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Fluorinated Poly(pentylene 4,4'-bibenzoate)s with Low Isotropization Temperatures and Unique Phase Transition Behavior

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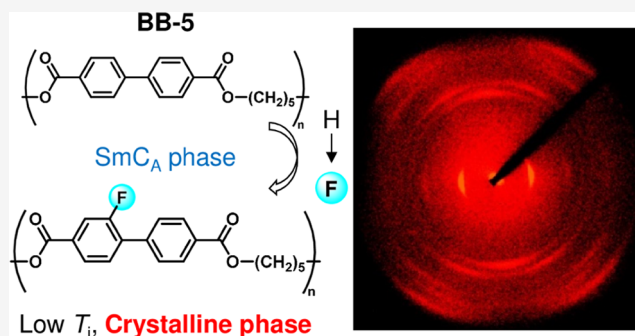


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ABSTRACT: Semiflexible main-chain thermotropic liquid crystalline polyesters (TLCPs) are applied in functional materials like heat dissipation sheets because of their good processability and ability to form nanostructures. Poly(pentylene 4,4'-bibenzoate) (**BB-5**) is a commonly investigated biphenyl-based TLCP that forms a SmC_A phase. The functional application of **BB-5** is limited by its high isotropization temperature (T_i). This study aims to lower the T_i of **BB-5** and enhance the processability of poly(pentylene 4,4''-terphenyl dicarboxylate) (**T-5**) compounds, which contain photo/electronic functional mesogens. Six fluorinated **BB-5** (**F-BB-5**) compounds are synthesized with T_i values at least 35 °C lower than that of **BB-5**. Fluorinated **T-5** (**F-T-5**) compounds show isotropic phases before thermal decompositions, unlike **T-5**. Moreover, **F-BB-5** and **F-T-5** form not only amorphous and SmC_A phases but also crystalline and higher-order smectic phases, which is unexpected from low-molecular-weight liquid crystals. Mesogenic fluorination of semiflexible TLCPs results in low T_i and even unique morphologies, opening up their new possibilities.

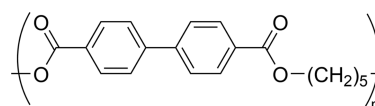


INTRODUCTION

Aromatic main-chain thermotropic liquid crystalline polyesters (TLCPs) have been extensively investigated owing to their versatile nano-orientation and excellent functionality that are derived from their nanostructures.^{1–5} They are used as (super)engineering plastics with high thermal stability, chemical resistance, and dimensional stability; low gas permeability; and excellent mechanical properties.^{6–8} However, wholly aromatic TLCPs often undergo thermal degradation prior to isotropization.⁹ Various aromatic polyesters have been developed to overcome this problem through copolymerization^{10–15} and the use of semiflexible TLCPs,^{16–18} in which flexible spacers are incorporated into the main chain.

Semiflexible TLCPs exhibit superior properties compared with those of wholly aromatic TLCPs, including melting points that are lower than degradation temperatures, low melt viscosities, and the presence of different phases, such as crystalline, smectic (Sm) A, SmC, SmC_A, and nematic phases.¹⁹ One of the most extensively investigated semiflexible TLCPs is poly(pentylene 4,4'-bibenzoate) (**BB-5**), which is a biphenyl-based polyester (Chart 1). Our group reported **BB-5** for the first time in 1988 and identified its LC structure.^{20,21} **BB-5** forms not only the SmC_A phase in a wide temperature range (86–196 °C) but also folded-chain lamellar structures. The phase transitions of **BB-5** that are associated with the odd–even effect of alkyl spacers have also been reported.²² Moreover, introducing methyl groups into the aliphatic spacer

Chart 1



of **BB-5** produces the SmC_A and SmA phase without crystallization.^{23,24} Various characteristics of **BB-5**, such as its nanostructure and thermally conductivity, were enhanced because of these properties.²⁵ However, **BB-5** suffers from a high isotropization temperature (T_i : 212 °C upon heating and 196 °C upon cooling), crystallization at 86 °C upon cooling, and the absence of various morphologies.

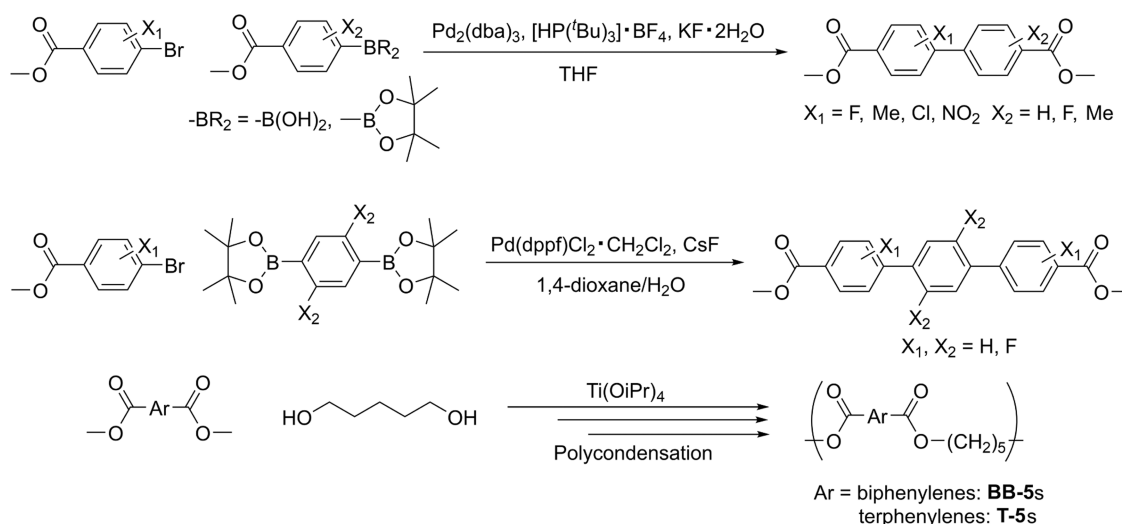
In recent years, nanostructures of block-copolymerized semiflexible TLCPs have been studied for their potential use in nano-applications because of their ability to be aligned by external fields, such as magnetic and electric fields or shear flow.^{26–29} However, new mesogenic strategies are required for the further performance improvement of these compounds, such as achieving better electrical/optical properties, lowering

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Scheme 1. Synthesis Methods of the Poly(pentylene 4,4'-bibenzoate) (BB-5) and Poly(pentylene 4,4'-terphenyl dicarboxylate) (T-5) Derivatives

processing temperatures, and generating higher-order smectic phases.^{30,31} Consequently, this study focuses on the lateral fluorine substitution of mesogenic biphenyl moieties. In low-molecular-weight liquid crystals, the lateral fluorine substitution of mesogens disrupts their molecular packing and reduces the number of intermolecular attractive forces, thus lowering the phase transition temperature and smectic phase stability.³² For LC molecules with small-aspect-ratio mesogens, such as biphenyl-based mesogens, lateral substitution sometimes suppresses liquid crystallinity.³³ However, the effects of small substituents, such as fluorine, in the mesogenic biphenyl moieties of semiflexible TLCPs on their liquid crystalline behavior have not been examined in detail,³⁴ although other laterally substituted semiflexible TLCPs have been reported previously.^{35,36}

In this study, we explore the effect of the lateral introduction of one or two fluorine groups on the phase transition behavior of BB-5. In addition, the effects of the methyl, chloro, and nitro groups were investigated. Note that nitro-substituted biphenyls sometimes form cubic phases.³⁷ Fluorinated BB-5s (F-BB-5s) have T_i that are at least 34.3 °C lower than that of BB-5 upon heating. F-BB-5s with SmC_A phases do not crystallize during the cooling process, and some of them exhibit lower T_i than those of conventional BB-5s. Lower melting and isotropic points allow not only a reduction in the processing temperature but also the use of more rigid calamitic molecules as mesogens. Based on these results, we have developed laterally fluorinated poly(pentylene 4,4'-terphenyl dicarboxylate)s T-5s (F-T-5s), which are semiflexible polyesters with terphenyl groups serving as mesogens, to obtain a smectic phase in the processable temperature range. For T-5, only the odd–even effect of the aliphatic spacer has been reported;³⁸ phase structure analysis and luminescence properties of T-5 have not been studied in detail. Moreover, our new F-BB-5s and F-T-5s exhibit unique phase transitions, which cannot be predicted from the properties of their low-molecular-weight liquid crystals.^{39,40} F-BB-5s contain not only the SmC_A and amorphous phases but also crystalline and higher-order smectic phases, and F-T-5s form the SmC_A , crystalline, and higher-order smectic phases. These results can be derived from polymer orientations.

EXPERIMENTAL SECTION

General Procedure for Biphenyl Monomers.⁴¹ Aryl bromide (1.0 equiv), aryl boronic acid (pinacol ester) (1.1–1.2 equiv), tris(dibenzylideneacetone)dipalladium(0) ($\text{Pd}_2(\text{dba})_3$) (0.5 mol %), tri-*tert*-butylphosphonium tetrafluoroborate (1.2 mol %), and potassium fluoride dihydride (3.3 equiv) were dissolved in tetrahydrofuran (THF). The mixture was heated at 50 °C for more than 4 h under an argon atmosphere, then filtrated through celite by diethyl ether. The filtrate was evaporated under reduced pressure. The residue was chromatographed over silica gel and recrystallized to give the target compound.

General Procedure for Terphenyl Monomers.⁴² Aryl bromide (2.1 equiv), aryl diboronic acid pinacol ester (1.0 equiv), [1,1-Bis(diphenylphosphino)ferrocene]palladium(0) ($\text{Pd}(\text{dppf})\text{Cl}_2 \cdot \text{CH}_2\text{Cl}_2$) (10 mol %), and cesium fluoride (6.0 equiv) were dissolved in 2:1 (v/v) 1,4-dioxane/water. The mixture was refluxed at 110 °C overnight under an argon atmosphere. Water was added, and the mixture was extracted with dichloromethane. The organic layer was washed with water three times, dried over MgSO_4 , filtrated, and evaporated under reduced pressure. The residue was chromatographed over silica gel and recrystallized to give the target compound.

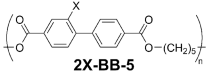
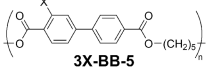
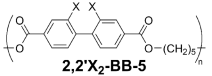
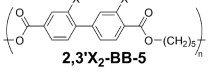
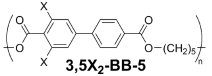
General Procedure for Melt Polycondensation.²⁰ Dimethyl ester monomer (1.0 equiv), 1,5-pentanediol (1.5 equiv), and tetrakis(isopropyl orthothitanate) ($\text{Ti}(\text{OiPr})_4$) (catalyst amount) were set in a test tube. The test tube was replaced with nitrogen for 10 min and melt-polymerized under flowing nitrogen at 220 °C for more than 4 h. The polymer was dissolved in chloroform and precipitated into methanol to give the target polymer.

Measurements. ^1H -NMR and ^{13}C -NMR spectra were recorded on BRUKER 500 (500 MHz) and JEOL 400 (100 MHz) spectrometers, respectively, for CDCl_3 solution using tetramethylsilane (TMS) as an internal standard. ^1H -NMR spectra are reported as follows: chemical shift (δ ppm), multiplicity (s = singlet, d = doublet, t = triplet, q = quartet, m = multiplet), integration, and coupling constants in units of Hz. ^{13}C -NMR spectra are reported as chemical shifts in ppm.

High-resolution (HR) EI mass spectra were recorded on a double-focusing mass spectrometer JEOL JMS-700, measured at Tokyo Institute of Technology Open Facility Center. This center is independent of our laboratory to ensure fairness.

Size exclusion chromatography (SEC) was performed using a JASCO system (PU-4185, AS-2055, CL-NETII/ADC, CO-4060, UV-4075, RI-4035) equipped with PS gel columns (SHODEX KF-405LHQ) with CHCl_3 as the eluent at a flowing rate of 0.2 mL min^{-1} at 40 °C after calibration with polystyrene standards.

Table 1. Thermal Properties and Number-Average Molecular Weights (M_n) of the Poly(pentylene 4,4'-bibenzoate)s (BB-5s) Determined Using DSC and SEC Measurements

Structure	Entry	T_g (°C)	Phase transition behavior ^a	ΔH_i (kJ mol ⁻¹)	M_n (g mol ⁻¹)
	BB-5 (1)	43	Cry (173 °C) SmC_A (212 °C) Iso	- ^b	- ^f
	2F-BB-5 (2)	-	Cry (131.9, 145.0 °C) Iso	5.78	3000
 2X-BB-5	2Me-BB-5 (3)	44.2	Amorphous	-	11000
	2Cl-BB-5 (4)	19.9	Amorphous	-	2400
	2NO₂-BB-5 (5)	53.0	Amorphous	-	5800
	3F-BB-5 (6)	25.3	SmC_A (177.7 °C) Iso	4.10	14500
 3X-BB-5	3Me-BB-5 (7)	6.3	No peak, but L-Sm	- ^c	1200
	3Cl-BB-5 (8)	40.1	Amorphous	-	3500
	3NO₂-BB-5 (9)	50.2	Amorphous	-	5100
	2,2'F₂-BB-5 (10)	36.5	Amorphous	-	8500
 2,2'X₂-BB-5	2,2'Me₂-BB-5 (11)	41.7	Amorphous	-	10300
	2,3'F₂-BB-5 (12)	11.4	No peak, but Cry or H-Sm	- ^c	1700
	2,3'X₂-BB-5				
	2,3'Me₂-BB-5 (13)	17.6	Amorphous	-	3200
 3,3'X₂-BB-5	3,3'F₂-BB-5 (14)	32.4	Cry or H-Sm (141.1, 149.6 °C) Iso	5.09 ^d	4400
	3,3'Me₂-BB-5 (15)	13.0	Amorphous	-	3700
	3,5F₂-BB-5 (16)	37.5	SmC_A (87.3 °C) Iso	- ^e	4900
 3,5X₂-BB-5					

^aPhase transition behavior. ^b ΔH_i of BB-5. ^c ΔH_i of 3Me-BB-5 and 2,3'F₂-BB-5. ^d ΔH_i of 3,3'F₂-BB-5. ^e ΔH_i of 3,5F₂-BB-5. ^f M_n of BB-5.

Polarizing optical microscopy (POM) was performed using a Leica DM2500P microscopy with a METTLER TOREDO FP82HT Hot stage controlled with the METTLER TOREDO Central Processor of the FP900 System at a rate of 10 °C min⁻¹.

Differential scanning calorimetry (DSC) was performed using PerkinElmer DSC 8500 equipment at a scanning rate of 10 °C min⁻¹ under a flow of dry nitrogen. Thermo-gravimetric analysis (TGA) was performed using a Rigaku Thermo Plus EVO2 series TG-DTA 8122 at a heating rate of 20 °C min⁻¹ under a flow of dry nitrogen. The initial mass of the samples was 5–7 mg.

Fiber samples were prepared by spinning the isotropic melt and quenching to room temperature. Press samples were prepared by sandwiching samples between two stainless used steel sheets (95 mm × 95 mm) with 30 kN force under vacuum, annealed at an appropriate temperature for 1 h, with a Manual Hydraulic Vacuum-Heating Press (Imoto Machinery Co., Ltd., Kyoto, Japan). Wide-angle X-ray diffraction (WAXD) patterns were obtained using a Bruker D8 DISCOVER equipped with a Vantec-500 detector and Cu K α radiation.

RESULTS AND DISCUSSION

We systematically designed and synthesized substituted bibenzoate monomers with fluorine atoms at the inner and/or outer lateral positions (Scheme 1) to investigate the relationship between the mesogenic structure and morphology of a substituted BB-5. For comparison, we also prepared bibenzoate monomers with methyl, chloro, or nitro groups at the lateral positions. Biphenyl bibenzoate monomers were prepared by Suzuki–Miyaura cross-coupling reactions from the corresponding bromobenzene and phenylboronic acid derivatives. Sixteen BB-5 derivatives (Table 1) were synthesized by

the Ti(OiPr)₄-catalyzed melt polycondensation of a corresponding monomer and 1,5-pentanediol (Scheme 1). Moreover, we developed fluorine-substituted terphenyl-based polyester T-5s to produce large-aspect-ratio mesogens with optical, luminescent, and electrical properties derived from the extended-conjugation length. All dimethyl ester monomers were measured by ¹H- and ¹³C-NMR except for 3,2',5',3'F₄-monomer that could not be measured by ¹³C-NMR due to its poor solubility in deuterated chloroform. ¹H-NMR and SEC measurements were performed on the chloroform-soluble polymers. DSC measurements were performed on the BB-5 derivatives at a rate of 10 °C min⁻¹ (Figures 1a and S1–S15). POM observations were performed on all polymers except BB-5 at a rate of 10 °C min⁻¹ (Figures 1c and S16–S25). WAXD measurements were performed on polymers that showed phase transition behaviors in the POM measurements (Figures 1d, 2a, and Figures S26–37). TGA measurements were performed on four of the T-5 derivatives at a rate of 20 °C min⁻¹ (Figures S38–41).

Laterally Substituted BB-5s. In previous studies, we demonstrated the specificity of BB-5 (1) compared to BB-6.²¹ BB-6 has a SmA phase and approximately parallel mesogens and alkyl chains. This characteristic is similar to the low-molar-mass SmA phase.⁴³ In contrast, 1 forms the SmC_A phase by tilting mesogens in the polymer chains by approximately 25° with respect to the chain axis. The mesogens adjacent along the chain tilt in opposite directions. This zigzag conformation is due to the effect of connecting mesogens via most extended spacers, which do not occur in the low-molar-mass SmC phase.

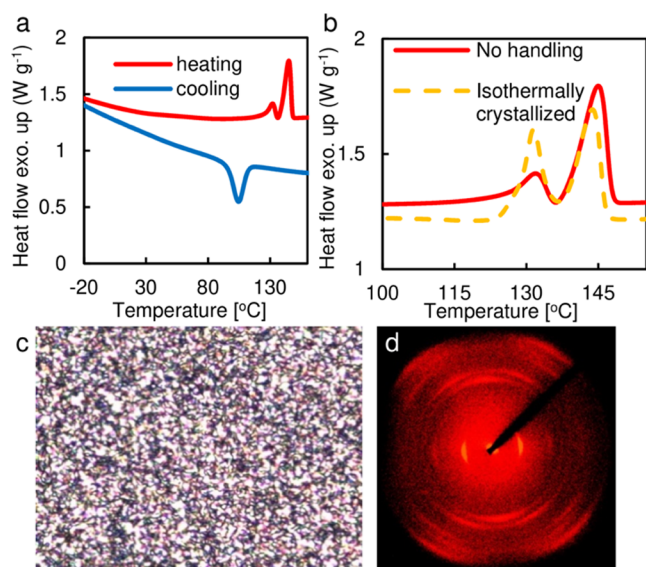


Figure 1. DSC thermograms of (a) **2** and (b) its unhandled and isothermally crystallized derivatives measured at a rate of $10\text{ }^{\circ}\text{C min}^{-1}$. (c) POM image of **2** being cooled at $95\text{ }^{\circ}\text{C}$. (d) WAXD pattern for the fiber sample of **2** annealed at $135\text{ }^{\circ}\text{C}$ for 1 h. The fiber axis lies in the horizontal direction.

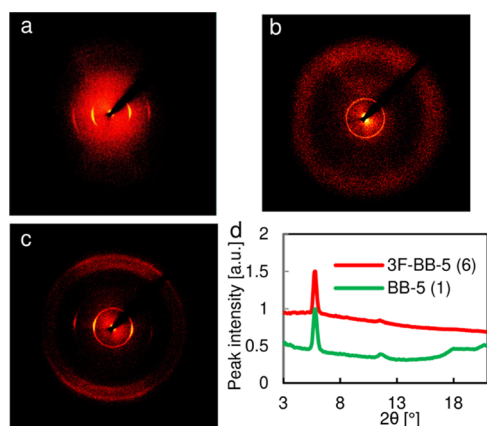


Figure 2. WAXD patterns for the (a) fiber sample of **6** annealed at $60\text{ }^{\circ}\text{C}$ for 1 h, (b) bulk sample of **6** measured at $150\text{ }^{\circ}\text{C}$, and (c) fiber sample of **1** annealed at $190\text{ }^{\circ}\text{C}$ for 1 h. The fiber axis lies in the horizontal direction. (d) WAXD intensity profiles measured for **1** and **6**. The circles show secondary reflections.

Alkylene 4,4'-biphenylates do not have liquid crystallinity,⁴⁴ while **1** forms a liquid crystalline phase because of the extension of the aspect ratio to the ester group. In the condensed phase, the mesogenic ester groups of **BB-5** become coplanar because of the immobilization of the polymer and the aspect ratio increases, resulting in the development of liquid crystalline properties. The phase transition behavior of **1** in ref **22** is as follows: during the heating process, **1** melts at $173\text{ }^{\circ}\text{C}$ to form SmC_A , which transforms into the isotropic liquid at $212\text{ }^{\circ}\text{C}$. In this study, **F-BB-5s** have T_i values that are at least $34\text{ }^{\circ}\text{C}$ lower than that of **BB-5**. Furthermore, the morphologies, such as the amorphous, SmC_A , crystalline, and higher-order smectic phases, depend on the position and number of fluorine atoms. In contrast, almost all of the other group (methyl, chloro, and nitro)-substituted **BB-5s** are amorphous. The glass-transition temperatures (T_g), phase transition behaviors, transition enthalpies (to isotropic) (ΔH_i),

and number-average molecular weights (M_n) of each **BB-5** derivative during heating are listed in Table 1, and the d -spacings are listed in Table S1. The liquid crystallinity of polyesters is independent of molecular weight, as previously reported.⁴⁵ Therefore, the relationship between the number-average molecular weight (M_n) and liquid crystallinity has not been considered.

The ΔH_i of **1** was reported in ref **22** (5.31 kJ mol^{-1}) and ref **20** (3.85 kJ mol^{-1}) during cooling. We resynthesized and evaluated **1** to compare its ΔH_i to that of the substituted **BB-5s** under the same measurement conditions as this study (Figure S1). However, the results differ from those in the previous study.²² Compound **1** has phase transition peaks at 170.1 and $198.2\text{ }^{\circ}\text{C}$ during the heating process, which overlapped. Although we could not compare the ΔH_i of the substituted **BB-5s** with that of **1**, the ΔH_i data are shown because it is a critical parameter in the liquid crystalline molecules.

2X-BB-5. **2F-BB-5 (2)** shows two phase transition peaks at 131.9 and $145.0\text{ }^{\circ}\text{C}$ upon heating with ΔH values of 1.71 and 5.78 kJ mol^{-1} , respectively (Figure 1a). The WAXD pattern shows some peaks in the wide-angle region (Figure 1d), and the d -spacing is $14.5\text{ }\text{\AA}$, which is $0.9\text{ }\text{\AA}$ shorter than that of **1**. Based on the many diffraction peaks in the WAXD pattern and the short d -spacing, it is concluded that **2** is a crystalline polymer. For a more detailed discussion of the phase transition behavior, we also considered the DSC measurement of an isothermally crystallized sample of **2** that was annealed at $115\text{ }^{\circ}\text{C}$ for 22 h at a heating rate of $10\text{ }^{\circ}\text{C min}^{-1}$ (Figure 1b). The higher-temperature peaks in the melting thermograms of the untreated and isothermally crystallized samples of **2** do not significantly differ. However, the lower-temperature peak of the isothermally crystallized sample is larger and sharper than that of the untreated sample. Moreover, the two phase transition temperatures are almost the same. These results indicate that **2** has two crystalline structures with different crystallization rates. However, the WAXD patterns of the fiber sample of **2** at $65\text{ }^{\circ}\text{C}$ (Figure S26) are similar to those at $135\text{ }^{\circ}\text{C}$. Therefore, we could only observe a crystal structure with a higher melting peak. **2Me-BB-5 (3)**, **2Cl-BB-5 (4)**, and **2NO₂-BB-5 (5)** exhibit no phase transition peaks in the DSC measurements. Moreover, **3** and **4** show no birefringence in the POM observations. Thus, **3** and **4** are amorphous. In contrast, **5** appears to form a phase, as observed by POM (Figure S16). However, most of the field of view is dark, and the WAXD pattern of the fiber sample includes only a circular halo in the wide-angle region (Figure S27), indicating that **5** is also amorphous.

3X-BB-5. **3F-BB-5 (6)** shows a phase transition at $177.7\text{ }^{\circ}\text{C}$ with a ΔH_i of 4.10 kJ mol^{-1} upon heating. The WAXD pattern for the fiber sample of **6** shows peaks at $2\theta = 5.8$ and 11.5° . In addition, no halo is observed in the wide-angle region (Figure 2a), whereas for the bulk sample of **6**, a halo is observed (Figure 2b). These primary and secondary reflections are associated with a layered structure with a d -spacing of $15.2\text{ }\text{\AA}$. This d -spacing coincides with that measured for **1** ($15.3\text{ }\text{\AA}$) (Figure 2c,d). Considering the odd–even effect of the alkyl spacer, it is reasonable to assume that **6** exhibits the SmC_A phase, although no halo is present in the wide-angle region in the fiber sample. The lack of halo observations is due to the lack of sample thickness. The T_i value of **6** is $34.3\text{ }^{\circ}\text{C}$ lower than that of **1** and only demonstrated a SmC_A phase. The absence of crystallization is advantageous for material

processing. The ΔH_i of **6** is 1.6 kJ mol^{-1} smaller than that of **2**. In general, the ΔH_i of the transition from the lower-order smectic to the isotropic phase is smaller than that of the crystalline/higher-order smectic to the isotropic phase, which supports that **2** is a crystalline polymer. **3Me-BB-5** (**7**) also shows some phases, as revealed by the DSC measurements (Figure S6) and POM observations (Figure S18). The oriented sample of **7** could not be prepared because of its low T_g . Thus, we performed a WAXD measurement for an unoriented sample of **7** sandwiched between thin glass films. The WAXD pattern demonstrates a primary reflection and a halo in the wide-angle region with a d -spacing of $15.0 \text{ \AA}$ (Figure S28). Therefore, **7** forms the SmC_A phase owing to its d -spacing. **3Cl-BB-5** (**8**) and **3NO₂-BB-5** (**9**) show no phase transition behaviors in either the DSC measurements or POM observations and are determined to be amorphous.

2,2'X₂-BB-5. Neither **2,2'F₂-BB-5** (**10**) nor **2,2'Me₂-BB-5** (**11**) show phase transition peaks or formations. Compounds **10** and **11** are amorphous.

2,3'X₂-BB-5. Although **2,3'F₂-BB-5** (**12**) shows no phase transition peaks on the DSC chart (Figure S11), the POM observation reveals that **12** forms some phase when it is left at room temperature (Figure S20). The WAXD pattern shows many peaks at $2\theta = 1.8, 5.9, 14.7, 16.2$, and 20.3° (Figures S30 and S31, Table S2), and the d -spacing of **12** was $15.0 \text{ \AA}$, $0.4 \text{ \AA}$ shorter than that of **1**. Therefore, **12** exhibits a crystalline or higher-order smectic phase. In contrast, **2,3'Me₂-BB-5** (**13**) is amorphous because its phase transition peak and birefringence are not observed.

3,3'X₂-BB-5. **3,3'F₂-BB-5** (**14**) demonstrates a double-melting behavior (Figure S13a). The WAXD pattern shows three peaks that are different from those of SmA , SmC , and SmC_A (Figures S32 and S33). To analyze the double-melting behavior, we performed the same measurements on **14** as on **2** (Figure S13b). The isothermally crystallized sample annealed at 125°C does not show a lower melting peak, indicating that the lower melting peak is derived from a thermally unstable organization with poor integrity upon rapid cooling. This phenomenon is observed in several (semi)crystalline polymers.⁴⁶ Moreover, the total ΔH_i of **14** is 0.99 kJ mol^{-1} larger than that of **6**, reinforcing the validity that **14** forms a three-dimensional ordered structure. Therefore, based on the similarity to **12**, it is assumed that **14** forms a crystalline or higher-order smectic phase. **3,3'Me₂-BB-5** (**15**) also has a phase transition peak at 32.7°C upon heating but the POM observation shows a dark region. Therefore, **15** is amorphous.

3,5X₂-BB-5. **3,5F₂-BB-5** (**16**) exhibits both endothermic and exothermic peaks upon heating (Figure S15). The T_i upon heating is 87.3°C , which is 124.7°C lower than that of **1**. The WAXD pattern is the same as that of **6**, with a d -spacing of $15.5 \text{ \AA}$ (Figure S34). Compound **16** has the SmC_A phase and the lowest T_i among **F-BB-5s**, with a T_g above room temperature, and does not crystallize.

The T_i values of **F-BB-5s** are lower than that of **1**. Compounds **6**, **10**, and **16** do not exhibit crystalline phases, unlike **1**. Moreover, **10** does not exhibit a smectic phase, suggesting that these polyesters are affected by the steric hindrance of fluorine. In contrast, **2** is a crystalline polymer, and both **12** and **14** forms crystalline or higher-order smectic phases. In biphenyl-type low-molecular-weight liquid crystals, the introduction of fluorine into lateral positions decreases the melting and isotropic points, which sometimes leads to a reduction in the liquid crystallinity. In other words, the

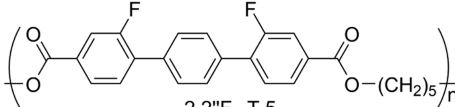
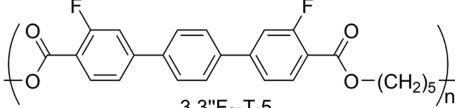
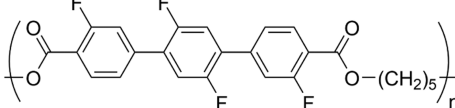
molecular orientation is reduced and the crystal structure formation is more complicated. However, in **BB-5**, the formation of higher-order structures predominates depending on the positions of the fluorine substitutions. Polymer immobilization plays a significant role in this process and provides unpredictable results for small liquid crystalline molecules.

Of the methyl-, chloro-, and nitro-substituted derivatives of **BB-5**, only **7** shows phase formation. However, the average molecular weight of **7** is 1200 (Table 1), which is considerably lower than that of the other polymers and may influence the phase formation. The aspect ratio of a biphenyl mesogen is insufficient when the substituents are larger than the methyl group, and most polymers are amorphous.

In general, the crystalline and higher-order smectic structures of organic compounds are highly dependent on thermodynamic factors. In addition, a detailed analysis of these polymers is complicated. Press samples of **12** and **14** prepared in this study are not uniaxially oriented, and their WAXD patterns show circular reflection peaks. Moreover, polymers rarely form tetragonal or hexagonal crystals, and the crystal lattice of only a few polymers can be determined by their d -spacings despite being uniaxially oriented. Therefore, a detailed analysis of the crystalline or higher-order smectic structures of **2**, **12**, and **14** is difficult. In this study, we do not focus on a detailed consideration of their structures and instead investigate the development of a new method for controlling the morphology of **BB-5**.

F-T-5s. The phase transition behavior and degradation temperatures at a 10% weight loss (T_{10}) of **T-5** (**17**) and **F-T-5s** are shown in Table 2. The POM images of these

Table 2. Thermal Properties of **T-5s**

 2,2'F₂-T-5		
 3,3'F₂-T-5		
 3,2',5',3'F₄-T-5		
entry	phase transition behavior ^a	T_{10} ($^\circ\text{C}$) ^b
T-5 (17)	Cry or Sm, decomposed	407.3
2,2'F₂-T-5 (18)	Cry or H-Sm (290.1°C) Iso	404.5
2,3'F₂-T-5 (19)	SmC _A (311.3°C) Iso	407.4
3,2',5',3'F₄-T-5 (20)	Cry (240.4°C) Iso	398.2

^aRecorded from POM observation at a heating rate of $10^\circ\text{C min}^{-1}$.

^bRecorded from TGA measurements at a heating rate of $20^\circ\text{C min}^{-1}$.

compounds are shown in Figures S23–S25 and their WAXD patterns are shown in Figures 3 and S35–S37. In the POM observations, **17** thermally decomposes before the isotropization, whereas all of **F-T-5s** exhibit T_i values during the heating process. All **T-5s** exhibit similar T_{10} values at approximately 404°C , which was higher than the T_i values of **F-T-5s**. Thus, lateral fluorine substituents in the mesogens allow the

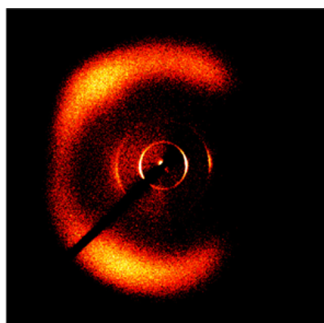


Figure 3. WAXD pattern for the press sample of 3,3''F₂-T-5 (**19**) annealed at 200 °C for 1 h, as viewed from the edge. (The right side of the WAXD pattern is dark owing to the malfunction of the measurement equipment).

processing of semiflexible TLCPs whose mesogens are electrically or optically functional molecules, such as terphenyl.

The WAXD pattern of 3,3''F₂-T-5 (**19**) shows a typical SmC_A pattern with *d*-spacing of 18.2 Å (Figures 3 and S37). Meanwhile, 2,2''F₂-T-5 (**18**) shows a pattern different from those of the SmA, SmC, and SmC_A phases (Figures S35 and S36). The *d*-spacing of **18** is 19.7 Å, which is 1.5 Å longer than that of **19**. However, **18** can have a crystalline or higher-order smectic phase for the following reasons: there are more than three peaks present in the WAXD pattern of **18**, its solubility in chloroform is relatively poor compared with that of **19**, or the sample of **18** is tough. 3,2',5',3''F₄-T-5 (**20**) is nearly insoluble in chloroform and the sample is also tough. Furthermore, **20** shows a phase formation that resembled needle growth, as observed in POM at 230 °C during cooling (Figure S25). These results indicate that **20** forms a crystalline phase.

Conformational Study of Fluorinated Mesogenic Moieties. The behavior of bibenzoate in the monomer and polymer was theoretically analyzed. We calculated the rotational energy of an ester group and the molecular orbital of bibenzoate with a dihedral angle (φ) between the ester and biphenyl groups (Figure 4a) of 0 and 90° using density functional theory at the ω B97XD/6-311G(d, p) level of theory.⁴⁷ The rotational barrier of $\varphi = 60^\circ$ is approximately 20 kJ mol⁻¹ and that of $\varphi = 90^\circ$ is approximately 30 kJ mol⁻¹. However, since dialkyl bibenzoates are relatively large, they

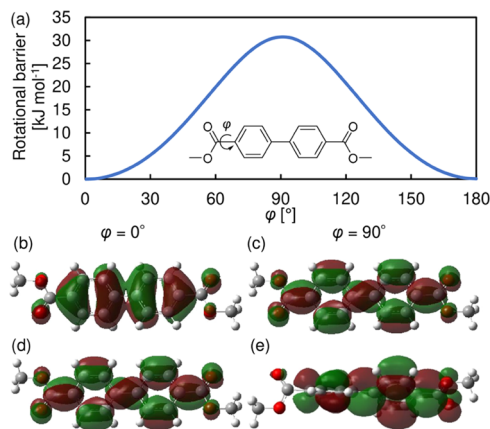


Figure 4. (a) Rotational barrier of bibenzoate. (b) HOMO and (c) LUMO of bibenzoate at $\varphi = 0^\circ$. (d) HOMO and (e) LUMO of bibenzoate at $\varphi = 90^\circ$.

have T_i values higher than 40 °C.⁴⁴ Thus, the ester group of bibenzoate can rotate in an isotropic phase. In addition, in $\varphi = 0^\circ$, the π -electron cloud extends from the biphenyl group to the ester group (Figure 4b,c), whereas in $\varphi = 90^\circ$, this π -electron cloud extension disappears (Figure 4d,e). These results suggest that, in a monomer, the π -conjugation of bibenzoate does not extend stably to ester groups because of their rotation or the sufficient existence of $\varphi =$ nonzero conformation, and the mesogen is only biphenyl. However, owing to polymerization, the existence probability of the parallel state between ester and biphenyl groups is increased, or ester groups are immobilized in the parallel state due to the strengthened viscosity in condensed phases. Therefore, the ester groups form pseudo-conjugates to which the mesogens are extended. Pseudo-conjugation of mesogens in the liquid crystal and crystalline states of main-chain liquid crystal polymers have been reported for several mesogens with ester and urethane linkages.^{48,49} However, the electronic state and structure of mesogens in the liquid crystalline state have not been discussed in detail. The bibenzoate monomer is not liquid crystalline,⁴⁴ only polymer **BB-5** is. The present calculations indicate that the pseudo-conjugation effect plays an important role in the liquid crystallinity of **BB-5**. Thus, it can be reasonably explained that the liquid crystalline nature of polybibenzoate appears due to polymeric effects.

For a more detailed discussion, we calculated the structural optimizations and rotational barriers of fluorinated biphenyls using the above theory, where (s) and (o) denote the same and opposite sides, respectively (Figures 5, S42, and S43). In liquid crystalline phases, mesogens are not completely confined and exhibit molecular motions, such as axis rotations. However, the molecular motions of the mesogenic ester groups are small in the crystalline and smectic phases of **BB-5**. Therefore, we hypothesized that the structures of the ester groups could be fixed and focused only on the biphenyl units.

The dihedral angles (ψ) of 3F-biphenyl and 3,5F₂-biphenyl are nearly identical to those of biphenyl, indicating that **6** and **16** exhibit SmC_A like **1**. 3,3''F₂(s)/(o)-biphenyl also exhibit a similar ψ to that of biphenyl. However, **14** forms a crystalline or higher-order smectic phase. Using the above theory, we also calculated the HOMO and dipole moments of bibenzoate (**A**) and 3,3''F₂-bibenzoates (**B–D**) in each conformation in the ground state (Table 3 and Figure S44).

The HOMOs of **B–D** are not significantly different from that of **A**, except for the electron clouds that extend to fluorine. The dipole moments differ depending on the conformation. **A**, **C**, and **D** only have the *z*-direction component, whereas **B** also has an all-direction component, with the *y*-direction component being the largest. The dipole moment values of **B** and **C** are one-and-a-half times larger than that of **A**, while that of **D** is one-third smaller than that of **A**. **D** has the lowest total energy among **B**, **C**, and **D**, and the energy difference between **B** and **D** is 3.65 kJ mol⁻¹ and that between **C** and **D** is 7.06 kJ mol⁻¹ (Table S4). These energy differences are so small that 3,3''F₂-bibenzoate can form all of the conformations of **B**, **C**, and **D**, even though the optimum structure of 3,3''F₂-bibenzoate is **D**. The large dipole moment of 3,3''F₂-bibenzoate, depending on the conformation, enhances the dipole–dipole interaction between the mesogens and would allow the formation of a higher-order structure. The ψ values of 2F-biphenyl and 2,3''F₂(s)/(o)-biphenyl are approximately 5° larger than that of biphenyl. The combination of the fact that the electronic interactions between the mesogens become

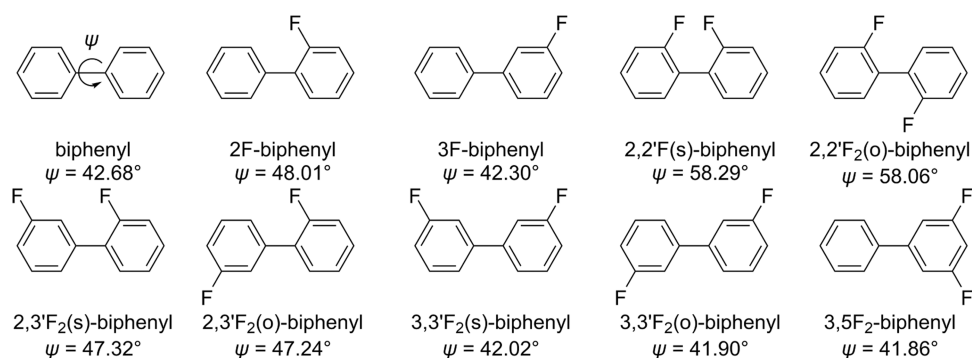


Figure 5. Chemical structures and calculated dihedral angles (ψ) of biphenyl and fluorinated biphenyls.

Table 3. Calculated Dipole Moments of Bibenzoate and 3,3'-F₂-Bibenzoate in Each Conformation^a

entry	x [D]	y [D]	z [D]	total [D]
A	0	0	1.22	1.22
B	-0.21	-2.11	-1.08	2.38
C	0	0	1.90	1.90
D	0	0	-0.37	0.37

^ax: short-axis direction; y: long-axis direction; z: perpendicular direction of π -plane.

not too small and the larger rotation barriers of $\psi = 0$ and 180° than that of biphenyl (Table S3) may allow **2** and **12** to form crystalline and higher-order smectic phases. The ψ of 2,2'-F₂(o)-biphenyl is 50.86° and that of 2,2'-F₂(s)-biphenyl is 58.29° . The large mesogenic twist results in very few electrical interactions. Consequently, **10** becomes an amorphous polymer. We also calculated the optimized conformation of F-T-Ss; the results have the same trend as to those of F-BB-Ss (Figure S45).

Summary for Morphologies. This study is summarized as follows (Figure 6). In condensed phases, such as liquid crystals and films, ester groups form pseudo-conjugates. As a result, the aspect ratio of the mesogen increases from biphenyl (monomer) to bibenzoate (polymer). Regarding the aspect ratio of BB-5, one or two lateral fluorine substitutions in the mesogen do not affect its orientation ability of BB-5. F-BB-5s possess lower T_i contents than BB-5. Lowering the T_i improves the processability of the polymer and the construction of nanostructures using a block polymer. Moreover, F-T-Ss with a 4,4''-terphenyl diester as the mesogen exhibit T_i values below their T_{10} , which is different from that of T-5. This result allows electrical and optical functional molecules to be treated as mesogens in semiflexible TLCPs.

Polymers tend to show crystalline or higher-order smectic phases (**2**, **12**) when fluorine is introduced into the mesogen on the inner side, and polymers are subject to smectic properties similar to BB-5 (**6**, **16**) when introduced on the outer side. This trend has also been observed for F-T-Ss (**18**, **19**, **20**). However, when the planarity of the mesogen is severely disrupted, the polymers become amorphous, such as

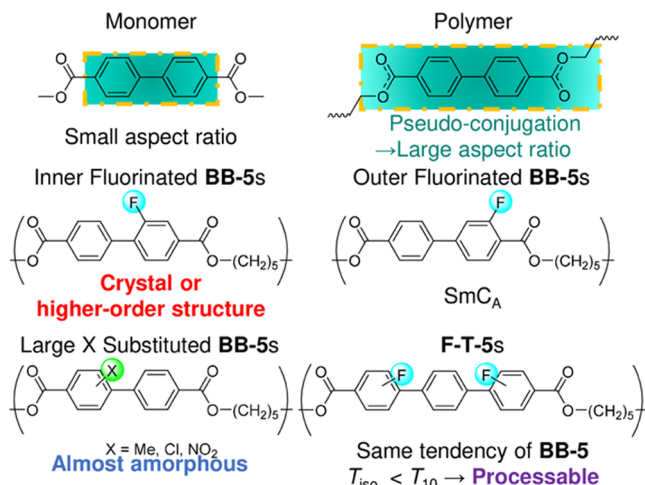


Figure 6. Summary of the properties of substituted poly(pentylene 4,4'-bibenzoate) (BB-5) and fluorinated poly(pentylene 4,4''-terphenyl dicarboxylate) (T-5).

10 with fluorine substitutions at the 2- and 2'-positions of the biphenyl mesogen. However, if the substitution sizes are larger than the methyl group, nearly all of the substituted BB-5s are amorphous because the substitution sizes are too large relative to the mesogen of BB-5. Some F-BB-5s form structures different from SmCA. However, we cannot provide a sufficient argument for this phenomenon. Moreover, the construction of a different phase structure expands the variety of polycrystals that can be formed from liquid crystal phases and contributes to the improvement of devices with higher-order structures, such as organic liquid crystal semiconductors.³⁰

CONCLUSIONS

We decreased the T_i of polybibenzoate BB-5 by at least 34.3°C by introducing fluorine groups at the appropriate lateral positions of biphenyl mesogens. The introduction of lateral fluorine substituents usually decreases the melting and isotropic points and smectic phase stabilities of low-molecular-weight liquid crystals. The same behavior is observed for F-BB-5s. In particular, 3,5F₂-BB-5 exhibits T_i at 87.3°C and contained only the SmCA phase. These properties overcome the major limitations of conventional BB-5 derivatives. We adjusted this strategy for terphenyl mesogen-based liquid crystalline T-5s by decreasing the T_i of F-BB-5s. F-T-5s exhibit T_i below their thermal degradation temperatures. Lateral fluorine substituents in mesogens enable the processing of T-5 and demonstrate the advantages of lateral

fluorination in semiflexible TLCPs. These results may facilitate the practical application of main-chain TLCPs with electrically and optically functional molecules serving as mesogens. They can also be used to overcome the disadvantages of the nanostructures formed by liquid crystalline block polymers and increase the choice of mesogens. In contrast to the T_i , we observe unexpected morphologies for F-BB-Ss and F-T-Ss. 3F-BB-5, 2,3'-F₂-BB-5, 3,3'-F₂-BB-5, 2,2''-F₂-T-5 and 3,2',5',3''-F₄-T-5 exhibit only crystalline or higher-order smectic phases. The introduction of fluorine into the inner lateral positions of the mesogens promotes the formation of crystalline or higher-order smectic phases. This feature differs from that of low-molecular-weight liquid crystals; however, the reason for this phenomenon remains unclear. It is hypothesized that the steric hindrance of fluorine causes small misalignments (distortions) in the mesogen structure compared with that of BB-5, and the spacers are adjusted to resolve this issue, resulting in a dense state. The unique behavior observed in this study is due to the immobilizing effect of the polymer.

In summary, the underlying mechanism of the formation of unique morphologies is elucidated, introduction of fluorine at appropriate positions is investigated, and new functional groups for TLCP improvement are developed. Lowering T_i and forming even crystalline or higher-order smectic phases have great advantages for improving functional materials/devices, such as heat sinks,⁵⁰ liquid crystalline organic semiconductors,³⁰ high-birefringence materials,^{51,52} and high-strength materials.⁵³ Thus, Liquid crystalline polyesters based on fluorinated oligophenylene mesogens have potential applications in a wide range of materials and devices.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.macromol.3c00773>.

Synthetic procedures in detail; ¹H- and ¹³C-NMR spectra; HRMS spectra; SEC curves; DSC thermograms; POM images; WAXD patterns; TG curves; and computational results (PDF)

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Notes

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