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Dark conglomerate phases of azobenzene derived bent-core mesogens – relationships between the molecular structure and mirror symmetry breaking in soft matter†

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New 4-bromoresorcinol based bent-core molecules with peripheral fluoro substituted azobenzene wings have been synthesized and the liquid crystalline self-assembly was investigated by differential scanning calorimetry (DSC), optical polarizing microscopy (POM), electro-optic studies and X-ray diffraction (XRD). A new type of optically isotropic mesophase composed of chiral domains with opposite handedness (dark conglomerate phases, DC phases) is observed, which for some homologues with medium alkyl chain length is stable down to ambient temperature. It is proposed that these DC phases are formed by helical twisted nano-domains of limited size and composed of the crystallized aromatic cores which are separated by the disordered alkyl chains. This structure is distinct from the previously known soft helical nano-filament phases (HNF phases, B₄ phases) formed by extended crystalline nano-filaments and also distinct from the fluid sponge phases composed of deformed fluid layers. Comparison with related bent-core molecules having H, F, Cl, I, CH₃ and CN groups in the 4-position at the resorcinol core, either with or without additional peripheral fluorines, provided information about the effects of these substituents on the tendency to form DC phases. Based on these relationships and by comparison with the minimum energy conformations obtained by DFT calculations a hypothesis is provided for the formation of DC phases depending on the molecular structure.

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1. Introduction

The observation of spontaneous development of macroscopic chirality in soft matter systems of achiral molecules is a contemporary research topic with great importance for the general understanding of spontaneous mirror symmetry breaking, and this might also be useful for numerous practical applications. This phenomenon is in recent years most often found in liquid crystalline phases of some molecules with an extended bent aromatic core (bent-core molecules).^{1,2} Helical superstructures in columns^{3,4} and self-assembly in helical filaments^{5–9} represent other prominent examples of mirror symmetry breaking at the nano-scale.¹⁰ However, in these 1D organizations chiral segregation occurs only locally, along the aggregates, whereas the lateral interaction between them is only weak, so that usually no macroscopic chirality could be achieved in the absence of a chiral dopant.² Only for achiral dibenzo[*g,p*]chrysenes with short peripheral chains macroscopic

spontaneous segregation of chiral conformers was observed in a columnar phase.⁴ In contrast, in polar smectic LC phases of bent-core liquid crystals (BCLCs) the spontaneous formation of conglomerates of macroscopically chiral domains is more frequently observed. Here chirality emerges in layers (smectic phases) which provides a stronger coupling and therefore can more easily transfer chirality over larger distances.¹ In addition, a geometric layer chirality arising from the combination of tilt and polar order in the polar smectic phases of bent-core molecules is assumed to play an important role in the development of macroscopic chirality in these smectic phases,¹¹ as it can couple with the molecular conformational chirality *via* diastereomeric interactions. On the other hand, in all cases of lamellar phases the packing of helical entities, ranging from helical conformers of single molecules *via* bent dimesogens to helical nano-filaments, is in competition with the formation of extended flat layers and thus leads to layer distortion.¹² In many cases this distortion is strong enough to remove the macroscopic anisotropic properties of liquid crystals, giving rise to isotropic mesophases which, due to the absence of birefringence, allow an easy identification of chirality by polarizing microscopy. These soft isotropic mesophases composed of a conglomerate of domains with opposite handedness are designated as dark conglomerate phases (DC phases). Depending on the local

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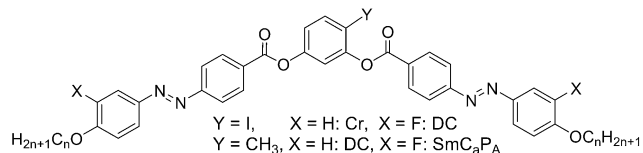
† Electronic supplementary information (ESI) available. See DOI: 10.1039/c4sm01255k



structure of these DC phases they can be classified as liquid crystalline sponge phases,^{13–19} formed by strongly deformed fluid layers or as helical nanofilament phases (HNF phases, also assigned as B₄ phases²⁰) having crystallized aromatic segments organized in helical nano-scale filaments which are separated by the disordered alkyl chain segments.^{21–27} Aside from these two known cases a number of additional DC phases with intermediate or with quite distinct nano-structures can be expected, only a few of them have very recently been observed.^{28–30} Moreover, conglomerates of chiral domains were also found in apolar SmC phases with only local polar domains (SmC_sP_R^[*] phases)^{31,32} and in nematic phases formed by some bent-core mesogens.^{33–35} In these cases of LC phases with reduced order and without long range layer chirality the formation of DC phases is also dependent on surface interactions. However, the relationships between the formation of the distinct types of DC phases and the required molecular structures are still unknown. Therefore, the search for new molecular structures capable of forming DC phases and the investigation of their general structure property relationships is of significant importance. Moreover, DC phases could represent interesting materials for a variety of applications, such as organic semiconductors, thin-film transistors and solar cells,³⁶ as thin-film polarizers,³⁷ as nonlinear optical materials³⁸ as well as for the detection and amplification of chirality,³⁹ and eventually also for separation of enantiomers and for enantioselective synthesis.⁴⁰

DC phases involving photoisomerizable azobenzene units were previously only reported for dimesogens composed of two azobenzene units connected by odd-numbered aliphatic spacers⁴¹ and for W-shaped molecules.⁴² Only recently, we have found that the azobenzene unit is a very useful building block for bent-core mesogens forming new types of HNF-like DC phases.^{43,44} Broad regions of these DC phases have been found for 4-iodoresorcinol based⁴³ as well as 4-methylresorcinol based bent-core mesogens with two azobenzene wings (see Scheme 1).⁴⁴ In contrast, related 4-chloro and 4-fluoro substituted compounds do not form any DC phases.

Fluorine substitution at the periphery of the bent aromatic core has a significant effect on the liquid crystalline behaviour of BCLCs,⁴⁵ for example it can change the switching from antiferroelectric to ferroelectric.^{1,46} In the case of the azobenzene based BCLCs it affects the formation of DC phases. Whereas F-substitution is required for the formation of DC phases of 4-iodoresorcinols with two azobenzene wings,⁴³ it removes the DC phase in the case of the analogous 4-methyl substituted compounds (Scheme 1).⁴⁴



Scheme 1 Effect of fluorine substitution (X = F vs. H) on the formation of DC phases in azobenzene-based BCLCs (abbreviations; DC = dark conglomerate phase, SmCaPA = anticlinic tilted antiferroelectric switching SmC phase, Cr = crystalline solid).^{43,44}

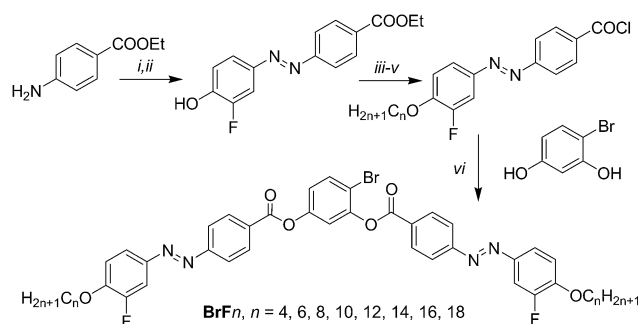
Here we report a new class of DC phase forming BCLCs based on 4-bromoresorcinol and having additional fluorine atoms at the periphery of the attached azobenzene wings, adjacent to the terminal alkyl chains. These compounds are compared with related compounds with F, Cl, I, CH₃ and CN substituents in the 4-position at the resorcinol core, either with or without additional peripheral fluorine substitution.^{31,32,43,44,47,48} The designation of the molecules reported herein follows the general notation **YXn**, where Y indicates the 4-substituent at the resorcinol core, X = F indicates the presence of peripheral fluorines, the absence of X indicates a non-fluorinated compound where X = H and *n* gives the alkyl chain length (see Scheme 1). Besides the series of 4-bromoresorcinol compounds **BrFn** also some additional compounds (**HF12**, **FF12**, **ClF12**, **MF12**, **IF16**) required for proper comparisons were newly synthesized. DFT calculations provide information about the effects of the different substituents Y on the minimum energy conformations, which is related to the experimentally determined tendency to form DC phases of different types. A hypothesis is provided for the prediction of formation of DC phase depending on the molecular structure.

2. Experimental

2.1 Synthesis

The synthesis of the target BCLCs **BrFn** was performed by acylation of 4-bromoresorcinol with two equivalents of the appropriate 4-(4-*n*-alkoxy-3-fluorophenylazo)benzoyl chloride in the presence of triethylamine as base and pyridine as acylation catalyst (Scheme 2).⁴⁹ Details of synthesis of the intermediates and final compounds as well as analytical data are reported in the ESI.[†]

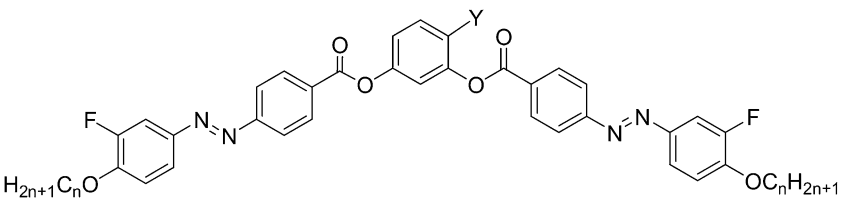
In analogy to the synthesis of compounds **BrFn** also the 4-substituted resorcinol based bent-core mesogens **HF12** (X = F, Y = H), **FF12** (X, Y = F), **ClF12** (X = F, Y = Cl), **MF12** (X = F, Y = CH₃) and **IF16** (X = F, Y = I) have been synthesized. The analytical data and other details of these new compounds (see Table 1) are also reported in the ESI.[†]



Scheme 2 Chemical structures and synthesis of compounds **BrFn**. *Reagents and conditions:* (i) NaNO₂, HCl, H₂O, 0 °C, 30 min; (ii) 1. 2-fluorophenol, NaOH, 0–5 °C, 3 h, 2. NaHCO₃; (iii) C_nH_{2n+1}Br, K₂CO₃, KI, 2-butanone, reflux, 18 h; (iv) 1. KOH, EtOH, reflux, 8 h, (2) H⁺; (v) SOCl₂, DMF, reflux, 1 h; (vi) TEA, pyridine, CH₂Cl₂, reflux, 6 h.



Table 1 Phase transition temperatures ($T/^\circ\text{C}$), mesophase types, and transition enthalpies [$\Delta H/\text{kJ mol}^{-1}$] of the synthesized compounds **BrFn**, **IF16**, **HF12**, **FF12**, **ClF12** and **MF12**^a



Compound	Y	n	1 st heating	1 st cooling
BrF4	Br	4	Cr 157 [57.7] Iso	Iso 117 [50.4] Cr
BrF6	Br	6	Cr 126 [45.9] Iso	Iso 96 [23.8] DC
BrF8	Br	8	Cr 119 [55.0] Iso	Iso 96 [25.9] DC
BrF10	Br	10	Cr 106 [19.0] Iso	Iso 99 [25.6] DC
BrF12	Br	12	Cr 106 [24.6] Iso	Iso 98 [25.7] DC
BrF14	Br	14	Cr 108 [67.2] Iso	Iso 99 [26.1] DC
BrF16	Br	16	Cr 100 [30.2] Iso	Iso 93 [28.5] ^b DC 89 [26.6] Cr
BrF18	Br	18	Cr 102 [30.5] Iso	Iso 89 [39.1] Cr
IF16	I	16	Cr 111 [20.7] Iso	Iso 104 DC 103 [28.5] ^c Cr
HF12	H	12	Cr ₁ 81 [21.2] Cr ₂ 138 [40.5] Iso	Iso 130 [16.5] Cr ₁ 103 [34.2] Cr ₂ 75 [21.6] Cr ₃
FF12	F	12	Cr 120 [60.2] Iso	Iso 116 [18.0] SmC _a P _A ^d 106 [45.0] Cr
ClF12	Cl	12	Cr 114 [51.3] Iso	Iso 95 [11.1] SmC _a P _A ^d 75 [37.1] Cr
MF12	CH ₃	12	Cr 90 [40.4] Iso	Iso 88 [13.9] SmC _a P _A ^d 72 [31.9] Cr

^a Peak temperatures from DSC with rate 10 K min⁻¹; for phase transitions in the second heating scan see Table S1; abbreviations: Cr = crystalline solid; DC = dark conglomerate phase; SmC_aP_A = antclinic tilted antiferroelectric SmC phase; Iso = isotropic liquid. ^b Obtained on cooling with 2 K min⁻¹. ^c Transition enthalpy value could not be determined for the DC–Cr transition due to overlapping. ^d The spontaneous polarization value (P_s) calculated in the SmC_aP_A phase for **FF12** is 460 nC cm⁻²; for **ClF12** is 480 nC cm⁻² and that for **MF12** is 640 nC cm⁻² (representative current response curves are shown in Fig. S11).

2.2 Methods

The mesophase behaviour and transition temperatures were measured using a Mettler FP-82 HT hot stage and a control unit in conjunction with a Nikon Optiphot-2 polarizing microscope. The associated enthalpies were obtained from DSC-thermograms which were recorded on a Perkin-Elmer DSC-7 with heating and cooling rates of 10 K min⁻¹. Electro-optical switching characteristics were examined in 6 μm polyimide coated ITO cells (EHC Japan) using the triangular-wave method. XRD patterns were recorded with a 2D detector (Vantec-500, Bruker). Ni filtered and pin hole collimated CuK α radiation was used. The exposure time was 15 min and the sample to detector distance was 8.95 and 26.7 cm for wide angle and small angle scattering experiments, respectively. Samples were aligned by slow cooling (rate: 1 K min⁻¹ to 0.1 K min⁻¹) of a small droplet on a glass plate and takes place at the sample–air interface. The samples were held on a temperature-controlled heating stage.

3. Results and discussion

3.1 Mesomorphic properties of compounds **BrFn** depending on terminal alkyl chain length

The phase sequences, transition temperatures ($^\circ\text{C}$) and associated phase transition enthalpies (kJ mol⁻¹) are summarized in Table 1 (see also Table S1† for data from the second heating scans). In the series of the bromosubstituted compounds **BrFn**

only the shortest homologue **BrF4** having a relatively high melting point at $T = 157^\circ\text{C}$ is non-mesomorphic. Also for the next homologue **BrF6** a direct transition from the crystalline state to the isotropic liquid takes place on heating at 126°C , but on cooling from the isotropic state a highly viscous optically isotropic mesophase appears at $T = 96^\circ\text{C}$, as indicated by microscopy between crossed polarizers. Rotating the analyzer by a small angle out of the precise 90° position leads to the appearance of dark and bright domains. Rotating the analyzer in the opposite direction reverses the dark and bright domains (see Fig. 1 for compound **BrF12**). Rotating the sample between crossed polarizers does not lead to any change and this indicates that the distinct regions represent chiral domains with opposite handedness, as typical for dark conglomerate phases (DC phases). No crystallization of this DC phase is observed upon cooling to room temperature. In the following heating cycles crystallization takes place either in the DC phase region or after transition of the DC phase to the isotropic liquid (see Fig. S1†). No crystallization is observed for any of the next homologues with $n = 8$ –14 (see Table 1 and Fig. 2b), even after storage for one year at room temperature.

All compounds **BrF8**–**BrF14** have very similar transition temperatures. The melting of the DC phases (second heating scans) takes place around 106°C and the formation of the DC phase on cooling occurs between 96 and 99°C (see Table 1 and Fig. 2a). Hence, there is a supercooling of this phase transition by about 8 K (peak temperatures), which is found nearly



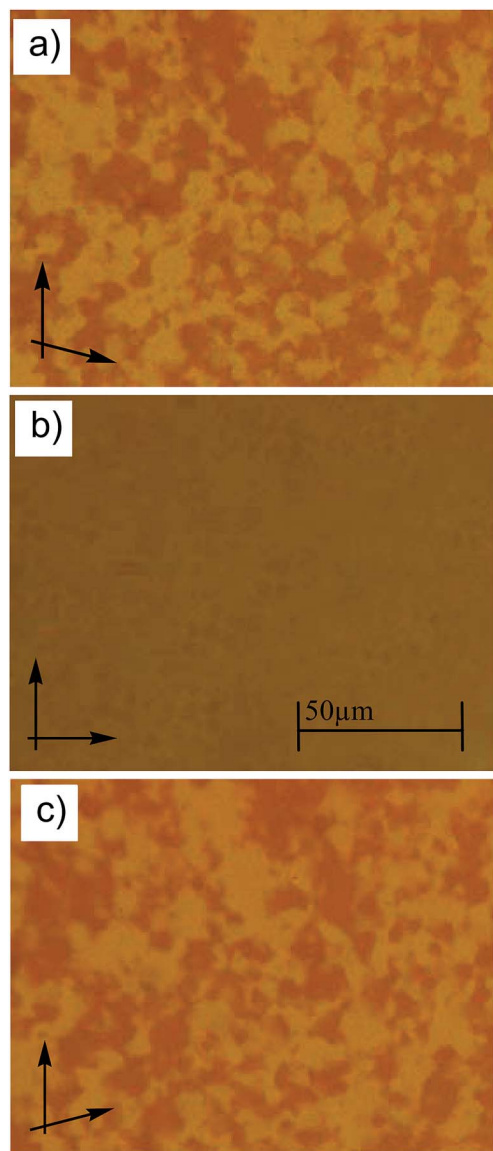


Fig. 1 Textures of the DC phase of compound **BrF12** at $T = 80\text{ }^{\circ}\text{C}$: (b) between crossed polarizers and (a) after rotating one polarizer by 7° from the crossed position in clock-wise direction and (c) in the anti-clockwise direction, showing dark and bright domains, indicating the presence of areas with opposite chirality sense.

independent of the scanning rate (2 or 10 K min^{-1}). The formation of the DC phases is associated with relatively high transition enthalpies, ranging between $\Delta H \sim 23.8$ and 28.5 kJ mol^{-1} , only slightly rising with growing chain length. Upon further chain elongation for **BrF16** with $n = 16$, the DC phase becomes unstable and immediately after its formation crystallization takes place. For the longest homologue **BrF18** ($n = 18$) the DC phase is completely removed and only a birefringent crystalline phase is observed.

In electro-optical experiments no current peak could be observed in the DC phases of any of the prepared materials **BrF6–BrF14** and also no birefringence is induced under an applied triangular wave voltage up to 200 Vpp in a $6\text{ }\mu\text{m}$ ITO cell. This behaviour is similar to those known for soft crystalline

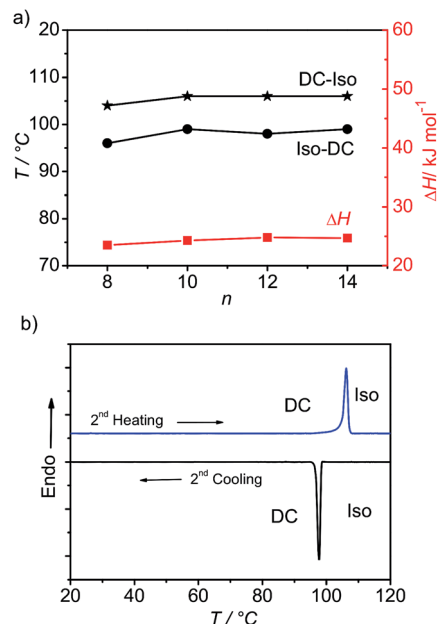


Fig. 2 (a) Dependence of DC–Iso transition temperatures (on heating and cooling scans with 10 K min^{-1}) and transition enthalpy values (on cooling from the isotropic liquid) of compounds **BrF8–BrF14** depending on alkyl chain length n and (b) DSC heating and cooling curves of compound **BrF12** with a rate of 10 K min^{-1} .

HNF phases (B_4 phases) and the DC phases of other related azobenzene based bent core mesogens,^{43,44} but it is clearly distinct from most of the fluid sponge-like smectic phases.

XRD investigation of the DC phases of compounds **BrF6**, **BrF8**, **BrF12** and **BrF14** (see Fig. 3a and b, 4 and S2–5†) shows an intense reflection with d -values in between half of the molecular length $L_{\text{mol}}/2$ and the full length L_{mol} (see Fig. 3c). For the determination of L_{mol} a 120° V-shaped conformation with all-*trans* alkyl chains was assumed (see Fig. S6†). The d/L_{mol} ratio was thus calculated to be in the range of 0.78–0.80 for all investigated compounds.

A linear increase of the d -values is observed with increasing length of the terminal chains from $d = 3.46\text{ nm}$ for **BrF6** to $d = 4.76\text{ nm}$ for **BrF14** (Fig. 3c). Thus the intense small angle scattering is assigned to a layer reflection. The d/L ratio around 0.78–0.80 would, according to $d/L_{\text{mol}} = \cos \beta$, allow a tilt angle of the molecules of around 38° . This relatively large difference between d and L_{mol} is similarly found for the DC phases of the related 4-iodo and 4-methylresorcinols **Ifn** and **Mn**,^{43,44} but distinguishes these DC phases from the previously reported HNF phases where d is usually close to the molecular length.^{24,25} No alignment could be achieved and therefore in the 2D patterns all scatterings form closed rings with uniform intensity distribution (see Fig. 3a). This is due to the disordered meso-scale structure, which is an inherent and very typical feature of all DC phases and leads to their optically isotropic appearance as well as to a broadening of the scattering; the domain size (determined using the Scherrer equation and assuming $K = 1$)⁵⁰ is in the range of 30–42 nm and grows with increasing alkyl chain length (see Fig. 3c).



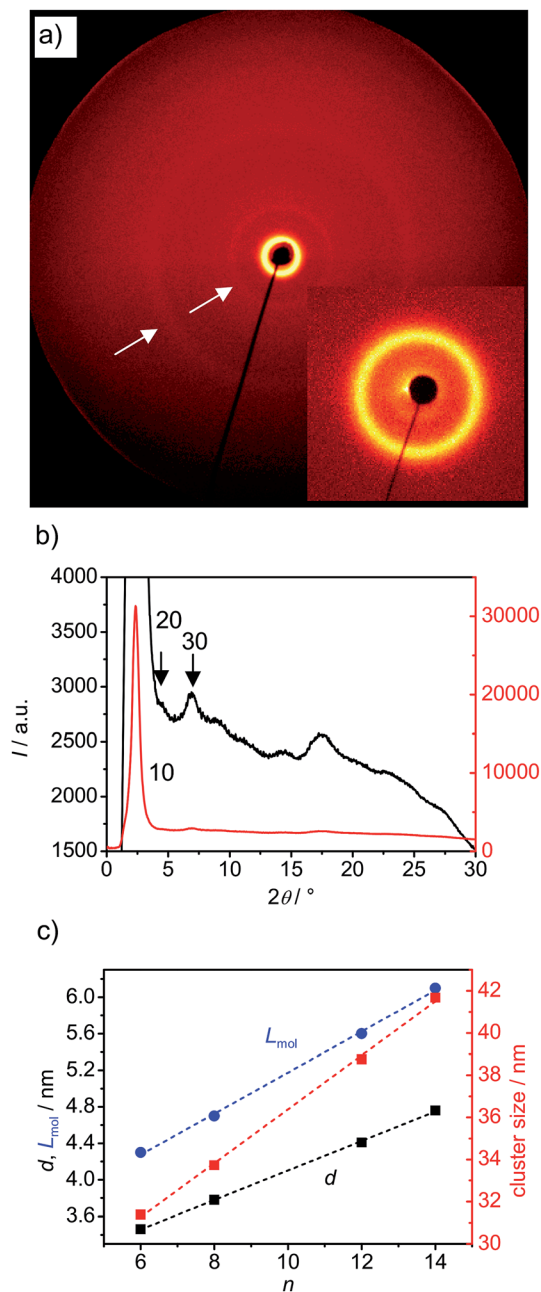


Fig. 3 (a) 2D XRD pattern of the DC phase of **BrF8** at $T = 90\text{ }^{\circ}\text{C}$, the inset shows the small angle region; (b) 2θ -scans over this XRD pattern; (c) dependence of d -values, molecular lengths (L_{mol} ; the used conformation is shown in Fig. S6†) and cluster size (determined using the Scherrer equation and assuming $K = 1$) in the DC phases of compounds **BrF6**–**BrF14** on the chain length.

The 2θ scan over the diffraction pattern of compound **BrF8** is shown in Fig. 3b, those of compounds **BrF6**, **BrF12** and **BrF14** are collated in Fig. 4. Beside the strong layer reflections very weak and broad scattering maxima are observed in the medium angle region ($2\theta = 5\text{--}9^{\circ}$) and in the wide angle region ($2\theta = 14\text{--}28^{\circ}$). This pattern excludes fluid sponge phases, which would show exclusively one completely diffuse wide angle scattering besides the layer reflection. The wide angle scattering is, similarly to the HNF phases, characterized by several scattering

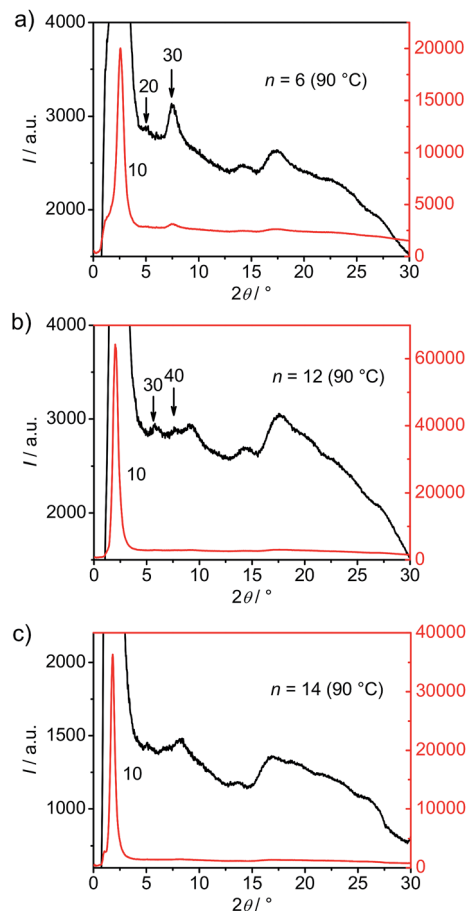


Fig. 4 Comparison of 2θ -scan over the XRD patterns of the DC phase of (a) **BrF6**, (b) **BrF12** and (c) **BrF14** at $T = 90\text{ }^{\circ}\text{C}$ (for more details, see Fig. S2–S5†).

maxima up to $2\theta = 28^{\circ}$. However, in contrast to typical patterns of HNF phases (B_4 phases, see Fig. 5b), the wide angle reflections are much broader and have significantly reduced intensity (compare red line in Fig. 5a with 5b, both shown in the logarithmic scale). For compounds **BrF8**–**BrF12** most reflections in the medium angle region can be indexed as harmonics (up to 4th order) of the layer reflection (see Fig. 4 and S2–S5†), as it is also the case for the previously reported **Mn** compounds⁴⁴ and usually found for HNF phases.^{18,24–26} Only the XRD pattern of compound **BrF14**, the compound with the longest alkyl chain among the compounds with stable DC phases, appears a bit different from the others, as the medium angle scatterings cannot be assigned as higher order layer reflections and also the shape of the wide angle scattering is a bit different (Fig. 4c).

Fig. 5a shows a comparison of the 2θ -scans of the DC phases of the related compound **M14** ($Y = \text{CH}_3$), **BrF14** ($Y = \text{Br}$) and **IF14** ($Y = \text{I}$) all having the same chain length. It clearly indicates that, though the position of the 10 reflection is nearly identical in all cases, the positions and intensities of the medium and wide angle scattering maxima are very distinct for the halogen substituted compounds **IF14** and **BrF14** compared to the methyl substituted compound **M14**. The scattering intensity of the iodo compounds **IFn** is the lowest, followed by the bromine

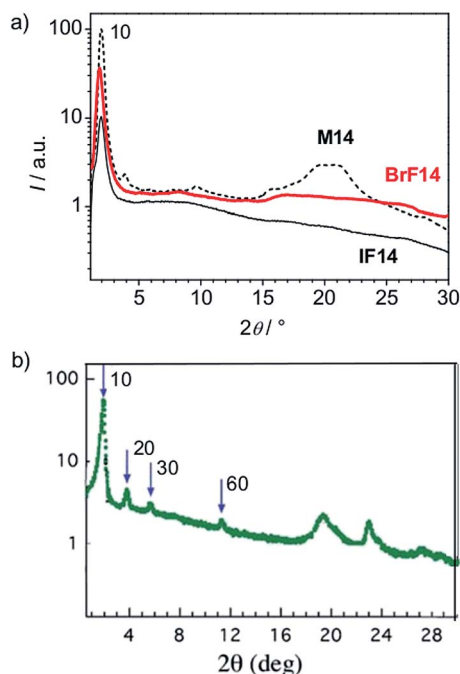


Fig. 5 (a) Comparison of the 2θ -scans of the DC phases of compounds **BrF14**, **IF14** and **M14** (base line is shifted for better visibility; the scattering intensity of the **IFn** compounds is so low that scattering by air contributes to the diffuse scattering in the medium angle range ($5\text{--}10^\circ$) of the XRD pattern of these I-substituted compounds) and (b) typical 2θ -scans of a HNF phase (B_4 phase) of a benzylideneaniline based bent-core mesogen (P-8-OPIMB) reproduced with modifications and permission from ref. 21, copyright RSC 2009.

substituted compounds **BrFn** and compounds **Mn**. Though having a higher intensity the wide angle scattering profile of the bromo compound **BrF14** is similar to **IF14**, indicating similarities in the molecular organizations in their DC phases.

3.2 Investigation of mixtures of compounds **BrFn** with 5-CB

It is known that HNF phases of benzylideneanilines can be diluted by nematic LC hosts to a high degree ($>95\%$) without loss of the chirality.⁵¹ This is a direct consequence of the helical nano-filament structure of these phases, allowing a swelling of the filaments by the nematic LC and transfer of chirality from the nano-filaments to the nematic LC. This effect is not observed for the sponge type liquid crystalline DC phases involving distorted fluid smectic layers. Therefore, investigation of mixtures with 4'-n-pentyl-4-cyanobiphenyl (5-CB) could be used as a tool providing additional information about the microstructure of the DC phases of compounds **BrFn**.

The 1 : 1 mixtures of **BrF8** or **BrF10** with 5CB do not show any DC phase. On heating these mixtures only a direct transition from the crystalline material to the isotropic liquid state occurs at 57°C and 54°C , respectively (Table 2). On cooling both of these mixtures form a nematic phase which crystallizes with formation of highly birefringent crystalline phases at $T \sim 36^\circ\text{C}$ and 33°C , respectively (see Table 2), but no DC phase is formed. The next homologue **BrF12** behaves differently; on heating this

Table 2 Phase transition temperatures and mesophase types of 1 : 1 mixtures of 5-CB and compounds **BrF8**–**BrF18** and comparison with related 4-methylresorcinol derivatives **M14**–**M18**^a (ref. 44)

Mixture	Heating $T/^\circ\text{C}$	Cooling $T/^\circ\text{C}$
BrF8 + 5-CB	Cr 57 Iso	Iso 43 N 36 Cr
BrF10 + 5-CB	Cr 54 Iso	Iso 42 N 33 Cr
BrF12 + 5-CB	Cr ^[*] 40 DC 55 Iso	Iso 53 DC 38 Cr ^[*]
BrF14 + 5-CB	Cr 38 DC 64 Iso	Iso 46 DC 35 Cr
BrF16 + 5-CB	Cr ₁ 33 Cr ₂ 69 Iso	Iso 63 Cr
BrF18 + 5-CB	Cr ₁ 42 Cr ₂ 75 Iso	Iso 65 Cr
M14 + 5-CB	Cr 85 Iso	Iso 54 N 53 Cr
M16 + 5-CB	Cr 64 DC 73 Iso	Iso 66 DC
M18 + 5-CB	DC 77 Iso	Iso 77 DC

^a Transition temperatures were taken from the observed textures using the polarized optical microscopy; abbreviations: N = nematic phase; Cr^[*] = crystalline phase composed of a conglomerate of chiral domains; for other abbreviations please see Table 1.

mixture a transition from the crystalline phase to the DC phase occurs at 40°C (see Fig. S7d–f†). On further heating a transition from the DC phase to the isotropic liquid takes place at 55°C . On cooling this mixture from the isotropic liquid the transition to the DC phase takes place at $T \sim 53^\circ\text{C}$ followed by crystallization at 38°C . Interestingly, this crystalline phase is nearly isotropic and composed of domains with opposite handedness (see Fig. S7a–c†). The mixture of the next homologue **BrF14** shows similar behaviour to that of **BrF12**, except that this weakly birefringent crystalline phase is achiral or the chiral domains are too small to be detected. Mixing the longer crystalline compounds **BrF16** and **BrF18** with 5-CB does not give any DC phases (see Table 2).

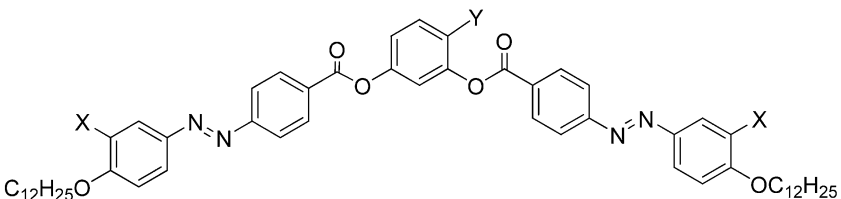
The significant destabilization of the DC phases by 5-CB confirms that the DC phases of compounds **BrFn** should be different from the classical HNF phases.^{21,24–26} It appears that these phases are formed by smaller nano-domains instead of long helical nano-filaments, in line with the results of the XRD studies showing relatively broad wide angle scatterings. Because no extended filaments are formed, no gel-like networks of these fibers are possible and therefore these compounds can take up only a limited amount of 5-CB, which is directly incorporated between the bent-core molecules and thus reduces the stability of the DC phases. The highest tendency for DC phase formation appears to be provided by the medium chain compounds **BrF12** and **BrF14**. Comparison with results obtained with related compounds indicate that the 5-CB mixtures of compounds **BrFn** behave very similar to the 5-CB mixtures of the iodo compounds **IFn**,⁴³ whereas for the 4-methylresorcinols **Mn** the DC phases in the mixtures have higher stability and their regions are shifted to longer alkyl chain lengths (Table 2), indicating slightly more stable aggregates.

3.3 Comparison of different types of 4-substituted resorcinol based BCLCs with azobenzene units

In Table 3 compound **BrF12** is compared with the non-substituted compound **H12** (ref. 52) and related BCLCs with



Table 3 Phase transition temperatures ($T/^{\circ}\text{C}$) and mesophase types as observed for different types of 4-substituted resorcinols (Y) with azo-benzene wings and the effect of peripheral F-atoms (X = H, F)^a



Compound	Y	cv^b/nm^3	μ^c/D	X	$T/^{\circ}\text{C}$	Ref.
H12	H	7		H	Cr 145 Iso	52
HF12	H			F	Cr 138 Iso	
F12	F	13	1.60	H	Cr 120 Iso	47
FF12	F			F	Cr 120 (SmCP _A 119) Iso	
Cl12	Cl	27	1.69	H	Cr ₁ 90 Cr ₂ 103 (N 97) Iso	47
ClF12	Cl			F	Cr 115 (SmCP _A 97) Iso	
Br12	Br	33	1.70	H	Cr (N 83) 93 Iso	47
BrF12	Br			F	DC 106 Iso	
I12	I	45	1.70	H	Cr 112 Iso	43
IF12	I			F	DC 111 Iso	43
M12	CH ₃	32	0.37	H	DC 102 Iso	44
MF12	CH ₃			F	Cr 90 SmCP _A 92 Iso	
CN12	CN	23	4.18	H	Cr 71 (SmC _s P _A 70) M ₁ 75 SmC _s P _F 83 SmC _s P _R ^[*] 96 SmC 127 N 137 Iso	32
CNF12	CN			F	Cr 104 SmC _s P _A 108 SmC _s P _R ^[*] 135 SmA 147 Iso	^d

^a Abbreviations: N = nematic phase; SmA = nonpolar smectic A phase; SmC = paraelectric smectic C phase; SmC_sP_R^[*] = synclitic tilted SmC phase composed of SmC_sP_F domains with randomized polar directions, homeotropically aligned samples of this phase are composed of a conglomerate of chiral domains; SmC_sP_F = polar SmC phase composed of enlarged ferroelectric domains; M₁ = highly viscous antiferroelectric switching SmC phase; SmCP_A = antiferroelectric switching SmC phase (B2 phase); SmC_sP_A = synclitic tilted antiferroelectric SmC phase (B2 phase); for other abbreviations see Table 1. ^b Crystal volumes of Immirzi.⁵³ ^c Dipole moments of the corresponding phenyl derivatives.⁵⁴ ^d Unpublished data.

other substituents in the 4-position of the resorcinol core, like halogens (Y = F, Cl, I), methyl and cyano. Comparison of the different azobenzene based BCLCs (all with $n = 12$) indicates that there is a tendency for reduction of the melting points by the substituents Y which becomes the more important the larger these substituents are. Thus LC phases can be observed for most of the 4-substituted compounds, whereas the non-substituted compounds **H12** and **HF12** represent relatively high melting solids. There appears to be a dependence of the phase type on the volume of Y (crystal volumes cv of Immirzi⁵³) and the polarity of the C–Y bond (dipole moments μ of the Ph–Y compounds⁵⁴). For molecules with X = H and having polar substituents Y at the resorcinol core no LC phase is observed (Y = F, I) or nematic phases are dominating (Y = Cl, Br, CN), whereas the nonpolar methyl group is obviously favourable for the formation of DC phases (compound **M12**). Introducing peripheral F-substituents increases the tendency for formation of polar SmC phases (SmCP_A) if the 4-substituent is small (Y = F, Cl, CN). Formation of DC phases is only supported by peripheral fluorination of BCLCs with large and polar substituents Y (Y = Br, I).

So, overall, large substituents in the 4-position at the resorcinol core appear to be favourable for DC phase formation, although there is an additional effect of peripheral F-substitution. Obviously the effect of peripheral fluorine substitution is the reverse for BCLCs with polar and non-polar substituents

Y. While for the methyl-substituted compounds the introduction of F removes the DC phase and a SmCP_A phase is formed, for compounds with large polar halogens (Br, I) introduction of peripheral F-substitution induces DC phases. The reason for this behaviour is not yet clear.

As can be deduced from the comparison in Fig. 2a and 6a, the transition enthalpy values of the DC–Iso transitions and the corresponding transition enthalpy values of the iodine substituted compounds **IF8–IF14** and for the bromine substituted analogues **BrF8–BrF14** are in the same range, once again confirming the similarity between the DC phases of the two series, already concluded from the similarity of the XRD patterns. Nevertheless, the DC–Iso transition temperatures and the related transition enthalpy values of compounds **IFn** are slightly higher (by ~ 3 to 10 K and ~ 2 to 6 kJ mol^{−1}, respectively) than those of their bromine substituted analogues **BrFn**. This might be due to the increased polarizability provided by iodine, increasing the attractive dispersion forces between the molecules. In line with increased intermolecular interactions a denser packing of compounds **IFn** could provide a stronger effect of molecular conformational chirality (helicity) on the aggregate stability and aggregate size. Also the observation that compounds **IF8–IF14** exhibit exclusively the DC phases and do not show any crystallization, while compounds **BrF8–BrF14** show crystalline phases in the first heating and thus form monotropic DC phases (though, the DC phases once formed



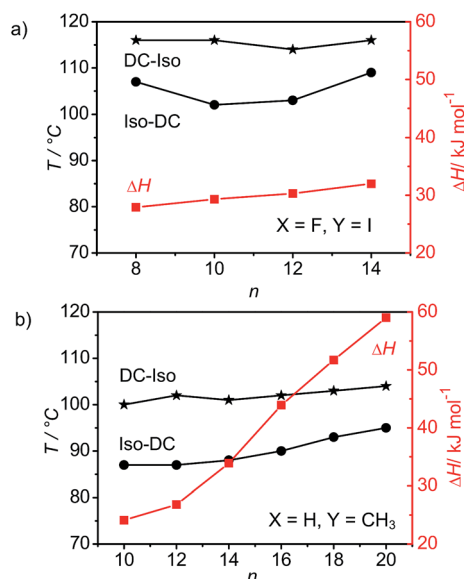


Fig. 6 Dependence of DC–Iso transition temperatures (on heating and cooling scans with 10 K min^{-1}) and transition enthalpy values (on cooling from the isotropic liquid) of two series of compounds related to **BrFn** (a) the 4-iodine-substituted compounds **IF8–IF14** (ref. 43) and (b) the 4-methyl-substituted compounds **M10–M20** (ref. 44) depending on the alkyl chain length n .

also do not crystallize) is in line with a stronger distortion of the long range molecular packing in the case of the iodo compounds. This could be attributed to the larger size of iodine compared to Br, which could provide a stronger helical twist of the molecular conformations.

Comparison of the series **BrFn** (Fig. 2a) with the methyl-substituted compounds **Mn** (Fig. 6b) with chain length $n = 10$ –14 indicates about 10 K higher DC–Iso transition temperatures for the compound having the more polar bromine substituent (and an additional peripheral F-substituent). The DC–Iso transition enthalpy values are comparable in both series, but for compound **Mn** there is a much stronger enthalpy increase upon chain elongation and in this case alkyl chain crystallization significantly contributes to the transition enthalpies.⁴⁴ There are clear difference also in the XRD patterns of the methyl substituted and the halogen substituted BCLCs (see Fig. 5a), indicating that the DC phases of compounds **Mn** are different from those of **BrFn** and **IFn**. The presence of only lower harmonics (up to the 4th) of the layer reflections (5th and 6th order harmonics of the layer reflections has been reported for typical B_4 phases, see for example Fig. 5b (ref. 21)) and the significantly increased broadness of the wide angle scatterings indicates a stronger distortion of layer structures and filament formation in the case of the halogen substituted compounds. This is also in line with the reduced resistance of the DC phases of compounds **BrFn** and **IFn** against dilution with the nematic phase of 5-CB (see Table 2).

DFT calculations of the model compounds **F1**, **Br1**, **I1**, **M1** and **CN1** with $n = 1$ were performed with the Gaussian 09 package,⁵⁵ using the B3LYP functional and LANL2DZ basis set using the solvation model IEFPCM in solvent chlorobenzene to

investigate the effects of the 4-substituents on the bending angle (γ) and on the twist of the wing groups in the lowest energy conformation. The results are summarized in Fig. 7 where two different projections are shown and in Table 4, providing numerical values of bending angles γ and twist angles α and β . In Fig. 7a the view is perpendicular to the resorcinol ring and in Fig. 7b the view is parallel to this ring along the wing in the 3-position. For all compounds, except **CN1**, the bending angle γ is not significantly affected by the 4-substituent and is around 120 – 123° in all cases (see Fig. 7a and Table 4). Only for **CN1** it is increased to about 130° due to the dipole interactions between the CN group and the adjacent C=O group, in line with previous investigations.⁵⁶ This contributes to the very different behaviour of the cyanoresorcinols **CNn** (ref. 31 and 32) compared to the other considered compounds (see Table 3).

The azobenzene units themselves are linear and the benzene rings in these wings as well as the C=O groups of the carboxylate groups are nearly coplanar in all cases. Different minimum energy conformations result from the twist of the planes of the azobenzene wings with respect to the resorcinol core and thus to each other, leading to chiral conformers for all investigated compounds. The angle α_1 , which describes the twist between the plane of the azobenzene wing in the 1-position with respect to the plane of the resorcinol core, is around 50° for all halogen substituted compounds. Larger differences can be found for the twist α_2 of the azobenzenes in the 3-position which changes between $\alpha_2 = 49$ and 109° (see Fig. 7b and Table 4). Overall, the effective twist between the planes of the two azobenzene wings β increases in the order $\text{F} < \text{Br} < \text{CH}_3 < \text{I}$ roughly in line with the growing size of the 4-substituent (see cv in Table 3; only the effect of the CH_3 group is larger than that of Br though their volumes are nearly equal). This is in line with the observed tendency of these molecules to form DC phases. Accordingly, no DC phase was found for any of the resorcinol and 4-fluororesorcinol derivatives, whereas DC phases were found for compounds with $\text{Y} = \text{Br}$, CH_3 and I. Although for $\text{Y} = \text{CN}$ the twist angle β is the largest, these molecules behave differently, due to the increased bending angle ($\gamma \sim 130^\circ$) and the possibility of antiparallel compensation of the very large dipoles of these groups. Therefore, these CN-substituted compounds do not form DC phases, but preferably form nematic and SmC phases. Nevertheless, the large twist in the low energy conformation might be responsible for the formation of chiral domains in some of the birefringent SmC phases of these compounds ($\text{SmC}_s\text{P}_R^{[*]}$).^{31,32}

Based on the combination of experimentally observed structure property relationships (Table 3) and results of DFT calculations the following relationships between the molecular structure and formation of DC phases can be hypothesized. As the azobenzene wings are completely flat the only source of conformational chirality is in this case the central part of the molecules, i.e. the connections between the resorcinol core and azobenzene wings. For this reason the substitution in position 4 at the resorcinol is important and has a significant effect on the molecular conformation and hence the ability to form DC phases. Due to the rigidity of the azobenzenes the



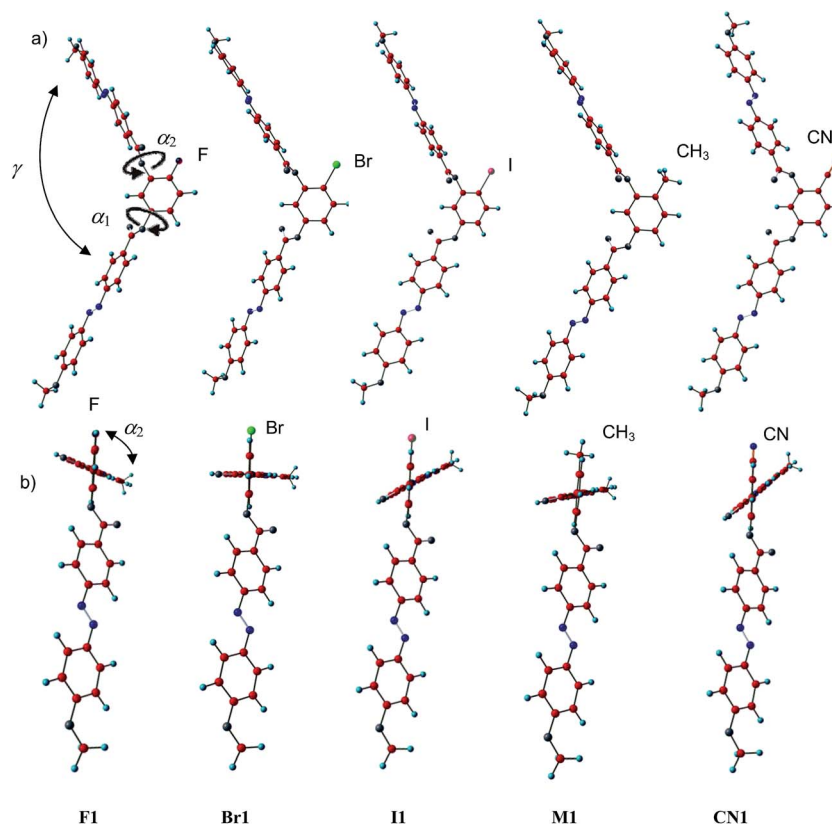


Fig. 7 Energy minimized molecular conformers of the model compounds ($n = 1$) as observed under two distinct perspectives, (a) perpendicular to the plane of the resorcinol core indicating changes of the bending angle γ and (b) along the azobenzene wing in the 3-position and parallel to the plane of the resorcinol core, indicating the distinct degrees of twist between the two planes of the azobenzene wings; the angles α_1 , α_2 and γ are explained for **F1**.

Table 4 Selected parameters of the energy minimized conformers obtained by DFT calculations of the distinct model compounds **Y1** ($n = 1$) depending on the 4-substituent **Y** at the resorcinol core^a

Y	$\alpha_1/^\circ$	$\alpha_2/^\circ$	$\beta/^\circ$	$\gamma/^\circ$
F	50.9	109.1	31.8	121.5
Br	48.8	89.8	41.4	120.1
I	49.8	58.1	72.1	123.4
CH ₃	53.6	72.4	54.0	119.9
CN	44.8	49.0	86.2	129.7

^a β = twist between the planes of the azobenzenes, calculated according to $\beta = (90 - \alpha_1) + (90 - \alpha_2)$, where α_1 and α_2 represent the twist angles of the planes of the azobenzenes with respect to the plane of the resorcinol unit, (α_1 azobenzene in the 1-position, α_2 in the 3 position); γ = bending angle which is calculated as the angle between the lines connecting the terminal ether oxygens at the azobenzenes with the carbon atoms 1 and 3 of the resorcinol core, respectively.

conformational chirality provided by the core has a strong effect on the overall molecular shape, providing a strong distortion to any packing in layers/filaments, thus leading to strongly distorted types of semicrystalline DC phases.⁵⁷ In contrast, other typical wing groups such as phenylbenzoates,² benzylideneanilines⁵⁸ and biphenyls² can themselves easily adapt twist conformations and can therefore contribute to the overall conformational chirality. In line with this, for such compounds

a bulky substituent in the central core unit is not required for DC phase formation. Moreover, as the helical twist can arise in the peripheral wing units the molecular shape is less distorted and such compounds can more easily adapt to the organization in twisted layers and extended helical filaments, predominately leading to the previously reported types of DC phases, the LC smectic sponges (phenyl benzoates) and the soft crystalline HNF phases (benzylidene anilines), respectively.

4. Conclusions and summary

Bent-core molecules derived from 4-bromoresorcinol with peripherally fluorinated azobenzene wings (**BrFn**) represent new materials forming optically isotropic DC phases composed of conglomerates of uniformly chiral domains, which can be observed for a chain length range $n = 8$ –14 as the only mesophase, whereas shorter and longer homologues form only birefringent crystalline phases. The DC phases of compounds with $n = 10$ –14 are stable down to ambient temperature and do not crystallize even after prolonged storage. For related compounds without the peripheral fluoro substitution (**Brn**) a DC phase was observed for only one homologue ($n = 10$) as a monotropic phase, whereas the other homologues form monotropic nematic and B_6 phases.⁴⁷ This indicates a strong DC phase stabilizing effect of peripheral F substitution in this case, which is reverse to the



4-methylresorcinols, for which DC phases were found for the non-fluorinated compounds **Mn** whereas the fluorinated compounds **MFn** form antiferroelectric switching SmCP_A phases (B2 phases).⁴⁴ XRD investigations and the relatively high viscosity of these DC phases indicate that they do not belong to the fluid sponge type smectic LC phases, but are more related to the soft crystalline HNF phases. However, in contrast to the classical HNF phases the helical nano-filaments appear to be not well developed, so that the helical twist gives rise to a strongly disordered structure formed by nanoscale twisted domains of crystallized aromatic cores separated by disordered alkyl chains. In this way the DC phases are similar to those formed by the related 4-iodoresorcinol derivatives,⁴³ but distinct from those formed by the previously reported B₄ type HNF phases of the benzylideneanilines.²⁴ It is thought that for the compounds **BrFn** the filaments are disrupted into smaller segments, as also proposed for the methyl substituted compounds **Mn**, and it appears that the degree of disruption increases in the order **Mn** < **BrFn** < **IFn**.

DFT calculations allowed the estimation of the effects of the substituents Y on the minimum energy conformers. This indicates that the azobenzenes themselves are flat and that mainly steric effects of the substituent Y lead to an increase of the twist between the planes of the azobenzene wings with increasing size of Y. This appears to be favourable for chiral segregation and layer distortion, due to the difficulty in the parallel packing of these highly twisted molecular species. The strong layer distortion is thought to be responsible for the preferred formation of twisted nano-domains instead of sponge-like deformed layers or extended helical nano-filaments.

The effect of polarity on the molecular conformation is the strongest for the CN substituted compounds, giving rise to a less bent (more rod-like) molecular shape, which appears to be unfavourable for formation of DC phases, but favours the formation of nematic and SmC phases. Nevertheless, the strong twist between the planes of the azobenzenes leads to a high “strength of chirality” of the energy minimum conformers which allows chiral segregation even in some of their polarization randomized SmC phases (SmC_sP_R^[*] phases).

In summary, the development of new types of DC phases depending on the molecular structure was investigated in a systematic way. Relationships between the molecular structure and formation of distinct types of DC phases were found and possible reasons for these relationships were hypothesized. We believe that this study will contribute to the improvement of the general knowledge about DC phases and the mechanisms of achiral symmetry breaking in soft matter as well as to the design of new molecules showing DC phases. BCLCs involving azo linkages are also very versatile materials due to their photochromic effects and high birefringence. Moreover, the *trans-cis* photoisomerization of the azobenzene wings in these BCLCs could lead to interesting perspectives for chirality switching and phase modulation by interaction with nonpolarized, linear or circular polarized light, *i.e.* to unique and potentially useful multifunctional chiral materials for application in high-density data storage systems, sensors, photonic switches and molecular logic gates,^{59,60} as well as with potential for discrimination of chiral physical forces and molecular species.

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References

- (a) R. A. Reddy and C. Tschierske, *J. Mater. Chem.*, 2006, **16**, 907; (b) H. Takezoe and Y. Takanishi, *Jpn. J. Appl. Phys.*, 2006, **45**, 597; (c) A. Eremin and A. Jakli, *Soft Matter*, 2013, **9**, 615.
- H. Takezoe, *Top. Curr. Chem.*, 2012, **318**, 303.
- (a) F. Vera, J. L. Serrano and T. Sierra, *Chem. Soc. Rev.*, 2009, **38**, 781; (b) M. Lehmann, M. Jahr, B. Donnio, R. Graf, S. Gemming and I. Popov, *Chem.-Eur. J.*, 2008, **14**, 4414; (c) C. Roche, H.-J. Sun, M. E. Prendergast, P. Leowanawat, B. E. Patridge, P. A. Heiney, F. Araoka, R. Graf, H. W. Spiess, X. Zeng, G. Ungar and V. Percec, *J. Am. Chem. Soc.*, 2014, **136**, 7169–7185; (d) M. H. C. J. V. Houtem, F. Benaskar, C. F. C. Fitié, R. Martín-Rapún, J. A. J. M. Vekemans and E. W. Meijer, *Org. Biomol. Chem.*, 2012, **10**, 5898; (e) R. Q. Albuquerque, A. Timme, R. Kress, J. Senker and H.-W. Schmidt, *Chem.-Eur. J.*, 2013, **19**, 1647.
- H. Nagayama, S. K. Varshney, M. Goto, F. Araoka, K. Ishikawa, V. Prasad and H. Takezoe, *Angew. Chem.*, 2010, **49**, 445.
- (a) D. Pijper and B. L. Feringa, *Soft Matter*, 2008, **4**, 1349; (b) K. Maeda and E. Yashima, *Top. Curr. Chem.*, 2006, **265**, 88; (c) C. C. Lee, C. Grenier, E. W. Meijer and A. P. H. J. Schenning, *Chem. Soc. Rev.*, 2009, **38**, 671; (d) W. Jin, Y. Yamamoto, T. Fukushima, N. Ishii, J. Kim, K. Kato, M. Takata and T. Aida, *J. Am. Chem. Soc.*, 2008, **130**, 9434.
- J. V. Selinger, M. S. Spector and J. M. Schnur, *J. Phys. Chem. B*, 2001, **105**, 1757.
- E. Barry, Z. Hensel, Z. Dogic, M. Shribak and R. Oldenbourg, *Phys. Rev. Lett.*, 2006, **96**, 018305.
- A. Lohr and F. Würthner, *Angew. Chem., Int. Ed.*, 2008, **47**, 1232.
- (a) E. Yashima, K. Maeda, H. Iida, Y. Furusho and K. Nagai, *Chem. Rev.*, 2009, **109**, 6102; (b) H. Jinnai, T. Kaneko, K. Matsunaga, C. Abetz and V. Abetz, *Soft Matter*, 2009, **5**, 2042.
- D. B. Amabilino, *Chirality at the Nanoscale*, VCH-Wiley Weinheim, 2009.
- D. R. Link, G. Natale, R. Shao, J. E. MacLennan, N. A. Clark, E. Körblová and D. M. Walba, *Science*, 1997, **278**, 1924.
- E. A. Matsumoto, G. P. Alexander and R. D. Kamien, *Phys. Rev. Lett.*, 2009, **103**, 257804.
- (a) G. Dantlgraber, A. Eremin, S. Diele, A. Hauser, H. Kresse, G. Pelzl and C. Tschierske, *Angew. Chem., Int. Ed.*, 2002, **41**, 2408; (b) G. Dantlgraber, S. Diele and C. Tschierske, *Chem. Commun.*, 2002, 2768; (c) C. Tschierske and G. Dantlgraber, *Pramana*, 2003, **61**, 455; (d) C. Keith, R. A. Reddy, A. Hauser, U. Baumeister and C. Tschierske, *J. Am. Chem. Soc.*, 2006, **128**, 3051; (e) C. Keith, R. A. Reddy, U. Baumeister, H. Hahn, H. Lang and C. Tschierske, *J. Mater. Chem.*, 2006, **16**, 3444; (f) H. Hahn, C. Keith,



- H. Lang, R. A. Reddy and C. Tschierske, *Adv. Mater.*, 2006, **18**, 2629; (g) C. Keith, G. Dantlgraber, R. A. Reddy, U. Baumeister, M. Prehm, H. Hahn, H. Lag and C. Tschierske, *J. Mater. Chem.*, 2007, **17**, 3796; (h) C. Keith, G. Dantlgraber, R. A. Reddy, U. Baumeister and C. Tschierske, *Chem. Mater.*, 2007, **19**, 694; (i) Y. Zhang, U. Baumeister, C. Tschierske, M. O'Callaghan and C. Walker, *Chem. Mater.*, 2010, **22**, 2869.
- 14 L. E. Hough, M. Spannuth, M. Nakata, D. A. Coleman, C. D. Jones, G. Dantlgraber, C. Tschierske, J. Watanabe, E. Körblova, D. M. Walba, J. E. MacLennan, M. A. Glaser and N. A. Clark, *Science*, 2009, **325**, 452.
- 15 G. Heppke, D. D. Parghi and H. Sawade, *Liq. Cryst.*, 2000, **27**, 313.
- 16 J. Thisayukta, Y. Nakayama, S. Kawauchi, H. Takezoe and J. Watanabe, *J. Am. Chem. Soc.*, 2000, **122**, 7441.
- 17 R. A. Reddy and B. K. Sadashiva, *Liq. Cryst.*, 2003, **30**, 1031.
- 18 A. Roy, M. Gupta, S. Radhika, B. K. Sadashiva and R. Pratibha, *Soft Matter*, 2012, **8**, 7207.
- 19 J. Ortega, C. L. Folcia, J. Etxebarria, N. Gimeno and M. B. Ros, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 2003, **68**, 11707.
- 20 T. Niori, T. Sekine, J. Watanabe, T. Furukawa and H. Takezoe, *J. Mater. Chem.*, 1996, **6**, 1231.
- 21 J. M. Perdiguerro, I. Alonso, C. L. Folcia, J. Etxebarria and J. Ortega, *J. Mater. Chem.*, 2009, **19**, 5161.
- 22 C. Zhang, N. Diorio, O. D. Lavrentovich and A. Jakli, *Nat. Commun.*, 2014, **5**, 3302.
- 23 (a) A. Jakli, Y.-M. Huang, K. Fodor-Csorba, A. Vajda, G. Galli, S. Diele and G. Pelzl, *Adv. Mater.*, 2003, **15**, 1606; (b) W. Weissflog, M. W. Schröder, S. Diele and G. Pelzl, *Adv. Mater.*, 2003, **15**, 630.
- 24 (a) L. E. Hough, H. T. Jung, D. Kröuerke, M. S. Heberling, M. Nakata, C. D. Jones, D. Chen, D. R. Link, J. Zasadzinski, G. Heppke, J. P. Rabe, W. Stocker, E. Körblova, D. M. Walba, M. A. Glaser and N. A. Clark, *Science*, 2009, **325**, 456; (b) D. M. Walba, L. Eshat, E. Körblova and R. K. Shoemaker, *Cryst. Growth Des.*, 2005, **5**, 2091; (c) D. Chen, J. E. MacLennan, R. Shao, D. K. Yoon, H. Wang, E. Körblova, D. M. Walba, M. A. Glaser and N. A. Clark, *J. Am. Chem. Soc.*, 2011, **133**, 12656.
- 25 (a) J. Thisayukta, H. Takezoe and J. Watanabe, *Jpn. J. Appl. Phys.*, 2001, **40**, 3277; (b) H. Niwano, M. Nakata, J. Thisayukta, D. R. Link, H. Takezoe and J. Watanabe, *J. Phys. Chem. B*, 2004, **108**, 14889; (c) H. Kurosu, M. Kawasaki, M. Hirose, M. Yamada, S. Kang, J. Thisayukta, M. Sone, H. Takezoe and J. Watanabe, *J. Phys. Chem. A*, 2004, **108**, 4674.
- 26 H. Kresse, J. Saltetnokova, H. Nadasi, W. Weissflog and A. Hauser, *Liq. Cryst.*, 2001, **28**, 1017.
- 27 Biphenyl based dimesogens: (a) E. Bialecka-Florjanczyk, I. Sledzinska, E. Górecka and J. Przedmojski, *Liq. Cryst.*, 2008, **35**, 401; (b) A. Zep, M. Salamonczyk, N. Vaupotić, D. Pocięcha and E. Gorecka, *Chem. Commun.*, 2013, **49**, 3119.
- 28 (a) H. Ocak, B. Bilgin-Eran, M. Prehm and C. Tschierske, *Soft Matter*, 2012, **8**, 7773; (b) H. Ocak, B. Bilgin-Eran, M. Prehm and C. Tschierske, *Soft Matter*, 2013, **9**, 4590.
- 29 E. Tsai, J. M. Richardson, E. Körblova, M. Nakata, D. Chen, Y. Shen, R. Shao, N. A. Clark and D. M. Walba, *Angew. Chem., Int. Ed.*, 2013, **52**, 5254.
- 30 G. B. Deepa, S. Radhika, B. K. Sadashiva and R. Pratibha, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 2013, **87**, 062508.
- 31 M. Alaasar, M. Prehm, M. Nagaraj, J. K. Vij and C. Tschierske, *Adv. Mater.*, 2013, **25**, 2186.
- 32 M. Alaasar, M. Prehm, K. May, A. Eremin and C. Tschierske, *Adv. Funct. Mater.*, 2014, **24**, 1703.
- 33 (a) G. Pelzl, A. Eremin, S. Diele, H. Kresse and W. Weissflog, *J. Mater. Chem.*, 2002, **12**, 2591; (b) T. Niori, J. Yamamoto and H. Yokoyama, *Mol. Cryst. Liq. Cryst.*, 2004, **409**, 475.
- 34 (a) V. Görtz and J. W. Goodby, *Chem. Commun.*, 2005, 3262; (b) V. Görtz, *Liq. Cryst. Today*, 2010, **19**, 37.
- 35 More recently helical superstructures, the so-called twist bent nematic phases, were reported for mesogenic dimers with odd-numbered spacers: (a) J. W. Emsley, M. Lelli, A. Lesage and G. R. Luckhurst, *J. Phys. Chem. B*, 2013, **117**, 6547; (b) V. Borshch, Y.-K. Kim, J. Xiang, M. Gao, A. J. Jakli, V. P. Panov, J. K. Vij, C. T. Imrie, M. G. Tamba, G. H. Mehl and O. D. Lavrentovich, *Nat. Commun.*, 2013, **4**, 2635; (c) D. Chen, J. H. Porada, J. B. Hooper, A. Klittnick, Y. Shen, M. R. Tuchband, E. Körblova, D. Bedrov, D. M. Walba, M. A. Glaser, J. E. MacLennan and N. A. Clark, *Proc. Natl. Acad. Sci. U. S. A.*, 2013, **110**, 15931.
- 36 (a) D. K. Yoon, Y. Yi, Y. Shen, E. D. Körblova, D. M. Walba, I. I. Smalyukh and N. A. Clark, *Adv. Mater.*, 2011, **23**, 1962; (b) D. Chen, C. Zhu, H. Wang, J. E. MacLennan, M. A. Glaser, E. Körblova, D. M. Walba, J. A. Rego, E. A. S. -Bustamante and N. A. Clark, *Soft Matter*, 2013, **9**, 462.
- 37 E. Peeters, J. Lub, J. A. M. Steebakkers and D. J. Broer, *Adv. Mater.*, 2006, **18**, 2412.
- 38 (a) I. C. Pintre, J. L. Serrano, M. B. Ros, J. M. Perdiguerro, I. Alonso, J. Ortega, C. L. Folcia, J. Etxebarria, R. Alicante and B. Villacampa, *J. Mater. Chem.*, 2010, **20**, 2965; (b) C. L. Folcia, I. Alonso, J. Etxebarria, I. Pintre and M. B. Ros, *Chem. Mater.*, 2006, **18**, 4617.
- 39 G. Lee, R. J. Carlton, F. Araoka, N. L. Abbott and H. Takezoe, *Adv. Mater.*, 2013, **25**, 245.
- 40 T. Ueda, S. Masuko, F. Araoka, K. Ishikawa and H. Takezoe, *Angew. Chem.*, 2013, **125**, 7001.
- 41 Dimesogen azobenzene: (a) S.-W. Choi, T. Izumi, Y. Hoshino, Y. Takanishi, K. Ishikawa, J. Watanabe and H. Takezoe, *Angew. Chem., Int. Ed.*, 2006, **45**, 1382; (b) G. Lee, R. J. Carlton, F. Araoka, N. L. Abbott and H. Takezoe, *Adv. Mater.*, 2013, **25**, 245; (c) A. Zep, K. Sitkowska, D. Pocięcha and E. Gorecka, *J. Mater. Chem. C*, 2014, **2**, 2323.
- 42 I. Miyake, Y. Takanishi, N. V. S. Rao, M. K. Paul, K. Ishikawa and H. Takezoe, *J. Mater. Chem.*, 2005, **15**, 4688.
- 43 M. Alaasar, M. Prehm and C. Tschierske, *Chem. Commun.*, 2013, **49**, 11062.
- 44 M. Alaasar, M. Prehm, M. Brautzsch and C. Tschierske, *J. Mater. Chem. C*, 2014, **2**, 5487.
- 45 There are numerous different effects of the F-substituents at the aromatic core, which influence each other and hence



specific effects are difficult to distinguish. Firstly, due to their slightly larger size compared to that of hydrogen, fluorine substituents change the steric intermolecular interactions and also have effects on the conformation of the adjacent alkyl chains. Secondly, fluorine has a strong influence on the intermolecular interactions as a result of its electronegative character, which affects the electron density of the aromatic ring leading to changes in the π - π stacking and π ...H-C electrostatic interactions between the aromatic rings, (M. O. Sinnokrot and C. D. Sherrill, *J. Am. Chem. Soc.*, 2004, **126**, 7690) as well as the polarization, surface charge distribution and polarizability of the whole conjugated π -system.

- 46 R. A. Reddy, V. A. Raghunathan and B. K. Sadashiva, *Chem. Mater.*, 2005, **17**, 274.
- 47 M. Alaasar, M. Prehm and C. Tschierske, *Liq. Cryst.*, 2013, **40**, 656.
- 48 I. Niezgoda, D. Pocięcha and Z. Galewski, *Thermochim. Acta*, 2014, **587**, 59.
- 49 M. Alaasar, M. Prehm and C. Tschierske, *Liq. Cryst.*, 2014, **41**, 126.
- 50 A. Guinier, *X-ray Diffraction*, Freeman, San Francisco, 1963.
- 51 T. Otani, F. Araoka, K. Ishikawa and H. Takezoe, *J. Am. Chem. Soc.*, 2009, **131**, 12368; F. Araoka, G. Sugiyama, K. Ishikawa and H. Takezoe, *Opt. Mater. Express*, 2011, **1**, 27.
- 52 N. G. Nagaveni, A. Roy and V. Prasad, *J. Mater. Chem.*, 2012, **22**, 8948.
- 53 A. Immirzi and B. Perini, *Acta Crystallogr., Sect. A: Cryst. Phys., Diffraction, Theor. Gen. Crystallogr.*, 1977, **33**, 216.
- 54 D. R. Lide, *CRC Handbook of Chemistry and Physics, Internet Version*, 2005, <http://www.hbcpnetbase.com>.
- 55 M. J. Frisch *et al.*, Gaussian 09, Revision A.1, Gaussian Inc., Wallingford CT, 2009.
- 56 (a) I. Wirth, S. Diele, A. Eremin, G. Pelzl, S. Grande, L. Kovalenko, N. Pancenko and W. Weissflog, *J. Mater. Chem.*, 2001, **11**, 1642; (b) L. Kövalenko, M. W. Schröder, R. A. Reddy, S. Diele, G. Pelzl and W. Weissflog, *Liq. Cryst.*, 2005, **32**, 857.
- 57 In contrast to the rigid resorcinol based bent-core mesogens, reported here, the azobenzene based dimesogens form HNF type DC phases,²⁷ presumably because in this case the inherent bent of the odd-numbered alkylene space provides helical conformational chirality, and because this kind of spacer is “softer” these molecules can still adopt helical nanofilaments even in the presence of the rigid azobenzene units.
- 58 Z. Fang and C. Cao, *J. Mol. Struct.*, 2013, **1036**, 447.
- 59 H. G. Heller and J. R. Langan, *J. Chem. Soc., Perkin Trans. 2*, 1981, 341.
- 60 P. Schmitz, H. Gruler and M. Eberhardt, *Mol. Cryst. Liq. Cryst.*, 1995, **262**, 129.

