

Cite this: *J. Mater. Chem. C*, 2013, **1**, 6667

## The influence of structure on the elastic, optical and dielectric properties of nematic phases formed from bent-core molecules

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The physical properties of the nematic phases formed by four bent-core oxadiazole based materials are reported. In particular, the splay ( $K_{11}$ ), twist ( $K_{22}$ ) and bend ( $K_{33}$ ) elastic constants, the birefringence and the dielectric anisotropy of the materials are described and the effect of chain length and the presence of fluoro-substituents at the outer phenylene group of the aromatic core structure on these parameters is determined. The birefringence and order parameter are found to be independent of the modification of molecular structure. The dielectric anisotropy is quite strongly dependent on molecular structure; the fluoro-substituted material has the largest magnitude of dielectric anisotropy while the alkyl-substituted compound has the smallest. Changes in the molecular length and fluoro-substitution in the bent-core materials are found to have little influence on the splay, twist and bend elastic constants at equivalent reduced temperatures. However, the material substituted with an alkyl terminal chain exhibits both smaller elastic constants and a less marked dependence on temperature than the alkoxy-substituted compounds. A possible insight into the behaviour of the elastic constants relevant to the formation of the dark conglomerate phase, which underlies the nematic phase in one of the compounds studied, is suggested by following the analysis proposed by Berreman and Meiboom. Importantly, using molecular field theory and atomistic modelling, we calculate elastic constants that are in excellent agreement with the experimental values. Our conclusion that the elasticity in the nematic phase formed from bent-core molecules is not strongly influenced by changes to the terminal chains or the presence of fluoro-substituents at the outer phenylene group of the aromatic core structure is in agreement with our previous work showing that the dominant parameter is the bend angle.

Received 8th August 2013

Accepted 3rd September 2013

DOI: 10.1039/c3tc31545b

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### 1 Introduction

Bent-core nematic liquid crystals (NLCs) are an exciting class of mesogenic systems. In addition to their potential to exhibit a biaxial nematic phase (a still somewhat contentious issue<sup>1,2</sup>) these systems are reported to possess extraordinary physical properties such as high Kerr constants,<sup>3</sup> large flexoelectricity,<sup>4</sup> and viscosity and elasticity behaviour quite different from that observed in calamitic NLCs.<sup>4–10</sup> Lately, there has been much interest in the particularly intriguing elastic properties of bent-core NLCs. We recently obtained excellent agreement between our experimentally determined elastic constants and calculations based on molecular-field theory and atomistic modelling in an oxadiazole bent-core NLC.<sup>6</sup> Subsequently, we established

that the bend angle of bent-core molecules plays an extremely important role in determining the elastic properties.<sup>10</sup> Our paper demonstrated that in case of bent-core NLCs with a bend angle of  $\sim 164^\circ$ , the bend ( $K_{33}$ ) elastic constant is larger than the splay ( $K_{11}$ ) elastic constant, analogous to the elastic behaviour in calamitic (rod-like) NLCs and in contrast to the expectation that all bent-core materials might have  $K_{33} < K_{11}$ . While the role of bend angle is relatively clear, the difficulty of measuring the physical properties of bent-core nematic liquid crystal materials has ensured that, to date there are no reports on the vital influence of other structural parameters on the elastic behaviour in these materials.

The Frank elastic constants are key physical parameters in nematic liquid crystals. The splay ( $K_{11}$ ), twist ( $K_{22}$ ) and bend ( $K_{33}$ ) elastic constants influence both the threshold voltage and the response times of nematic devices. Knowledge of the elastic constants allows an understanding of the microscopic structure of the ordered state, testing mean field theory, and providing insight into the director distortion in devices. Further, information about the elastic constants is needed in measurements of flexoelectricity, a parameter that is currently hotly debated in

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nematic systems formed from bent-core molecules. The influence of molecular structure variations in calamitic liquid crystals on the elastic behaviour is well-understood. Numerous reports describe the influence of factors such as the change in molecular length and width on the elastic properties of calamitic NLCs.<sup>11–15</sup>

This paper describes structure–property correlations determined in the nematic phases of four related bent-core materials. We consider the influence of the terminal chain length and the presence of fluoro-substituents at the outer phenylene group of the aromatic core structure on the optical, dielectric and elastic properties of the nematic phase and consider, where possible, the influence of the underlying phases on the elastic properties. We further examine theoretical fits to experimental data, together with calculations of the elastic constants from atomistic modelling.

## 2 Experimental and computational methods

### 2.1 Materials and devices

Four bent-core mesogens derived from a 1,3,4-oxadiazole-biphenyl (ODBP) core are shown with their phase transitions in Fig. 1. Although the absolute values of the transition temperatures are slightly higher than those quoted elsewhere,<sup>16–18</sup> the nematic range is the same and differences are attributed to the absolute accuracy of hot stages at these high temperatures. All temperatures are, therefore, quoted with respect to the nematic to isotropic phase transition temperature,  $T_{NI}$  correcting for differences in absolute temperature accuracy in different apparatus. The rationale to choosing compounds **1–4** is that oxadiazoles of similar structure are well-known materials (the first claims of nematic biaxiality were made for compound **1** (ref. 19 and 20)) which have the same core and different alkyl/alkoxy chain lengths, which is a key consideration in the present work. The fact that the materials also have distinct phases underlying the nematic phase offers an additional opportunity to study the behaviour of elasticity near the phase transitions. Compounds **1** and **2** exhibit an underlying smectic-C (SmC) phase, compound **3** an unidentified smectic phase and the 'dark conglomerate (DC)' phase is found in compound **4**.

All measurements employ devices with glass substrates and a liquid crystal layer approximately 5  $\mu\text{m}$  thick. The transparent

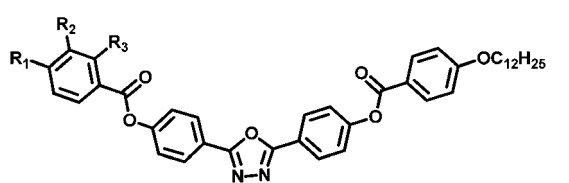
indium-tin-oxide electrode area is 30  $\text{mm}^2$  ( $20 \Omega \square^{-1}$ ) and a high temperature conducting glue (H21D, Epoxy Technology) is used to attach the wires to the electrode surfaces. The electrical properties of the devices were calibrated using air as a standard reference at all temperatures studied and the devices were capillary-filled at 190 °C. All compounds exhibit negative dielectric anisotropy ( $\Delta\epsilon$ ), across the entire nematic regime and as a consequence the measurement of the splay and bend elastic constants from Freedericksz transition requires homeotropic alignment, which is a challenge in bent-core materials. While excellent alignment was obtained using trichloro-octadecyl silane in heptane across the entire nematic regime for compounds **2** and **4**, the alignment degraded slightly at low temperatures for compounds **1** and **3**. We take careful account of this in our results. A homogenously aligned interdigitated electrode device (sometimes known as an in-plane switching, IPS device) is used to measure the twist elastic constant, with the rubbing direction parallel to the field. The IPS device thickness is also  $\sim 5 \mu\text{m}$ , with an electrode gap and width of 20  $\mu\text{m}$  and 10  $\mu\text{m}$  respectively. The devices were held in a Linkam THMS600 hot stage equipped with a TMS 93 controller which allows temperature control of the samples with a relative accuracy of  $\pm 0.1 \text{ K}$ .

### 2.2 Birefringence measurements

The birefringence of the high temperature nematic phases exhibited by the bent-core materials is measured by analysis of the reflection spectra from planar devices *via* the Berreman methodology.<sup>21</sup> Details of both the apparatus used and the fitting procedure are given in ref. 6 and 22. Reflection spectra are first taken for the empty device, allowing the thickness to be measured with an accuracy of  $\pm 0.02 \mu\text{m}$ , before filling with the liquid crystal. The ordinary and extraordinary refractive indices ( $n_o$  and  $n_e$ ) are determined from spectra obtained with the nematic director oriented perpendicular and parallel to the incident light polarization respectively. The birefringence,  $\Delta n = n_e - n_o$ , may then be calculated with an accuracy of  $\pm 0.001$ . All values of refractive index are quoted for light of wavelength 589 nm.

### 2.3 Measurement of the dielectric anisotropy and elastic constants

The methodology employed to determine the splay and bend elastic constants has also been described in detail elsewhere,<sup>6,10</sup> and is based on an analysis of the variation in capacitance of a device that results from the electric-field induced Freedericksz transition. For homeotropic devices, the parallel component of dielectric permittivity,  $\epsilon_{||}$  is observed at fields below the Freedericksz threshold while at high field strength, the director alignment approaches the direction perpendicular to the field. A detailed fit to the capacitance–voltage curve allows the perpendicular component of dielectric permittivity,  $\epsilon_{\perp}$  and hence the dielectric anisotropy,  $\Delta\epsilon$  to be deduced with an absolute accuracy that depends on the quality of the alignment, typically  $\sim 5\%$  for excellent homeotropic alignment. The relative accuracy of a particular data set which describes the



Compound **1**:  $R_1 = C_{12}H_{25}O$ ,  $R_2 = R_3 = H$   
 Compound **2**:  $R_1 = C_9H_{19}O$ ,  $R_2 = R_3 = F$   
 Compound **3**:  $R_1 = C_9H_{19}O$ ,  $R_2 = R_3 = H$   
 Compound **4**:  $R_1 = C_5H_{11}$ ,  $R_2 = R_3 = H$

Iso 217.7°C N 203.6°C SmC  
 Iso 221.4°C N 206.0°C SmC  
 Iso 236.6°C N 204.6°C SmX  
 Iso 239.6°C N 184.6°C DC

**Fig. 1** The chemical structure and phase transitions of 1,3,4-oxadiazole bent-core compounds **1–4** (phase transitions other than that directly below the nematic phase are omitted).



temperature dependence is however better than this,  $\sim 2\%$ . The threshold voltage is related to the bend modulus while both the splay and bend constants influence the steepness of the threshold behaviour and the fit also allows the splay and bend elastic constants,  $K_{11}$  and  $K_{33}$  to be determined. This methodology allows the elastic constants to be determined straightforwardly, with an accuracy of around 5%, but it does not take account of any contributions from flexoelectric effects.

Although measurement of the twist elastic constant,  $K_{22}$ , is intricate in comparison with measuring the splay and bend elastic constants in LCs, we have successfully utilized in-plane devices to measure  $K_{22}$  in a positive  $\Delta\epsilon$  bent-core NLC material.<sup>10</sup> In the present work, similar devices with the rubbing direction perpendicular to the length of the electrodes are used to measure  $K_{22}$ .<sup>23,24</sup> The voltage is applied initially parallel to the director and the electric-field induced Freedericksz transition occurs at a voltage  $V_{\text{th}}^{K_{22}}$  given by:

$$V_{\text{th}}^{K_{22}} = \frac{\pi l}{d} \sqrt{\frac{K_{22}}{\epsilon_0 \Delta\epsilon}} \quad (1)$$

where  $l$  is the electrode separation and  $d$  is the thickness of the device. The Freedericksz threshold is best measured from observations of the transmitted light intensity when the device is held on a polarizing microscope with the director initially aligned with the polarizer. We estimate that finding the elastic constant in this way gives an absolute uncertainty in the value of  $K_{22}$  of around 60%, but we expect that the relative accuracy of this method in a particular data set is much better than this ( $\sim 20\%$ ). We believe that we can derive useful information from this method despite its intrinsic uncertainties.

#### 2.4 Atomistic calculation of the order parameters and elastic constants

The approach to calculating the order parameters and elastic constants was presented in ref. 25 and application to bent-core materials has also been described in detail in ref. 6 and 10. Indeed, calculations of the splay ( $K_{11}$ ) and bend ( $K_{33}$ ) elastic constants for compound 4 have already been presented,<sup>6</sup> but here the study is extended to include compounds 2 and 3, and the twist elastic constant ( $K_{22}$ ) is also evaluated. All four compounds have the  $\text{OC}_{12}\text{H}_{25}$  group as one terminal chain. Compounds 1–3 have alkoxy chains as the second terminal group, while compound 4 has an alkyl chain with the least number of carbon atoms (5) at the other end of the molecule. Compounds 2 and 3 are similar in structure as they both have the same terminal alkoxy groups but differ in that the former has two fluoro-substituents in the outer phenylene group of one arm (see Fig. 1). In this paper, compounds 2 and 3 are chosen for the simulation work to investigate the sensitivity of the elastic constants of the bent-core material to the length of the lateral chains and to the presence of substituents in the aromatic rings.

As in ref. 6 the Rotational Isomeric State (RIS) approximation was used,<sup>26</sup> and Metropolis Monte Carlo (MC)<sup>27</sup> sampling of conformers was performed. Conformer geometry and energy were defined on the basis of quantum chemical calculations,

carried out for single molecules of molecular fragments in vacuum.<sup>28</sup> Thus, the following choices were made:

- Planar geometry of the central three rings (2,5-diphenyl-1,3,4-oxadiazole) was assumed.<sup>29</sup>

- In phenyl benzoate, a planar benzoate moiety was assumed and two states of the  $\text{C}_{\text{ar}}-\text{C}_{\text{ar}}-\text{O}-\text{CO}$  dihedral were considered ( $\pm 90^\circ$ ), which correspond to the middle of the highest probability region for this dihedral.<sup>6</sup>

- The benzoate group of phenyl-2,3-difluoro benzoate was also assumed to be planar, with the CO group on the opposite side of the fluorine (F) atoms; for the  $\text{C}_{\text{ar}}-\text{C}_{\text{ar}}-\text{O}-\text{CO}$  dihedral the same choice as for phenyl benzoate was taken. These assumptions were based on torsional potentials obtained by relaxed scans at the DFT/M062X/6-31+g\*\* level of the theory for the  $\text{C}_{\text{ar}}-\text{C}_{\text{ar}}-\text{O}-\text{CO}$  dihedral, and at the DFT/B3LYP/6-31+g\* level for the  $\text{C}_{\text{ar}}-\text{C}_{\text{ar}}-\text{CO}-\text{O}$  dihedral. The torsional potential for the latter has two minima in correspondence to the planar configurations, and the configuration having the CO group pointing in the same direction as the F atoms is higher in energy than the other by about 3 kJ mol<sup>-1</sup>. The torsional potential for the  $\text{C}_{\text{ar}}-\text{C}_{\text{ar}}-\text{O}-\text{CO}$  dihedral has the same general form as that of phenyl benzoate, but the barriers at  $0^\circ$  and  $180^\circ$  are significantly lower; thus, higher rotational freedom is expected for the  $\text{C}_{\text{ar}}-\text{O}$  bond.

- The  $\text{CH}_2-\text{CH}_2$  and the  $\text{O}-\text{CH}_2$  bonds of the alkyl and alkoxy chains were allowed to jump between the *trans* ( $180^\circ$ ), *gauche*+ ( $+65^\circ$ ) and *gauche*- ( $-65^\circ$ ) states, with relative energies defined as in ref. 6. Two possible equivalent states were assumed for the  $\text{C}_{\text{ar}}-\text{CH}_2$  bond, with the first  $\text{CH}_2-\text{CH}_2$  bond perpendicular to the phenyl ring, on one or the other side. Also for the  $\text{C}_{\text{ar}}-\text{O}$  bond, two equivalent states were assumed, but in this case with the  $\text{O}-\text{CH}_2$  bond lying on the same plane of the aromatic ring, on either side.<sup>6</sup> These two states are equivalent if the aromatic group is a benzoate, whereas in the case of 2,3-difluorobenzoate the conformation having the  $\text{O}-\text{CH}_2$  bond pointing opposite to the F atoms is more stable by about 3 kJ mol<sup>-1</sup> (DFT/B3LYP/6-31g\*\*).

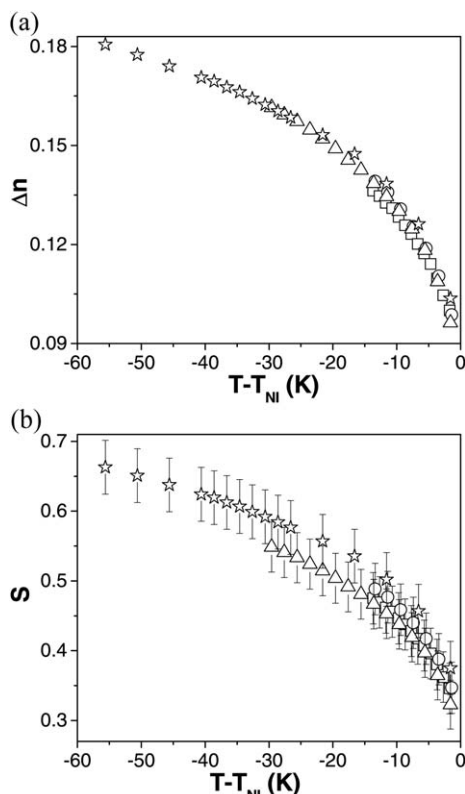
In MC sampling of conformers, the structures having pairs of atoms closer than a cut-off distance equal to  $0.82\sigma$ , where  $\sigma$  is the sum of their van der Waals radii, were discarded. Van der Waals radii equal to 0.185 nm (C), 0.15 nm (N and O), and 0.1 nm (H)<sup>30</sup> were assumed. In this way sterically hindered conformations were rejected, including those with adjacent *gauche*+ *gauche*- (or *gauche*- *gauche*+ ) pairs in the hydrocarbon chains. For each conformer, the molecular surface was generated by the fast molecular surface calculation library (MSMS),<sup>31</sup> assuming a rolling sphere radius equal to 0.3 nm and density of vertices equal to  $5 \text{ \AA}^{-2}$ . The same van der Waals radii used for the cut-off distance were assumed for this purpose.

## 3 Results and discussion

### 3.1 Birefringence and order parameter

Fig. 2(a) shows the behaviour of the birefringence,  $\Delta n$  as a function of reduced temperature ( $T - T_{\text{NI}}$ ) in compounds 1–4. The birefringence increases monotonically with decreasing temperature for all compounds, behaviour that is entirely consistent with that observed in nematic materials generally.





**Fig. 2** The birefringence,  $\Delta n$  (a) and order parameter,  $S$  (b) plotted as a function of reduced temperature,  $T - T_{NI}$  for compounds **1** (square), **2** (circle), **3** (triangle) and **4** (star).

Indeed, the birefringence behaviour of each of the four materials is indistinguishable at the same reduced temperature. All four compounds exhibit  $\Delta n \sim 0.10$  near  $T_{NI}$  increasing to  $\sim 0.14$  (compounds **1** and **2**),  $\sim 0.16$  (compound **3**) and  $\sim 0.18$  (compound **4**) as the low lying phases are approached (the differences are entirely attributable to the different nematic ranges of each of the compounds). It is clear that the birefringence of the materials is dominated by the oxadiazole-based core. Sathyaranayana *et al.* report similar values of birefringence in a different bent-core material.<sup>7</sup> On the other hand, a shape-persistent thiadiazole bent-core NLC exhibits a much larger birefringence (0.17 just below  $T_{NI}$ , gradually increasing to  $\sim 0.39$  at lower temperatures); the high birefringence is attributed to the four triple bonds in the thiadiazole mesogen.<sup>10</sup> An asymmetric L-shaped molecule has recently been reported to have rather high values of birefringence;  $\Delta n > 0.25$  at temperatures far from  $T_{NI}$ .<sup>32</sup>

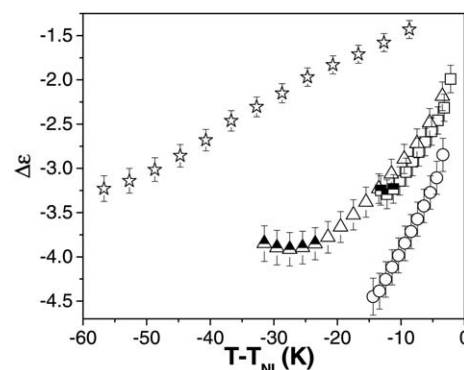
As mentioned, the similarity of the birefringence of compounds **1–4** at equivalent reduced temperatures indicates that neither changes in terminal chain length nor the presence of fluoro-substituents on the outer phenylene group of the aromatic core structure has a strong influence on this parameter. This observation can be compared with the behaviour observed in calamitic NLCs such as alkoxy-azoxybenzenes, alkyl cyano-biphenyls (*n*CB), alkylcyanophenyl cyclohexanes (*n*PCH) and alkylcyclohexyl isothiocyanato-benzenes (*n*CHBT). In these systems, a strong odd–even effect is seen for short chain lengths

and in general,  $\Delta n$  decreases with increasing molecular length.<sup>33–35</sup> However, the variation is not as marked at longer chain lengths, which offer a better comparison with compounds studied here. Indeed the most significant change is seen when the core structure is changed. Further, Avci *et al.* report the birefringence of two similar L-shaped molecules substituted with  $C_{12}$  and  $C_5$  terminal chains; values coincide to around 10 K below  $T_{NI}$ , but then appear to diverge at lower temperatures.<sup>32</sup>

The orientational order parameter,  $S$  can be determined from the birefringence data using the well-known Haller methodology<sup>36</sup> (details are given in ref. 6 and 10). Fig. 2(b) shows the order parameter as a function of  $T - T_{NI}$  for all four compounds.  $S$  increases from  $\sim 0.30$  near  $T_{NI}$  for all of them to  $\sim 0.48$  (compounds **1** and **2**),  $\sim 0.55$  (compound **3**) and  $\sim 0.66$  (compound **4**) near the low lying phases respectively. Excellent agreement is found between these measurements and the order parameter ( $P_{200}$ ) values obtained *via* Polarised Raman Spectroscopy (PRS).<sup>16</sup> Ref. 6 explicitly shows the excellent agreement of the two different approaches for compound **4**. Order parameter measurements carried out *via* NMR on a different bent-core material by Dong *et al.* also are in excellent agreement with the values obtained here.<sup>37</sup> The order parameter values obtained (Fig. 2(b)) at the same reduced temperatures, are very similar to those obtained for calamitic nematics ( $\sim 0.6$ –0.75)<sup>16</sup> but are slightly higher than in a thiadiazole<sup>10</sup> and another bent-core system.<sup>7</sup> It is clear that the variation in molecular structure between compounds **1–4** has a negligible influence on the order parameter. Dong *et al.* also illustrated (*via* NMR) that  $S$  does not change on increasing the chain length in two carboxylate derived bent-core materials (with ten and eleven C atoms).<sup>37</sup>

### 3.2 Dielectric anisotropy and elastic constants

The magnitude of dielectric anisotropy,  $|\Delta\epsilon|$  determines the interaction strength of the LC with an electric field, and is therefore a vital parameter that influences the threshold voltage of a material. Fig. 3 shows  $\Delta\epsilon$  as a function of  $T - T_{NI}$  for compounds **1–4**. The magnitude of the dielectric anisotropy is found to increase with decreasing temperature in all compounds as would be expected and as is also observed in



**Fig. 3** The dielectric anisotropy,  $\Delta\epsilon$  as a function of reduced temperature,  $T - T_{NI}$  for compounds **1** (square), **2** (circle), **3** (triangle) and **4** (star). The few half filled symbols for compounds **1** and **3** represent the slight deterioration of alignment at low temperatures.



calamitic NLCs.<sup>38</sup> There is almost no difference in  $|\Delta\epsilon|$  between compounds **1** ( $C_{12}$ ) and **3** ( $C_9$ ) at similar reduced temperatures; again significant changes with chain length are seen in calamitic NLCs only for relatively short chains.<sup>35,39</sup> Compound **2** ( $C_9$ ) exhibits the largest absolute value of  $\Delta\epsilon$  varying from  $-2.8$  ( $T - T_{NI} = -3.4$  K) to  $-4.4$  ( $T - T_{NI} = -14.4$  K) and the effect can be attributed to the conjugative mesomeric effect of the phenyl ether oxygen on the lateral fluoro substituents, as is seen for calamitic NLCs.<sup>40</sup> Compound **4**, which has one terminal alkyl chain replacing the alkoxy chain, has the smallest dielectric anisotropy, varying from just  $-1.4$  ( $T - T_{NI} = -8.7$  K) to  $-3.2$  ( $T - T_{NI} = -56.7$  K); this is approximately half of the value measured in compounds **1–3**. While such a variation could be due to the different terminal chain in compound **4**, it is to be remembered that the alkyl chain is only  $C_5$ , *i.e.*, it is somewhat shorter than in the other materials. A comparison of  $\Delta\epsilon$  for alkyl (e.g. 8CB) and alkoxy (8OCB) compounds in calamitic NLCs shows little difference in  $\Delta\epsilon$ ,<sup>39,41</sup> while  $|\Delta\epsilon|$  is slightly smaller in 8CB than 5CB. However, we suggest that in the oxadiazole materials substituting an alkyl group in place of an alkoxy terminal chain reduces the lateral dipole of the molecule, hence reducing the dielectric anisotropy.

As already mentioned, although excellent homeotropic alignment was obtained in all compounds, the alignment quality deteriorated for compounds **1** and **3** close to the underlying phase transition. The data points affected in Fig. 3 have been denoted by half-filled symbols; the deterioration was noticed within  $\sim 3$  K of the underlying SmC phase in compound **1** and within  $\sim 10$  K of the underlying phase for compound **3**. We have included these data points partly because of the unexpected deterioration of the alignment (which was reproducible in several different cells), but also to illustrate the influence that small changes in alignment can have on the apparent values of the physical parameters being measured. In the case of both compounds **1** and **3**, the effect of the reduction in alignment quality is, unsurprisingly, to reduce the dielectric anisotropy measured.

To present a clear view of the elastic behaviour of compounds **1–4** with respect to the molecular structure, the splay ( $K_{11}$ ), twist ( $K_{22}$ ) and bend ( $K_{33}$ ) elastic constants are shown separately in Fig. 4(a)–(c) respectively. Note that data are not included unless the alignment quality is excellent, specifically; it makes no sense to deduce elastic constants for the half-filled data points in Fig. 3. Fig. 4(a) shows that  $K_{11}$  increases monotonically with decreasing temperature for all materials. The data for compounds **1–3** are almost indistinguishable, taking values of  $\sim 3.5$  pN close to  $T_{NI}$  and increasing to  $\sim 8$  pN within  $10$  K of the low-temperature transition. Both the magnitude and temperature variation of the splay elastic constant is different in compound **4**, varying much more gradually from  $\sim 3$  pN close to the transition to  $\sim 8.7$  pN at  $\sim 57$  K below it. Fig. 4(b) also shows a monotonic increase in  $K_{22}$  for all compounds as the temperature is reduced. The magnitude of twist is rather small for the materials ( $\sim 1$  pN), and again, compounds **1–3** exhibit almost indistinguishable  $K_{22}$  values at equivalent reduced temperatures. Both the magnitude and temperature dependence of  $K_{22}$  are much smaller in compound

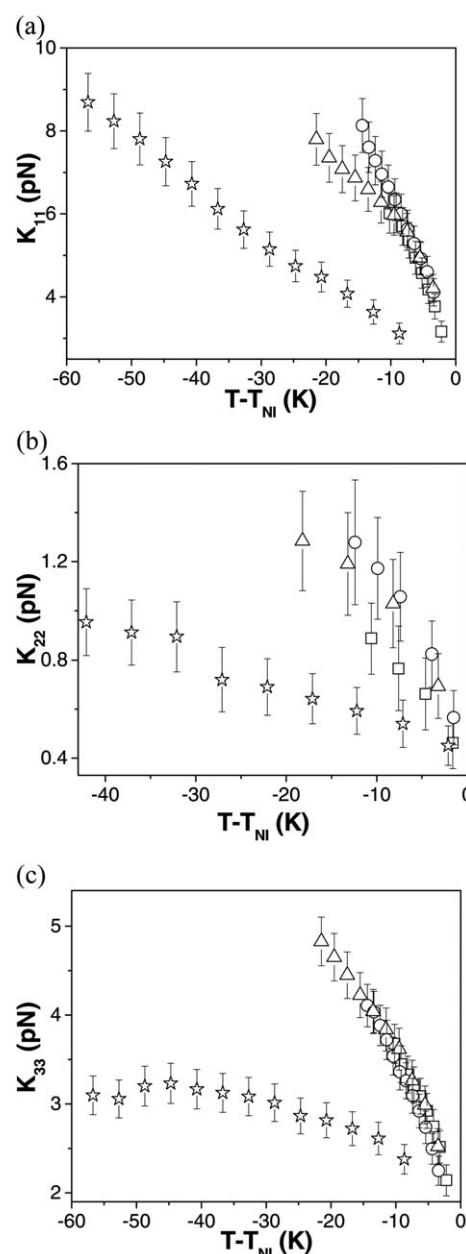


Fig. 4 The splay,  $K_{11}$  (a) twist,  $K_{22}$  (b) and bend,  $K_{33}$  (c) elastic constants as a function of reduced temperature,  $T - T_{NI}$  for compounds **1** (square), **2** (circle), **3** (triangle) and **4** (star).

**4** than in the other materials. Fig. 4(c) displays the behaviour of  $K_{33}$  which generally is seen to increase with decreasing temperature. Once again, the behaviour of  $K_{33}$  is almost indistinguishable for compounds **1–3**, changing from  $\sim 2$  pN close to  $T_{NI}$ , and taking values  $\sim 4$  pN,  $10$  K below the transition. The behaviour of  $K_{33}$  in compound **4** is again very different from the rest of the compounds, increasing only slightly from  $2.4$  pN ( $T - T_{NI} = -8.7$  K) to  $3.2$  pN ( $T - T_{NI} = -44.7$  K) and decreasing marginally thereafter. The behaviour of  $K_{11}$  and  $K_{33}$  for compound **4** have been described in detail previously and are found to be in excellent agreement with the calculations obtained from molecular-field theory and atomistic modelling.<sup>6</sup>

Clearly, overall the elastic behaviour is almost indistinguishable between compounds **1–3**, while the alkyl substituted compound **4** shows significantly lower values of all three elastic constants, together with weaker temperature dependence. For all of the materials we find  $K_{11} > K_{33} > K_{22}$ , analogous to the behaviour of most other bent-core NLCs (a thiadiazole bent-core NLC with a bend angle of  $\sim 164^\circ$  exhibits  $K_{33} > K_{11} > K_{22}$  (ref. 10) in common with calamitic NLCs). Indeed, by extracting data from ref. 32 with differently substituted L-shaped molecules, it can also be seen that neither the length of the terminal chains ( $C_{12}$  or  $C_5$ ) nor the presence of fluoro-substituents in the aromatic core structure significantly changes  $K_{11}$  or  $K_{33}$  in the investigated series of compounds.

We now consider the temperature dependence of the elastic constants near the low lying phases for all compounds (Fig. 4(a)–(c)). The simplest materials to consider are compounds **2** and **4**, for which excellent homeotropic alignment was achieved across the nematic temperature range. The splay and bend elastic constants of compound **4** have been discussed in detail before (ref. 6); the transition to the underlying DC phase is first order, and no pretransitional divergence of the elastic constants is anticipated. Fig. 4(a) and (c) shows that neither splay nor bend exhibit pretransitional divergence in the nematic phase. The small decrease in the bend constant at lower temperatures is potentially of interest with respect to the discussion of the twist-bend nematic phases.<sup>42</sup> No pretransitional divergence is found in either  $K_{11}$  or  $K_{33}$  for compound **2** which has a low lying SmC phase (there are too few data points to draw conclusions about pretransitional behaviour of  $K_{22}$ ). Such behaviour is in agreement with that reported by Tadapatri *et al.* who report no pretransitional divergence in either  $K_{11}$  or  $K_{33}$  for a symmetric bent-core benzoate compound ( $OC_{12}$  on both terminal ends) with a low lying SmC phase.<sup>8</sup> In contrast, Sathyaranayana *et al.* report a strong pretransitional divergence in  $K_{33}$  for a bent-core material with a low lying SmC phase.<sup>7</sup> Findon and Gleeson<sup>43</sup> measured  $K_{11}$  in the nematic and SmC phase of a calamitic mixture, with a continuous variation across the transition.

Any discussion of the pretransitional behaviour in compounds **1** and **3** is complicated by the fact that both materials suffered from deterioration in the otherwise excellent homeotropic alignment as the lower-lying phase was approached. Indeed it is interesting that the deterioration phenomenon was observed in several different cells filled with these two materials, occurring over approximately the same temperature range in every case ( $\sim 3$  K and  $\sim 10$  K for compounds **1** and **3** respectively), but at different temperatures for each compound, ruling out thermal deterioration of the alignment layer. While we cannot, therefore make any comparison of the pretransitional elastic behaviour in these compounds, we can note that such a reproducible change in alignment is unusual and may be of interest given current discussions around anomalous behaviour and texture observations in nematic materials.

### 3.3 Elastic constants and order parameter

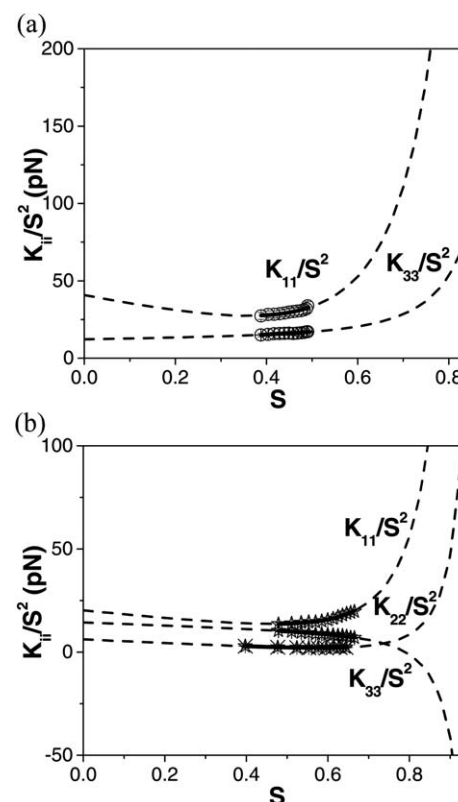
A comparison between the elastic constants and the order parameter not only throws light on the molecular mean field

theory but also helps in understanding the microscopic structure of the ordered state. The elastic constants of the few bent-core NLCs<sup>6,7,10</sup> and mixtures of calamitic and bent-core NLCs<sup>41,44</sup> reported to date do not strictly obey the mean field theory approximation which suggests a square dependence of the order parameter ( $K_{ii} \propto S^2$ ). Indeed, it has been shown for two different bent-core materials<sup>6,10</sup> that the most appropriate way to describe the relationship is to utilize a third order dependency as advocated by Berreman and Meiboom.<sup>45</sup> The relationship is given by:

$$\frac{K_{ii}}{S^2} = K_i^{(2)} + K_i^{(3)} S + K_i^{(4)} \left( \frac{S}{1-S} \right)^2, \quad (2)$$

where  $K_i^{(2)–(4)}$  are second, third and fourth order elastic coefficients obtained from fitting which could, in principle, be temperature dependent, and the subscript  $i$  is 1, 2 or 3 for splay, twist and bend respectively. The second order fitting parameter  $K_i^{(2)}$  is the value of  $K_{ii}/S^2$  when extrapolated to zero order parameter while the coefficients  $K_i^{(3,4)}$  define the behaviour of elastic constants with changing order parameter. In particular, the term in (2) including  $K_i^{(4)}$  accounts for pretransitional divergence of the elastic constants as a low-lying smectic or crystalline phase is approached (high values of order parameter).

Fig. 5(a) and (b) show  $K_{ii}/S^2$  plotted as a function of order parameter for compounds **2** and **4** respectively. The fits to



**Fig. 5** The dependence of  $K_{ii}/S^2$  ( $i = 1, 2$  or 3 for splay, twist and bend respectively) as a function of order parameter for: (a) compound **2**,  $K_{11}/S^2$  (open circles) and  $K_{33}/S^2$  (circles with a plus sign). (b) The same quantities with star symbols for compound **4** and  $K_{22}/S^2$  (star with a cross sign). The solid lines show fits of eqn (2) to the data and the dashed lines extrapolate the fits to low and high order parameter regimes (outside the region of stability of the nematic phase).

**Table 1** Fit parameters obtained from eqn (2) for compounds **2** and **4**

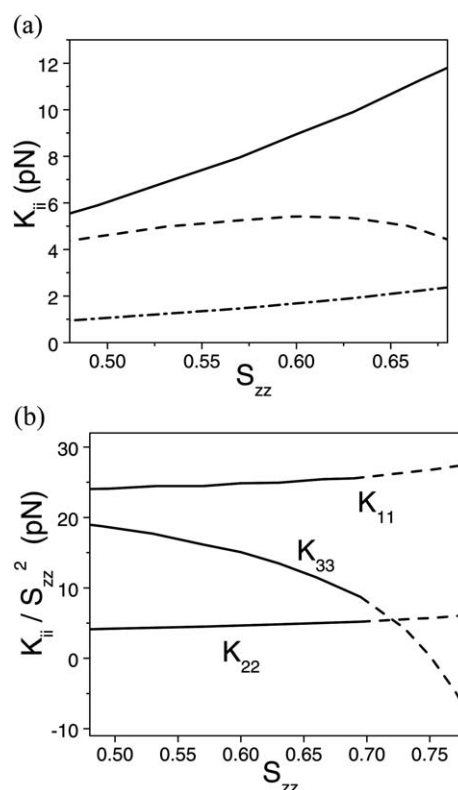
Compounds	$K_1^{(2)}$ (pN)	$K_1^{(3)}$ (pN)	$K_1^{(4)}$ (pN)	$K_2^{(2)}$ (pN)	$K_2^{(3)}$ (pN)	$K_2^{(4)}$ (pN)	$K_3^{(2)}$ (pN)	$K_3^{(3)}$ (pN)	$K_3^{(4)}$ (pN)
<b>2</b>	$41 \pm 13$	$-55 \pm 41$	$20 \pm 8$	—	—	—	$12 \pm 5$	$5 \pm 16$	$2.3 \pm 3.1$
<b>4</b>	$20 \pm 2$	$-19 \pm 5$	$3.2 \pm 0.3$	$6.2 \pm 0.7$	$-9.1 \pm 1.6$	$0.6 \pm 0.1$	$14 \pm 1.5$	$-7.5 \pm 3.3$	$-0.6 \pm 0.2$

eqn (2), shown as solid lines, have been made using the assumption that  $K_i^{(2-4)}$  are temperature independent and the requirement that the elastic constants converge at  $S = 0$  is relaxed. The dotted lines in Fig. 5(a) and (b) are the extrapolated values of  $K_{ii}/S^2$  calculated using the fitting parameters  $K_i^{(2-4)}$  which are given in Table 1. Note that fits are not made to the  $K_{22}$  data for compound **2** as there are only 5 experimental data points. Further, as the elastic constants could not be measured close to the underlying phase for compounds **1** and **3**, fits to the data are not included as they cannot be used to give any insight to the pretransitional behaviour of the materials.

Berreman and Meiboom successfully applied the theory to three standard calamitic NLCs (MBBA, PAA, E7).<sup>45</sup> Their fits all diverge to large (positive) values of elastic constant at high order (low temperature), with the splay curves diverging at higher temperatures (lower order) than those for twist and bend, consistent with a virtual low-lying smectic phase. While we recognise that conclusions drawn from theoretical fits to experimental data depend on both the quality and quantity of the data points, it is nonetheless interesting to apply such an approach to the elastic constants of compounds **2** and **4**. Fig. 5(a) indicates that both  $K_{11}/S^2$  and  $K_{33}/S^2$  for compound **2** diverge to larger values at high  $S$ . The splay curve again diverges at a lower order than the bend, consistent with the occurrence of an underlying SmC phase in this material. The behaviour of compound **4** is different with both  $K_{11}/S^2$  and  $K_{22}/S^2$  diverging to larger values at high  $S$  while  $K_{33}/S^2$  diverges to negative values. Interestingly, again the splay data begin to diverge at lower order (higher temperatures) than twist or bend even though the lower temperature phase is a DC phase. Further, the bend data clearly diverge to negative values at high order, a point of potential relevance to the twist-bend phases predicted by Dozov *et al.*<sup>46</sup> Currently, little is known about the behaviour of the elastic constants in the DC phase which is the low-lying phase in this material. However, the phase is known to be sponge-like with a local SmC structure and the formation relies on a small (or negative) saddle-splay elastic constant,  $K_{24}$ .<sup>47</sup> The divergence of  $K_{11}/S^2$  at lower order than  $K_{33}/S^2$  is again consistent with the formation of a local SmC structure. Perhaps more interestingly, the Erickson inequalities<sup>48</sup> suggest that  $|K_{24}| < K_{22}$  and as  $K_{22}$  takes extremely small values in compound **4** (<1 pN) we can note that the formation of the DC phase is consistent with the elastic behaviour measured. The very low and negatively diverging values of  $K_{33}$  may also be relevant to the formation of the DC phase. Clearly, using eqn (2) to analyse the behaviour of the bent-core materials gives an interesting possible insight into the formation of the underlying phase, though the negative divergence of  $K_{33}/S^2$  on approaching the DC phase must be confirmed in other systems before firm conclusions can be drawn.

### 3.4 Atomistic calculations of the elastic constants

It has already been demonstrated for one material (compound **4**) that excellent qualitative and good quantitative agreement (within 2 pN) is found between the experimentally determined splay and bend elastic constants and calculations carried out *via* simulation and atomistic modelling.<sup>6</sup> Indeed, Fig. 6(a) shows the result of calculations of all three elastic constants of compound **4** as a function of the order parameter  $S_{zz}$ , with  $z$  being the axis passing through the carbon atoms of the oxadiazole ring. In agreement with experiment we find  $K_{22} < K_{33} < K_{11}$ , with very small  $K_{22}$ . Remarkably, the trend of the calculated elastic constants as a function of the order parameter is also in agreement with experiment:  $K_{11}$  and  $K_{22}$  increase with increasing order, whereas  $K_{33}$  first increases and then decreases at high ordering. In the theoretical expressions for the elastic constants<sup>25</sup> there is an implicit dependence on the molecular



**Fig. 6** (a) The elastic constants obtained by atomistic calculations for compound **4**, as a function of the order parameter ( $S_{zz}$ ):  $K_{11}$  (solid line),  $K_{22}$  (dash-dotted line) and  $K_{33}$  (dashed line). (b) Ratio of elastic constants ( $K_{ii}$ ) to the squared order parameter ( $S_{zz}$ ), as obtained from atomistic calculations for compound **4**. Dashed lines are used for points beyond the experimental nematic region.  $S_{zz}$  is the order parameter for the axis passing through the carbon atoms of the oxadiazole ring.



order parameter, which involves higher order terms beyond those proportional to  $S_{zz}^2$ . Fig. 6(b) shows the ratios  $K_{ii}/S_{zz}^2$ , for illustrative purposes a wide range of order parameters is considered, including values beyond the experimental ones. It may be worth pointing out that the branches of the curves in Fig. 6(b) and 5(b) at high ordering have a different meaning: in the former case they represent the elastic constants of a hypothetical phase with nematic order extending to very high ordering, whereas in the latter case the fitting curves are meant to reproduce the whole experimental behaviour, including extra-nematic effects. The results reported in Fig. 6(b) show that the splay and twist elastic constants calculated for the nematic phase of compound 4 are not strictly proportional to  $S_{zz}^2$ , but the ratios  $K_{11}/S_{zz}^2$  and  $K_{22}/S_{zz}^2$  exhibit a relatively small increase with increasing order. On the contrary, a strong dependence on the order parameter is predicted for the ratio  $K_{33}/S_{zz}^2$ , which would diverge to negative values at very high ordering. This trend agrees with the behaviour shown in Fig. 5(b).

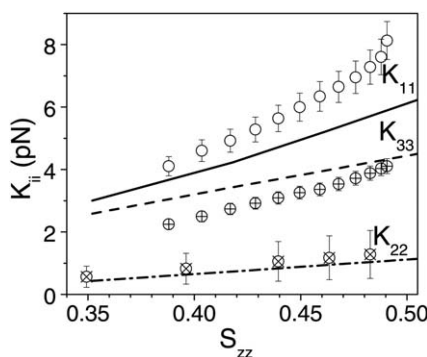
Calculations of the elastic constants were also carried out for compounds 2 and 3. Both differ from compound 4 for the replacement of the pentyl with a longer nonyloxy chain. In compound 2 there are also two additional F atoms in the phenyl ring attached to the nonyloxy chain. In the framework of the molecular field model used here, the F atoms are expected to affect the elastic constants if they can induce a change of the conformational preferences, thus a change of the average molecular shape. Quantum chemical calculations show that the main effects of the F atoms are those of lowering the energy barriers opposing the rotation of the CO–O–Car–C<sub>ar</sub> bond and stabilizing the conformations having both the CO group and the O–CH<sub>2</sub> bond pointing in the same direction, opposite to the F atoms. We have found that neither these variations, nor those deriving from the change of the lateral chain significantly affect the average molecular shape: the plots of the elastic constants as a function of the  $S_{zz}$  order parameter calculated for compounds 2 and 3 are very similar to each other and also not very different from that shown in Fig. 6(a) for compound 4. The

small influence of changes in the lateral arms on the elastic constants is in line with the experimental findings for compounds 1–4. Fig. 7 explicitly shows excellent agreement (within 2 pN) between the experimental and calculated elastic constants for compound 2. It is to be noted that though the calculations do not point to any difference in the elastic constants for compound 4 from the rest, the experimental values are slightly lower for compound 4 than the other compounds which could most probably be due to the pentyl/nonyloxy chain difference. Previous calculations for single molecular conformers pointed to the strong sensitivity of elastic constants of bent-core nematics to the conformation of the lateral chains.<sup>6</sup> This is not in contradiction to the negligible effect of the terminal chains in compounds 1–4, evidenced here both by experiments and calculations, since the internal flexibility leads to an average molecular shape where the effects of single conformations are washed out.

## 4 Conclusions

This paper describes the temperature dependence of the splay, twist and bend elastic constants along with the dielectric anisotropy, birefringence and order parameter measurements in four related bent-core oxadiazole compounds. The birefringence,  $\Delta n$  as well as the order parameter,  $S$  for all compounds is found to be independent of the change in molecular length or the presence of fluoro-substituents at the outer phenylene group of the aromatic core structure. Excellent agreement is found for the order parameter values derived from Haller method with those reported from PRS<sup>16</sup> in all the compounds. The dielectric anisotropy is, however, found to depend strongly on the molecular structure, with the presence of fluoro-substituents at the outer phenylene group of the aromatic core structure (compound 2) resulting in the highest magnitude of dielectric anisotropy while the pentyl-substituted (compound 4) shows the smallest magnitude.

All of the bent-core compounds studied show  $K_{22} < K_{33} < K_{11}$  analogous to other bent-core compounds with comparable bend angles. The dependence of the elastic constants on temperature is found to be indistinguishable for compounds 1, 2 and 3, all of which have alkoxy terminal chain substitution. The temperature variation of the elastic constants is much less marked in compound 4 than the other materials. A third-order dependence of the elastic constants on order parameter was found to give excellent fits to the experimental data for compounds 2 and 4, but was not used to analyse the elastic behaviour of the other two compounds, both of which showed an unusual but reproducible deterioration in alignment on approaching the underlying higher order phase. Extrapolation of the elastic behaviour was carried out using the fitting parameters obtained for compounds 2 and 4. The insight offered by such an analysis is consistent with what is known about the structure of the DC phase and role of the saddle-splay constant,  $K_{24}$  in its formation. It is possible that the bend elastic constant may also tend to diverge to negative values on approaching the DC phase, a suggestion that merits further experimental study. We have demonstrated that molecular-field theory and atomistic modelling can be used to calculate the



**Fig. 7** The experimentally deduced splay,  $K_{11}$  (open circles), twist,  $K_{22}$  (circles with a cross) and bend,  $K_{33}$  (circles with a plus sign) elastic constants as a function of order parameter,  $S$  for compound 2. Note that the error bars in this figure include the absolute uncertainty in the measurement which is most significant for measurements of  $K_{22}$ . The elastic constants calculated by atomistic simulations for compound 2 are denoted by  $K_{11}$  (solid line),  $K_{22}$  (dash-dotted line) and  $K_{33}$  (dashed line).

temperature dependence of the elastic constants for the materials. The calculations show remarkable agreement with experimentally determined values. Experiments and computation agree in finding that the elastic constants of compounds **1–4** are only weakly affected by changes in the terminal chains. The differences between compound **4** and the other compounds, detected experimentally, are probably beyond the predictive capability of the atomistic calculations. Both approaches show that there is no effect of the presence of fluoro-substituents on the outer phenylene group of the aromatic core structure (compound **2**) on the elasticity of the LC materials.

Both the analysis of experimental data and theoretical predictions agree in showing that the elastic constants of bent-core NLCs exhibit significant deviations from the square dependence on the order parameter, which is generally assumed for calamitic NLCs.

In summary, the combination of careful experimentation with calculations has begun to reveal some of the relationships between the molecular structure and the physical properties of nematic phases formed from bent-core molecules. Further, this work has offered a tantalising glimpse into the behaviour of the elastic constants on approaching an underlying DC phase.

## Acknowledgements

This work was supported by the Engineering and Physical Sciences Research Council (EPSRC) under Project no. EP/G023093/1. CG acknowledges the PhD School of Materials Science and Engineering (University of Padova) and Merck Chemicals Ltd. for financial support. HL is grateful to Merck Chemicals Ltd. and the EPSRC for a CASE Award. VG gratefully acknowledges the Royal Society for funding through a Dorothy Hodgkin Fellowship.

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