Review

Liquid Crystal Dimers and Smectic Phases from the Intercalated to the Twist-Bend

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Abstract: In this review we consider the relationships between molecular structure and the tendency of liquid crystal dimers to exhibit smectic phases, and show how our application of these led to the recent discovery of the twist-bend, heliconical smectic phases. Liquid crystal dimers consist of molecules containing two mesogenic groups linked through a flexible spacer, and even- and odd-membered dimers differ in terms of their average molecular shapes. The former tend to be linear whereas the latter are bent, and this difference in shape drives very different smectic behaviour. For symmetric dimers, in which the two mesogenic groups are identical, smectic phase formation may be understood in terms of a microphase separation into distinct sublayers consisting of terminal chains, mesogenic units and spacers, and monolayer smectic phases are observed. By contrast, intercalated smectic phases were discovered for nonsymmetric dimers in which the two mesogenic units differ. In these phases, the ratio of the layer spacing to the molecular length is typically around 0.5 indicating that unlike segments of the molecules overlap. The formation of intercalated phases is driven by a favourable interaction between the different liquid crystal groups. If an odd-membered dimer possesses sufficient molecular curvature, then the twist-bend nematic phase may be seen in which spontaneous chirality is observed for a system consisting of achiral molecules. Combining the empirical relationships developed for smectogenic dimers, and more recently for twist-bend nematogenic dimers, we show how dimers were designed to show the new twist-bend, heliconical smectic phases. These have been designated SmC_{TW} phases in which the director is tilted with respect to the layer plane, and the tilt direction describes a helix on passing between layers. We describe three variants of the SmC_{TB} phase, and in each the origin of the symmetry breaking is attributed to the anomalously low-bend elastic constant arising from the bent molecular structures.

Keywords: liquid crystal dimers; intercalated; interdigitated; twist-bend nematic; twist-bend smectic; chirality; resonant soft X-ray scattering

1. Overview

Over the last decade arguably the hottest topic in liquid crystals science has been the twist-bend nematic, N_{TB}, phase following its discovery in 2011 [1], some ten years after its prediction by Dozov [2]. We will return to the N_{TB} phase later, but widely overlooked in Dozov’s seminal work was the prediction of twist-bend smectic phases, and in this review, we trace the discovery of these phases for liquid crystal dimers [3]. Although the aim of this Special Issue is to provide an overview of the state-of-the-art of current UK liquid crystals research, the work we describe would not have been possible without a close collaboration between the Universities of Aberdeen and Warsaw. Indeed, the very essence of liquid crystals research is the need for a multidisciplinary approach, and science should know no borders. In keeping with this Special Issue’s aim, however, we have attempted to focus primarily on the contribution to the overall work made in Aberdeen, but note that this story is fundamentally one of collaboration.
At the root of this work are liquid crystal dimers, and we begin in Section 2 with a very brief description of the characteristic behaviour of this fascinating class of low molar mass liquid crystals. In Section 3, we consider the smectic behaviour of symmetric dimers, and follow this in Section 4 by describing the discovery of the intercalated smectic phases for nonsymmetric dimers. We then return to describe the \( N_{TB} \) phase in Section 5 and briefly consider the types of liquid crystal dimer known to exhibit the phase. Pulling together the themes developed in Sections 3–5, we describe the discovery of the newest class of smectic phases in liquid crystal dimers, the twist-bend smectic phases in Section 6. We finish this review with an outlook of what may be achieved in this area in the future.

2. Liquid Crystal Dimers

Liquid crystal dimers consist of molecules containing two semi-rigid mesogenic moieties linked though a flexible spacer normally, but not always, an alkyl chain, and can be divided broadly into two classes. In symmetric dimers the two mesogenic units are the same, whereas in nonsymmetric dimers they differ. Detailed reviews of structure-property relationships in liquid crystal dimers can be found elsewhere [4–6], whereas for the purpose of this overview we need to consider only what may be referred to as their archetypal behaviour. Figure 1a shows how the melting points and nematic-isotropic transition temperatures, \( T_{NI} \), vary as the length of the spacer is increased for a series of symmetric liquid crystal dimers, the \( \alpha, \omega \)-bis(4-nitroazobenzene-4′-oxy)alkanes, [7]

![Image of molecular structure](image)

and these are referred to using the acronym BNABO\( \text{H}_n \) in which \( n \) refers to the number of methylene units in the flexible spacer. It can be seen in Figure 1a that \( T_{NI} \) initially exhibits a strong alternation as the parity of the spacer is varied, but that this attenuates on increasing spacer length. In this alternation it is the even members of the series that show the higher values of \( T_{NI} \). It is interesting to note that the melting points for this particular series also alternate on increasing the spacer length, and again the even members show the higher values, but this is somewhat less regular behaviour than that seen for \( T_{NI} \), and is not observed for all dimer series. The dependence of \( T_{NI} \) on spacer length and parity in dimers is most often attributed to molecular shape when considering the spacer in its all-\( \text{trans} \) conformation. For an even-membered spacer, the two mesogenic units are more or less parallel, whereas for an odd-membered spacer they are inclined to each other and the molecule is bent (Figure 1b). The linear shape of an even-membered dimer is more compatible with the molecular organisation found in the nematic phase than the bent-shape of an odd-membered dimer, and this accounts for the higher values of \( T_{NI} \) seen for the former. Although intuitively pleasing, this interpretation does not account for the pronounced alternation also observed in the nematic-isotropic entropy change on increasing the spacer length, and to do so the inherent flexibility of the spacer must be accounted for by considering a wide range of conformations and not solely the all-\( \text{trans} \) form [5]. For our purposes, however, it is sufficient to remember that, on average, an even-membered dimer is essentially linear whereas an odd-membered dimer is bent, and that this difference in average shape decreases as the spacer length increases given the increasing number of conformations available to the spacer.
The interest in liquid crystal dimers can be traced back to the early 1980s and the suggestion by Griffin and Britt that they may be used as model compounds for the technologically important semi-flexible main chain liquid crystal polymers [8] whereas their discovery, many decades earlier, by Vöhringer had been largely overlooked [9]. The dimeric molecular architecture represented a marked deviation in the design of a low mass liquid crystal. The overwhelming majority of low molar mass liquid crystals up to that point consisted of molecules containing a single semi-rigid core attached to which were one or two flexible alkyl chains. In essence, the interactions between the cores accounted for the liquid crystal behaviour and the terminal chains used to reduce the melting point and drive smectic phases. In a dimer, this structure is inverted and now the core of the molecule is flexible, and we have seen already the importance of this in controlling their shape and hence, transitional behaviour.

The majority of dimers reported in the 1980s showed only nematic behaviour and Griffin and Britt attributed this to their inherent molecular flexibility suppressing smectic phase formation [9]. With hindsight, the absence of smectic behaviour was surprising. It was well-known that molecular inhomogeneity, such as the chemically distinct regions found in a dimer, drives the formation of smectic phases, and there appeared to be no fundamental reason for dimers not to exhibit smectic behaviour. It was almost a decade later, however, before the first family of liquid crystal dimers were reported that showed rich smectic polymorphism, the α,ω-bis(4-n-alkylanilinebenzylidene-4'-oxy)alkanes [10]:

\[
\text{H}_{2m+1}C_{m}\underbrace{\text{N}}_{\text{m}}\overbrace{\text{O(\text{CH}_2)_n\text{O}}}^{\text{m}}\overbrace{\text{N}}_{\text{m}}\underbrace{\text{C}_{n}\text{H}_{2m+1}}_{\text{m}}
\]

and these are referred to using the acronym \(m\text{OnO.m}\) in which \(n\) and \(m\) refer to the number of carbon atoms in the spacer and terminal chains, respectively. The strategy underpinning the design of this family of dimers was straightforward and centred upon the need to be able to readily vary the lengths of both the spacer and the terminal alkyl chains, a condition met by the \(m\text{OnO.m}\) molecular architecture. In addition, they may be considered to be the dimeric analogues of the \(N\)-(4-n-alkoxybenzylidene)-4'-n-alkylanilines known to be a rich source of smectic phases [11].

The versatility in the synthetic approach used to prepare the \(m\text{OnO.m}\) series allowed for the transitional behaviour of 132 members of the family to be reported [10]. These included varying the spacer length, \(n = 1\text{--}12\), and the terminal alkyl chain lengths, \(m = 0\text{--}10\). These dimers did indeed exhibit rich smectic polymorphism including smectic A and C.
These observations are now established as being rather general for symmetric dimers with parity of the spacer was varied (Figure 2) as seen for T₅ polymer also promoted smectic behaviour. It is also interesting to note that the SmA-I transition temperatures also exhibited a pronounced odd-even effect as the length and parity of the spacer was varied (Figure 2) as seen for T₅NI (Figure 1a). This indicates that the bent odd-membered dimers experience greater difficulty in packing into smectic structures than their linear, even-membered counterparts. Increasing the terminal chain length for a given spacer promotes nematic behaviour and this is shown for the m.O₅O₅ series in Figure 3, and as would be expected.

The study of the m.OₙOₕₕ series revealed that increasing the spacer length for a given terminal chain length promotes nematic behaviour and this is shown for the 5.OₙO₅ series in Figure 2. This was surprising and contravened the very general observations at the time that increasing the length of an alkyl chain promoted smectic behaviour in low molar mass mesogens, and that increasing the length of the spacer in a semi-flexible main chain polymer also promoted smectic behaviour. It is also interesting to note that the SmA-I phase, the smectic layers have a periodic modulation analogous to that found in the Sm C° ribbon phase. This modulated hexatic phase is only observed for dimers with an odd parity spacer, suggesting that the average bent molecular shape drives its formation. This is a theme that runs through this overview.

The dependence of the transition temperatures on the number of methylene groups, n, in the flexible spacer for the 5.OₙO₅ series [10]. The broken line joins the melting points. Filled squares denote T_{SmAI}; unfilled squares T_{SmAN}; circles T_{NI}; triangles T_{SmASmB}.

A simple empirical relationship emerged from the study of the m.OₙOₕₕ family of compounds relating the observation of smectic behaviour to the relative lengths of the terminal chains, m, to that of the spacers, n [10]. Thus, for smectic behaviour to be observed m > 0.5 n. In addition, all the smectic phases observed for these dimers possessed a monolayer structure, i.e., the layer spacing, d, corresponded to the full molecular length, l. These observations are now established as being rather general for symmetric dimers with only a small number of exceptions known [5], and imply that in the smectic layer the mesogenic units, spacers and terminal chains may be considered to microphase separate into distinct sublayers as sketched in Figure 4a. An alternative packing arrangement of the dimers in which the terminal chains and spacers are randomly mixed to give an
intercalated structure (Figure 4b), although entropically favoured, was not observed, and this was attributed to an unfavourable interaction between the terminal chains and spacers offsetting the favourable entropic term [13].

![Graph](image)

**Figure 3.** The dependence of the transition temperatures on the number of carbon atoms, \( m \), in the terminal chains for the \( m\text{OSO}_m \) series [10]. The broken line joins the melting points. Filled squares denote \( T_{\text{SmA}} \); unfilled squares \( T_{\text{SmAN}} \); circles \( T_{\text{NI}} \); unfilled diamonds \( T_{\text{SmASmC}} \); filled diamonds \( T_{\text{SmAG}} \); filled triangles \( T_{\text{SmASmF}} \); unfilled triangles \( T_{\text{SmCSmF}} \).

![Sketch](image)

**Figure 4.** A sketch of (a) the monolayer smectic A phase \((d \sim l)\) exhibited by symmetric liquid crystal dimers, and (b) the intercalated smectic A phase \((d \sim l/2)\) observed only rarely for symmetric liquid crystal dimers.

It is noteworthy that replacing the ether links by methylene links between the spacer and mesogenic units in a dimer has a significant effect on its transitional properties. This is
highlighted by a comparison of the behaviour of the \( \alpha,\omega \)-bis(4-\( n \)-alkyloxyanilinebenzylidene-4′-yl)alkanes [14,15],

![Chemical structure](image)

referred by the acronym \( mO-n-Om \) in which the hyphen is used to reflect the reversal of the Schiff’s base link compared to the \( mO-OnO.m \) series, with that of the corresponding ether-linked materials, the \( mO-OnO-Om \) series. Surprisingly, for short chain lengths \((m = 1, 2)\), the \( mO-5-Om \) series exhibited a nematic phase and at lower temperatures, a second mesophase that exhibited a fan-like optical texture in coexistence with regions of schlieren texture and this was assigned as an anticlinic, intercalated smectic C phase although its monotropic nature precluded an unambiguous identification using X-ray diffraction (XRD). This behaviour was noted to be in contrast to that seen for the corresponding \( mO-OnO-Om \) series for which smectic behaviour was observed only if the length of the terminal chains was greater than half that of the spacer, as described earlier for the \( mO-OnO.m \) series. By comparison, the even-membered \( mO-n-Om \) \((n = 4, 6)\) series with short terminal chains showed solely nematic behaviour as expected. For long terminal chain lengths \((n = 9, 10)\), the \( mO-5-Om \) series exhibited a G/J soft crystal phase with a modulated layer structure, whereas the corresponding members of the \( mO-6-Om \) series a G/J soft crystal phase was observed but with a simple layer structure. This difference in behaviour is similar to that seen for the corresponding members of the \( mO-OnO-Om \) series. It is important to note that the switch from an ether- to a methylene-linked spacer accentuates the bent shape of an odd-membered dimer as we will see later. The behaviour of the \( mO-OnO-Om \) series reinforced the view that the difference in shape between odd and even-membered dimers accounts, at least in part, for the differing smectic behaviour observed, with the bent odd-members having a stronger tendency to pack into tilted, alternating lamellar phases. The particularly surprising behaviour seen for \( mO-5-Om \) with \( m = 1 \) and 2, was later shown in fact to be a twist-bend nematic phase and we return to this in Section 5 [16].

4. Nonsymmetric Dimers and Intercalated Smectic Phases

We have seen that symmetric dimers have a strong tendency to form smectic phases having monolayer structures, and that this may reflect an unfavourable interaction between the spacers and terminal chains inhibiting intercalated arrangements. The question now arose: what if we overcome this unfavourable interaction by designing nonsymmetric dimers that exhibited a favourable specific interaction between the unlike mesogenic units, and could this drive the formation of intercalated smectic phases? To investigate this intriguing possibility, the \( \alpha \)-(4-cyanobiphenyl-4′-yloxy)-\( \omega \)-(4-\( n \)-alkylanilinebenzylidene-4′-oxy)alkanes were studied [17,18]:

![Chemical structure](image)

The acronym used to describe these dimers is \( \text{CBO}nO.m \), in which \( n \) and \( m \) refer to the number of carbon atoms in the spacer and terminal chains, respectively. This general structure was chosen because it was known that mixtures of nematogenic 4-\( n \)-alkoxy cyanobiphenyls and \( N \)-(4-\( n \)-alkyloxybenzylidene)-4′-\( n \)-alkylanilines showed induced smectic behaviour implying a specific favourable interaction between the unlike mesogenic units [19,20]. The \( \text{CBO}nO.m \) family of dimers showed new patterns of liquid crystal behaviour: for example, Figure 5 shows the dependence of the transition temperatures on the length of the terminal chain for the \( \text{CBO}4O.m \) series and it is striking that smectic
behaviour is observed for short and long terminal chains, but for intermediate chain lengths only a nematic phase is seen. This behaviour contravened a very general observation for conventional low molar mass mesogens that, for a given series, the smectic A-nematic transition temperature simply increases on increasing the length of a terminal chain and at some point, nematic behaviour is extinguished. To understand the behaviour seen in Figure 5, we must consider how the structure of the smectic phase changes as we increase $m$. For short chain lengths, the ratio of the smectic layer spacing to molecular length, $d/l$, is about 0.5 indicating an intercalated arrangement of the dimers in which differing fragments of the molecules overlap (Figure 6a), the SmA$_c$ phase. The driving force for this arrangement was attributed to a specific interaction between the unlike mesogenic units suggested to be an electrostatic quadrupolar interaction between groups having quadrupole moments of opposite signs [21]. We should note that as sketched in Figure 6a the intercalated smectic A phase would appear to be polar, but we assumed that such molecular groupings would be randomly arranged at the macroscopic level such that ferroelectric properties would not be exhibited. This view was later supported by the failure to detect macroscopic polarisation in these phases. Returning to Figure 5, $d/l$ for the smectic phase shown by the dimers with long terminal chains is around 1.8, implying an interdigitated arrangement of the dimers (Figure 6b), the SmA$_d$ phase, and such an arrangement is stabilised by the electrostatic interaction between the polar and polarizable cyanobiphenyl groups while the smectic phase results from the molecular inhomogeneity arising from the long terminal chains. The cross-over from an intercalated to an interdigitated structure arises from insufficient space between the layers of mesogenic units, governed by the length of the spacer, to accommodate the terminal chains and so the layers are pushed apart reducing the specific interaction between the unlike groups. The disappearance of smectic behaviour for intermediate chain lengths reflects a competition between these two incompatible arrangements, neither of which wins and hence, only a nematic phase is observed.

![Figure 5](image_url)

**Figure 5.** The dependence of the transition temperatures on the number of carbon atoms, $m$, in the terminal chain for the CBO$_4$O$_m$ series [18]. The broken line joins the melting points. Filled squares denote $T_{\text{SmA}_c}$; unfilled squares $T_{\text{SmA}_d}$; circles $T_{\text{NI}}$; unfilled diamonds $T_{\text{SmC}_m}$.
Not only was the intercalated smectic A phase shown by the CBO4O.10 series [18]. Qualitatively similar behaviour to the CBO4O.10 series is seen such that smectic phases are observed for short and long spacers, and exclusively nematic behaviour for intermediate spacer lengths. In contrast to the CBO4O.10 series, however, intercalated smectic phases are observed for long spacer lengths and interdigitated phases for short spacers. This is wholly consistent with the view that the terminal chain must be accommodated within the space between the layers that is governed by the length of the spacer.

If we now consider how the phase behaviour changes on varying the length of the spacer while holding the terminal chain constant, Figure 7 shows the behaviour of the CBO4O.10 series [18]. The broken line joins the melting points. Filled squares denote T_{SmAI}; unfilled squares T_{SmAN}; circles T_{NI}; unfilled diamonds T_{SmCSmA}.

![Figure 5](image1.png)

Figure 5. The dependence of the transition temperatures on the number of carbon atoms, n, in the flexible spacer for the CBO4O.10 series [18]. The broken line joins the melting points. Filled squares denote T_{SmAI}; unfilled squares T_{SmAN}; circles T_{NI}; unfilled diamonds T_{SmCSmA}.

![Figure 6](image2.png)

Figure 6. Sketches of (a) the intercalated and (b) the interdigitated smectic A phases shown by nonsymmetric liquid crystal dimers.

![Figure 7](image3.png)

Figure 7. The dependence of the transition temperatures on the number of methylene groups, n, in the flexible spacer for the CBO4O.10 series [18]. The broken line joins the melting points. Filled squares denote T_{SmAI}; unfilled squares T_{SmAN}; circles T_{NI}; unfilled diamonds T_{SmCSmA}.
Not only was the intercalated smectic A phase shown by the CBO\textit{m}O\textit{m} series but other intercalated variants including the smectic C (Sm\textit{C}_\textit{c}) and I phases, as well as intercalated soft crystal phases were observed. It is important to note that tilted intercalated smectic phases were observed only for dimers containing long odd-membered spacers, and it was suggested that this reflects the difficulty that these bent dimers experience packing into an intercalated arrangement. Of particular interest was the intercalated smectic C phase shown, for example, by CBO9O.6. The anticlinic structure of the phase was identified on the basis of the observation of a schlieren optical texture containing both \( s = \pm 1/2 \) and \( s = \pm 1 \) defects [22], and the value of \( d/l \) was established using X-ray diffraction to be about 0.5 indicating an intercalated arrangement of the molecules. A sketch of the molecular arrangement proposed for the intercalated smectic C phase is shown in Figure 8 in which the tilt direction alternates between the layers such that the global tilt angle is zero but locally, within a layer, is non-zero. This structure was later confirmed using electron spin resonance spectroscopy that revealed within the intercalated smectic C phase two distinct directors with azimuthal tilt directions differing by 180° [23]. Within a layer the tilt angle was estimated to be about 18°, fully consistent with the bent geometry of these odd-membered dimers in which the mesogenic units make an angle of around 15° with the spacer. The intercalation of the dimers has been widely accepted to account for the measured values of \( d/l \) of around 0.5 although a number of issues remain to be fully resolved as recently discussed elsewhere [24].

![Figure 8](image)

**Figure 8.** Sketch of the molecular arrangement in the intercalated smectic C phase. The arrow indicates the local tilt direction.

In Figure 5, we saw how the transition temperatures varied on increasing the terminal chain length for the CBO4O\textit{m} series and this surprising pattern of behaviour was understood in terms of the formation of intercalated smectic phases for short terminal lengths and interdigitated smectic phases for long terminal chains. The behaviour of the CBO5O\textit{m} series was very different; see Figure 9. On cooling the nematic phase shown by the members with \( m = 1-9 \), what appeared to be a smectic phase formed, and the values of the transition temperature between this phase and the nematic phase decreased essentially linearly on increasing \( m \). The structure of the lower temperature phase could not be established using X-ray diffraction due to its monotropic nature. The behaviour seen in Figure 9 was noted as being unusual and it was also noted that the CBO3O\textit{m} series showed broadly similar behaviour. These observations were thought to reflect the bent shape of these dimers, but its physical significance was not apparent. As we will see in the next section, however, the behaviour may now be accounted for in terms of the twist-bend nematic phase. Smectic behaviour was observed for long homologues in both these odd-membered series and we will return to this later.
This series showed a tilted soft crystal phase, possibly a G/J phase composed of finite thickness which pack to give macroscopic left-handed domains [28]. The formation of helical nanofilaments in these structures is understood in terms of the formation of intercalated smectic phases for short terminal spacers. This supported the suggestion that molecular shape was an important factor in smectic phase formation. This tendency for bent odd-membered dimers to pack into tilted, intercalated smectic phases to alleviate the difficulties in packing into orthogonal arrangements has similarities to the behaviour of bent-core mesogens [26]. This view was further reinforced by the behaviour of a series of 4-decyloxy-4′-hydroxybiphenyl esters of α,ω-alkanedicarboxylic acids reported by Białecka-Florjańczyk et al. [27]:

\[
\begin{align*}
\text{C}_{10}\text{H}_{21}\text{O} & \text{-} \text{O} \cdot \text{(CH}_2)_n \text{O} \cdot \text{O} \cdot \text{(CH}_2)_n \text{OC}_{10}\text{H}_{21}
\end{align*}
\]

The even members of this series showed a tilted soft crystal phase, possibly a G/J phase, whereas the odd members exhibited the B4 phase previously only observed for bent-core mesogenic materials. The B4 phase is thought to consist of bundles of twisted, rope-like smectic ribbons of finite thickness which pack to give macroscopic left- and right-handed domains [28]. The formation of helical nanofilaments in these structures is thought to be driven by the instability of the flat layers arising from the mismatch in the projections of the two crystal lattices associated with each arm of the bent core [29]. The elastic strain required to connect the two lattices may be relieved, at least in part, by bending the layers with saddle-splay curvatures. A similar model is clearly applicable to bent odd-membered dimers although would necessarily have to also include the inherent flexibility associated with the spacer. The observation of such behaviour for this particular series may be attributed to the strong lateral interactions known to exist between biphenyl fragments, a view supported by the observation of the soft crystal phase for the even members. Again, we see that molecular shape is a key factor in the pronounced differences in behaviour between odd and even-membered dimers, and the similarities between bent odd-membered dimers and the semi-rigid bent-core mesogens will be further discussed later.
5. The Twist Bend Nematic Phase

We now return to the prediction of the twist-bend nematic, N\textsubscript{TB}, phase by Dozov [2] and its subsequent discovery by Cestari et al. [1]. At the root of Dozov’s prediction was the assertion that bent molecules have a natural disposition to pack into bent structures, but that pure bend cannot fill space and so is forbidden. Instead, bend must be accompanied by other deformations of the director; either twist or splay. In the case of twist, this gives rise to the twist-bend nematic phase in which the director adopts a helical structure in which it is tilted with respect to the helical axis (Figure 10). Perhaps the most intriguing aspect of Dozov’s work was the prediction of spontaneous chirality in a fluid system composed of achiral molecules having no positional order. The helices formed may be either left- or right-handed and equal amounts of both are expected.

![Figure 10](image)

**Figure 10.** A schematic representation of the twist-bend nematic, N\textsubscript{TB}, phase.

Odd-membered liquid crystal dimers with their bent molecular structures (Figure 1b) provided the ideal testbed for Dozov’s prediction, and some ten years later, Cestari et al. reported the discovery of the N\textsubscript{TB} phase in 1,7-bis(4-cyanobiphenyl-4’-yl)heptane (CB7CB) [1]:

\[
\text{NC} - \text{C}_n - \text{O} - (\text{CH}_2)_p - \text{C}_n - \text{CN}
\]

The assignment of the N\textsubscript{TB} phase was confirmed using techniques including freeze fracture transmission electron microscopy [30] and resonant soft X-ray scattering (RSoXS) [31]. The events surrounding this discovery are reviewed by Dunmur in another contribution to this collection of papers [32]. A striking feature of the N\textsubscript{TB} phase is that the helical pitch is very short, typically around 10 nm, corresponding to around 3–4 molecular lengths. Nematic-nematic transitions had been reported previously for other odd-membered dimers that, with the benefit of hindsight, are examples of N\textsubscript{N} transitions [33,34]. A common feature of these dimers is the use of methylene links to connect the spacer to the semi-rigid mesogenic units. By contrast, the majority of dimers reported prior to the discovery of the N\textsubscript{TB} phase contained ether-linked spacers, i.e., O(CH\textsubscript{2})\textsubscript{n}O, although interest in methylene-linked dimers had been triggered earlier by a prediction made by Ferrarini et al. [35,36] that systems containing a high concentration of bent conformers in the isotropic phase should exhibit a nematic-nematic transition. Again, short odd-membered methylene-linked dimers provided a testbed for these predictions. It was found that switching O(CH\textsubscript{2})\textsubscript{n}O for (CH\textsubscript{2})\textsubscript{n+2} saw the values of T\textsubscript{NI} fall for both odd- and even-membered spacers but the reduction was greater for odd-membered spacers. This resulted in a more pronounced alternation in T\textsubscript{NI} for the methylene-linked dimers on varying the spacer length and parity compared...
to that seen for their ether-linked counterparts [14,15]. By contrast, the entropy change associated with the nematic-isotropic transition increased for even-membered spacers but decreased for odd-membered spacers on replacing O(CH$_2$)$_n$O with (CH$_2$)$_{n+2}$. These experimental observations were in complete agreement with the predictions of a model described by Luckhurst and co-workers in which the only difference between the dimers is the bond angle between the para-axis of the mesogenic unit and the first bond in the spacer [37,38]. For an ether-linked dimer this angle is about 126.4° whereas for a methylene-linked dimer it is around 113.5°. The change in this angle means that an odd-membered methylene-linked dimer is more bent than the corresponding ether-linked dimer (Figure 11). The discovery of the N$_{TB}$ phase reignited interest in the methylene-linked dimers and previously reported dimers were shown to exhibit the new phase [16]. In addition, odd-membered ether-linked dimers were also shown to exhibit the N$_{TB}$ phase [39] and led to the reassignment of smectic phases as N$_{TB}$ phases as shown, for example, in Figure 9 for the CB6O.m series [18,40].

![Figure 11. A comparison of the shapes of odd-membered methylene- and ether-linked dimers: (a) CB7CB and (b) CBO5OCB.](image)

There is now a large collection of odd-membered dimers known to exhibit the N$_{TB}$ phase, and structure-property studies have focussed on the nature of the link between the spacer and mesogenic units [41–49], the length and parity of the spacer [39,50–53], the structure of the mesogenic units [54–64], and the chemical nature of the terminal groups [65–68]. Although the majority of twist-bend nematogens are odd-membered dimers, other examples include trimers and higher oligomers [69–74], hydrogen-bonded supramolecular systems [75–79], rigid-bent core systems [80,81] and polymers [82]. A recent overview of structure-property relationships in twist-bend nematogens may be found elsewhere [83].

The overarching structural requirement for the observation of the twist-bend nematic phase is molecular curvature. This is in complete agreement not only with Dozov’s model [2], but also with Maier-Saupe theory for V-shaped molecules that predicts the N$_{TB}$ phase will be formed by just a narrow range of molecular curvatures with the N-N$_{TB}$ transition temperature being particularly sensitive to the molecular bend angle [84]. The uniformity of molecular curvature is also critical in driving the formation of the N$_{TB}$ phase as revealed by the behaviour of the azobenzene-based dimer, CB6OABOBu [85]:

\[
\text{NC} \quad \begin{array}{c}
\includegraphics[width=2cm]{azobenzene_structure}
\end{array} \quad \text{N} \quad \begin{array}{c}
\text{OC}_2\text{H}_9
\end{array}
\]

This dimer shows an isothermal N$_{TB}$-N transition when illuminated using UV-light and this may be attributed to the trans-cis photoisomerization of the azo-linkage. This transition is reversible on removing the light source, driven by the thermal cis—trans relaxation. The cis isomer is more strongly bent than the trans isomer (Figure 12) and a priori it may have been expected that increasing the concentration of the more bent isomer would enhance the stability of the N$_{TB}$ phase. It is clear from this and other studies [86,87] that quite the opposite is true. This counter-intuitive observation was accounted for in terms of the shapes of the two isomers. The molecular curvature of the trans isomer is governed by the geometry of the spacer and hence, is spatially uniform. In contrast, the cis isomer contains two centres of bend, namely the spacer and the azobenzene fragment, which results in a change in bend polarity along the molecule. This spatially varying bend is not compatible with the local structure of the N$_{TB}$ phase.
In contrast, the TB-12-x members of the CB6O.m series, in which the hexyloxy spacer ensured the necessary molecular curvature for the NTB phase, resulted in a change in bend polarity along the molecule. This spatially varying bend is not compatible with the local structure of the NTB phase. Ironically, Dozov highlighted the challenge in obtaining the NTB phase for bent mesogens, but also noted that the same arguments could give rise to a similar symmetry breaking even in apolar banana smectics [2]. This aspect of the study was largely overlooked, and the search for these twist-bend smectic phases became a focus of our work. At the outset of our studies, the great majority of the NTB phases reported either vitrified or crystallised on cooling, and only rarely had smectic-NTB or B-NTB transitions been reported [58,88–90]. In designing materials to potentially exhibit twist-bend smectic phases we identified three key structural criteria: (i) a uniform molecular curvature compatible with the NTB phase, and, based on our previous studies of dimeric smectogens, (ii) a specific favourable interaction between the mesogenic units to promote smectic behaviour, and (iii) a terminal chain that could be readily varied in length to control the packing of the molecules and to promote molecular inhomogeneity.

The first question to address is how does the more pronounced molecular curvature, required to promote the formation of twist-bend phases, affect the tendency of dimers to exhibit smectic phases? Figure 13 shows the dependence of the transition temperatures on the length of the terminal chain for the CB6O.m series [40,91]:

\[
\text{NC-} \begin{array}{c}
\text{ether-bridged diester-} \\
\text{N}
\end{array} \begin{array}{c}
\text{in which the hexyloxy spacer ensured the necessary molecular curvature for the NTB} \\
\text{phase to be observed [42]. Our initial study of this series included varying the terminal} \\
\text{chain length, } m, \text{ from } m = 1–10, [40] \text{ and in a subsequent study we reported the} \\
\text{transitional behaviour of the longer members with } m = 11–18 [91]. The phase behaviour of the} \\
\text{CB6O.m series with } m = 1–10 [40] \text{ may be compared to that of the CBO5O.m series shown in} \\
\text{Figure 9 [18]. The bent nature of both series reduces their smectic tendencies compared to, for} \\
\text{example, the linear CBO4O.m series shown in Figure 5 [18]. For the CBO5O.m series, NTB} \\
\text{phases are observed for } m = 1–7. NTB behaviour is extinguished at } m = 8 \text{ and} \\
\text{smectic-nematic transitions are observed for } m = 8–10. \text{ For the more bent CB6O.m series,} \\
\text{NTB phases are observed for } m = 1–10, \text{ and smectic behaviour emerges at } m = 10 \text{ which} \\
\text{exhibits a Sm-NTB transition to be discussed later [40]. This reduction in the smectic} \\
\text{tendencies on increasing molecular bend may be quantified by comparing the scaled transition} \\
\text{temperature, } T_{\text{smN}}/T_{\text{NI}}, \text{ for CBO5O.10 of 0.924, with } T_{\text{smNTB}}/T_{\text{NI}} \text{ for CB6O.10 of 0.909.} \\
\text{It is interesting to note that the value of } T_{\text{NTBN}}/T_{\text{NI}} \text{ for CB6O.10 is 0.925, i.e., essentially} \\
\text{the same as } T_{\text{smN}}/T_{\text{NI}} \text{ for CBO5O.10. It appears, therefore, that increasing molecular} \\
\text{curvature increases the tendency to exhibit the NTB phase at the expense of smectic behaviour.} \\
\text{Ironically, Dozov highlighted the challenge in obtaining the NTB phase would be to suppress} \\
\text{the formation of smectic phases in bent-core systems [2] in which symmetry breaking had} \\
\text{been attributed to specific polar interactions [92]. For odd-membered dimers the origin of} 
\text{the symmetry breaking is quite different and may be attributed to anomalously low values of} \\
\text{the bend elastic constant arising from the bent molecular geometry. Thus, the inherent} 
\text{flexibility and bent shape of these odd-membered dimers suppresses smectic behaviour,}
\]

![Figure 12. The shapes of the (a) trans and (b) cis isomers of CB6OABOBu.](image)

6. The Twist-Bend Smectic Phases

Dozov, in his seminal work, not only predicted the existence of the NTB phase for bent mesogens, but also noted that the same arguments could give rise to a similar symmetry breaking even in apolar banana smectics [2]. This aspect of the study was largely overlooked, and the search for these twist-bend smectic phases became a focus of our work. At the outset of our studies, the great majority of the NTB phases reported either vitrified or crystallised on cooling, and only rarely had smectic-NTB or B-NTB transitions been reported [58,88–90]. In designing materials to potentially exhibit twist-bend smectic phases we identified three key structural criteria: (i) a uniform molecular curvature compatible with the NTB phase, and, based on our previous studies of dimeric smectogens, (ii) a specific favourable interaction between the mesogenic units to promote smectic behaviour, and (iii) a terminal chain that could be readily varied in length to control the packing of the molecules and to promote molecular inhomogeneity.

The first question to address is how does the more pronounced molecular curvature, required to promote the formation of twist-bend phases, affect the tendency of dimers to exhibit smectic phases? Figure 13 shows the dependence of the transition temperatures on the length of the terminal chain for the CB6O.m series [40,91]:

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\text{NTB phases are observed for } m = 1–10, \text{ and smectic behaviour emerges at } m = 10 \text{ which} \\
\text{exhibits a Sm-NTB transition to be discussed later [40]. This reduction in the smectic} \\
\text{tendencies on increasing molecular bend may be quantified by comparing the scaled transition} \\
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\text{It is interesting to note that the value of } T_{\text{NTBN}}/T_{\text{NI}} \text{ for CB6O.10 is 0.925, i.e., essentially} \\
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\text{the formation of smectic phases in bent-core systems [2] in which symmetry breaking had} \\
\text{been attributed to specific polar interactions [92]. For odd-membered dimers the origin of} 
\text{the symmetry breaking is quite different and may be attributed to anomalously low values of} 
\text{the bend elastic constant arising from the bent molecular geometry. Thus, the inherent} 
\text{flexibility and bent shape of these odd-membered dimers suppresses smectic behaviour,}
\]
and the challenge was to design odd-membered dimers in which the tendency to form smectic phases, rather than the N\textsubscript{TB} phase, was enhanced.

![Graph showing the dependence of the transition temperatures on the number of carbon atoms, m, in the terminal chain for the CB6O\textit{m} series [40,91]. The melting points are connected by the broken lines. Unfilled circles denote T\textsubscript{NI}; filled circles T\textsubscript{N}\text{TB}; filled squares T\textsubscript{SmA}; unfilled squares T\textsubscript{SmAN}; unfilled diamonds T\textsubscript{SmCTB-SH}SmA; filled diamonds T\textsubscript{SmCTB-SH}N; unfilled triangles T\textsubscript{SmCTB-DH}SmCTB-SH; filled triangles T\textsubscript{SmCTB-DH}N; crosses T\textsubscript{SmX}SmCTB-DH.](image)

**Figure 13.** The dependence of the transition temperatures on the number of carbon atoms, \(m\), in the terminal chain for the CB6O\textit{m} series [40,91]. The melting points are connected by the broken lines. Unfilled circles denote \(T_{NI}\); filled circles \(T_{N}\text{TB}\); filled squares \(T_{SmA}\); unfilled squares \(T_{SmAN}\); unfilled diamonds \(T_{SmCTB-SH}\text{SmA}\); filled diamonds \(T_{SmCTB-SH}N\); unfilled triangles \(T_{SmCTB-DH}\text{SmCTB-SH}\); filled triangles \(T_{SmCTB-DH}N\); crosses \(T_{SmX}\text{SmCTB-DH}\).

In order to achieve this goal, we increased the interactions between the mesogenic moieties by increasing the structural anisotropy of the benzylideneaniline moiety in the CB6O\textit{m} series to obtain the CB6O\textit{IBeOm} series [3]:

![Molecular structure of CB6OIBeOm](image)

Again, we chose the hexyloxy spacer to impart the required molecular curvature and varied the terminal chain length, \(m = 1–10\), to increase molecular inhomogeneity and further promote smectic behaviour. For short terminal chain lengths, \(m = 1–6\), a transition between the conventional and twist-bend nematic phases was observed, whereas for \(m \geq 7\), the \(N_{TB}\) phase was extinguished and up to four lamellar phases were found below the \(N\) phase (Figure 14). The layer spacings in all the smectic phases corresponded to approximately the molecular length. The lowest temperature lamellar phase, seen for most of the homologues, \(m = 2, 4–10\), is a hexatic-type smectic phase, designated HexI, in which the molecules tilt towards the apex of the local in-plane hexagons. The three higher temperature smectic phases showed liquid-like ordering of the molecules within the layers, and the changes in optical texture seen on cooling from the nematic phase are shown in Figure 15a. The highest temperature smectic phase was optically uniaxial and assigned as a \(SmA\) phase. On cooling, a schlieren texture developed, and at the transition there was no change in layer spacing, only the layer thermal expansion coefficient differed between the phases, both negative in sign, suggesting both phases to be orthogonal. The observed schlieren texture suggested that this phase was the biaxial smectic \(A, SmA_b\), phase, in which molecular rotation around the long axis is, to some degree, frozen and this assignment is consistent with the increase in
layer spacing on cooling. The absence of a polar response under an electric field indicated that in the SmA\textsubscript{b} phase the dipole moments of the molecules must be locally compensated. On cooling the SmA\textsubscript{b} phase into the lowest temperature liquid-like smectic phase, the layer spacing decreased continuously suggesting a transition to a tilted phase, but surprisingly a homeotropic optical texture was restored (Figure 15a) ruling out the possibility of a simple all-in-one plane synclinic or anticlinic SmC phase. The observed optical uniaxiality of this tilted phase strongly suggested an averaging of the molecular orientations arising from the formation of a helical structure. This view was supported using soft X-ray resonant scattering which revealed that, at the transition from the SmA phase, a resonant Bragg peak developed corresponding to a pitch length of around 150 Å decreasing to around 117 Å at the transition to the HexI phase (Figure 15b). The pitch length was incommensurate with the layer thickness, corresponding to around 3–4 smectic layer distances. This heliconical smectic C phase was designated as the SmC\textsubscript{TB} phase and the molecular arrangement within the phase is shown schematically in Figure 16. The SmC\textsubscript{TB} phase does not appear optically active suggesting that essentially free molecular rotation occurs around the long molecular axis such that the biaxiality is too weak to give rise to detectable layer chirality, and it is known that optical activity is negligible in chiral smectic phases in which the pitch length is much shorter than the wavelength of light as is the case here. A resonant peak was also detected in the HexI phase corresponding to a pitch length of around 100 Å, around 2.2 smectic layer spacings, and this was essentially temperature independent (Figure 15b). The HexI phase showed a weakly birefringent texture (Figure 15a) with optically active domains and gave a circular dichroism signal around the absorption band of the material. This optical activity was attributed to the strongly inhomogeneous electron distribution across the layer and the tilting of the biaxial molecules, i.e., layer chirality (Figure 17); this originates given that the bent molecules are tilted with respect to the layer normal and changing the direction of the bend vector with respect to the tilt plane results in a non-superimposable mirror image. The morphology of the HexI phase was studied using atomic force microscopy revealing uniformly oriented entangled filaments with an average diameter of around 50 nm. These filaments had a uniform twist sense over areas of micron dimensions. The HexI phase, therefore, exhibits structural chirality at different length-scales, namely layer chirality, nanoscale helices and mesoscopic helical filaments, and strongly resembles the B\textsubscript{4} phase but this has an internal crystalline structure [28,93] whereas the HexI phase does not. It should be noted that although all the smectic phases are single layer type with \( d \sim l \), in the HexI phase, an additional, very weak diffraction signal corresponding to \( 2l \) periodicity was detected [94], evidencing a tendency towards bilayer packing driven by a small imbalance in the density of the cyano groups at consecutive layer interfaces. Based on the RSoXS pattern it was deduced that the structure of HexI phase is anticlinic with a longer helix superimposed on that of the bilayer unit. A comparison of the calculated and experimental RSoXS patterns suggests that the layer chirality is coupled to the handedness of the longer superimposed helix.
Figure 14. The dependence of the transition temperatures on the length of the terminal chain, \(m\), for the CB6OIBeOn series [3]. The N-I transition temperatures have not been shown in order to enlarge the other phase regions. The melting points have been omitted for the sake of clarity. Filled circles denote \(T_{NTBN}\); unfilled squares \(T_{SmAN}\); unfilled diamonds \(T_{SmAbSmA}\); filled diamonds \(T_{SmC_{TB-\alpha}SmAb}\); crosses \(T_{HexI}\).

Figure 15. (a) The optical textures observed between crossed polarisers for CB6OIBeO8 on cooling in an homeotropic cell. A glass bead has been used to indicate that the textures represent the same area of the slide; (b) temperature dependence of the resonant soft X-ray diffraction signal for CB6OIBeO7 recorded on cooling (taken from [3]).
The heliconical SmC_{TB} phase was the first unambiguous example of a short pitch helical structure consisting of achiral dimers arranged into a lamellar phase. The driving force for the spontaneous formation of this short-pitch length heliconical structure is presumably the anomalously low-bend elastic constant which may be attributed to the bent molecules. Such a view is supported by the striped optical texture observed in the SmC_{TB} phase which, by analogy to the interpretation of the striped texture observed for the N_{TB} phase [95], suggests that the bend elastic constant is very low. In chiral systems, the SmCα* phase, found between the SmA* and SmC* phases, has a similarly short pitch length incommensurate with the layer spacing, typically between 5 and 8 smectic layers, and has a small tilt angle [96,97]. In this structure, the helix formation relieves the frustration in systems in which next nearest interactions favour antiparallel tilt. Given the similarity in structure between the SmC_{TB} phase shown by CB6OIBeOm series and the SmCα* phase, we designate this SmC_{TB} variant as the SmC_{TB-α} phase. Polar bent core molecules had also been reported to show a short pitch heliconical smectic phase designated Sm(CP)_{hel} and this is thought to be associated with the growth of polar order at the anticlinic-synclinc transition in smectic phases with a small tilt angle [98–100]. Again, the pitch length was incommensurate with the layer spacing, and typically around three-layer thicknesses. Earlier, liquid crystal phases containing helices of both handedness had been reported for...
bent-core liquid crystals by Sekine et al. [101], although this was later assigned as a B4 phase, and technically crystalline in nature.

We saw earlier that CB6O.10 was the shortest member of the CB6O.m series to exhibit a smectic phase (Figure 13) and this was subsequently studied in detail and assigned as a SmC\textsubscript{TB} phase [94]. The SmC\textsubscript{TB}-N\textsubscript{TB} transition was clearly observed using DSC, but was not associated with changes in optical textures. This suggested that in both phases there was a similar averaging of molecular orientation due to the formation of a helix. Only a small change in birefringence was seen at the SmC\textsubscript{TB}-N\textsubscript{TB} transition, evidencing that the conical angle in both phases is similar, and estimated to be 15°. Non-resonant XRD measurements revealed the layer spacing in the smectic phase to be almost two molecular lengths suggesting a bilayer arrangement with the molecules arranged head-to-head within a layer. A resonant signal was evident in the RSoXS pattern of the N\textsubscript{TB} phase associated with the helix, and showed that the pitch length decreased with decreasing temperature (Figure 18). At the N\textsubscript{TB}-SmC\textsubscript{TB} transition, the resonant signal (q\textsubscript{4} in Figure 18) locks at approximately four molecular lengths and this persists for several degrees below the phase transition. This signal was purely resonant indicating it was associated with the helical structure. A signal associated with the bilayer structure was also observed but off-scale in Figure 18. The continuous evolution of structure at the N\textsubscript{TB}-SmC\textsubscript{TB} transition suggested that in the smectic phase just below the transition, the molecules in adjacent layers are azimuthally rotated by exactly 90° on the tilt cone and this may be described as an ideal clock structure (see Figure 19a). A model used to estimate the intensities of the RSoXS signals suggested that deeper in the phase, the structure may be described as a distorted clock with δ between 60–70° (Figure 19b). In such an arrangement the helical pitch is commensurate with two smectic bilayer spacings; in other words, approximately four molecular lengths. On cooling the SmC\textsubscript{TB} phase, the resonant signal (q\textsubscript{4}) split symmetrically into two resonant signals, (q\textsubscript{4} + q\textsubscript{m}) and (q\textsubscript{4} − q\textsubscript{m}), with the former much more intense. This splitting increased with decreasing temperature. A very weak signal at 2q\textsubscript{m} was also observed.

![Figure 18. The resonant soft-X-ray scattering pattern for CB6O.10 (taken from [94]).](image-url)
The RSoXS pattern of either a smectic phase in which the basic four-layer unit repeats with a regular distribution of the molecules on the tilt cone (Figure 19a,b), or the all-in-one-plane bilayer structure with alternating synclinic and anticlinic interfaces between the constituent molecular layers, should contain only the signal \( q_4 \) and its harmonics. The splitting of the \( q_4 \) signal in Figure 18 indicated that an additional modulation, with a longer periodicity, is superimposed over the basic four-layer unit. It was noted that the pattern shown in Figure 18 was very similar to that seen for the SmC*\( \text{FI}_2 \) phase observed over narrow temperature ranges between anticlinic and synclinic SmC* phases in chiral rod-like materials [102]. The SmC*\( \text{FI}_2 \) has also a four-layer unit cell in which the molecules in consecutive layers form a distorted helix such that the azimuthal angle, \( \delta \), between the projections of the long molecular axis in adjacent layers onto the smectic plane is no longer 90° (Figure 19b). In addition, the azimuthal angle increases by some amount, \( \varepsilon \), with respect to the adjacent layer causing the four-layer unit cell to rotate giving an additional helical structure (Figure 19c). The smectic phase shown by CB6O.10 is thought to be similar to this structure, but it should be stressed that given the molecules are achiral, the driving force for its formation must be different than in case of the SmC*\( \text{FI}_2 \) phase. We termed this phase SmC\( \text{TB-DH} \) where DH indicates double helix and show the structure as a distorted clock in Figure 19c and schematically in Figure 20. An important difference to the structure of the SmC*\( \text{FI}_2 \) Phase is that in the SmC\( \text{TB-DH} \) phase, the modulation superimposed on the basic four-layer unit is much stronger, and similar to the basic modulation, is on the nanometre scale. A model used to predict the RSoXS signal intensities has shown that the handedness of the short four-layer structure determines that of the longer helical modulation. The angle \( \delta \) is essentially temperature independent whereas \( \varepsilon \) is strongly temperature dependent giving rise to the evolution of the helical structure shown in Figure 20. Close to the \( N_{\text{TB}} \)-SmC\( \text{TB-DH} \) transition, we see an almost ideal clock four-layer structure (Figure 19a) but as \( \varepsilon \) increases on decreasing temperature, the structure evolves towards an anticlinic arrangement. We should also note that the bent nature of the molecules introduces another level of structural chirality, the sign of which is defined by the relative orientation of three vectors: the layer normal, the tilt projection onto the smectic layer, and the molecular bend direction. This structural chirality is referred to as layer chirality and was described earlier (Figure 17) [92]. Presumably the sign of the layer chirality is coupled to the handedness of the four-layer basic unit and hence to the longer, superimposed helix. We have seen earlier that in the \( N_{\text{TB}} \) phase steric interactions drive the formation of the helical structure, and the pitch length decreases on reducing the temperature. On cooling into the SmC\( \text{TB-DH} \) phase an ideal four-layer clock helical structure forms, in which the azimuthal tilt angle changes by 90° on passing from layer to layer. As the temperature is reduced, the competition increases between the tendency to twist and the entropy-driven tendency for molecules to remain in one tilt plane which allows for easier molecular movement along the layer.

Figure 19. Schematic representations of the SmC\( \text{TB} \) phases using the clock model in which the ellipse represents the direction of the tilt projected onto the layer plane. (a) The ideal clock in which molecules rotate on the tilt cone by exactly 90° between consecutive layers. (b) The distorted clock in which the azimuthal angle between the projections of the long molecular axis in adjacent layers onto the smectic plane is no longer 90°. (c) A change in azimuthal angle to the adjacent layer causing the four-layer unit cell to rotate giving an additional helical structure.
normal, and thus, the structure continuously evolves from the four-layer clock to an almost anticlinic bilayer arrangement. By comparison, in chiral materials a series of discontinuous phase transitions have been observed rather than a continuous evolution of structure [96].

A study of the longer homologues of the CB6O.m series, \((m = 12, 14, 16, 18)\) revealed a sequence of four smectic phases: SmA—SmC\(_{\text{TB-SH}}\)—SmC\(_{\text{TB-DH}}\)—SmX below the I or N phase (Figure 13) [91]. The lowest temperature phase, SmX, was not studied in detail, but its X-ray diffraction pattern indicated that it is a hexatic-type smectic phase. The SmA phase was optically uniaxial (Figure 21) and the layer spacing corresponded to two molecular lengths indicating a bilayer structure in which the cyano groups are concentrated at alternating interfaces, as seen for CB6O.10. X-ray diffraction revealed that the SmA, SmC\(_{\text{TB-SH}}\), and SmC\(_{\text{TB-DH}}\) phases are liquid-like in terms of in-plane ordering. The SmC\(_{\text{TB-SH}}\) phase was optically biaxial whereas the SmC\(_{\text{TB-DH}}\) is optically uniaxial (Figure 21). The RSoXS pattern of the SmC\(_{\text{TB-SH}}\) phase contained a resonant peak corresponding to four molecular layers and its position was temperature independent, and using non-resonant XRD, a bilayer structure was observed. SH is used to indicate that the SmC\(_{\text{TB-SH}}\) phase is characterised by having a single helix. On entering the SmC\(_{\text{TB-DH}}\) phase, symmetric satellites of the resonant peak developed indicating an additional modulation superimposed on the basic four-layer helix. Unlike the continuous development of this peak splitting seen in the RSoXS pattern for CB6O.10, the satellites’ peaks change discontinuously with temperature, and this is thought to be associated with surface interactions. The SmC\(_{\text{TB-SH}}\)—SmC\(_{\text{TB-DH}}\) phase transition is not associated with a change in layer spacing as measured using non-resonant XRD, implying that the tilt angle does not change. The optical biaxiality of the SmC\(_{\text{TB-SH}}\) phase suggests a distorted clock structure for its four-layer unit cell (Figure 19b). In the optically uniaxial SmC\(_{\text{TB-DH}}\) phase an additional shift in azimuthal angle, \(\varepsilon\), between consecutive layers gives rise to a longer pitch-length helical modulation superimposed on the basic 4-layer helical structure. This additional helix has a pitch length corresponding to around 16 layers deep into the SmC\(_{\text{TB-DH}}\) phase but as the temperature approaches the SmC\(_{\text{TB-SH}}\)—SmC\(_{\text{TB-DH}}\) transition, the pitch length jumps to around 46-layer distances. As we saw for CB6O.10, this change in the pitch length reflects the evolution of the structure from a bilayer towards a four-layer arrangement with alternating synclinic and anticlinic interfaces. The equal intensities of the satellite peaks in the RSoXS pattern for the SmC\(_{\text{TB-DH}}\) phase imply that the angle \(\delta\) must be small such that the basic four-layer unit is a strongly distorted clock in

![Figure 20. A schematic representation of the evolution of the four-layer structure in the SmC\(_{\text{TB-DH}}\) phase on increasing \(\varepsilon\), the difference between the azimuthal angle between adjacent layers. For clarity the tilt angle has been increased. (Taken from [94]).](image-url)
which alternating interfaces approach synclinic and anticlinic arrangements. A study of the SmC$_{TB-DH}$–SmC$_{TB-SH}$ transition using optical microscopy revealed selective reflection colours covering the whole optical spectrum at temperatures very close to the transition, which indicated unwinding of the additional helix of the SmC$_{TB-DH}$ phase (Figure 22) [91]. The selective reflection of light occurs when the pitch of the helix, as it unwinds, becomes comparable to the wavelength of visible light, and this was the first observation of such behaviour for achiral liquid crystals.

![Figure 21. Optical textures of CB6O.14 observed between crossed polarizers in a 3 μm thick cell with homeotropic anchoring (taken from [91]).](image1)

![Figure 22. The optical texture shown by CB6O.12 taken in a 3 μm thick cell with homeotropic anchoring at the SmC$_{TB-DH}$–SmC$_{TB-SH}$ transition showing selective reflection. The colours appear simultaneously because of a small temperature gradient in the sample. ( Taken from [91]).](image2)

The corresponding dimers containing a thioether-linked terminated chain, CB6O.Sm$_m$ [103]

also show the SmC$_{TB-DH}$–SmC$_{TB-SH}$ transition for $m > 13$, but the monotropic nature of these phases prevented their detailed study. This series of bent dimers also showed the helical filament B$_4$ phase described earlier.

To investigate the effects of increasing the spacer length on the formation of the heliconical SmC$_{TB}$ phases, we studied the properties of the 1-(4-cyanobiphenyl-4′-yl)-10-(4-alkylaniline-benzylidene-4′-oxy)decanes (CB10O.$m$) [104]:

![Chemical structure image](image3)
with the terminal chain length, \( m = 1\text{–}10, 12, 14, 16 \text{ and } 18 \). The CB10O.m series shows a rich phase polymorphism including the N and N\(_{TB}\) phases, and six different smectic phases (Figure 23). All the homologues showed the conventional N phase. For the shortest members, \( m = 1\text{–}3 \), the N\(_{TB}\) phase was seen. An intercalated SmC\(_A\) phase emerged for CB10O.3, and for \( m = 4\text{–}6 \) the N\(_{TB}\) phase was extinguished and a direct SmC\(_A\)-N transition observed. The N\(_{TB}\) phase remerged with CB10O.7, and the SmC\(_A\) phase extinguished after \( m = 8 \). The homologues with \( m = 9 \) and 10 were exclusively nematogenic, showing both N and N\(_{TB}\) phases. Smectic behaviour re-emerged at \( m = 12 \), and the longer homologues exhibited heliconical SmC\(_{TB}\) phases. This pattern of behaviour has clear similarities to that seen for the CBO4O.m series (Figure 5) for which smectic phases were observed for short and long terminal chain lengths and solely nematic behaviour for intermediate chain lengths [18]. For the CBO4O.m series, this was interpreted in terms of the change in the structure of the smectic phases on increasing chain length from being interdigitated to intercalated (Figure 6). A similar explanation accounts for the behaviour seen for the CB10O.m series except that with increasing \( m \) we now see a switch from intercalated to interdigitated bilayer smectic phases. Three homologues (\( m = 12, 14, 16 \)) show SmC\(_{TB\text{-}SH}\) and SmC\(_{TB\text{-}DH}\) phases whereas for the longest homologue only the SmC\(_{TB\text{-}SH}\) phase is seen, presumably the transition to the SmC\(_{TB\text{-}DH}\) phase is precluded by the formation of the underlying SmY phase. As described for the CB6O.m series, the SmC\(_{TB\text{-}SH}\) phase is optically biaxial implying a strongly distorted clock arrangement (Figure 19b) whereas the SmC\(_{TB\text{-}DH}\) phase is optically uniaxial, given the additional modulation superimposed on the basic four-layer structure leading to space-averaging of the azimuthal positions of the molecules along the layer normal (Figure 20). It is interesting to note that there is no apparent change in layer spacing at the SmC\(_{TB\text{-}SH}\)-SmC\(_{TB\text{-}DH}\) transition implying the tilt angle is similar in both. The striking difference in the nature of the smectic phases shown by \( m = 3\text{–}8 \), and \( m \geq 12 \) is revealed in the behaviour of an approximately equimolar mixture of homologues with \( m = 6 \) and 16 that exhibited the N\(_{TB}\) phase over a broad temperature range although neither individual component does. This shows that the intercalated and interdigitated smectic phases are incompatible and destabilised in the mixture, revealing the underlying N\(_{TB}\) phase.

![Figure 23. The dependence of the transition temperatures on the length of the terminal chain, \( m \), for the CB10O.m series [104]. The melting points have been omitted for the sake of clarity. The dependence of the transition temperatures on the length of the terminal chain, \( m \), for the CB10O.m series [104]. The melting points have been omitted for the sake of clarity. Unfilled circles denoted T\(_{NI}\); filled circles T\(_{N\text{TB}}\); crosses T\(_{Sm\text{CN} \text{/N}_{TB}}\); dashes T\(_{SmX}\); squares T\(_{Sm\text{AN}}\); unfilled diamonds T\(_{Sm\text{C}_{TB\text{-}SH}}\); filled diamonds T\(_{Sm\text{C}_{TB\text{-}DH}Sm\text{C}_{TB\text{-}SH}}\); stars T\(_{Sm\text{Y}}\).](image-url)
7. Summary and Outlook

In this review we have discussed the relationships between the smectic behaviour of liquid crystal dimers and their molecular structures. These are dominated by the average shape of the molecule and markedly different behaviour is seen for bent-odd membered dimers than for linear even-membered dimers. For symmetric dimers there is a strong preference for the formation of monolayer smectic phases driven apparently by the incompatibility between the mixing of the spacers and terminal chains. This may be overcome for nonsymmetric dimers for which intercalated and interdigitated smectic phases are observed depending on the relative lengths of the spacers and terminal chains. The stronger tendency of odd-membered dimers to exhibit tilted phases is attributed to the difficulty associated with the packing of their bent shape into orthogonal phases. By adjusting the molecular curvature, it is possible to drive the formation of spontaneously chiral twist-bend smectic phases, and to date, we have reported three variants of the SmC\(_{\alpha}\) phase. In the SmC\(_{\text{TB}}\) phase, the helical pitch length is incommensurate with the layer spacing and the phase is optically uniaxial. In the SmC\(_{\text{TB-SH}}\) phase the helical pitch corresponds to approximately four molecular lengths, and may be described by the distorted clock model and the phase is optically biaxial. The SmC\(_{\text{TB-DH}}\) phase has an additional helical modulation described by the rotation of the basic four-layer units by a temperature dependent angle, and thus is optically uniaxial. To date, the twist-bend arrangement has only been reported for either monolayer or bilayer smectic phases, and not for intercalated structures. We anticipate, however, the discovery of further variants of the SmC\(_{\alpha}\) phase as a wider expanse of molecular structure space is explored. More widely, the spontaneous formation of chiral structures with multiple different levels of chirality by achiral molecules is an area of intense global interest and of fundamental importance in the physical and biological sciences. The study of liquid crystal dimers has an exciting and vibrant future!

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