Luminescence Tuning of Fluorinated Bistolanes via Electronic or Aggregated-Structure Control

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Abstract: Organic luminescent materials have a wide range of practical applications, but the understanding of the relationship between molecular structure and luminescent behavior is lacking. Herein, we synthesized fluorinated bistolanes with an electron-donating alkoxy substituent at one terminal and an electron-withdrawing substituent at the other to realize systematic control of the electron-density distribution. Evaluation of the phase transition behavior revealed that most of the fluorinated bistolanes showed liquid-crystalline (LC) behavior, with the phase transition temperature depending on the terminal substituents. Additionally, the fluorinated luminophores displayed intense photoluminescence (PL) in solution and in their crystal phases. Remarkably, the PL color shifted dramatically depending on the dipole moment (μ||) along the long molecular axis; thus, PL tuning can be achieved through electronic modulation by precise control of the μ|| of the luminophore. Interestingly, in the LC phases under thermal conditions, the maximum PL band shifted by 0.210 eV upon phase transition from the crystal to smectic A LC phases, indicating that PL tuning can also be achieved by controlling the aggregated structure. These results offer a new molecular design for easily tunable PL materials using the molecular properties or external stimuli for promising applications, including light-emitting displays and PL sensing materials.

Keywords: fluorine; π-conjugated molecules; photoluminescence; liquid crystals; electron-density distribution

1. Introduction

Organic luminescent materials can be broadly classified into two groups: solution-state and solid-state luminophores. Solution-state luminophores, such as rhodamine and fluorescein, are applicable in biomedical imaging and optical probes [1–4], whereas solid-state luminophores are typically employed in lighting devices, light-emitting diodes, and electroluminescence devices [5–9]. To realize practical applications, significant efforts have been devoted to the discovery of novel luminescent materials, and as a result, various kinds of luminescent molecules have been developed [10,11]. However, there have been few systematic studies on the relationship between molecular structure and luminescent behavior, even though luminescence tuning is a key feature for practical light-emitting applications.

Over the past few decades, numerous fluorine-containing organic materials have been developed because chemical reactivity and material properties are dramatically influenced by the unique characteristics of fluorine, e.g., the largest electronegativity of all elements, the second smallest atom...
next to hydrogen, and the strong dissociation energy of C–F bonds [12,13]. Our group has extensively studied efficient synthetic protocols for organofluorine molecules containing CF3 groups [14,15] and CF2CF2 fragments [16,17]. More recently, we have also developed a significant interest in functional materials, e.g., liquid-crystalline (LC) and photoluminescence (PL) materials, based on fluorinated π-conjugated molecules such as pentafluorophenyl-containing tolane [18] and bistolane molecules [19–23]. Intriguingly, bistolanes 1aA and 1bA (Figure 1a) exhibited blue-light PL in solution as well as in crystalline (Cr) phases [23]. Remarkably, 1aA and 1bA in the LC phases under thermal conditions also displayed PL behavior, with the PL intensity and color reversibly changing during the thermal Cr ⇄ LC phase transition.

Previous studies by our research group on fluorinated bistolane-type luminophores have demonstrated that the PL behavior depends on the electron-density distribution or aggregated structure. Herein, we describe the molecular design and synthesis of these fluorinated bistolane-type luminophores as well as their LC and PL behaviors. Additionally, the effect of precisely modulating the electron-density distribution or aggregated structure on the PL behavior is discussed.

2. Materials and Methods

2.1. General

1H and 13C NMR spectra were recorded using a Bruker AVANCE III 400 NMR spectrometer (1H: 400 MHz and 13C: 100 MHz, Bruker Corporation, Rheinstetten, Germany) in chloroform-d (CDCl3) solution, and the chemical shifts are reported in parts per million (ppm) using the residual proton in the NMR solvent (δH = 7.26 ppm for CHCl3 and δC = 77 ppm for CDCl3). 19F NMR (376 MHz) spectra were recorded using a Bruker AVANCE III 400 NMR spectrometer in CDCl3 solution with CFCl3 (δF = 0 ppm) as an internal standard. Infrared (IR) spectra were acquired via the KBr method using a JASCO FT/IR-4100 type A spectrometer (JASCO Corporation, Tokyo, Japan); all spectra are reported in wavenumbers (cm⁻¹). High-resolution mass spectroscopy (HRMS) was performed using a JEOL JMS-700MS spectrometer (JEOL Ltd., Tokyo, Japan) via fast atom bombardment (FAB). All reactions were performed using dried glassware with a magnetic stirrer bar. All chemicals were of reagent grade and, if necessary, were purified in the usual manner prior to use. Column chromatography was conducted on silica gel (Wako-gel@60N, 38–100 μm, FUJIFILM Wako Pure Chemical Corporation, Osaka, Japan) and thin layer chromatography (TLC) was performed on silica gel TLC plates (Merck, Silica gel 60F254, FUJIFILM Wako Pure Chemical Corporation, Osaka, Japan).
2.2. Synthesis

Fluorinated bistolanes 1aB–D and 1bB–D were synthesized according to the synthetic procedure shown in Scheme 1. Acetoxy (AcO)-substituted bistolanes 1aB and 1bB were prepared by the Pd(0)-catalyzed Sonogashira cross-coupling reaction of 4-[2-(4-alkoxyphenyl)ethyn-1-yl]phenylacetylene 3a or 3b, readily accessible from the corresponding (4-alkoxyphenyl)acetylene 2a or 2b [21,23], with 4-acetoxy-2,3,5,6-tetrafluoriodobenzene (B). Derivatives 1aC or 1bC with a trifluoromethyl (CF3) group and 1aD or 1bD with a cyano (CN) group were prepared by the nucleophilic addition of a lithium acetylide, generated from 3a or 3b and n-butyllithium, to perfluorotoluene (C) or perfluorobenzonitrile (D), followed by elimination of lithium fluoride. The following sections provide typical synthetic procedures for the preparation of bistolanes 1aB–D with a methoxy (CH3O) substituent.

Scheme 1. Synthetic procedures for fluorinated bistolanes 1aB–D and 1bB–D.

2.3. Preparation of AcO-Substituted Bistolane 1aB

In a 50 mL two-necked round-bottomed flask were placed Cl2Pd(PPh3)2 (0.051 g, 0.050 mmol), PPh3 (0.015 g, 0.050 mmol), CuI (0.031 g, 0.10 mmol), 4-[2-(4-methoxyphenyl)ethyn-1-yl]phenylacetylene (3a, 0.24 g, 1.0 mmol), and 4-acetoxy-2,3,5,6-tetrafluoriodobenzene (B, 0.41 g, 1.2 mmol) in Et3N (10 mL) and THF (5 mL). The mixture was stirred at reflux temperature overnight. The precipitate formed in the reaction mixture was separated by atmospheric filtration and the filtrate was poured into saturated aqueous NH4Cl solution (20 mL). The crude product was extracted using AcOEt (20 mL, three times) and washed with brine (20 mL, once). The organic layer was dried over anhydrous Na2SO4, filtered, and concentrated using a rotary evaporator. The resultant crude product was purified by silica-gel column chromatography using hexane/AcOEt (20:1, v/v) as an eluent, providing coupling product 1aB (0.21 g, 0.48 mmol) in 48% yield as a white solid. This product was recrystallized by slow evaporation from CH2Cl2/MeOH (1:1, v/v) to obtain 1aB in crystalline form.

2.3.1. 1-Acetoxy-2,3,5,6-tetrafluoro-4-[2-[4-(4-methoxyphenyl)ethyn-1-yl]phenyl]ethyn-1-yl]benzene (1aB)

Yield: 48% (white solid); mp: 159 °C determined by differential scanning calorimetry (DSC); 1H NMR (CDCl3): δ 2.42 (s, 3H), 3.84 (s, 3H), 6.88–6.91 (m, 2H), 7.47–7.57 (m, 6H); 13C NMR (CDCl3): δ 20.0, 25.4, 75.3 (t, J = 4.0 Hz), 87.8, 92.3, 101.6 (t, J = 3.8 Hz), 102.3 (t, J = 18.0 Hz), 114.0, 114.9, 121.0, 125.1, 129.3 (t, J = 14.7, 2.9 Hz), 131.6, 132.0, 133.3, 139.5–142.2 (m, CAr-F, 1C), 145.6–148.4 (m, CAr-F, 1C), 160.1, 166.6; 19F NMR (CDCl3): δ = 137.34 (dd, J = 23.9, 9.9 Hz, 2F), -153.66 (dd, J = 23.9, 9.9 Hz, 2F); IR (KBr): ν 3262, 2210, 1792, 1596, 1491, 1491, 1394, 1315 cm−1; HRMS (FAB+) m/z [M]+ calcd for C25H14O3F4: 438.0879; found: 438.0881.
2.3.2. 1-Acetoxy-2,3,5,6-tetrafluoro-4-[2-[4-(4-hexyloxyphenyl)ethyn-1-yl]phenyl]ethyn-1-yl]benzene (1bB)

Yield: 54% (white solid); mp: 143 °C determined by DSC; $^1$H NMR (CDCl$_3$): $\delta$ 0.91 (t, $J = 7.0$ Hz, 3H), 1.32–1.37 (m, 4H), 1.43–1.50 (m, 2H), 1.76–1.83 (m, 2H), 3.42 (s, 3H), 3.98 (t, $J = 6.6$ Hz, 2H), 6.87–6.89 (m, 2H), 7.45–7.48 (m, 2H), 7.50–7.56 (m, 4H); $^{13}$C NMR (CDCl$_3$): $\delta$ 14.2, 20.1, 22.7, 25.8, 29.3, 31.7, 68.3, 75.3 (t, $J = 4.3$ Hz), 87.7, 92.5, 101.6 (t, $J = 4.3$ Hz), 102.4 (t, $J = 18.1$ Hz), 114.8, 121.0, 125.2, 129.3 (tt, $J = 17.6, 2.9$ Hz), 131.6, 132.0, 133.3, 139.5–142.0 (m, C$_{Ar}$-F, 1C), 145.7–148.3 (m, C$_{Ar}$-F, 1C), 159.7, 166.6 (one sp-carbon was overlapped with another alkyne carbon); $^{19}$F NMR (CDCl$_3$): $\delta$ –137.90 (dd, $J = 23.1, 9.8$ Hz, 2F), –154.22 (dd, $J = 23.1, 9.8$ Hz, 2F); IR (KBr): $\nu$ 2962, 2860, 2367, 2208, 1795, 1597, 1495, 1405 cm$^{-1}$; HRMS (FAB$^+$) $m/z$ [M]$^+$ calcd for C$_{30}$H$_{23}$OF$_3$: 508.1662; found: 508.1655.

2.4. Preparation of CF$_3$-Substituted Bistolane 1aC

In a 50 mL two-necked round-bottomed flask was placed 4-[2-(4-methoxyphenyl)ethyn-1-yl]phenylacetylene (3a, 0.46 g, 2.0 mmol) in THF (30 mL). The solution was cooled to 0 °C and then n-BuLi (1.6 mol L$^{-1}$ hexane solution, 1.3 mL, 2.0 mmol) was added. The mixture was stirred at 0 °C for 0.5 h. Then, perfluorotoluene (C, 0.42 mL, 3.0 mmol) was added dropwise at 0 °C and the resultant solution was continuously stirred at room temperature overnight. The reaction mixture was poured into saturated aqueous NH$_4$Cl solution (40 mL). The crude product was extracted using AcOEt (40 mL, three times) and washed with brine (40 mL, once). The organic layer was dried over anhydrous Na$_2$SO$_4$, filtered, and concentrated using a rotary evaporator. The resultant crude product was purified by silica-gel column chromatography using hexane/AcOEt (20:1, v/v) as an eluent, yielding coupling product 1aC (0.32 g, 0.72 mmol) in 36% yield as a white solid. This product was recrystallized by slow evaporation from CH$_2$Cl$_2$/MeOH (1:1, v/v) to obtain 1aC in crystalline form.

2.4.1. 1-Trifluoromethyl-2,3,5,6-tetrafluoro-4-[2-[4-(4-methoxyphenyl)ethyn-1-yl]phenyl]ethyn-1-yl]benzene (1aC)

Yield: 36% (white solid); mp: 168 °C determined by DSC; $^1$H NMR (CDCl$_3$): $\delta$ 3.84 (s, 3H), 6.88–6.91 (m, 2H), 7.47–7.50 (m, 6H); $^{13}$C NMR (CDCl$_3$): $\delta$ 55.5, 74.9 (t, $J = 3.7$ Hz), 87.7, 92.8, 104.7, 108.8–109.3 (m), 109.7–109.8 (m, 1C), 114.3, 114.9, 120.7 (q, $J = 273.6$ Hz), 120.3, 125.9, 131.7, 132.2, 133.4, 142.9–145.7 (m), 145.8–148.3 (m), 160.2; $^{19}$F NMR (CDCl$_3$): $\delta$ –56.67 (t, $J = 23.2$ Hz, 3F), –135.09 (dd, $J = 18.1, 18.1, 5.7$ Hz, 2F), –141.27 (m, 2F); IR (KBr): $\nu$ 2360, 2210, 1660, 1643, 1595, 1493, 1404, 1345 cm$^{-1}$; HRMS (FAB$^+$) $m/z$ [M]$^+$ calcd for C$_{24}$H$_{17}$OF$_7$: 448.0698; found: 448.0699.

2.4.2. 1-Trifluoromethyl-2,3,5,6-tetrafluoro-4-[2-[4-(4-hexyloxyphenyl)ethyn-1-yl]phenyl]ethyn-1-yl]benzene (1bC)

Yield: 32% (white solid); mp: 133 °C determined by DSC; $^1$H NMR (CDCl$_3$): $\delta$ 0.91 (t, $J = 7.0$ Hz, 3H), 1.33–1.37 (m, 4H), 1.43–1.50 (m, 2H), 1.76–1.83 (m, 2H), 3.42 (s, 3H), 3.98 (t, $J = 6.6$ Hz, 2H), 6.87–6.89 (m, 2H), 7.45–7.48 (m, 2H), 7.52–7.59 (m, 4H); $^{13}$C NMR (CDCl$_3$): $\delta$ 14.1, 22.7, 25.8, 29.3, 31.7, 68.3, 74.9 (t, $J = 7.2$ Hz), 87.6, 93.0, 104.7 (t, $J = 4.2$ Hz), 108.9–109.2 (m, 1C), 114.6, 114.8, 120.7 (q, $J = 273.6$ Hz), 120.2, 125.9, 131.6, 132.2, 133.3, 143.0–145.6 (m, 1C), 145.8–148.3 (m, 1C), 159.8 (one sp-carbon was overlapped with another alkyne carbon); $^{19}$F NMR (CDCl$_3$): $\delta$ –56.77 (t, $J = 23.2$ Hz, 3F), –135.13 (dd, $J = 18.2, 18.2, 5.9$ Hz, 2F), –141.39 (m, 2F); IR (KBr): $\nu$ 2957, 2984, 2360, 2211, 1659, 1606, 1594 cm$^{-1}$; HRMS (FAB$^+$) $m/z$ [M]$^+$ calcd for C$_{29}$H$_{21}$OF$_7$: 518.1481; found: 518.1484.

2.5. Preparation of CN-Substituted Bistolane 1aD

In a 50 mL two-necked round-bottomed flask was placed 4-[2-(4-methoxyphenyl)ethyn-1-yl]phenylacetylene (3a, 0.47 g, 2.0 mmol) in THF (30 mL). The solution was cooled to 0 °C and then n-BuLi (1.6 mol L$^{-1}$ hexane solution, 1.3 mL, 2.0 mmol) was added. The mixture was stirred at 0 °C for 0.5 h. Then, perfluorobenzonitrile (D, 0.30 mL,
2.4 mmol) was added dropwise at 0 °C, and the resultant solution was continuously stirred at room temperature overnight. The reaction mixture was poured into saturated aqueous NH₄Cl solution (40 mL). The crude product was extracted using AcOEt (40 mL, three times) and washed with brine (40 mL, once). The organic layer was dried over anhydrous Na₂SO₄, filtered, and concentrated using a rotary evaporator. The resultant crude product was purified by silica-gel column chromatography using hexane/AcOEt (20:1, v/v) as an eluent, yielding coupling product 1aD (0.26 g, 0.64 mmol) in 32% yield as a yellow solid. This product was recrystallized by slow evaporation from CH₂Cl₂/MeOH (1:1, v/v) to obtain 1aD in crystalline form.

2.5.1. 1-Cyano-2,3,5,6-tetrafluoro-4-[2-[4-[2-(4-methoxyphenyl)ethyn-1-yl]phenyl]ethyn-1-yl]benzene (1aD)

Yield: 32% (yellow solid); mp: 195 °C determined by DSC; ¹H NMR (CDCl₃): δ 3.84 (s, 3H), 6.87–6.92 (m, 2H), 7.47–7.50 (m, 6H), 7.52–7.61 (m, 4H); ¹³C NMR (CDCl₃): δ 55.5, 75.2, 87.7, 93.1, 93.3, 107.1 (dt, J = 84.9, 4.9 Hz), 110.5 (d, J = 2.3 Hz), 111.2 (t, J = 2.8 Hz), 114.3, 114.9, 120.0, 126.3, 131.7, 132.3, 133.4, 145.3–146.0 (m), 147.8–148.7 (m), 160.2; ¹⁹F NMR (CDCl₃): δ = –133.19 (ddd, J = 18.0, 18.0, 6.7 Hz, 2F), –133.90 (ddd, J = 18.0, 17.9, 6.1 Hz, 2F); IR (KBr): ν 2351, 2207, 1643, 1595, 1487, 1478 cm⁻¹; HRMS (FAB⁺) m/z [M⁺] calcd for C₂₄H₁₁NOF₄: 405.0777; found: 405.0780.

2.5.2. 1-Cyano-2,3,5,6-tetrafluoro-4-[2-[4-[2-(4-hexyloxyphenyl)ethyn-1-yl]phenyl]ethyn-1-yl]benzene (1bD)

Yield: 17% (yellow solid); mp: 154 °C determined by DSC; ¹H NMR (CDCl₃): δ 0.91 (t, J = 6.9 Hz, 3H), 1.32–1.37 (m, 4H), 1.44–1.48 (m, 2H), 1.79 (quin, J = 7.0 Hz, 2H), 3.98 (t, J = 6.6 Hz, 2H), 6.87–6.90 (m, 2H), 7.45–7.48 (m, 2H), 7.52–7.59 (m, 4H); ¹³C NMR (CDCl₃): δ 14.2, 22.7, 25.8, 29.3, 31.7, 68.3, 75.5, 87.9, 93.3 (m), 93.3, 106.6–108.3 (m), 110.0, 111.3 (m), 114.6, 114.8, 119.9, 126.3, 131.7, 132.3, 133.4, 143.5–143.8 (m), 144.9–145.1 (m), 159.91; ¹⁹F NMR (CDCl₃): δ = –133.24 (ddd, J = 19.5, 17.8, 6.3 Hz, 2F), –133.94 (ddd, J = 19.5, 17.8, 6.3 Hz, 2F); IR (KBr): ν 2944, 2872, 2210, 1645, 1606, 1596, 1520, 1474, 1416 cm⁻¹; HRMS (FAB⁺) m/z [M⁺] calcd for C₂₉H₂₁NOF₄: 475.1559; found: 475.1549.

2.6. X-ray Crystallography

Single crystals of 1aB and 1aC were obtained by a double purification protocol using column chromatography, followed by recrystallization (CH₂Cl₂/MeOH = 1:1, v/v or CH₂Cl₂/hexane = 1:1, v/v). The obtained single crystals were mounted on a glass fiber. X-ray diffraction data were recorded using a Rigaku XtaLab mini diffractometer (Rigaku Corporation, Tokyo, Japan) or a Rigaku Saturn 724 diffractometer (Rigaku Corporation, Tokyo, Japan) equipped with a VariMax Mo optic system using MoKα (λ = 0.71075 Å). The reflection data were integrated, scaled, and averaged using CrysAlisPro (ver. 1.171.38.46, Rigaku Oxford Diffraction, Rigaku Corporation, Tokyo, Japan, 2015). Empirical absorption corrections were applied using the SCALE3 ABSPACK scaling algorithm (CrysAlisPro, Rigaku Corporation, Tokyo, Japan). The structures were solved by a direct method (SHELXT-2014/S) and refined using a full-matrix least-squares method on F² for all reflections (SHELXL-2014/7) [24]. The crystallographic data were deposited into the Cambridge Crystallographic Data Centre (CCDC) database (CCDC 1908239 for 1aB and 1908240 for 1aC). These data can be obtained free of charge from the CCDC via www.ccdc.cam.ac.uk/data_request/cif.

2.7. Computation

All density functional theory (DFT) computations were performed using the Gaussian 09 (Revision C.01) package [25]. Geometry optimizations were executed using the CAM-B3LYP hybrid functional [26,27] and the 6-31+G(d) basis set with an implicit solvation model, namely, the conductor-like polarizable continuum model (CPCM) [28–30], for CH₂Cl₂. The vertical excitation energies and dipole moments of the optimized structures were calculated using the time-dependent
(TD)-DFT method at the same level as in the theory. To obtain the optimized structures in the excited ($S_1$) state, geometry optimizations were carried out using the TD-DFT method.

2.8. Phase Transition Behavior

The phase transition behavior of the fluorinated bistolanes was observed by polarizing optical microscopy (POM) using an Olympus BX53 microscope (Olympus Corporation, Tokyo, Japan) equipped with a cooling and heating stage (Linkam Scientific Instruments, 10002L, Surrey, UK). The thermodynamic behavior was determined using differential scanning calorimeter (DSC, Shimadzu, DSC-60 Plus, Kyoto, Japan) at heating and cooling rates of 5.0 °C min$^{-1}$ under a N$_2$ atmosphere.

2.9. Photophysical Behavior

Ultraviolet-visible (UV-vis) absorption spectra were recorded using a JASCO V-500 (JASCO Corporation, Tokyo, Japan) absorption spectrometer. Steady-state PL spectra were obtained using a JASCO FP-8500 (JASCO Corporation, Tokyo, Japan) or a Hitachi F-7000 fluorometer (Hitachi Ltd., Tokyo, Japan). The absolute PL quantum yields were recorded using a calibrated integrating sphere with a Hamamatsu Photonics C11347-01 Absolute PL Quantum Yield Measurement System (Hamamatsu Photonics KK, Hamamatsu, Japan).

3. Results and Discussion

3.1. Molecular Design

Using DFT and TD-DFT calculations in the Gaussian 09 suite of programs, we examined the effect of the terminal substituent in each designed molecule on the electron-density distribution over the entire molecule. The geometries of 1aB–D in the ground ($S_0$) and excited ($S_1$) states were optimized at the CAM-B3LYP/6-31+G(d) level of theory with the implicit solvation CPCPM model for CH$_2$Cl$_2$. Figure 2 shows the structures of 1aB–D and their calculated electrostatic potential maps, dipole moments along the long molecular axis (μ||), isodensity surfaces, and orbital energies for the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO).

![Figure 2](image_url)

Figure 2. Chemical structures of 1aB–D and their electrostatic potential maps, dipole moments (μ||) along the long molecular axis, isodensity surfaces with the color-bar (isovalue = 0.0004), and orbital energies for the HOMO and LUMO in the $S_1$ state.

The electrostatic potential maps of 1aB–D reveal that the electron density at the fluorinated aromatic ring decreases in the following order: 1aB > 1aC > 1aD. The dipole moment (μ||) along the...
long molecular axis can be used to assess the electron-density distribution over the entire molecule quantitatively. AcO-substituted 1aB had the smallest $\mu_{||}$ value (3.48 D), and this value increased when the AcO group was replaced by a CF$_3$ or CN group ($\mu_{||}$ = 9.63 D for 1aB and 12.2 D for 1aC). These results clearly confirm that the incorporation of AcO, CF$_3$, and CN groups into the fluorinated bistolane scaffold can be used to systematically change the electron-density distribution over the entire molecule. The electron-density distribution also affected the molecular orbitals, with the orbital lobes in the HOMO largely localized on the relatively electron-rich aromatic rings, whereas the LUMO lobes were relatively localized on the electron-deficient aromatic rings. The orbital separation between the HOMO and LUMO in the S$_1$ state may contribute to unique PL behavior through intramolecular charge transfer [31]. The energy differences ($\Delta E_{H,L}$) between HOMO and LUMO levels were 5.08 eV for 1aB, 4.89 eV for 1aC, and 4.67 eV for 1aD. Thus, 1aD is expected to show a PL band in the long-wavelength region, with the PL wavelength shifting towards shorter wavelengths as the transition energy increases, i.e., 1aB $>$ 1aC. Theoretical calculations for 1bB–D, bearing long alkoxy substituents, also provided similar $\mu_{||}$ and HOMO/LUMO distributions (see Supplementary Materials), assuming that the alkoxy-substituent does not affect in the electron density distribution. Consequently, systematic control of the electron-density distribution induced by modulating the electronic character of the terminal substituent should realize luminescence tuning for the fluorinated luminophores. According to our previous study, modulation of terminal substituents also influences the LC behavior [21,23], not only through changes in the electron-density distribution but also in the steric bulkiness of the substituent. Thus, systematic modulation of the terminal substituent in 1aB–D may also allow control of the LC behavior.

3.2. Synthesis and Crystal Structure

Based on the aforementioned molecular design, fluorinated bistolanes 1aB–C with a CH$_3$O group and 1bB–D with a hexyloxy (C$_6$H$_{13}$O) moiety were synthesized according to the procedure shown in Scheme 1. AcO-substituted 1aB was obtained in 48% yield by the Pd(0)-catalyzed Sonogashira cross-coupling reaction of 3a, readily accessible from 2a [21,23], with B. In contrast, CF$_3$-substituted 1aC and CN-substituted 1aD were obtained in 36% or 32% yield, respectively, by nucleophilic aromatic substitution of the in situ generated lithium acetylide derived from 3a with perfluorotoluene and perfluorobenzonitrile, respectively [32]. In a similar manner, C$_6$H$_{13}$O-substituted analogue 1bB was prepared in 54% yield through a Sonogashira cross-coupling reaction and 1bC and 1bD were prepared in 32% and 17% yields, respectively, through nucleophilic substitution reactions. Each fluorinated bistolane was purified by column chromatography and recrystallization from a 1:1 solution of CH$_2$Cl$_2$ (good solvent) and MeOH or hexane (poor solvent). The structure of each compound was confirmed by $^1$H, $^{13}$C, and $^{19}$F NMR, IR, and HRMS and the spectra indicated that the purity was sufficient to evaluate the LC and PL behaviors.

Among the fluorinated bistolanes, AcO-substituted 1aB and CF$_3$-substituted 1aC furnished single crystals appropriate for X-ray crystallographic analysis. 1aB crystalized with a monoclinic crystal structure in the P 2$_1$ space group with two molecular units in the unit cell. As shown in Figure 3a, the three aromatic rings were found to be almost coplanar, although there may be a rapid equilibrium between the coplanar and twisted conformations through free rotation of the C$_{sp^2}$–C$_{sp^2}$ single bonds [33]. The packing structure (Figure 3b) revealed several close contacts that were shorter than the sums of the van der Waals radii (carbon: 170 pm, hydrogen: 120 pm, and fluorine: 148 pm) [34]: $H_{skyl}$···$C_{sp^2}$ (288 pm), $H_{aryl}$···$C_{sp^2}$ (284 pm), $H_{aryl}$···$F$ (253 pm), $H_{skyl}$···$F$ (246 pm), $H_{aryl}$···$O$ (257 pm). Thus, the crystal lattice of 1aB contains two CH–π interactions and three hydrogen bonding interactions. In contrast, 1aC crystalized with a monoclinic crystal structure in the P 2$_1$/c space group with four molecular units in the unit cell. As shown in Figure 3c, each 1aC molecule was also found to be coplanar with several short contacts (Figure 3d), such as $C_{sp^2}$···$C_{sp^2}$ (332 pm), $H_{aryl}$···$F$ (251 pm), $C_{sp^2}$···$F$ (314 pm), and $F$···$F$ (265 pm). Thus, 1aC also formed a tight packing structure with one π–π interaction, one hydrogen bonding interaction, and one lone-pair–π interaction [35,36].
Figure 3. Molecular and crystal structures of (a) 1aB and (c) 1aC. Packing structures of (b) 1aB and (d) 1aC. The light-blue lines in the packing structures indicate close contacts shorter than the sum of van der Waals radii (C: 170 pm, H: 120 pm, F: 148 pm).

3.3. Phase Transition Behavior

The phase transition behavior of fluorinated bistolanes 1aB–D and 1bB–D was investigated using POM and DSC. Figure 4 shows the phase transition behavior observed during the 2nd heating cooling processes and the microphotographic images of the LC phases.

Figure 4. Phase transition behavior of (a) 1aB–D and (b) 1bB–D during the 2nd heating [H] and cooling [C] processes. Abbreviations: Cr, crystal; SmA, smectic A; N, nematic; and Iso, isotropic phases. (c) Microphotographic images recorded by POM during the cooling process.
Among 1aB–D with a methoxy group at one molecular terminal and an electron-withdrawing substituent at the other, AcO-substituted 1aB exhibited no LC behavior, as evidenced by the dramatic change in the POM images from nonfluidic bright-viewing to fluidic dark-viewing fields during both the heating and cooling processes. The steric hindrance of the AcO group and the dipole moment (μ⊥) along the short molecular axis induced by the carbonyl moiety in the AcO group likely make it difficult to order structures in the condensed phase, resulting in a direct phase transition to the isotropic (Iso) phase. In contrast, the POM images of 1aC with a CF3 group and 1aD with a CN group displayed fluidic bright-viewing fields between the crystal (Cr) and Iso phases; POM and DSC measurements only revealed a single LC phase for 1aC and 1aD. As the microphotographic images of 1aC and 1aD display a four-brushed Schlieren texture, which is typical of nematic (N) LC phases, the single LC phases of 1aC and 1aD are assigned as N phases. The melting temperature (Tm, defined as the phase transition temperature between the Cr and LC phases) and the clearing temperature (Tc, defined as the phase transition temperature between the LC and Iso phases) are much higher for 1aD than for 1aC (Figure 3a). This behavior is consistent with the linear and rigid structure of the CN group, which induces tight packing in an antiparallel direction owing to the large μ⊥ value [37–39].

Changing the length of the flexible chain at the molecular terminal by replacing CH2 with C6H13O affected the LC behavior. 1bB–D all showed LC phases, and the Tm and Tc values for 1bB–D were dramatically decreased compared with the corresponding values for 1aB–D. Typically, such stabilization of LC phases can be attributed to the introduction of a longer flexible unit [23]. The POM and DSC measurements revealed that AcO-substituted 1bB and CN-substituted 1bD displayed only a N LC phase owing to the steric bulkiness of AcO and the rigid linear rigid structure of CN with a large μ⊥ value. However, CF3-substituted 1bC exhibited both SmA and N LC phases, analogous to those previously reported for 1bA. Thus, a suitable choice of substituent at the molecular terminal is important for achieving precise control of the LC behavior, including the LC phase, phase transition temperature, and LC temperature range.

### 3.4. PL Behavior in the Solution Phase

Next, we examined the photophysical behavior of the fluorinated bistolanes. First, we investigated the absorption and PL behavior in CH2Cl2 solution (1.0 × 10−5 mol L−1 for absorption and 1.0 × 10−6 mol L−1 for PL). Figure 5a,b shows the absorption and PL spectra, respectively, of 1aB–D (solid lines) and 1bB–D (dashed lines). Figure 5c shows photographs of the PL colors and the corresponding Commission Internationale d’Eclairage (CIE) chromaticity diagram. The photophysical data are listed in Table 1.

![Table 1. Photophysical data for 1aB–D and 1bB–D in CH2Cl2 solution.](image-url)

|        | λabs (nm) | 1 (ε) | Ground (S0) State | λPL (nm) | ΦPL | Excited (S1) State |
|--------|-----------|-------|-------------------|----------|-----|-------------------|
|        | μ∥ (D)    | HOMO/LUMO (eV) | μ∥ (D) | HOMO/LUMO (eV) |        |       |
| 1aB    | 333 (55.2) | 1.91 | -7.09/-1.30      | 410 (0.85) | 3.48 | 6.74/-1.66        |
| 1aC    | 341 (48.1) | 7.22 | -7.13/-1.58      | 444 (0.99) | 9.63 | 6.83/-1.94        |
| 1aD    | 356 (46.7) | 9.67 | -7.15/-1.87      | 488 (0.91) | 12.2 | 6.86/-2.19        |
| 1bB    | 335 (54.5) | 2.27 | -7.07/-1.29      | 414 (0.86) | 3.93 | 6.72/-1.65        |
| 1bC    | 343 (52.5) | 7.51 | -7.12/-1.58      | 451 (0.95) | 10.0 | 6.81/-1.94        |
| 1bD    | 360 (44.8) | 9.95 | -7.13/-1.86      | 493 (0.87) | 12.6 | 6.84/-2.19        |

1 Concentration of the solution: 1.0 × 10−5 mol L−1. 2 Molar extinction coefficient (10−3 L mol−1 cm−1). 3 Obtained by DFT calculations at the CAM-B3LYP/6-31+G(d) level of theory. 4 Concentration of the solution: 1.0 × 10−6 mol L−1. 5 Measured using a calibrated integrating sphere system. 6 Obtained by TD-DFT calculations at the CAM-B3LYP/6-31+G(d) level of theory.

In CH2Cl2 solution, CH3O-Substituted 1aB–D each exhibited a single absorption band and the maximum absorption wavelength (λabs) varied significantly depending on the electron-withdrawing substituent: 333 nm (AcO-substituted 1aB), 341 nm (CF3-substituted 1aC), and 356 nm (CN-substituted 1aD) (Figure 5a), in which the λmax in the calculated UV-Vis spectra obtained from TD-DFT calculation
observed in similar tendency: 338 nm (AcO-substituted 1aB), 347 nm (CF3-substituted 1aC), and 360 nm (CN-substituted 1aD) (Figure S21). As listed in Table 1, the values in the S0 state were calculated to be 1aB (1.91 D) < 1aC (7.22 D) < 1aD (9.67 D), indicating a strong correlation between the \( \lambda_{abs} \) and \( \mu_{||} \) values. Upon excitation at \( \lambda_{abs} \), 1aB–D in CH2Cl2 each displayed a single PL band at a maximum PL wavelength (\( \lambda_{PL} \)) of 410–488 nm with a high PL efficiency (\( \Phi_{PL} = 0.85–0.91 \)) (Figure 5b). As shown in Figure 5c, 1aB–D emitted deep-blue, blue PL, light-green PL, respectively, and the PL colors can also be quantitatively evaluated using the CIE chromaticity diagram. Similar to \( \lambda_{abs} \), \( \lambda_{PL} \) also exhibited a shift towards longer wavelengths depending on the \( \mu_{||} \) value. Replacing the CH3O substituent with the longer C6H12O unit did not change the absorption or PL spectra significantly owing to the similar \( \mu_{||} \) values. These results clearly demonstrate that the electron-density distribution induced by the electron-withdrawing substituent is an important parameter for controlling the photophysical properties in the solution phase.

![Figure 5](image_url)

**Figure 5.** (a) Absorption spectra of 1aB–D (solid lines) and 1bB–D (dashed lines) in CH2Cl2 (1.0 \( \times \) 10^{-5} mol L^{-1}); (b) photoluminescence (PL) spectra of 1aB–D (solid lines) and 1bB–D (dashed lines) in CH2Cl2 (1.0 \( \times \) 10^{-6} mol L^{-1}); (c) photographs under ultraviolet (UV) irradiation (\( \lambda_{ex} = 365 \) nm) and CIE chromaticity diagram for the PL in CH2Cl2 solution.

TD-DFT calculations revealed that the HOMO→LUMO transition corresponded with the lowest-energy excitation with a large oscillator strength. Considering that the energy gap between the HOMO and LUMO (\( \Delta E_{H-L} \)) in the S0 state decreases in the order 1aB (5.69 eV) > 1aC (5.55 eV) > 1aD (5.28 eV), increasing the \( \mu_{||} \) value results in a narrower \( \Delta E_{H-L} \) and a shift of \( \lambda_{abs} \) towards longer wavelengths: 1aB (338 nm) < 1aC (347 nm) < 1aD (360 nm). \( \Delta E_{H-L} \) in the S1 state exhibited a similar tendency (1aB > 1aC > 1aD), corresponding to the order of the \( \mu_{||} \) values. Thus, these results clearly indicate that the electronic transitions for absorption and PL can be controlled by tuning \( \mu_{||} \).
The PL colors of luminophores with large $\mu_{\parallel}$ values have often been reported to show obvious variations depending on the solvent polarity [40]. Therefore, we tested the PL sensing properties of CF$_3$-substituted 1bC as a representative fluorinated bistolane in various solvents. Figure 6 shows the observed PL behavior, and the photophysical data are summarized in Table 2.

**Figure 6.** (a) PL spectra of 1bC in various solvents (concentration: $1.0 \times 10^{-6}$ mol L$^{-1}$); (b) photographs under UV irradiation ($\lambda_{ex}$ = 365 nm) and CIE chromaticity diagram for the PL in various solvents.

**Table 2.** Photophysical properties of 1bC in various solvents.

| Solvent (c) | Toluene (2.38) | THF (7.58) | CH$_2$Cl$_2$ (8.93) | MeCN (35.9) | DMF (36.7) |
|-------------|----------------|------------|---------------------|--------------|------------|
| $\lambda_{PL}$ (nm)$^2$ | 409 | 482 | 451 | 518 | 523 |
| $\Phi_{PL}$ $^3$ | 0.79 | 0.55 | 0.95 | 0.58 | 0.40 |

$^1$ $\epsilon$: Dielectric constant at 25 °C. $^2$ Concentration of the solution: $1.0 \times 10^{-6}$ mol L$^{-1}$. $^3$ Measured using a calibrated integrating sphere system.

Irradiating 1bC at 343 nm in less polar toluene ($\epsilon = 2.38$, where $\epsilon$ is a dielectric constant at 25 °C and an indicator of solvent polarity [40]) produced deep-blue PL with a single PL band at $\lambda_{PL} = 409$ nm, which was shifted towards shorter wavelengths by 0.282 eV compared with the $\lambda_{PL}$ in CH$_2$Cl$_2$ ($\epsilon = 8.93$). In contrast, dramatic shifts towards longer wavelengths were observed when more polar solvents, e.g., MeCN ($\epsilon = 35.9$) and DMF ($\epsilon = 36.7$), were employed; based on the CIE diagram, greenish-yellow PL was observed in both cases. Although THF ($\epsilon = 7.58$) is less polar than CH$_2$Cl$_2$, the PL band ($\lambda_{PL} = 482$ nm) was shifted towards longer wavelengths by 0.177 eV. The unique PL behavior in THF is likely due to the oxygen atom, which allows THF to easily solvate 1bC with its large $\mu_{\parallel}$ value through intermolecular interactions, such as hydrogen bonding, and may promote the nonradiative deactivation from the S$_1$ to the S$_0$ state [41–43]. Consequently, it was successfully demonstrated that the fluorinated bistolanes can be applied as PL sensing materials for solvent polarity.

### 3.5. PL Behavior in the Crystal Phase

To our delight, we found that 1aB–D and 1bB–D crystals prepared by recrystallization via a slow evaporation technique exhibited PL. Figure 7 shows the obtained PL spectra acquired, and the photophysical data are listed in Table 3.
behavior in THF is likely due to the oxygen atom, which allows THF to easily solvate with its large $\mu_{||}$ value through intermolecular interactions, such as hydrogen bonding, and may promote the nonradiative deactivation from the S$_1$ to the S$_0$ state [41–43]. Consequently, it was successfully demonstrated that the fluorinated bistolanes can be applied as PL sensing materials for solvent polarity.

3.5. PL Behavior in the Crystal Phase

To our delight, we found that 1aB–D and 1bB–D crystals prepared by recrystallization via a slow evaporation technique exhibited PL. Figure 7 shows the obtained PL spectra acquired, and the photophysical data are listed in Table 3.

![Figure 7. PL spectra of (a) 1aB–D and (b) 1bB–D crystals; (c) CIE chromaticity diagram calculated from the corresponding PL spectra.](image)

Table 3. Photophysical properties of 1aB–D and 1bB–D crystals.

| Compound | $\lambda_{PL}$ (nm) $^1$ | $\Phi_{PL}$ $^2$ |
|----------|------------------------|-----------------|
| 1aB      | 420                    | 0.48            |
| 1aC      | 459                    | 0.80            |
| 1aD      | 500                    | 0.36            |
| 1bB      | 414 (418)              | 0.17            |
| 1bC      | 479 (443)              | 0.73            |
| 1bD      | 526 (527)              | 0.31            |

1 Upon excitation at 358 nm for 1aB, 361 nm for 1aC, 400 nm for 1aD, and 320 nm for 1bB–D. The values in parentheses are the PL maximum wavelengths in the LC state upon excitation at 320 nm. 2 Measured using a calibrated integrating sphere system.

As shown in Figure 7a,b, $\lambda_{PL}$ changed dramatically depending on the electron-withdrawing substituent, even in crystals: 414–420 nm (deep-blue PL) for AcO-substituted 1aB and 1bB, 459–479 nm (blue to light-blue PL) for CF$_3$-substituted 1aC and 1bC, and 500–526 nm (yellowish-green PL) for CN-substituted 1aD and 1bD. Compared with the $\lambda_{PL}$ values in CH$_2$Cl$_2$ solution, those observed for the crystals were slightly shifted towards longer wavelengths, which may be due to quick deactivation via nonradiative processes owing to intermolecular interactions in the condensed phases. As discussed above, exchanging the CH$_3$O group for a longer C$_6$H$_{13}$O moiety did not have a significant effect on the electronic structure; however, an intriguing band shift of 0.113 eV for 1aC vs. 1bC and 0.123 eV for 1aD vs. 1bD was observed. According to our previous reports [21,23], this change in the PL behavior likely originates from a dramatic change in the aggregated structure induced by the difference in alkoxy chain length. Similar to the trend observed for the PL in CH$_2$Cl$_2$ solution, a relatively strong correlation was observed between $\mu_{||}$ and $\lambda_{PL}$ in the crystals. These results demonstrate that the PL behavior of the
fluorinated bistolanes can also be modulated in crystal by controlling $\mu_\parallel$ via the electron-withdrawing character of the terminal substituent.

3.6. PL Behavior in the LC Phase

Finally, owing to the fluorinated bistolanes possessing LC phases and unique PL characteristics, we examined the PL behavior in the LC phase under thermal conditions. As representative examples, frozen LC samples of 1bB, 1bC, and 1bD were obtained by rapid cooling in a dry-ice bath after a suitable LC phase was observed. The obtained PL spectra are shown in Figure 8.

![Figure 8. PL spectra of the Cr and LC phases of (a) 1bB, (b) 1bC, and (c) 1bD upon excitation at 320 nm.](image)

When the N LC samples of AcO-substituted 1bB and CN-substituted 1bD were irradiated at 320 nm light, single PL bands were observed at $\lambda_{PL} = 418$ and 527 nm, respectively, which were almost identical to those in the Cr phase (Figure 8a,c). Interestingly, as shown in Figure 8b, the initial Cr sample of CF$_3$-substituted 1bC showed light-blue PL ($\lambda_{PL} = 479$ nm), but a significant shift to shorter wavelengths was observed after the phase transition to the SmA phase, leading to deep-blue PL ($\lambda_{PL} = 443$ nm), which is almost identical to that in CH$_2$Cl$_2$ solution. It is anticipated that rearrangement of the molecular aggregates during the Cr $\rightarrow$ SmA phase transition results in loose molecular packing owing to thermally activated molecular motion and monomeric emission characteristics. Accordingly, it can be concluded that the PL of fluorinated bistolanes can be controlled not only by tuning the electron-density distribution but also by modulating molecular aggregation through thermal phase transitions, which is a promising approach for realizing functional PL sensing materials, such as PL thermometers.

4. Conclusions

In conclusion, we designed and synthesized fluorinated bistolanes with an alkoxy substituent at one molecular terminal and various electron-withdrawing substituent at the other terminal via a four-step protocol. The phase transition temperatures of the fluorinated bistolanes were found to depend significantly on the terminal substituent, with the LC behavior affected by both the steric and
electronic nature of the substituent. The PL behavior in solution clearly revealed that both $\lambda_{PL}$ and the PL color changed dramatically depending on the $\mu_{||}$ value. The present fluorinated bistolanes also showed relatively strong PL in the crystal phase, with the PL behavior modulated not only by $\mu_{||}$ but also by molecular aggregation. Notably, the PL emission of a CF$_3$-substituted fluorinated bistolane was significantly shifted towards shorter wavelengths by 0.210 eV via a phase transition from the Cr to the SmA LC phase. As the PL behavior of the fluorinated bistolanes can be tuned by the electron-density distribution as well as the aggregated structure in condensed phases, this molecular design approach offers a useful method for fabricating PL materials that can be easily tuned by using molecular properties or external stimuli.

Supplementary Materials: The following are available online at http://www.mdpi.com/2076-3417/9/9/1905/s1, Figures S1–S18: NMR spectra, Table S1: crystallographic data, Figure S19: DSC thermograms, Tables S2–S6: phase transition behavior and thermodynamic parameter, Figure S20: photophysical spectra in solution, Figure S21: comparison between calculated and experimental UV-vis spectra, Figure S22: photophysical spectra in crystal, Figure S23: PL spectra in LC phases, Figures S24–S34: HOMO and LUMO diagram, Tables S7–S18: cartesian coordinate for calculated molecules.

Author Contributions: Conceptualization, S.Y.; methodology, S.Y. and M.M.; investigation, S.Y., M.M., T.A., T.K. (Tosio Kubota) and T.K. (Tsutomu Konno); writing—original draft preparation, S.Y., M.M. and T.K. (Tsutomu Konno); writing—review and editing, S.Y., M.M., T.A., T.K. (Tosio Kubota) and T.K. (Tsutomu Konno); visualization, S.Y. and M.M.; supervision, S.Y.; project administration, S.Y.; funding acquisition, S.Y.

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