The nematic liquid crystal (NLC) phase is the least ordered but most commonly used liquid crystal state, forming the basis of almost all commercial liquid crystal devices. Nematic liquid crystals are characterized by long-range orientational order, and the combination of their fluidity, optical anisotropy, and responsiveness to electric fields makes them attractive electo-optic materials. Most nematic phases are formed from either rodlike (calamitic) or disklike (discotic) molecules, but recently, systems formed by molecules with a bent-core structure have emerged as an exciting new class of nematogens. Indeed, most of the studies focus on mixtures of calamitic and bent-core carboxylate molecules and has been explained through the mutual alignment of both types of molecules [16]. However, the presence of a biaxial nematic phase was questioned by Vanarakas et al., suggesting that the phase could consist of uniaxially distributed biaxial clusters [11]. Fracescangeli et al. further explained the splitting of small angle x-ray diffraction patterns, taken to be a signature of a biaxial nematic phase, as the presence of skewed smectic fluctuations (cybotactic groups) [12].

The behavior and performance of NLC devices is dominated by the viscoelastic properties of the material; the elastic constants influence the threshold voltage for switching, whereas, the elastic constant ratio dictates the steepness of the electro-optic response and, therefore, limits to the complexity of the device. More fundamentally, there is considerable interest in studying the behavior of elastic constants because of the insight they offer in areas, such as testing continuum theory, understanding director distortion in devices, and providing insight into the microscopic structure of the ordered state and order parameter fluctuations [13]. The behavior of the elastic constants associated with the splay $K_{11}$, twist $K_{22}$, and bend $K_{33}$ deformations of the director in calamitic liquid crystals is well understood, although interestingly, it has emerged over the past decade that materials that include bent-core nematics are atypical [14,15]. Dodge et al. showed that the bend elastic constant (measured both by Freedrickz transition and light scattering experiments) reduced to half the original value in a mixture of calamitic and bent-core benzoate molecules [15]. Anomalous temperature dependence of $K_{11}$ and $K_{33}$ has been observed in mixtures of calamitic and bent-core carboxylate molecules and has been explained through the mutual alignment of both types of molecules [16]. Indeed, most of the studies focus on mixtures of calamitic and bent-core structures [14–18] with just a few that consider pure bent-core nematic systems [4,19,20]. The few reports of the elastic properties of purely bent-core nematic materials to date find that $K_{11} > K_{33}$ (the converse is true for calamitic nematic materials). Sathyanarayana et al. demonstrate that the temperature dependence of the elastic constants in a bent-core material is very different from that observed in calamitic systems and relate their findings to a coupling of the bent molecular shape with the bend distortions of the...
unusual bent-core nematic systems [23]. This paper offers
phases, predictions that are now being applied to a few
resulting in the occurrence of new lower symmetry nematic
chains. Importantly, we have carried out molecular-field atomistic calculations of the elastic constants
reported to date where the underlying phase is not a tilted
chain, and we note that the nature of the low-temperature nematic
regime in C5-Ph-ODBP-Ph-OC12 is, therefore, still an open
question.
C5-Ph-ODBP-Ph-OC12 exhibits negative dielectric anisotropy $\Delta \varepsilon$ across the entire nematic phase regime [7],
and as a consequence, minor modifications are required for both the experimental geometry and the fitting programs
that are more commonly used to measure elastic constants in materials with positive $\Delta \varepsilon$. We, therefore, checked
our method using the negative calamitic nematic material $N$-p-methoxybenzylidene-p-butylaniline (MBBA), which has
phase transitions: isotropic 42 °C N, Fig. 1(b).
All measurements employ devices with glass substrates
and a liquid crystal layer ~5 μm thick. The transparent
(Indium-tin-oxide) electrode area is 30 mm$^2$ (20 Ω□), 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 for C5-Ph-ODBP-Ph-OC12
and at ~30 °C for MBBA. Either planar or homeotropic
alignment was required depending on the experiment carried
out; the former was achieved using a high-temperature (SE130)
polyimide layer, and the latter can be obtained using trichloro-
cta-decyl silane in heptane. Excellent monodomain alignment
was confirmed by use of polarizing microscopy, although we
note that there was no indication of any biaxial nematic regime
via conoscopy in the homeotropic device. The devices were
held in a Linkam THMS600 hot stage equipped with a TMS 93
controller, which allowed temperature control of the samples
with a relative accuracy of ±0.1 °C.
The following subsection presents, in detail, the techniques
employed in the measurement of birefringence and elastic
constants followed by the calculation details of atomistic
modeling.

A. Birefringence measurements
The birefringence measurements are undertaken via an
analysis of the reflection spectra from planar aligned devices
using the Berreman methodology [26]. This technique is
particularly useful for accessing refractive indices at high
temperatures where Abbé refractometers cannot be used. The
details of both the apparatus used and the fitting procedure
are given in Ref. [27]; the fitting program implements an
optical model for the propagation of electromagnetic waves in
media consisting of isotropic and anisotropic layers. Reflection
spectra are first taken for the empty device, allowing the
thickness to be measured with an accuracy of ±0.02 μm,
before filling at 200 °C. The ordinary $n_e$, and extraordinary
$n_o$ refractive indices are determined from spectra obtained
with the nematic director oriented perpendicular and parallel

II. EXPERIMENTAL AND COMPUTATIONAL METHODS
The bent-core mesogen studied is derived from a 1,3,4-
oxadiazole-biphenyl core (ODBP) and is shown in Fig. 1(a).
The phase transitions of C5-Ph-ODBP-Ph-OC12 were mea-
sured to be isotropic 239.6 °C N 184.6 °C DC, where N is
the nematic phase and DC is the dark conglomerate phase;
although the absolute values are higher than those quoted
elsewhere [6,24], the nematic range is the same, and the
difference is 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 $T_{NI}$. This material is particularly interesting as
studies using polarized Raman spectroscopy and small-angle
x-ray scattering [24] and electroconvection [6,7] have indi-
cated anomalies in the low-temperature region of the nematic
phase. Specifically, below $T - T_{NI} \sim -30$ K, nonzero
biaxial order parameters have been measured, and changes
in the nature of electroconvection are also apparent. The x-ray
FIG. 1. Chemical structures of (a) C5-Ph-ODBP-Ph-OC12 and
(b) MBBA.

data [24] show clear lobes in the small-angle scattering pattern.
However, the angular peak separation of the lobes (~70°) is
temperature independent, and the perpendicular correlation
length is only ~1 molecular length. The x-ray data [24] are,
therefore, not consistent with the occurrence of cybotactic
clusters [25]. Consequently, although the available data for C5-
Ph-ODBP-Ph-OC12 do not confirm phase biaxiality, neither
are the x-ray data consistent with a cybotactic nematic phase,
and we note that the nature of the low-temperature nematic
regime in C5-Ph-ODBP-Ph-OC12 is, therefore, still an open
question.

director. Majumdar et al., however, suggest that nanoscale
molecular clusters with tilted smectic correlations cause the
softening of $K_{22}$ and $K_{33}$ relative to $K_{11}$, Salter et al. [21]
also show that $K_{11} > K_{33}$ and find an unusually low ratio of
$K_{33}/K_{11} \sim 0.5$ in a bent-core nematic material, although they
find no anomalous flexoelectric behavior in contrast to Ref. [1].
It is clear that bent-core NLCs exhibit elastic properties that
appear to be not readily explained by models developed
for calamitic systems and that their unusual properties offer
the potential for novel electro-optic devices. Indeed, Dozov
[22] predicted pathological elasticity in bent-core materials
resulting in the occurrence of new lower symmetry nematic
phases, predictions that are now being applied to a few
unusual bent-core nematic systems [23]. This paper offers
importance contributions to the understanding of elasticity in
bent-core nematic systems as it reports the splay and bend
elastic constants across the entire nematic range (DC) phase.
Importantly, we have carried out molecular-field atomistic calculations of the elastic constants
for the molecule studied, and we compare these calculations
to the experimental data.

All measurements employ devices with glass substrates
and a liquid crystal layer ~5 μm thick. The transparent
(Indium-tin-oxide) electrode area is 30 mm$^2$ (20 Ω□), 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 for C5-Ph-ODBP-Ph-OC12
and at ~30 °C for MBBA. Either planar or homeotropic
alignment was required depending on the experiment carried
out; the former was achieved using a high-temperature (SE130)
polyimide layer, and the latter can be obtained using trichloro-
cta-decyl silane in heptane. Excellent monodomain alignment
was confirmed by use of polarizing microscopy, although we
note that there was no indication of any biaxial nematic regime
via conoscopy in the homeotropic device. The devices were
held in a Linkam THMS600 hot stage equipped with a TMS 93
controller, which allowed temperature control of the samples
with a relative accuracy of ±0.1 °C.

The following subsection presents, in detail, the techniques
employed in the measurement of birefringence and elastic
constants followed by the calculation details of atomistic
modeling.

A. Birefringence measurements
The birefringence measurements are undertaken via an
analysis of the reflection spectra from planar aligned devices
using the Berreman methodology [26]. This technique is
particularly useful for accessing refractive indices at high
temperatures where Abbé refractometers cannot be used. The
details of both the apparatus used and the fitting procedure
are given in Ref. [27]; the fitting program implements an
optical model for the propagation of electromagnetic waves in
media consisting of isotropic and anisotropic layers. Reflection
spectra are first taken for the empty device, allowing the
thickness to be measured with an accuracy of ±0.02 μm,
before filling at 200 °C. The ordinary $n_e$, and extraordinary
$n_o$ refractive indices are determined from spectra obtained
with the nematic director oriented perpendicular and parallel

FIG. 1. Chemical structures of (a) C5-Ph-ODBP-Ph-OC12 and
(b) MBBA.

S. KAUR et al.

PHYSICAL REVIEW E 86, 041703 (2012)

041703-2
to the incident light polarization, respectively. The average refractive index \( \bar{n} = \sqrt{(2n_x^2 + n_y^2)/3} \) and the birefringence \( \Delta n = n_x - n_o \) may then be calculated with an accuracy of \( \pm 0.001 \). All values of refractive index are quoted for light of wavelength 589 nm.

**B. Measurement of the elastic constants \( K_{11} \) and \( K_{33} \)**

The elastic constants are determined from the variation in the material dielectric constant through the electric-field induced Freedrickz transition. Homeotropic alignment is required since C5-Ph-ODBP-Ph-OC12 has negative \( \Delta \varepsilon \) across the whole nematic regime, and in this geometry, the director experiences an electric torque on the application of the electric field perpendicular to the plane of the device. Obtaining homeotropic alignment for bent-core nematic materials is a particular challenge, although we have obtained excellent results for this material using trichloro-octa-decyl silane in heptane. The apparatus includes an Agilent Precision LCR Meter E4980A, which measures the dielectric permittivity across the frequency range 20 Hz–2 MHz using a 0.05 V_ac probe voltage where necessary in the presence of an ac electric field of up to 20 V. The parallel component of the dielectric permittivity \( \varepsilon || \) is observed at fields below the Freedrickz threshold, whereas, at increasing field strength, above the transition, the director begins to align perpendicular to the field, thus, allowing the perpendicular component of the dielectric permittivity \( \varepsilon _\perp \) to be deduced. The bend elastic modulus \( K_{33} \) is directly related to the threshold voltage \( V_{th} \) by

\[
V_{th} = \pi (K_{33}/\varepsilon_0 \Delta \varepsilon)^{1/2}.
\]

The splay elastic constant \( K_{11} \) influences the steepness of the variation in capacitance above the threshold and can be deduced using a computer fitting program, which employs the following equation:

\[
\frac{V}{V_{th}} = 2\pi^{-1}(1 + \gamma \sin^2 \theta_m)^{1/2} \int_0^{\pi/2} \left( \frac{F_k}{F_x F_y} \right)^{1/2} d\theta,
\]

where \( F_k(\theta) = 1 + k \sin^2 \theta_m, F_x(\theta) = 1 - \sin^2 \theta_m, F_y(\theta) = 1 + \gamma \sin^2 \theta_m, k = (K_{11}/K_{33}) - 1, \gamma = (\varepsilon \perp/\varepsilon ||) - 1, \) and \( \theta_m \) is the director angle in the center of the liquid crystal layer [28]. This methodology allows the elastic constants to be deduced with an accuracy of around 5%. We note that no account has been taken of any possible contribution due to flexoelectricity in the sample.

**C. Calculation of elastic constants and definition of the molecular geometry**

The elastic constants of C5-Ph-ODBP-Ph-OC12 were calculated assuming a uniaxial nematic phase, according to the surface interaction model, wherein, the orientational mean field experienced by a molecule in the nematic phase \( U^{\sigma \sigma} \) is parametrized according to the anisotropy of the molecular surface. It is expressed as

\[
U^{\sigma \sigma}(\Omega) = \varepsilon k_B T \int_S dS P_2(n \cdot s),
\]

where \( \Omega \) are the Euler angles, which define the molecular orientation, \( k_B \) is the Boltzmann constant, \( T \) is the temperature, \( P_2 \) is the second Legendre polynomial, \( n \) and \( s \) are unit vectors parallel to the director and perpendicular to the surface element \( dS \), respectively. The integral is over the molecular surface \( S \), and \( \varepsilon \) is a parameter that specifies the orienting strength. It is related to the reduced temperature and the molecular order parameters. In this framework, molecular expressions for the elastic constants are obtained as linear combinations of Cartesian tensor components, each defined as an integral over the molecular surface [29].

C5-Ph-ODBP-Ph-OC12 is a highly flexible molecule, and the prediction of the elastic constants and orientational order parameters requires calculation of an average over all the conformational degrees of freedom of the core and the lateral wings. The rotational isomeric state (RIS) approximation was used [30], according to which, only the conformations corresponding to the minima of the torsional potential (the so called conformers) are considered. RIS conformers were collected by Monte Carlo (MC) sampling. The bonds of C5-Ph-ODBP-Ph-OC12 that were allowed to rotate are shown in Fig. 2; the corresponding dihedrals are denoted as \( \chi_i \) (\( i = 0 - 15 \)). At each MC move, a random number of bonds was allowed to jump from one to another state, and then, the new conformer was accepted on the basis of its torsional energy according to the Metropolis criterion [31].

In our calculations, planar geometry of the central three ring moiety was assumed in agreement with the results of density-functional theory calculations for 2,5-diphenyl-1,3,4-oxadiazole reported in Ref. [32]; the torsional potential reported there for the bond between phenyl and oxadiazole exhibits two minima for planar structures, separated by barriers of about 30 kJ/mol corresponding to the perpendicular arrangement. Planar or nearly planar geometry of 2,5-diphenyl-1,3,4-oxadiazole was also found in crystals [33]. The geometry choices for the phenyl benzoate groups are less obvious since the potential energy surface of this group has been the object of several investigations with some differences in the results [34–36]. There is general agreement about the preference for the planar geometry of the benzoate unit due to the presence of high energy barriers, which oppose the rotation of the bond between the aromatic carbon (\( C_{ar} \)) and the COO moiety (\( C_{ar}^{-}\text{COO} \)). Thus, in our MC calculations, the benzoate unit was frozen in the planar geometry. Some discrepancies exist in the literature on the torsional potential for the \( C_{ar}^{-}\text{ar}^{-}\text{O-CO} \) dihedral. Energy barriers at 0° and 180° (planar arrangement) are found, ranging from about 3 kJ mol\(^{-1} \) to more than 10 kJ mol\(^{-1} \). A smaller, in several cases, even insignificant barrier is found for the perpendicular arrangement of the phenyl rings, whose height strongly depends on the level of the theory. Thus,
despite some uncertainty about the details, all calculations point to a wide distribution of the C\textsubscript{ar}-C\textsubscript{ar}-O-CO dihedral in the range of \(\pm (30^\circ - 150^\circ)\). For the sake of simplicity, in our calculations, we assumed for \(\chi\)\textsubscript{16} and for \(\chi\)\textsubscript{17} two values \(\pm 90^\circ\) in the middle of their highest probability region. The torsional potential for the C\textsubscript{ar}-CH\textsubscript{2} bond exhibits two equivalent minima, corresponding to the geometry with the first CH\textsubscript{2}-CH\textsubscript{2} bond of the alkyl chain lying on a plane perpendicular to the aromatic plane [37]; therefore, we assumed two equivalent states with \(\chi\) equal to \(\pm 90^\circ\). Also, the torsional potential for the C\textsubscript{ar}-O bond has two equivalent minima, but in this case, they correspond to the O-CH\textsubscript{2} bond lying on the same plane of the aromatic ring [37]; the values \(\chi\)\textsubscript{4} = 0\(^\circ\) and \(\chi\)\textsubscript{4} = 180\(^\circ\) were used in our calculations. For the CH\textsubscript{2}-CH\textsubscript{2} and the O-CH\textsubscript{2} bonds in the lateral wings, three states were taken: the trans conformation (dihedral equal to 180\(^\circ\)) and the gauche +, gauche - pair (\(\pm 65^\circ\)). These three states are not energetically equivalent, and this was taken into account for sampling the conformer space according to the Metropolis criterion. The torsional energy of the J conformer was calculated as

\[
V^J = \sum_{i=0,2,5,15} \left( \delta^J_{i,g} + \delta^J_{i,g'} \right) \Delta V_i + \sum_{i=0,1,5,14} \left( \delta^J_{i,g} + \delta^J_{i+1,g} + \delta^J_{i,g} - \delta^J_{i+1,g} \right) \Delta V_{i,i+1},
\]

where the index \(i\) indicates the bond and \(\delta\) is the Kronecker symbol (\(\delta^J_{i,g} = 1\) if, in the \(J\) conformer, the \(i\) bond is in the \(g\) state). The first sum accounts for the energy difference between the gauche and the trans states; \(\Delta V_i = 2.2\) kJ mol\(^{-1}\) was assumed for all the chain dihedrals with the exception of \(\chi_0\) for which \(\Delta V_0 = -2.85\) kJ mol\(^{-1}\) was used [37]. The second sum in Eq. (4) introduces the energy stabilization associated with the presence of adjacent gauche + gauche + (or gauche - gauche -) pairs, and \(\Delta V_{i,i+1} = -0.75\) kJ mol\(^{-1}\) was assumed [37]. In sampling the conformers of C5-Ph-ODBP-Ph-OC12, structures having pairs of atoms closer than a cutoff distance equal to 0.82\(\sigma\), where \(\sigma\) is the sum of their van der Waals radii, were discarded. Van der Waals radii for C\textsubscript{ar} were taken from [38] and the same van der Waals radii used for this purpose.

III. RESULTS

A. Birefringence and order parameter of C5-Ph-ODBP-Ph-OC12

Figure 3 shows the variation in birefringence in the nematic phase of C5-Ph-ODBP-Ph-OC12 as a function of reduced temperature. The birefringence increases monotonically with decreasing temperature as expected and as observed for calamitic systems. The inset to Fig. 3 shows explicitly the ordinary, extraordinary, and average refractive indices in the nematic phase as well as the isotropic refractive index. The average refractive index is slightly lower than the isotropic phase value and, although a similarly small difference is often observed in calamitic materials, it is more common for the isotropic refractive index to take lower values than the mean value. The only comparable data published for bent-core liquid crystals is for an oxazole-derived system measured at infrared wavelengths (1.5 \(\mu\)m) where there is no difference reported between the mean value of the refractive index in the nematic phase and the isotropic value [40].

The birefringence of C5-Ph-ODBP-Ph-OC12 varies from \(\sim -0.1\) to \(\sim 0.18\) across the temperature range and is around a factor of 2 higher (near \(T_{NI}\)) than that reported by Sathyarayana et al. for a different bent-core material [19]. In the case of mixtures of rodlike and bent-core molecules, \(\Delta n\) has generally been found to decrease with the increasing concentration of bent-core molecules [16,18]. The orientational order parameter \(S\) can be deduced from the refractive index data following the well-known method of Haller [41]:

\[
S = \left( \frac{n^2 - n^2_o}{n^2 - 1} \right) \left\{ \frac{\tilde{\alpha}}{\Delta \alpha} \right\},
\]

where \(\tilde{n}^2 = \frac{1}{2} (n^2_e + 2n^2_o), \tilde{\alpha} = \frac{1}{3} (\alpha_{||} + 2\alpha_{\perp}), \) and \(\Delta \alpha = \alpha_{||} - \alpha_{\perp}.\) The \(\alpha_{||}\) and \(\alpha_{\perp}\) give the polarizability of individual molecules [42]. The temperature dependence of \(S\) can be written as

\[
S = \left( 1 - \frac{T}{T^*} \right)^\beta,
\]

where \(T^*\) and \(\beta\) are adjustable fitting parameters and \(T\) is the temperature measured in Kelvin. The fitting parameters \([T^* = 512.82\) K, \((\tilde{\alpha}/\Delta \alpha) = 1.89, \) and \(\beta = 0.19\)] are deduced by equating Eqs. (5) and (6). Although our value of \(\beta\) is much less than 0.5, which is predicted by mean-field theory, it is in good agreement with measurements of many other systems; Haller observed 13 different liquid crystals that exhibit \(\beta \sim 0.2 \pm 0.03,\) whereas, recent papers by Sathyarayana et al. and Kundu et al. find \(\beta \sim 0.2\) for both calamitic materials and mixtures of calamitic and bent-core materials [18,19,43]. Figure 4 shows the order parameter as a function
of reduced temperature deduced from refractive index data together with the value obtained for the same material from Raman spectroscopy in which $P_{200}$ is equivalent to $S$ [24]. There is excellent agreement between these order parameter measurements as well as with those made on a different bent-core nematic material via NMR by Dong [44], Dong et al. [45], and Weissflog et al. [46]. Order parameters also deduced from refractive index data for a different material [19] are slightly lower than those in Fig. 4 but exhibit a similar temperature dependence.

**B. Elastic constants of C5-Ph-ODBP-Ph-OC12 determined experimentally**

First, the elastic constants of the calamitic liquid crystal MBBA are given in Table I. These data are within 5% of those reported elsewhere [47], validating our methodology. The dielectric anisotropy of C5-Ph-ODBP-Ph-OC12 is found to decrease from $-1.4$ at $T - T_{NI} = -8.7$ K to $-3.2$ at $T - T_{NI} = -56.7$ K. Figure 5 shows the splay $K_{11}$ and bend $K_{33}$ elastic constants of C5-Ph-ODBP-Ph-OC12 as a function of reduced temperature. $K_{11}$ increases monotonically with decreasing temperature, more than doubling over the temperature range, whereas, $K_{33}$ is almost temperature independent. Examining the bend elastic constant in more detail, it can be seen to increase slightly as the temperature decreases until below a reduced temperature of $T - T_{NI} \sim -45$ K, a very small decrease is suggested by the data. It is clear that there is no evidence of pretransitional divergence of either $K_{11}$ or $K_{33}$. Both Sathyanarayana et al. [19] and Tadapatri et al. [20] report elastic constants in bent-core materials with underlying SmC phases, both also describing an approximately linear temperature dependence of $K_{11}$, varying between $\sim 2$–$17$ and $\sim 2$–$28$ pN, respectively. The behavior of $K_{33}$ appears to be quite different in each case, being almost temperature independent here, increasing continuously from $\sim 1$ to 5 pN in Ref. [20], and showing a strong pretransitional divergence in Ref. [19]. The inset of Fig. 5 shows that the ratio of $K_{33}/K_{11}$ of C5-Ph-ODBP-Ph-OC12 decreases from 0.8 to 0.3 as the temperature decreases. Values as low as 0.52 have been observed in long-chain calamitic nematics [48], although it is more usual in calamitic systems that $K_{11} < K_{33}$.

Discussions of the temperature dependence of the elastic constants in bent-core nematic materials have employed a comparison of the data with the order parameter. As a simple interpretation of mean-field theory (discussed in more detail in the following section) predicts a dependence of the elastic constants on $S^2$, several authors have examined the relationship $K_{ij} \propto \Delta n^2$. In order to facilitate a comparison of our data with that in Refs. [19,20], Fig. 6 show plots of $K_{11}$ and $K_{33}$ (inset) as a function of birefringence $\Delta n$ and order parameter $S$. Using this approach, the data for $K_{11}$ appear to behave differently in the low- and high-temperature regimes, changing at $T - T_{NI} \sim -30$ K; there is approximately a factor of 2 difference in the exponents associated with each region, which are $\sim 2.5$ and 4.5 for the high- and low-temperature regimes, respectively (a single exponent does not fit the data over the whole nematic regime). Interestingly, the change in behavior occurs at approximately the same temperature where anomalous behavior has been reported for this compound via Raman spectroscopy and electroconvection measurements [7,24]. The anomaly in $K_{11}$ observed in the high- and low-temperature regimes is not clear from either the splay elastic constant or the order parameter data alone. In comparison, examining the relationship $K_{ij} \propto \Delta n^x$ has resulted in values of $x = 2.9$ for the $K_{11}$ data of Sathyanarayana et al. [19]

![FIG. 4. Order parameter $S$ as a function of reduced temperature $T - T_{NI}$ obtained from open squares: refractive index and open circles: Raman data.](image)

![FIG. 5. Open squares: splay $K_{11}$ and open circles: bend $K_{33}$ elastic constants as a function of reduced temperature $T - T_{NI}$ in the nematic phase. The inset shows the temperature variation in open triangles: $K_{33}/K_{11}$.](image)
and values of $x = 2.5$ for mixtures of rod and bent-core nematic materials [16]. The inset of Fig. 6 shows that the bend elastic constant $K_{33}$ exhibits an approximately linear dependence on $\Delta n$ until at $T - T_{NI} = -44.7 \ K (\Delta n \approx 0.174)$ and, thereafter, decreases. Analogous results are observed in high concentration mixtures of bent-core molecules in 5CB where $K_{33}$ increases with increasing $\Delta n$ and attains a maximum at a certain $\Delta n$ value before decreasing [18]. Kundu et al. also observed a decrease in $K_{33}$ with decreasing temperature in a $17M$ concentration of bent-core molecules in 8OCB [16].

IV. DISCUSSION

The behavior of the elastic moduli of C5-Ph-ODBP-Ph-OC12 can be summarized as follows:

(1) In common with other bent-core nematic materials $K_{11} > K_{33}$, differing from calamitic nematic systems.

(2) The values of both elastic constants are comparable to but slightly lower than in other bent-core systems.

(3) The birefringence takes values of around 0.18, and the order parameters deduced from both refractive index data and Raman scattering give values between 0.35 and 0.70, comparing well with order parameter data for other bent-core nematic materials.

(4) The temperature dependence of $K_{11}$ is analogous to other bent-core materials, increasing approximately linearly with reducing temperature. However, its relationship with the order parameter shows two distinct regimes with exponents of around 2.5 and 4.5 at high and low temperatures, respectively.

(5) The behavior of $K_{33}$ is quite distinct for this system; it is almost temperature independent, taking a maximum value of $\sim 3$ pN. There is some evidence of a decrease in $K_{33}$ at low temperatures in the nematic regime, but there is no obvious pretransitional divergence.

The elastic constants clearly behave differently from calamitic systems, and although there are some similarities with other bent-core systems, there are also differences. In the subsequent sections, we first consider the elastic behavior in the context of existing physical models followed by computational investigation of our system.

A. Examining existing physical models

We organize the discussion to consider first the absence of critical divergence on approaching the underlying phase as this is perhaps the simplest result to understand. We then consider the relationship of the elastic constants with the order parameter and, finally, the values of the elastic constants and their ratios.

According to de Gennes [49] and Chen and Lubensky [50], all three nematic elastic constants diverge at the SmC-N phase transition. Findon and Gleeson [51] report that, for a calamitic nematic mixture, $K_{11}$ increases continuously from $\sim 4 \ pN$ close to the nematic to isotropic transition to $\sim 30 \ pN$ close to the SmC transition. Although there is some evidence of the variation becoming slightly more rapid close to the transition, there is no clear pretransitional behavior, and $K_{11}$ is continuous across the transition into the SmC phase where it takes values ranging from $\sim 30$ to $100 \ pN$. The temperature dependence of $K_{11}$ reported here and by others for different bent-core materials [19–21] is in broad agreement, and it increases approximately linearly across the nematic phase range as the temperature reduces. On the other hand, Sathyaranayana et al. report a strong pretransitional divergence in $K_{33}$, observed neither by us nor in Refs. [20,21]. In both cases (Refs. [19] and [20]), the same general behavior might be expected as the underlying phase is SmC. However, in C5-Ph-ODBP-Ph-OC12, there is a first-order transition to an underlying DC phase, and although the elastic behavior in that phase has not yet been considered either theoretically or experimentally, a first-order transition normally severely curtails pretransitional phenomena. Therefore, the lack of pretransitional divergence in the elastic constants reported here is as expected. Similar behavior might also be expected for the material in Ref. [21] as the underlying phase is a crystal phase, but measurements are reported only for temperatures $7 \ K$ below the isotropic phase transition, well away from the nematic to crystal transition.

We next consider the theoretical relationship between elastic constants and order parameter. Nehring and Saupe [52], Saupe [53], and Khoo and Wu [54] showed that a mean-field approach suggests that $K_{ii}$ are related to the order parameter $S$ by

$$K_{ii} = C_{ii} V_n^{-7/3} S^2,$$

where $C_{ii} = (3\gamma_2/L)(L m^{-1} \gamma_2)^{1/3}$ and $C_{ii}$ are reduced elastic constants that depend on molecular properties; $V_n$ is the molar volume, $L$ is the molecular length, $m$ is the number of molecules in a steric unit, $A^0 = 1.3 \times 10^{-8} \ erg \ cm^6$, and $\gamma_2 = \gamma_2 = x/z$ and $\gamma_2 \gamma_3 = (x/z)^2$, where $x$, $y$, and $z$ are the average molecular lengths in directions across and along the molecular long axis. For calamitic materials with small alkyl chains, $C_{ii}$ are found to be temperature independent, but as the chain length increases, $C_{ii}$ have considerable temperature dependence, and to account for the decrease in $K_{33}/K_{11}$ with increasing chain length (discussed further below), the increase in $C_{11}$ is found to be more than for
one molecule long. Given this and the nature of our underlying phase transition, we suggest that an explanation other than clustering must be sought to explain the elastic behavior of this and potentially other bent-core nematic systems.

The second-order dependence of the elastic constants on the order parameter [Eq. (7)] follows the arguments presented by other authors who have studied elastic constants in bent-core nematic systems, and it is tempting to suggest that the anomalous behavior observed in the exponents (Fig. 6) in the two different temperature regimes could suggest the emergence of biaxiality. However, a more detailed and less commonly used mean-field treatment of the elastic constants by Berreman and Meiboom [57] indicates that the correct expression to use includes a third-order dependence given by

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

where \( K_i^{(2)-(4)} \) are fitting parameters (which could, in principle, be temperature dependent). The ratio \( K_{ii}/S^2 \) is shown as a function of order parameter \( S \) in Fig. 7(b), and the solid lines show fits of Eq. (8) to our data assuming no temperature dependence in \( K_i^{(2)-(4)} \) (measured in pN), which are found to be \( K_i^{(2)} = 20.2 \pm 2.1, K_i^{(3)} = -19.2 \pm 4.6, K_i^{(4)} = 3.2 \pm 0.3, K_i^{(2)} = 14.4 \pm 1.5, K_i^{(3)} = -7.5 \pm 3.3, \) and \( K_i^{(4)} = -0.6 \pm 0.2 \). It is clear that excellent fits are obtained, although only if the condition that \( K_{11} \) and \( K_{33} \) converge at \( S = 0 [57] \) is relaxed.

We next consider the ratio \( C_{33}/K_{11} \) in our bent-core nematic liquid crystal. It has been shown that there is a correlation between \( C_{33}/K_{11} \) and \( L/W \) where \( L \) and \( W \) are the length and breadth of the molecule [48,58]. For short-chain calamitic nematic systems, this naturally leads to \( K_{33}/K_{11} > 1 \) as the molecular length is greater than the breadth. However, there are several systems where the ratio is modified. In the case of the homologous series of \( p,p'-\text{di}-n-\text{alkyl} \) and \( p,p'-\text{di}-n-\text{alkoxy} \) azobenzenes, both \( K_{11} \) and \( K_{33} \) become greater with increasing chain length, whereas, the ratio \( C_{33}/K_{11} \) decreases along the series [55] due to an increase in the number of conformations adopted by the flexible chain. In hockey-shaped molecules [59] as well as bent-core systems, the molecular shape and population of conformers can certainly lead to situations where \( W_{\text{eff}} > L \) (\( W_{\text{eff}} \) is the effective width of the molecule). Finally, there are several papers that demonstrate that the lower-temperature regimes of bent-core systems include large cybotactic SmC or SmA clusters, which can explain many of the unique features of bent-core nematic systems and which could also lead to a situation where \( W_{\text{eff}} > L \). Thus, there are several reasons why one might expect \( K_{11} > C_{33} \) in bent-core systems, so it is not surprising that we and others [16,19–21] report such a relationship. We note that, in the case of C5-Ph-ODBP-Ph-OC12, there is no x-ray evidence for pretransitional SmA or SmC clusters in the nematic regime [24], indeed, these would be unexpected given the first-order transition to a DC phase below the nematic phase.

In C5-Ph-ODBP-Ph-OC12, \( K_{11} > C_{33} \) is common with other bent-core nematic systems but contrasts the behavior found in most calamitic liquid crystals. Interestingly, almost 40 years ago, Gruler [60] and Helfrich [61] each presented
models that predicted a decrease in the bend elastic constant in the case of banana-shaped molecules. Helfrich employed phenomenological and molecular considerations to argue that

\[ K_{33} = \frac{\tilde{K}_{33}}{1 + \beta^2 N_0^{1/3} \left( \frac{\tilde{K}_{33}}{2k_BT} \right)}, \]

where \( \tilde{K}_{33} \) is the bend elastic modulus without molecular adjustment to the strain, \( K_{33} \) is the modulus adjusted for the bend structure, \( \beta \) is the molecular curvature (180° − \( \theta \) bend), and \( N_0 \) is the molecular density. The above formula does not incorporate an internal field correction, the elongated molecular shape, or the degree of order, which are taken into consideration by Gruler [60]. Gramsbergen and de Jeu reported a mere 5% adjustment of \( K_{33} \) for a bent-core thiadiazole compound using the above formula [62]. In C5-Ph-ODBP-Ph-OC12, \( K_{33} \) is \( \sim 3 \times 10^{-12} \) N (from Fig. 5), \( \beta = 0.7 \) rad, \( N_0 = 10^{27} \) m\(^{-3} \), \( k_BT \) is 655.78 J, \( T = 475 \) K, and we get \( \tilde{K}_{33} = 3.38 \times 10^{-12} \) N, which is again a rather small change (\( \sim 10\% \)). Indeed, it seems that this adjustment cannot account for any of the values of \( K_{33} \) reported; Table II summarizes the influence of this correction on selected materials.

Clearly, the formula suggested by Helfrich offers a much smaller shift in the bend elastic constant than would be needed to return to the situation for calamitic liquid crystals where \( K_{11} < K_{33} \). However, Gruler presented a molecular viewpoint of the elastic theory and obtained a much more significant effective change of 10 pN in \( K_{33} \) for bent molecules [60]. This suggests that the inclusion of intrinsic molecular deformations in the calculations for bend elastic constants is important. Indeed, in summarizing the discussion so far, it is clear that the elastic behavior of C5-Ph-ODBP-Ph-OC12 and other bent-core nematic systems is significantly different from that of calamitic systems, and we next consider whether a more sophisticated approach based on mean-field theory, which makes use of atomistic modeling is more suited to understanding these systems.

Recent calculations of this kind by Cestari and Ferrarini on typical mesogenic systems show that the high rigidity of para-azoxyanisole compared with 4-alkyl-4′-cyanobiphenyls (5 and 8CBs) in which contributions of various conformers (due to the slightly bent shape) are important, results in lower values of elastic constants [63]. The number of possible molecular conformers increases with the increase in length of the flexible chain, and the authors emphasized that the presence of a V-shaped structure has a marked effect in lowering the bend elastic constant. A recent case study on two mesogenic odd dimers with a certain degree of bend in the all-trans state conformation by Cestari et al. predicted a low bend elastic constant, which can even become negative at higher-order parameters [64] and could be a prerequisite to the twist-bend nematic phase [23]. Nearly a decade earlier, Dozov also proposed that the natural tendency of banana-shaped molecules to induce bend distortions could result in negative \( K_{33} \) values [22]. The next section presents the computational modeling of C5-Ph-ODBP-Ph-OC12 and demonstrates how the elastic constants behave in the case of a bent-core system.

### B. Calculation of the elastic constants of C5-Ph-ODBP-Ph-OC12

C5-Ph-ODBP-Ph-OC12 can assume markedly different shapes as a consequence of rotations around the bonds highlighted in Fig. 2. In our calculations, this was taken into account by averaging over the conformations of the lateral wings and of the aromatic core as explained in Sec. II.C. Based on considerations of the relevant torsional potentials, two conformations were assumed for the core, which differ from each other for having the carbonyl groups on opposite sides (I) and on the same side (II) of the oxadiazole ring as shown in Fig. 8(a). In these conformations, the angle between the \( para \) axes of the B and D phenyls is equal to 138°, and the angle between the A and E phenyls takes the same value. The calculated splay and bend elastic constants are reported in Fig. 8(b) as a function of the orientational order parameter \( S_{2z} \) (\( z \) is the axis passing through the carbon atoms of the oxadiazole ring). We can see that, in agreement with the experiment and in contrast to the usual behavior of rodlike calamitics, \( K_{11} > K_{33} \) is predicted. Also, in line with what is found experimentally, the calculated \( K_{11} \) steadily increases with increasing order (it increases slowly with ordering in

**TABLE II. Influence of the Helfrich correction on the values of the bend elastic constant \( K_{33} \) in selected bent-core nematic materials.**

<table>
<thead>
<tr>
<th>Material/reference</th>
<th>( \beta ) (rad)</th>
<th>( T ) (K)</th>
<th>( K_{33} ) (pN)</th>
<th>( \tilde{K}_{33} ) (pN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C5-Ph-ODBP-Ph-OC12</td>
<td>0.7</td>
<td>475.0</td>
<td>3.0</td>
<td>3.4</td>
</tr>
<tr>
<td>Reference [19]</td>
<td>0.82</td>
<td>419.5</td>
<td>4.0</td>
<td>5.2</td>
</tr>
<tr>
<td>Reference [4]</td>
<td>∼ 1.0</td>
<td>347</td>
<td>0.88</td>
<td>0.97</td>
</tr>
</tbody>
</table>

![Fig. 8](image-url)
other conformer, for a large molecule, such as C5-Ph-ODBP-Ph-OC12. For the oxadiazole ring.

order parameter for the axis passing through the carbon atoms of the two single conformers (shown in the insets) as a function of the

considered to explain the fact that $K_{11} > K_{33}$, but none could fully explain the behavior of splay and bend in the bent-core materials. The relationship between the elastic constants and the order parameter is found to be in excellent agreement with the equation suggested by Berreman and Meiboom [57], which includes third-order terms, provided that the condition of convergence of splay and bend at zero-order parameter is relaxed. Elastic constants were also determined from molecular-field theory and atomistic modeling, and notably good agreement was found with the experimental results. The value of $K_{11}$ is found to be almost independent of the molecular conformation, whereas, $K_{33}$ is highly geometry dependent, and in particular, it is extremely sensitive to changes in the molecular curvature. Importantly, the calculations show that what has been considered to be anomalous behavior of the elastic constants can, in fact, be predicted by molecular-field theories of nematics, provided that the bent molecular shape is suitably taken into account. An important distinctive feature evidenced by both our experimental results and the calculations is that $K_{33}$ can reduce to extremely low values well below the nematic to isotropic transition temperature, which is due to the difficulty in reconciling a bent molecular shape with a uniform director at high-order parameter values. This feature, which could be of interest for applications where low elastic constants are required, is also highly relevant to recent observations of splay-bend nematic phases in dimeric liquid crystals. Indeed, the low cost of bend deformations could well be important in the formation of domains of opposite chirality, recently observed in the nematic phase of bent-core bis-(phenyl)oxadiazole derivatives [66]. Finally, we note that this analysis of the splay and bend elastic constants of C5-Ph-ODBP-Ph-OC12 gives no insight into any possible biaxial nature of the nematic phase of this system, although it is clear that cybotactic clusters need not be considered to be present to explain the observed physical properties.

\[ K_{33} \text{ becomes huge and prevails. It was already shown that, even for conventional NLCs, the molecular curvature may have a strong influence on the bend elastic constant} \ [63]. \text{This effect is enhanced by the peculiar structure of bent-core NLCs. Other systems whose bend elasticity was found to be particularly sensitive to conformational changes are odd mesogenic dimers as mentioned earlier. For those systems, } K_{11} > K_{33} \text{ was again predicted, along with a nearly temperature independent } K_{33}, \text{ decreasing at lower temperatures} [64]. \text{Such theoretical predictions were subsequently confirmed by the experiment} [65]. \]

FIG. 9. (Color online) (a) and (b) Elastic constants calculated for the two single conformers (shown in the insets) as a function of the order parameter for the axis passing through the carbon atoms of the oxadiazole ring.

calamitic NLCs), whereas, $K_{33}$ remains nearly constant and tends to decrease at high ordering (it increases with order in calamitic NLCs). Both $K_{11}$ and $K_{33}$ are slightly higher than the experimental values, but strict quantitative agreement is probably beyond the present capability because of the detailed knowledge of the molecular geometry that is needed for this purpose.

To get an insight into the relationship between molecular structure and elastic constants, it can be useful to look at Figs. 9(a) and 9(b), which show the results of calculations for a couple of distinct conformers of C5-Ph-ODBP-Ph-OC12 (shown in the insets): They both have the core in the I configuration and the all-$trans$ chains and only differ in the value of the $\chi_3$ dihedral, which is equal to $+90^\circ$ in one case and to $-90^\circ$ in the other. We can see that the rotation around a single bond, changing the curvature of the molecular structure, has dramatic effects on the bend elastic constant. For the less bent conformer, a more conventional behavior is predicted, $K_{33} > K_{11}$ is found, and both elastic constants are large as expected for a large molecule, such as C5-Ph-ODBP-Ph-OC12. For the other conformer, $K_{11}$ remains large, but $K_{33}$ takes negative values, which would be incompatible with the existence of a uniform nematic phase because it implies a tendency for spontaneous bending of the director [22]. We can explain the unusual behavior of $K_{33}$ reported in Fig. 8(b) as the result of different contributions, which lead, on average, to small $K_{33}$ values with a weak temperature dependence: Elongated conformers, which would impart the nematic phase with high bending stiffness, are simultaneously present with bent conformers, which would promote bending of the mesophase director. At very high ordering, the negative contribution to $K_{33}$ becomes huge and prevails. It was already shown that, even for conventional NLCs, the molecular curvature may have a strong influence on the bend elastic constant [63]. This effect is enhanced by the peculiar structure of bent-core NLCs. Other systems whose bend elasticity was found to be particularly sensitive to conformational changes are odd mesogenic dimers as mentioned earlier. For those systems, $K_{11} > K_{33}$ was again predicted, along with a nearly temperature independent $K_{33}$, decreasing at lower temperatures [64]. Such theoretical predictions were subsequently confirmed by the experiment [65].

V. CONCLUSION

In conclusion, splay and bend elastic constants along with birefringence and order parameter measurements of a bent-core oxadiazole NLC have been reported. The order parameter derived from the refractive indices is in excellent agreement with that obtained from Raman spectroscopy. The splay elastic constant is found to be higher than the bend elastic constant in common with other bent-core systems. It is interesting to note that no pretransitional behavior in either $K_{11}$ or $K_{33}$ is observed close to the transition to the underlying DC phase. Various existing models have been considered to explain the fact that $K_{11} > K_{33}$, but none could fully explain the behavior of splay and bend in the bent-core materials. The relationship between the elastic constants and the order parameter is found to be in excellent agreement with the equation suggested by Berreman and Meiboom [57], which includes third-order terms, provided that the condition of convergence of splay and bend at zero-order parameter is relaxed. Elastic constants were also determined from molecular-field theory and atomistic modeling, and notably good agreement was found with the experimental results. The value of $K_{11}$ is found to be almost independent of the molecular conformation, whereas, $K_{33}$ is highly geometry dependent, and in particular, it is extremely sensitive to changes in the molecular curvature. Importantly, the calculations show that what has been considered to be anomalous behavior of the elastic constants can, in fact, be predicted by molecular-field theories of nematics, provided that the bent molecular shape is suitably taken into account. An important distinctive feature evidenced by both our experimental results and the calculations is that $K_{33}$ can reduce to extremely low values well below the nematic to isotropic transition temperature, which is due to the difficulty in reconciling a bent molecular shape with a uniform director at high-order parameter values. This feature, which could be of interest for applications where low elastic constants are required, is also highly relevant to recent observations of splay-bend nematic phases in dimeric liquid crystals. Indeed, the low cost of bend deformations could well be important in the formation of domains of opposite chirality, recently observed in the nematic phase of bent-core bis-(phenyl)oxadiazole derivatives [66]. Finally, we note that this analysis of the splay and bend elastic constants of C5-Ph-ODBP-Ph-OC12 gives no insight into any possible biaxial nature of the nematic phase of this system, although it is clear that cybotactic clusters need not be considered to be present to explain the observed physical properties.
ACKNOWLEDGMENTS

This work was supported by the Engineering and Physical Sciences Research Council under Project No. EP/G023093/1. C.G. acknowledges the Ph.D. School of Materials Science and Engineering (University of Padova) and Merck Chemicals Ltd. for financial support. We thank D. Allender for useful discussions.