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The design and investigation of de Vries liquid crystals based on 5-1 phenyl-pyrimidine and (R,R)-2,3-epoxyhexoxy backbone 2

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Calamitic liquid crystals (LCs) based on 5-phenyl-pyrimidine derivatives have been 15 16 designed, synthesized and characterized. The 5-phenyl pyrimidine core was functionalized with chiral (R,R)-2,3-epoxyhexoxy chain on one side and either siloxane or perfluoro 17 18 terminated chains on the opposite side. The one involving perfluorinated chain shows SmA* phase over a wide temperature range of 82 °C, whereas the siloxane analogue exhibits both 19 SmA^* and SmC^* phases over broad range of temperatures and a weak first order SmA^* -20 21 SmC^* transition is observed. For the siloxane analogue, the reduction factor for the layer 22 shrinkage R (relative to its thickness at the SmA* - SmC* transition temperature, T_{AC}) is ~ 0.373 and layer shrinkage is 1.7% both at a temperature of 13 °C below the T_{AC} . This 23 compound is considered to be having "de Vries smectic" characteristics with the de Vries 24 25 coefficient, C_{deVries} of ~ 0.86 on the scale of zero (the maximum layer shrinkage) to 1 (zero layer shrinkage). A three-parameter mean-field model is introduced for the orientational 26 27 distribution function (ODF) to reproduce the electro-optic properties. This model explains the experimental results and leads to the ODF which exhibits a cross-over from the sugar-loaf to 28 29 diffuse-cone ODF some 3 °C below T_{AC}.

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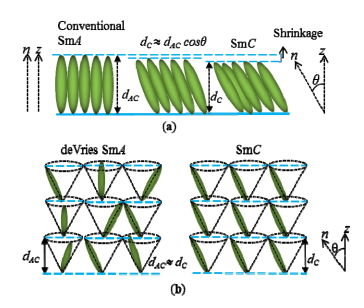
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34 I. INTRODUCTION

35 Chiral smectic liquid crystals (LCs) having mesophases close to the room temperature 36 are extremely useful for the next generation of displays and electro-optical devices. The 37 devices based on ferroelectric smectics with extremely fast switching time (μ s) compared to 38 the currently used slowly switching nematics (ms) are very promising [1]. However to obtain the defect free alignment in an LC cell over a broad range of temperatures, a pre-requisite of 39 40 zero or a small layer shrinkage for practical applications is required and the achievement of 41 which is still an open challenge. In principle, one is able to align the material very well in the 42 nematic or smeetic A phases. However on a phase transition from SmA to SmC, a tilt emerges 43 whereby the molecular long axis *n* tilts relative to the layer normal z by an angle, θ , that varies with temperature (Fig. 1a). Consequently, the layer spacing (d_c) in the SmC phase decreases 44 45 by a factor of $cos\theta$ provided liquid crystalline molecules are close enough to being considered as rigid rods. The layer shrinkage arising from the tilt together with surface anchoring of 46 47 molecules leads to chevron structure/s formed in the cell. These structures with opposite fold 48 directions create zigzag defects at the interface in between the two oppositely folded 49 cheverons. The emergence of these defects are impediments to a successful 50 commercialization of devices based on ferroelectric LCs [2-4]. The solution is therefore focussed on to developing chiral LCs with a minimal or zero layer-shrinkage at the SmA*-51 52 SmC^* transition temperature and within the temperature range of SmC^* [4]. It is normally the case that where the layer shrinkage is much lower than the scaling factor, $\cos\theta$, then the 53 54 smectic phases of such compounds are known as 'de Vries smectics' (Fig. 1b).

A dozen or so compounds have been identified to have de Vries characteristics to the 55 56 varying degrees of perfection. According to the semi-qualitative studies carried out so far, the 57 calamitic LCs that combine low orientational order parameter but large lamellar orderings are likely to exhibit better de Vries type characteristics [5-9]. Most of the de Vries smectics 58 59 contain trisiloxane/carbosilane backbones and perfluorinated side chains on either ends of the 60 core; these terminations strongly promote lamellar orderings [10]. In this case, the alkane 61 chains were replaced by an epoxy group (a precursor in the synthesis of the difluoro material). In order to obtain a large P_S in the SmC* phase, it is pertinent to prepare highly 62 63 enantiomeric enriched chiral molecules to avoid the cancelation effects of the oppositely

handed enantiomers. In the literature, excellent protocols are given for preparing epoxides 64 65 from allylic alcohols with large enantiomeric excesses [11]. However to our knowledge, materials with enantiomeric enriched 5-phenyl pyrimidine epoxides have not yet been 66 reported to have de Vries like behavior [12] before. The epoxide group may induce large 67 electrical polarization. A use of 5-phenyl pyrimidine though as an aromatic core is 68 69 extensively used in the synthesis of compounds that exhibit de Vries-like characteristics in 70 chiral and non-chiral mesogens [13-15]. With this concept in mind, two new chiral materials 71 containing epoxides in their structures have been designed, synthesized and investigated.



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FIG. 1 Schematic representation of the molecular arrangements in the SmA and SmC mesophases as per (a) conventional rigid-rod model and, (b) de Vries diffuse cone model [16]. z is the layer normal, n is the molecular long axis orientation, θ is the angle between n and z, d_C and d_A are the layer spacing in SmC and SmA, respectively.

In this article, we report synthesis and properties of the two 5-phenyl pyrimidine 77 derivatives that contain the chiral (R,R)-2,3-epoxyhexoxy chain in conjunction with 78 79 trisiloxane (adpc042) or perfluorinated (DR257) terminal chains (see appendix for the synthetic procedure). Both moieties are there0fore structurally related to each other, both use 80 81 the same aromatic core and the epoxide chiral chain. In the DR257, siloxane group is replaced with perfluorinated butane as aliphatic linkage of six carbons with the aromatic core. 82 83 A modification of the chain ending could have a significant effect on the properties and with 84 the objective of finding its effect on the electro-optic properties, we decided to explore this 85 further to establish a well-desired structure-property relationship. The mesophases formed by these compounds were characterized by differential scanning calorimetry (DSC), polarized 86

optical microscopy (POM), x-ray diffraction (XRD), electro-optics and dielectric spectroscopy. Characterization of the LCs shows that the perfluorinated epoxide DR257 exhibits a stable SmA^* phase over a broad range of temperatures whereas the siloxane terminated analogue adpc042 exhibits both SmA^* and SmC^* phases. In this trisiloxane compound we achieve a layer shrinkage of 1.7 %, at a temperature of -13 °C below the SmA^* - SmC^* transition temperature. Since adpc042 LC exhibits both SmA^* and SmC^* phases, properties of this compound are detailed below.

94 II. EXPERIMENT

The phase sequence and the transition temperatures were found by DSC thermograms 95 96 (Perkin-Elmer DSC-7 calorimeter). The experiment was carried out under cooling at a rate of 10 °C min⁻¹. The electro-optical studies of the two LCs were conducted using planar-aligned 97 98 cells with parallel rubbed polyimide alignment layer KSRP-XX/D611P6NSS05. These cells 99 were purchased from EHC, Japan. LC cells were studied by POM and electro-optics. 100 Olympus BX 52, equipped with a rotating table was used. The hot-stage was fixed on to the 101 rotating table. The hot stage was connected to a temperature controller, Eurotherm 2604. 102 XRD measurements were carried out on Bruker D8 diffractometer using Cu-K_a source with a 103 wavelength 0.154 nm. Dielectric spectroscopic studies were performed using an Alpha High 104 Resolution Dielectric Analyser (Novocontrol GmbH, Germany) in the frequency range 1 Hz -105 10 MHz. Indium Tin Oxide (ITO) coated glass substrates were used to make cells for studies 106 of the real and imaginary parts of the dielectric permittivity. Substrates were coated with RN 107 1175 polymer (Nissan Chemicals, Japan) for obtaining the planar alignment. The cell 108 thickness (d) was controlled by Mylar spacers. UV-VIS spectrometer (Avaspec-2048) was 109 used to measure d through interference fringes. The ITO sheet resistance of the substrates (20 110 Ω/\Box) is low enough to shift the peak frequency for the resistance of the ITO in series with the 111 cell, beyond the experimental window of measurements. The dielectric spectra were analysed using Novocontrol WINDETA program. 112

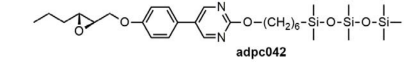
113 III. RESULTS AND DISCUSSIONS

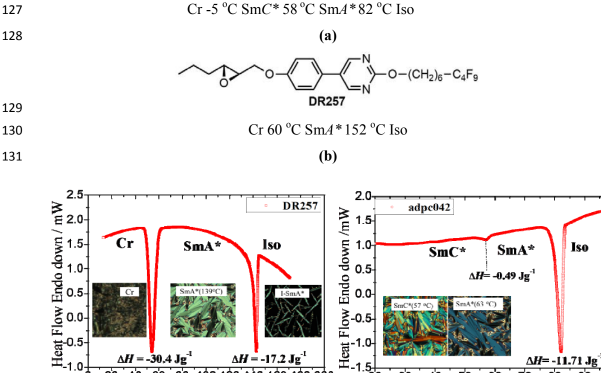
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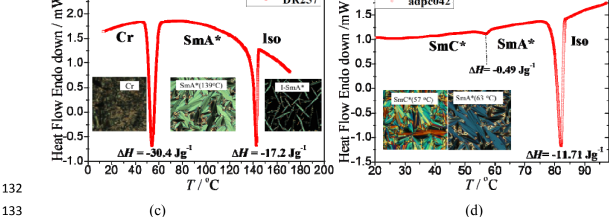
A. Differential Scanning Calorimetry and Polarizing Optical Microscopic Studies

115 The molecular structure and the transition temperatures of siloxane pyrimidine 116 adpc042 and the fluorinated pyrimidine DR257 are given in Figs. 2 (a) and (b) (synthetic

procedures given in the appendix). Result of a DSC thermogram of DR257 is given in Fig. 117 2c. Here two transitions with the enthalpies, ΔH of ~17.2 Jg⁻¹ and ~30.4 Jg⁻¹ at temperatures 118 of ~ 143 °C and ~56 °C are recorded. To identify the mesophases, textures of a 9 μ m planar 119 120 cell filled with DR257 are recorded with POM. Images corresponding to the two mesophases 121 are shown in the inset of Fig. 2c. On cooling from the isotropic temperature, the mesophase 122 grows from the typical 'batonnet' structures to the focal conic texture. Both focal conic and 123 dark homeotropic textures obtained by POM are the characteristics of SmA* phase [17]. Textures confirm that the phase transition from SmA* to a crystalline state is at a temperature 124 125 of 56 °C.







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FIG. 2 Molecular structures and the DSC thermograms of adpc042 [(a) and (d)] and 134 DR257 [(b) and (c)] are given. The phase transition temperatures are obtained under the 135 cooling cycle with a quasi-equilibrium condition at the rate of $\sim 1^{\circ}$ C min⁻¹ using POM. 136 The cooling curves of the DSC thermograms are obtained at the rate of 10 $^{\circ}$ C min⁻¹. 137 138 Iso= isotropic phase, Cr= crystalline state. Insets in these two figures are the POM 139 images of the mesophases seen under the crossed polarizers. Textures are recorded for a 140 9 µm planar-aligned cells in the cooling run.

141 A representative the DSC plot of adpc042 LC under cooling exhibits two peaks (Fig. 142 2d). Both DR257 and adpc042 show the first order Iso -SmA* transition with ΔH of ~11.71 Jg⁻¹ and 17.2 Jg⁻¹ at a temperature of ~ 82 °C and ~ 145 °C. This is confirmed by POM where 143 the focal conic fan shaped texture from a 9 µm planar-aligned cell (Fig. 2d inset) is recorded. 144 On lowering the temperature, the sample undergoes a weakly first order SmA^* - SmC^* 145 transition with ΔH of ~0.49 Jg⁻¹ (0.069 kcal mol⁻¹) at 58 °C [12, 18]. For a conventional 146 SmA^* - SmC^* transition where the tilt angle stays at almost zero value in the SmA^* phase 147 down to the transition temperature T_{AC} , the DSC thermogram exhibits a second order 148 149 transition with a step in the baseline without a peak for the enthalpy of transition [19]. The 150 SmC^* phase of adpc042 is distinguishable from SmA^* by POM where a fan shaped texture 151 from a planar cell is altered to a broken fan shaped one (inset Fig. 2d). The dark homeotropic 152 texture is changed to Schlieren texture on transition from SmA^* to SmC^* . The entire 153 replacement of trisiloxane in adpc042 by tetra fluorocarbon chain in DR257, radically alters the mesomorphic behaviour, the temperature range of SmA* is increased from 23 °C to 89 °C, 154 whereas SmC^* disappears. An increased temperature range of the SmA^* phase in DR257 155 156 could be due to a large dipole moment associated with the fluorinated tail [20]. The siloxane 157 groups in adpc042 enhance the thermal stability of the compound as well as lower the phase 158 transition temperatures. These features may be of great importance from a point of view of 159 the molecular design and applications.

160 The POM images of a planar-aligned cell of thickness 9 μ m filled with adpc042 161 recorded at 63 °C (~19 °C below the Iso- Sm*A** transition temperature and 5 °C above the 162 T_{AC}), are given in Fig. 3. The rubbing direction R_d is fixed at an angle, $\alpha = \sim 15^\circ$ to the 163 polarizer **P**.

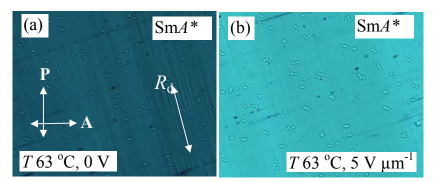


FIG. 3 POM images of Sm A^* phase at a temperature of 63 °C in a 9 µm planar-aligned cell filled with adpc042 LC: (a) 0 V and (b) 5 V µm⁻¹ (square wave AC electric field of 110 Hz).

168 The electric field treatment of the LC cell gives rise to a uniform mono-domain 169 texture (Fig. 3). An application of the external electric field across a cell in well-aligned 170 SmA^* phase produces a change in the color of the texture (compare Figs. 3a and 3b), due to 171 an increase in the birefringence Δn with the field, a characteristic of the de Vries behavior. In 172 this phase, the molecules are tilted but are azimuthally disordered. The electric field produces 173 a bias in the azimuthal distribution and the molecules tilt in a particular direction determined 174 by the polarity of the field. The resulting SmA^* is reduced to the same symmetry as SmC^* 175 phase through azimuthal ordering of the already tilted liquid crystalline molecules.

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B. X-ray diffraction(XRD)

Figure 4(a) shows the temperature dependence of the smectic layer spacing 177 determined by the XRD experiment. On cooling the sample below the isotropic phase, the 178 layer spacing in SmA* phase initially increases linearly with decreasing temperature. On 179 approaching the SmA*- SmC* transition temperature, the thickness reverses its trend within 180 181 SmA* from an increase to a decrease due to an emergence of the molecular tilt even in the SmA* phase. On further cooling of the sample cell, the layer thickness in the SmC* phase 182 183 continues to decrease much more slowly than expected from a conventional smectic, in which the layer thickness scales as, $\cos \theta$. Figure 4b shows the simulated molecular 184 185 structures of the compound adpc042 using Hyperchem Program. The simulated length l, of the molecule is ~ 31 Å. A possible molecular arrangement in layers is shown in Fig. 4b such 186 187 that the total length of a pair of molecules is larger than the smectic layer thickness.

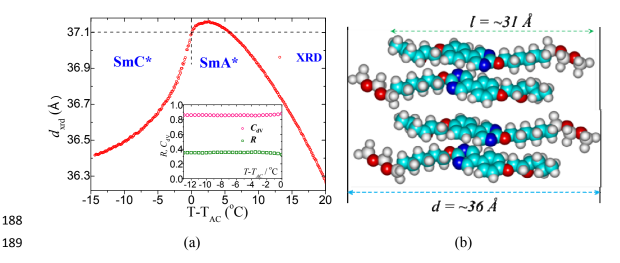


FIG. 4 (a) Temperature dependence of the smectic layer spacing determined by XRD (\blacktriangle) relative to the layer thickness at the Sm*A*^{*} to Sm*C*^{*} transition temperature. The inset shows plots of the de Vries coefficient $C_{deVries}$ () and the reduction factor R (\square) vs. the

reduced temperature $(T-T_{AC})$. De Vries characteristics are defined in terms of $C_{deVries}$ and *R*. Measurements are carried out on a compound adpc042, (b) The simulated molecular structures of the compound adpc042.

A layer shrinkage of 1.7% is calculated from the XRD at a temperature of 13 °C below the Sm A^* - Sm C^* transition temperature T_{AC} . The layer-shrinkage is small. The behaviour is similar to that in Sm A^* phase where the molecular directors in Sm A^* phase are distributed on to a cone where the azimuthal angle is degenerated. At the Sm A^* to Sm C^* transition, the tilt directions condense to within narrower limits, hence the disorder in the azimuthal angle disappears in the Sm C^* phase. This process in itself requires no change in the layer thickness.

203 The reduction factor *R* of a smectic LC is defined as,

204
$$R = \frac{\delta(T)}{\theta_{opt}(T=T_{AC})} = \frac{\cos^{-1}(d_C(T)/d_{AC}(T=T_{AC}))}{\theta_{opt}(T=T_{AC})}$$
(1)

205 where δ (T) governs the layer thickness at a temperature T within SmC* phase relative to the layer thickness at the SmA*- SmC* transition temperature, d_{AC} , on assuming that the rigid 206 rod model is applicable to the LC under investigation [21]. θ_{opt} is the optical tilt angle 207 determined by the POM (Fig. 5). According to Eq. (1), an ideal de Vries smectic with 208 $d_c(T) \approx d_{AC}(T = T_{AC}$ (Fig. 1b) produces a defect-free bookshelf geometry in SmC* 209 phase with a reduction factor R = 0, *i.e.* the short-range molecular tilt order at a lower 210 temperature in the SmA* phase becomes long range close to the SmA*- SmC* transition 211 temperature, where the maximum de Vries cone angle is equal to the saturated optical tilt 212 angle at a temperature close to T_{AC} . On entering the SmC* from SmA* phase, the azimuthal 213 214 ordering of the molecular directors on the cone localizes these on to a single orientation. In 215 this case, the layer contraction is absent and then $d_C/d_{AC} \approx 1$, and therefore $R \approx 0$. Li *et al.* 216 suggested another method of characterizing de Vriesness from measurements of the layer 217 thickness from XRD in terms of the de Vries co-efficient, C_{deVries} [22]. The magnitude of this varies from '0' (classic rigid rod model with a maximum layer shrinkage) to 1 (ideal de Vries 218 219 $SmA^* - SmC^*$ transition with zero layer shrinkage) such that

220
$$C_{deVries} = 1 - \left[\frac{(d_{AC} - d_C)}{(d_{AC}(1 - \cos \theta))}\right]$$
 (2)

The material adpc042 gives rise to $R \approx 0.373$ at 13 °C below the SmA* - SmC* transition 221 temperature. Understandably, R is away from zero but still low, hence the material is close to 222 being a 'de Vries smectic'. An estimated value of $C_{deVries}$ is ~0.86 (Fig. 3 inset) at 13 °C 223 below the SmA^* - SmC^* transition temperature. The inset plot of Fig. 4 shows that both R 224 and $C_{deVries}$ are almost independent of temperature throughout the entire temperature range of 225 the SmC* phase. The lowest reported value of R so far is 0.17 for the chiral de Vries 226 227 compound QL32-6 that exhibits a maximum layer contraction of only 0.2% at 3 °C below the 228 SmA*-SmC* transitions with an optical tilt angle of 20[°] [13].

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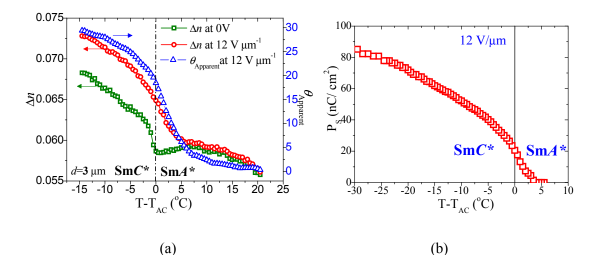
C. Electro-optical studies

The birefringence Δn and the apparent optical tilt angle $\theta_{Apparent}$ are determined by 230 231 recording the intensity of a transmitted beam of light through a LC cell by varying the 232 positions of the polarizer and the analyzer. The procedure involves first keeping P fixed, and 233 then rotating the analyzer automatically by various angles. This procedure is repeated for at least three different positions of the polarizer. From the data, Δn and $\theta_{Apparent}$ are 234 235 calculated. This procedure is different from that given by Park et al. [23], who record the 236 transmitted intensity by rotating the sample and also record the transmitted intensity for the P and A (i) parallel and (ii) crossed to each other. The experiment is conducted by applying a 237 triangular signal of frequency 46 Hz and an amplitude of 12 $V_{0-peak} \ \mu m^{-1}$. Frequency of the 238 field is chosen so as to allow sufficient time for the electro-optic switching to occur but to 239 240 prevent ionic-conductivity contributing to the switching current. An amplitude of the voltage 241 applied to the LC sample adpc042 is large enough so as to make the tilt angle increase slowly with field and then eventually to saturate by the field while avoiding a risk of damage to the 242 243 sample by the applied electric field.

Figure 5 shows results of the birefringence (Δn) as a function of temperature with and without electric field applied across the cell. On cooling the sample in the absence of field from the isotropic to Sm*A** phase, the magnitude of Δn first increases slightly, but then starts to decrease within Sm*A** phase. This decrease in Δn with temperature is due to the de Vries tilt appearing in the Sm*A** phase and a distribution of the in-layer directors occurring on to the cone. On further cooling the sample in the absence of electric field, Δn suddenly increases

at the SmA^{*} - SmC^{*} phase transition temperature T_{AC} followed by slow increase with a 250 decrease in temperature. An application of the electric field ($E=12 \text{ V} \mu \text{m}^{-1}$) gives rise to a 251 continuous rise in Δn in the entire temperature range of the SmA^{*} and SmC^{*} phases. This 252 field induced increase in Δn is consistent with a change in the interference colors observed by 253 254 POM (Fig. 3). The behavior is typical of the diffuse-cone model of SmA^* phase with a weak first order SmA* - SmC* phase transition (Fig. 2b). In the conventional SmA* of a LC, one 255 256 may expect a small increase in Δn to occur at the phase transition from a uniaxial SmA* (with 257 zero tilt angle at $T = T_{AC}$) to a biaxial SmC* phase.

For adpc042, the measured value of the spontaneous polarization P_S [24] is plotted as 258 259 a function of the reduced temperature in Fig. 5(b). Measurements are carried out on a 5 μ m 260 planar cell under the application of a square wave, frequency 110 Hz and large enough 261 amplitude of 12 V/ μ m. The field is such that the helical structure for temperatures within SmC* is unwound. It is interesting to note that the field induced polarization is much higher 262 conventional ferroelectric liquid crystals. P_S increases with a reduction in 263 than for 264 temperature below the SmC*-SmA* transition. The liquid crystalline material under study yields $P_{S} \sim 82.5 \text{ nC cm}^{-2}$ for $T = (T_{AC} - 30)$ °C. 265



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FIG. 5. Results given here are for adpc042. (a) The temperature dependence of 268 the apparent tilt angle ($\theta_{Apparent}$) and the birefringence Δn by applying a maximum 269 field of (12 V μ m⁻¹, red open circles); Δn in the absence of the electric field 270 (green square). Blue triangles denote the apparent tilt angle $\theta_{Apparent}$ for a field of 271 12 V/ μ m. Measurements are carried out on a 3 μ m planar-aligned cell filled with 272 adpc042 LC. (b) Spontaneous polarization P_S measured under the cooling process 273 274 from the isotropic temperature plotted as a function of (T-T_{AC}). The measurements are carried out on a 5 µm planar cell by applying a square wave AC 275 voltage of 12 $[V_{peak-peak}/\mu m]$ at a frequency of 110 Hz. 276

The measured values of $\theta_{Apparent}$ as a function of the electric field for different temperatures in the Sm*A** phase are plotted in Fig. 6a. The co-ordinate system and the definition of the various angles are given in Fig. 6b. At a higher temperatures in the Sm*A** phase, the magnitude of $\theta_{Apparent}$ is low and it increases linearly with the applied field. For temperatures closer to the T_{AC} , $\theta_{Apparent}$ becomes nonlinear and it continues to increase slowly up to an electric field of 12 V μ m⁻¹ and eventually the tilt angle tends to saturate with the electric field.

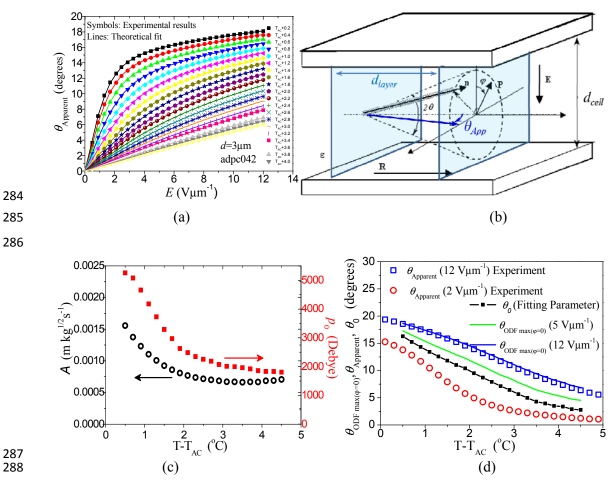


FIG. 6. (a) Variation of the field induced apparent optical tilt $\theta_{Apparent}$ (symbols) and 289 the corresponding fitted values (solid lines) for selected temperatures. Measurements 290 291 are carried out in a 3 µm planar cell filled with adpc042 LC. (b) The schematic of the dynamics with the laboratory co-ordinate system (c) Temperature dependence of the 292 model parameters A and p_0 found from the experiments, (d) temperature dependencies 293 294 of the measured apparent tilt angle $\theta_{Apparent}$ (open symbols), the cone angle θ_0 (the line 295 with symbols) apositions of the distribution function of the maxima (lines) for the various electric field strengths applied across a planar-aligned cell. 296

297 Several approaches exist in the literature for modeling the unusual electro-optic 298 characteristics of the de Vries smectics [25-30]. The Langevin-Debye model, proposed by 299 Fukuda [25] in a different context, was used by Clark et al. [27] to explain the electro-optical 300 properties of de Vries smectic LCs. This model assumes that for a fixed temperature and zero 301 electric field, the molecular directors in the SmA* phase are tilted but azimuthally distributed 302 on to a cone. In this model, the free energy is expressed as $U = -pE\cos\varphi$, where p is the local 303 dipole moment. However, this model does not correctly explain the dependence of the 304 induced apparent tilt angle ($\theta_{Apparent}$ for the electric field) for temperatures rather closer to the SmA^*-SmC^* transition temperature T_{AC} . In 2013, the Boulder group [28] modified this 305 model where they added an additional term involving the square of the electric field in the 306 expression for the free energy. This is expressed as $U = -p_0 E \sin \theta \cos \varphi (1 + \alpha E \cos \varphi)$. 307 Here, α is the phenomenological scaling factor and $p_0 \sin\theta$ is the dipole moment of the 308 309 domain correlated in the molecular tilt created by the condensation of azimuthal angles φ . 310 The first term $-p_0 Esin\theta \cos\varphi$ corresponds to the interaction of the dipole with the field. The second term $-\alpha p_0 E^2 \sin\theta \cos^2 \varphi$ includes the tilt susceptibility that increases with the 311 312 square of the field E and it leads to a sigmoidal response in both Δn and $\theta_{Apparent}$ with E. In this model, the field induced $\theta_{Apparent}$ varies between the values inferred from the Δn at zero 313 field (θ_{\min}) to the maximum electric field (θ_{\max}). These limiting values of θ_{\min} and θ_{\max} 314 315 themselves are temperature independent but the actual values within these limits nevertheless 316 are temperature dependent. Therefore, we propose a somewhat different model. This 317 involves three parameters for explaining the electro-optic response where the mean-field free 318 energy is expressed as:

$$U = -p_0 E \sin\theta \, \cos\varphi + A^2 \sin^2(\theta - \theta_0) \tag{3}$$

320 Here the first term, $p_0 E \sin \theta \cos \varphi = \mathbf{P} \cdot \mathbf{E}$, describes the usual dipole interaction energy with the field. $|\mathbf{P}|$, $[= (p_0 \sin \theta)]$, is the magnitude of the dipole moment of a tilt correlated 321 322 domain. This first term of Eq. (3) is linear in the external electric field E. The second term in this equation defines the cone distribution with a cone aperture angle of $2\theta_0$ and a distribution 323 width proportional to $\sqrt{k_BT}/A$. Here k_B is the Boltzmann constant, T is the absolute 324 temperature and A is the phenomenological co-efficient for the zero field molecular 325 distribution. Note that for the case, $\theta_0 < \sqrt{k_B T} / A$, the model produces "sugar-loaf" ODF (Fig. 326 7(a)), while a larger θ_0 combined with a narrower distribution width correspond to the 327 328 "volcano" or "diffuse-cone" distribution (Fig. 7(a)). Thus both scenarios discussed in [31] are 329 reproduced with the same expression for the free energy. Figure 7b illustrates the trend of $[(k_BT/A)^{1/2} - \theta_0]$, as a function of the reduced temperature (T-T_{AC}) for adpc042. One can see 330

that at 2 to 3 degrees below the phase transition, this function crosses zero. This is a cross-over between the sugar-loaf and the diffuse-cone ODFs.

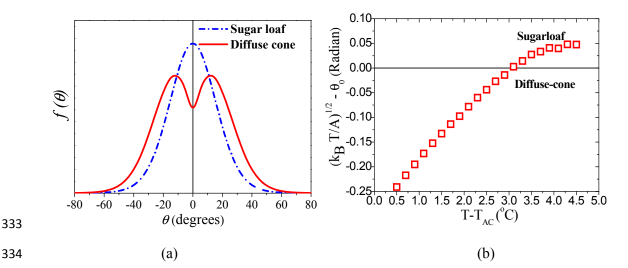


FIG. 7. (a) 2D schematic representation of sugar-loaf (dash-dot lines in blue colour) and diffuse-cone (solid lines in red colour) ODFs. (b) $[(k_BT/A)^{1/2} - \theta_0]$ plotted as a function of the reduced temperature (T-T_{AC}). Measurements are carried out on a 3 µm planar cell filled with adpc042.

For the range of the electric fields used, we observe no inflection point in the field dependence of $\theta_{Apparent}$ *i.e.* no change in the slope of the derivative of the angle at the point. Therefore, we can achieve a good fit of the experimental data to the model without using terms of higher powers of *E*.

In order to simulate the behaviour of $\theta_{Apparent}$, we use the mean-field approach combined with equations for the optical response [28]. An average $\langle Y \rangle$ over the orientational distribution can be written as $\langle Y \rangle = \int_0^{2\pi} \int_0^{\pi/2} Y(\theta, \varphi) f(\theta, \varphi) \sin \theta d\theta d\varphi$, where the mean field ODF $f(\theta, \varphi)$ is expressed as:

347
$$f(\theta,\varphi) = \exp[-U/k_B T] / \int_0^{2\pi} \int_0^{\pi/2} \exp[-U/k_B T] \sin\theta d\theta d\varphi$$
(4)

348 The apparent electro-optical tilt angle θ_{Apparent} is given by:

349
$$\tan 2\theta_{Apparent} = \frac{\langle \sin 2\theta \cos \varphi \rangle}{\langle \cos^2 \theta - \sin^2 \theta \cos^2 \varphi \rangle}$$
(5)

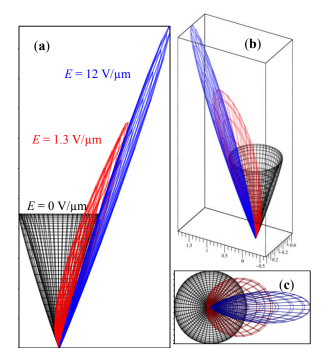
On fitting the experimentally obtained voltage dependencies of $\theta_{Apparent}$ to the above model, we obtain values of p_{0} , A and θ_{0} each as a function of temperature (Fig. 6c, and 6d). All the parameters tend to increase on approaching T_{AC} as the cell is cooled from the isotropic state to the Sm A^* phase. Such an increase is in agreement with the previous models [27, 28].

Onusing the dipole moment of the correlated domain as $p_0 \approx 5000$ D at 0.5 °C above 354 the SmC^* - SmA^* phase transition temperature, we can estimate the effective size of a 355 domain using the procedure similar to that adopted by Shen et al. [28]. The effective 356 357 molecular dipole moment can be estimated from the polarization density measurements performed on a planar-aligned cell in SmC* phase. In this case P_S is saturated at 69 nC/cm²; 358 $\theta_{Apparent} = 30^{\circ}$ at $(T_{AC} - 19)^{\circ}C$, with the molar mass, M = 590.3 g/mol, density $\rho = 1.2$ 359 g/cm³, the dipole moment is found as $\mu_{eff} = \frac{P_S M}{\rho N_A sin \theta_{Apparent}} \approx 0.34$ D. Therefore, the tilt-360 correlated domain in our experiment contains approximately 6, 000 to 15,000 molecules. The 361 effective domain correlation length $\sqrt[3]{\frac{p_0 sin \theta_{Apparent}}{P_S}}$, at a temperature close to the SmA* -362 SmC^* phase transition, is of the order of 30 nm [30]. 363

364 Parameter A clearly favours the "volcano"- type distribution as illustrated by Fig. 8. A comparison of the cone angle θ_0 that fits the experimentally obtained $\theta_{Apparent}$ for different 365 fields is shown in Fig. 6d. Note that for higher field strengths, $\theta_{Apparent}$ exceeds θ_0 . This is 366 367 indicative of a large electro-clinic effect. The solid lines in Fig. 6d show angular positions of the maximum in the ODF simulated for the two electric field strengths (5 and 12 V μ m⁻¹) 368 here. The experimental values of $\theta_{Apparent}$ for 12 Vµm⁻¹ are slightly lower than for an angle for 369 which θ_{ODF} is a maximum. This is expected from a highly distorted difuse-cone ODF, the 370 371 distortion is brought about by the applied electric field.

Figures 8a-d show a 3D illustration of the ODF function obtained for a temperature of ($T_{AC} + 1$) °C. The ODF is normalized as given below such that its volume for different fields is a constant: $f_{3D} = \frac{f(\theta, \varphi)}{\sqrt[3]{\int_0^{2\pi} \int_0^{\pi/2} f(\theta, \varphi)^3 \sin \theta d\theta d\varphi}}$. Figure 8c shows the cross-section of the ODF

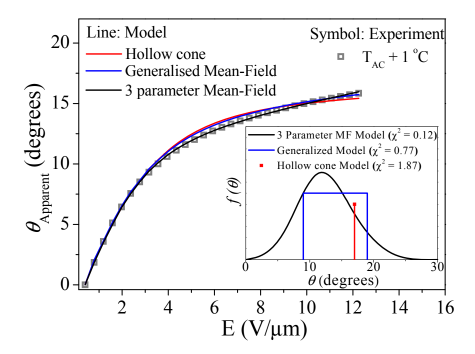
in the plane of the LC cell where the maximum of the ODF *vs.* apparent tilt angle is plotted. The electro-clinic effect for the following electric fields are given in Fig. 8: zero, moderate (1.3 V/ μ m) ("cone unwinding"), and for large fields; 12 V/ μ m (tilt increases but φ is condensed within a narrow range of values). Note that the modeled ODF shown in Fig. 8 corresponds to the core part of the mesogen that exhibits birefringence at visible wavelengths of light.



381

FIG. 8. The ODF of the smectic compound, adpc042, at a temperature of $T = (T_{AC} + 1)$ °C for different electric field strengths (0 V, 1.3 Vµm⁻¹, and 12 Vµm⁻¹). (a-c) are the axonometric projections of $f_{3D}(\theta, \varphi)$.

385 To test different models, we fit experimental data of $\theta_{Apparent}$ as a function of the electric field at a temperature 1 °C above the SmA*- SmC* phase transition temperature, T_{AC} (Fig. 9), to 386 387 the following models: the fixed-angle hollow cone [27], the generalized mean-field model 388 [28] and our 3-parameter mean-field model. For low fields data fitted to the various models is 389 quite adequate. However for the medium and high electric fields, the 3-parameter model shows a closer fit to the experimental data. This is supported by drawing a comparison 390 between sum of the squares of the residuals (χ^2) for the various fits as shown in the inset of 391 Fig. 9. Though, the three models provide reasonable fit to the data, our proposed model has no 392 393 intrinsic assumption of either the sugar-loaf or the diffuse cone type distribution, i.e. it is not 394 limited to the assumption of a certain ODF in the SmA* phase. Therefore testing and 395 discrimination among the models is carried out automatically during the fitting procedure. 396 Moreover, the model produces a continuous function for the ODF, having a better physical 397 significance.



399

FIG. 9. The experimental, $\theta_{Apparent}$ (\Box) *vs* electric field E shown at a temperature of 1 ^oC above the Sm*A**- Sm*C** phase transition temperature, T_{AC}, are fitted to various models: The hollow cone, the generalized mean Field (MF) and the 3 parameter MF one. A schematic representation of the ODFs for the corresponding three models is given in the inset.

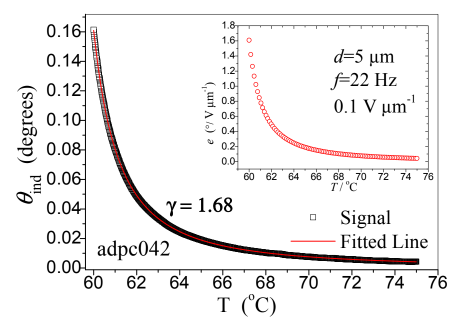
The de Vries LCs normally exhibit large values of the electro-clinic coefficient (denoted by *e*) [19, 32, 33]. The electro-clinic response is recorded for the material adpc042 in a planar-aligned cell. The experiment is carried out by keeping the smectic layer normal at an angle of 22.5° with respect to the polarizer. The light intensity *I*, transmitted through the LC cell, in the absence of the field *E* is given by:

410
$$I = I_0 \sin^2(2\alpha) \sin^2(\pi \Delta n d/\lambda)$$
(6)

411 I_0 is the incident intensity, α is the angle between the molecular director and the polarizer, Δn 412 is the birefringence, *d* is the cell thickness, and λ is the wavelength of the incident light. 413 When an electric field is applied across the cell, the transmitted intensity varies linearly with 414 the induced tilt angle $\delta \alpha = \theta_{ind}$. Differentiating Eq. (6) with respect to α is given as:

415
$$\delta I = 2I_0 \sin (4\alpha) \sin^2(\pi \Delta n d/\lambda) \theta_{ind}$$
(7)

416 From Eqns. (6) and (7) and for $\alpha = 22.5^{\circ}$ we obtain $\theta_{ind} = \delta I/4I$. Figure 10 shows θ_{ind} as a 417 function of temperature over the temperature range of SmA* phase. The magnitude of θ_{ind} increases with decreasing temperature. On approaching, T_{AC} , from the high temperature side of Sm*A**, magnitude of θ_{ind} diverges and this corresponds to the divergence of the correlation length of the tilt domain and here the azimuthal angle is condensed to lie within narrower limits.



422

432

FIG. 10. The temperature dependence of electro-clinic response θ_{ind} of adpc042 (\Box): experimental value, red line (—): fitting to Eq. (7). The inset figure is the electroclinic co-efficient *e* (red open circle) calculated using the Eq. $e = \theta_{ind}/E$ as a function of temperature. Experiments were conducted under cooling within the temperature range of Sm*A** phase up to a temperature of $T = (T_{AC} + 1)$ °C by the application of 0.1 V_{0-peak} µm⁻¹ (planar aligned cell, *d*=5 µm sine wave AC voltage of 22 Hz).

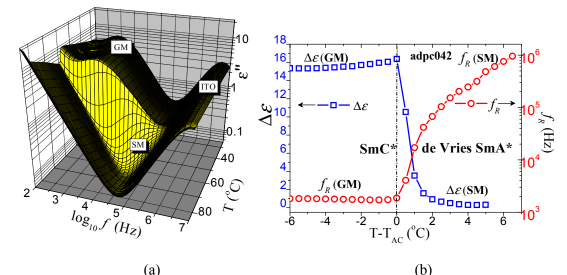
430 The amplitude of θ_{ind} with temperature can be expressed by the power law equation as 431 follows [34]

$$\theta_{ind} = \frac{c}{(T - T_{AC})^{\gamma}} \tag{8}$$

Here *C* is the scaling constant, T_{AC} is the Sm*A**-Sm*C** transition temperature and γ is the critical exponent. For the smectic material under investigation $\gamma = 1.68$. For a conventional Sm*A**-Sm*C** transition, γ is 1.33 [35]. Values of γ greater than 1.33 reflect the short-range correlation of molecular directors in three dimensions, again a characteristic of the de Vries smectics. The inset plot in Fig. 10 shows the temperature dependence of the electroclinic coefficient *e*, calculated from the definition of $e = \theta_{ind}/E$.

439 **D. Dielectric Spectroscopy**

Figure 11a shows a three dimensional plot of temperature dependent dielectric loss 440 spectra (ε'') of a planar aligned cell filled with the material adpc042 LC. Note that the 441 442 temperature independent high frequency process is due to the resistance of ITO in series with 443 the capacitance of the filled cell. The relaxation process in the SmA^* phase corresponds to 444 fluctuation of the tilt angle and this mode is known as the soft mode (SM). This arises mainly 445 as the system approaches T_{AC} the elastic constant controlling the tilt fluctuations decreases or 446 gets softer. The low frequency dielectric relaxations in the SmC^* phase is associated with the 447 Goldstone (GM) mode in which the molecular director is subjected to continual symmetry 448 breaking.



449

450 (a) (b) 451 **FIG. 11.** (a) A 3-D plot of temperature dependent dielectric loss spectra (ε ") of the 452 compound adpc042 (cell thickness, $d=10 \ \mu$ m) in a planar-aligned cell. The dielectric 453 measurements are carried out under cooling. Temperature stabilization is 0.05 °C and the 454 applied voltage is 0.1 V_{rms} (b) Dielectric strength $\Delta\varepsilon$ (open blue squares \Box) and the 455 corresponding relaxation frequency f_R (red open circles) as a function of temperature. 456 SM and GM refer to the soft and Goldstone modes, respectively.

457 The dielectric spectra are analysed using the Novocontrol WINFIT programme. The 458 temperature dependent dielectric strength $\Delta \varepsilon$ and the relaxation frequency f_R are obtained by 459 fitting the dielectric spectra to the Havriliak-Negami Eq. [36]:

460
$$\varepsilon^{*}(\omega) = \varepsilon^{'} - i\varepsilon^{''} = \varepsilon_{\infty} + \sum_{j=1}^{n} \frac{\Delta \varepsilon_{j}}{\left[1 + \left(i\omega\tau_{j}\right)^{\alpha_{j}}\right]^{\beta_{j}}} - \frac{i\sigma_{dc}}{\varepsilon_{0}\omega}$$
(9)

where, ε_{∞} is the high frequency permittivity depending on the electronic and atomic 461 462 polarizability, j is the number of relaxation processes which varies from 1 to n, $\omega = 2\pi f$ is the angular frequency in radians, ε_0 is the free space permittivity, τ_i is relaxation time of the j^{th} 463 process, $\Delta \varepsilon_i$ is the dielectric strength and α_i and β_i are the symmetric and asymmetric 464 broadening parameters of the j^{th} dielectric relaxation process related to the distribution of 465 relaxation times. The term $(-i\sigma_{dc}/\epsilon_0\omega)$ gives the dielectric loss due to the ionic conduction at 466 and is dominant at lower frequencies due to the inverse ω term. The relaxation frequency f_i of 467 the j^{th} process is related to τ_i as [37]: 468

469
$$f_{j} = \frac{1}{2\pi\tau_{j}} \left[\sin\left(\frac{\alpha_{j}\pi}{2+2\beta_{j}}\right) \right]^{1/\alpha_{j}} \left[\sin\left(\frac{\alpha_{j}\beta_{j}\pi}{2+2\beta_{j}}\right) \right]^{-1/\alpha_{j}}$$
(10)

The temperature dependencies of $\Delta \varepsilon$ and f_R are shown in Fig. 11b. The amplitude of $\Delta \varepsilon$ shows a pronounced increase when the system approaches T_{AC} from the Sm A^* phase. The maximum value of $\Delta \varepsilon$ at T_{AC} is 16.8. The corresponding f_R is decreasing in the Sm A^* phase with a sharper trend in its lower temperature range. Remarkably strong soft mode dielectric absorption is observed in the dielectric spectra of de Vries LCs in contrast to the compounds that exhibit a conventional Sm A^* [38-41].

476 IV. CONCLUSION

We have designed and synthesised two 5-phenyl pyrimidine derivatives with siloxane and fluorocarbon chain terminations, both with a chiral (R,R)-2,3-epoxyhexoxy side chain. Mesophases formed by these compounds are investigated using a number of techniques: DSC, POM, XRD, electro-optical, and dielectric spectroscopy.

481 We find that the maximum layer contraction for the siloxane terminated adpc042 at a temperature of 13 °C below the Sm A^* - Sm C^* transition is 1.7% and a reduction factor R is 482 483 0.37 on a scale of 1 to 0. This compound is considered to be a good "de Vries smectic" with 484 the de Vries coefficient $C_{deVries}$ of 0.86 on the scale of 0 to 1. The soft mode dielectric relaxation strength $\Delta \varepsilon$ shows a critical behaviour when the system approaches SmA* -SmC* 485 486 transition from the high temperature side. The critical exponent of the electro-clinic response is found to be $\gamma = 1.68$. Value of the exponent is much greater than 1.33, found for most 487 488 conventional smectics. Temperature dependence of the correlation length suggests that the material has de-Vries characteristics. 489

490 We introduce a three parameter MF model for describing the ODF of the molecular 491 director in SmA* phase with a view of fitting the field-induced experimental apparent tilt angle $\theta_{Apparent}$ data as a function of the electric field. Close to the SmA* - SmC* transition 492 temperature, the birefringence in SmA* phase is seen to decrease with a reduction in 493 temperature. This is due to the emergence of a non-zero value of cone angle θ_0 , the 494 magnitude of which increases to 17° as the SmA*-SmC*phase transition is approached. The 495 proposed model fits the experimental data on apparent tilt angle and leads to the diffuse-cone 496 ODF, at temperatures close to the phase transition temperatures. However θ_0 decreases with 497 498 an increase in temperature and the ODF displays a cross over from the diffuse-cone to the sugar loaf at ~ $3.5 \,{}^{0}$ C above T_{AC}. 499

500 Acknowledgements

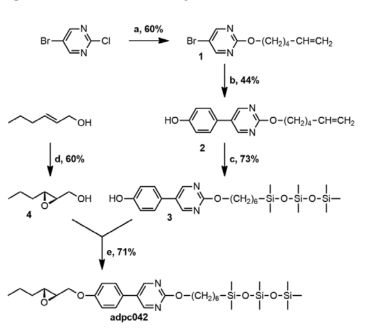
Work was supported by 13/US/I2866 from the Science Foundation Ireland as part of the US– Ireland Research and Development Partnership program jointly administered with the United States National Science Foundation under grant number NSF-DMR-1410649. Financial support for the Belfast group was from the Department for Employment and Learning under with grant code USI 056. X-ray measurements were done in Warsawa, under the EU Cost ICI 1208 2014-17 project. We thank Professor Satyendra Kumar for co-ordinating the US-Ireland project.

508 APPENDIX: SYNTHETIC PROCEDURE

All reagents were purchased from Sigma Aldrich, Fluorochem, Alfa Aesar, ABCR, Synthonix and used without any further purification. Solvents were purchased from Sigma Aldrich, DMF was purchased pre-dried, THF was dried using a sodium/benzophenone still under N₂ and DCM was dried using CaH₂. All reactions were generally carried out under argon using oven-dried glassware. TLC plates were performed on Merck silica gel 60 F_{254} and were visualized using a 254 nm light source. Flash column chromatography was performed on Fluorochem silica gel 60 (40-63 micron).

IR spectra were recorded using a Perkin Elmer Spectrum Two FT-IR spectrometer. ¹H and
¹³C spectra were recorded at 25°C (CDCl₃ as solvent and TMS as reference) using a Bruker
400 MHz Ultrashield (Avance 400). HRMS spectra were recorded using a Waters – TOF
Electrospray micromass LCT premier. Optical rotations were recorded using a polarimeter
Perkin Elmer: model 341 Polarimeter.

521 The synthesis of adpc042 was carried out through the scheme shown in FIG. 12.



522

523

FIG. 12. General scheme for the preparation of adpc042

Further details of synthesis and characterization of each stage of intermediates are givenbelow:

526 The chemical structure of compound 1 is given in Fig. 13.

$$\mathsf{Br} \xrightarrow{\mathsf{N}} \mathsf{O} \text{-} (\mathsf{CH}_2)_4 \text{-} \mathsf{CH} \text{=} \mathsf{CH}_2$$

527 528

FIG. 13. Chemical structure of compound 1

529 5-Hexen-1-ol (0.62 g, 6.2 mmol) was dissolved in dry toluene (15 mL) and sealed in a 530 schlenk under argon. Sodium (0.16 g, 7.00 mmol) was added as a solid and the solution was 531 stirred at 50 °C overnight. 5-Bromo-2-chloropyrimidine (1.00 g, 5.17 mmol) was added and 532 the solution stirred for 8 hours at 50 °C. The solution was filtered, water (20 mL) was added 533 and the crude extracted with ethyl acetate (3 x 30 mL). The product was purified by column 534 chromatography (ethyl acetate: hexane, 1:9, $R_f = 0.45$) to yield a colourless oil (0.80 g, 3.11 535 mmol, 60%).

¹H NMR (400 MHz, CDCl₃) δ : 8.51 (s, 2H), 5.81 (ddt, J = 6.67, 6.67, 10.18, 16.91, H), 4.98

538 ¹³C NMR (101 MHz, CDCl₃) δ: 164.15 (C), 159.75 (2CH), 138.59 (CH), 115.01 (CH₂),

539 111.79 (C), 68.42 (CH₂), 33.55 (CH₂), 28.36 (CH₂), 25.36 (CH₂).

540 **IR (film):** = 3076, 2936, 1640, 1570, 1432, 1332, 1176, 1122, 1024, 912, 794 cm⁻¹.

541 **HRMS (EI):** m/z calcd for $C_{20}H_{27}N_4O_2Br_2 [2M + H^+] 513.0501$, found: 513.0513.

The chemical structure of the compound 2 in Fig. 10 is given in Fig.14 and the synthesis and

543 characterization are detailed below:

544 545

FIG. 14. Chemical structure of compound 2

546 4-hydroxyphenylboronic acid (0.32 g, 2.33 mmol) and 1 (0.50 g, 1.94 mmol) were sealed in a schlenk under argon. Toluene (12 mL) was added and the solution degassed for 5 minutes. 547 548 Degassed methanol (7 mL) and potassium carbonate dissolved in degassed water (3 mL) were added to the schlenk. Tetrakis(triphenylphosphine)palladium(0) (0.11 g, 0.10 mmol) 549 550 was added as a solid and the solution was refluxed overnight. The crude was concentrated and water (30 mL) was added before extraction with DCM (3 x 30 mL). The organic phase 551 was dried with magnesium sulphate and concentrated in vacuo. The product was purified by 552 column chromatography (ethyl acetate: DCM, 15:85, where $R_f = 0.57$ in ethyl acetate: 553 554 hexane, 1:1) to yield a white solid (0.23 g, 0.85 mmol, 44%).

¹H NMR (400 MHz, CDCl₃) δ : 8.66 (s, 2H), 7.90 (s, 1H), 7.37 (d, J = 8.72, 2H), 7.03 (d, J = 8.71), 5.79 (ddt, J = 6.65, 6.65, 10.18, 16.91, 1H), 4.97 (m, 2H), 4.42 (t, J = 6.58, 2H), 2.11 (m, 2H), 1.85 (m, 2H), 1.58 (m, 2H).

¹³C NMR (101 MHz, CDCl₃) δ: 164.22 (C), 157.17 (C), 156.95 (2CH), 138.61 (CH), 128.48

559 (C), 127.98 (2CH), 126.19 (C), 116.73 (2CH), 114.97 (CH₂), 68.06 (CH₂), 33.55 (CH₂), 560 28.48 (CH₂), 25.38 (CH₂).

561 **IR (film):** = 3099, 3022, 2950, 1602, 1558, 1434, 1326, 1270, 1180, 1072, 924, 834 cm⁻¹.

HRMS (EI): The molecular weight for $C_{16}H_{19}N_2O_2$ [M + H⁺] was calculated as 271.1447

and found to be 271.1446.

The chemical structure of compound **3** is given in Fig. 15; synthesis and characterization are detailed below:

$$HO \longrightarrow N \longrightarrow O - (CH_2)_6 - Si - O - Si -$$

568 **2** (0.22 g, 0.81 mmol) was dissolved in dry THF (12 mL) under argon. 1,1,1,3,3,5,5-569 heptamethyltrisiloxane (0.35 g, 1.59 mmol) and platinum(0)-1,3-divinyl-1,1,3,3-570 tetramethyldisiloxane (0.20 mL of 0.1M solution, 0.02 mmol) were added to the flask and the 571 solution was stirred for 2 hours. The crude is concentrated and purified by column 572 chromatography (ethyl acetate: hexane, 2:8, where $R_f = 0.30$ in ethyl acetate: hexane, 3:7) to 573 yield a colourless wax (0.29 g, 0.59 mmol, 73%).

¹H NMR (400 MHz, CDCl₃) δ : 8.65 (s, 2H), 7.39 (d, J = 8.70), 6.96 (d, J = 8.72, 2H), 5.37

(s, 1H), 4.39 (t, J = 6.72), 1.83 (m, 2H), 1.42 (m, 6H), 0.54 (m, 2H), 0.08 (s, 9H), 0.06 (s, 6H,), 0.02 (s, 6H).

- ¹³C NMR (101 MHz, CDCl₃) δ: 164.71 (C), 157.08 (CH), 156.18 (C), 128.11 (CH), 128.08
- 578 (C) 127.30 (C), 116.48 (CH), 68.19 (CH₂), 33.36 (CH₂), 29.07 (CH₂), 25.91 (CH₂), 23.39
- 579 (CH₂), 18.47 (CH₂), 2.04 (3CH₃), 1.50 (2CH₃), 0.41 (3CH₃).
- 580 **IR (film):** = 2958, 1600, 1442, 1332, 1258, 1048, 840 cm⁻¹.
- 581 **HRMS (EI):** m/z calcd for $C_{23}H_{41}N_2O_4Si_3$ [M + H⁺] 493.2374, found: 493.2379.

The chemical structure of compound 4 is given in Figure 16 and other details are followed below:

584

585

FIG. 16. Chemical structure of compound 4

~____О

586 Trans-2-hexen-1-ol (1.00 g, 9.98 mmol), Shi Epoxidation Diketal Catalyst (0.77 g, 3.00 587 mmol) and Tetrabutylammonium hydrogensulfate (0.06 g, 0.18 mmol) were dissolved in 588 dimethoxymethane/acetonitrile (100 mL, 2:1). Acetic acid (0.35 mL) was added to potassium 589 carbonate (70 mL of 0.1M solution) and this was added to the reaction which was then cooled to -10 °C. Oxone (8.48 g, 13.78 mmol) was dissolved in ethylenediaminetetraacetic acid 590 disodium salt (50 mL of 0.004M solution in water) and potassium carbonate (8.00 g, 57.91 591 592 mmol) was dissolved in water (50mL). The oxone and the carbonate solution were added to 593 the reaction flask dropwise over 3 hours while keeping the temperature below 0 °C. Water 594 was added and the organic phase extracted with DCM (3 x 50 mL) before drying with 595 magnesium sulphate and concentrating the solution. The epoxide was purified by column chromatography (hexane: diethyl ether, 2:1 -> 100% ether once product appears, $R_f = 0.53$ in 596

597 DCM: ethyl acetate, 1:1) to yield a colourless oil (0.70 g, 5.99 mmol, 60%). The column was 598 compacted using hexane: diethyl ether, 2:1 and 1% triethylamine to neutralize the silica.

¹H NMR (400 MHz, CDCl₃) δ: 3.91 (ddd, J = 2.58, 5.58, 12.53, 1H), 3.63 (ddd, J = 4.30,

600 7.29, 12.52), 2.96 (td, J = 2.36, 5.50, 5.65, 1H), 2.92 (m, 1H), 1.74 (m), 1.50 (m, 3H), 0.96 (t, 1), 1.50 (m, 2H), 0.96 (t, 2H), 0.96

601 J = 7.26, 3H).

602 ¹³C NMR (101 MHz, CDCl₃) δ: 61.93 (CH₂, 58.55 (CH), 56.01 (CH), 33.78 (CH₂), 19.47

- 603 (CH₂), 14.10 (CH₃).
- 604 **IR (film):** = 3418, 2962, 2874, 1650, 1464, 1382, 1220, 1046, 900, 849 cm⁻¹.
- 605 $[\alpha]_D^{20}$: +38.75 (c 0.022, CHCl₃).

The chemical structure of adpc042 is given in Fig. 17 and other details are followed below:

608

FIG. 17. Chemical structure of compound adpc042

3 (0.56 g, 1.14 mmol), **4** (0.12 g, 1.03 mmol) and triphenylphosphine (0.33 g, 1.26 mmol) were dissolved in dry THF (20 mL) under argon. Diethyl azodicarboxylate (0.22 g, 1.24 mmol) was dissolved in dry THF (8 mL) under argon before being added dropwise to the reaction flask. The flask was stirred overnight and then concentrated. The product was purified by column chromatography (ethyl acetate: hexane, 1:9, where $R_f = 0.60$ in ethyl acetate: hexane, 3:7) to yield a white wax (0.43 g, 0.73 mmol, 71%).

615 ¹H NMR (400 MHz, CDCl₃) δ : 8.65 (s, 2H), 7.43 (d, J = 8.81, 2H), 7.02 (d, J = 8.81, 2H), 616 4.38 (t, J = 6.71, 2H), 4.23 (dd, J = 3.34, 11.08, 1H), 4.02 (dd, J = 5.52, 11.08), 3.13 (ddd, J =617 2.29, 3.27, 5.53, 1H), 2.98 (td, J = 2.20, 5.53, 5.64, 1H), 1.83 (m, 2H), 1.65-1.30 (m, 10H), 618 0.99 (t, J = 7.28), 0.54 (m, 2H), 0.08 (s, 9H), 0.06 (s, 6H), 0.01 (s, 6H).

619 ¹³C NMR (101 MHz, CDCl₃) δ: 164.76 (C), 158.92 (C), 157.10 (2CH), 127.95 (C), 127.89

620 (2CH), 127.68 (C), 115.71 (2CH), 68.82 (CH₂), 68.15 (CH₂), 56.66 (CH), 56.22 (CH), 33.84

621 (CH₂), 33.36 (CH₂), 29.06 (CH₂), 25.91 (CH₂), 23.39 (CH₂), 19.44 (CH₂), 18.46 (CH₂), 14.10

622 (CH₃), 2.04 (3CH₃), 1.50 (2CH₃), 0.40 (2(CH₃).

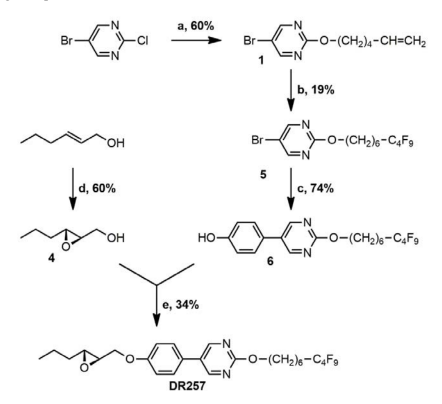
623 **IR (film):** = 2958, 1598, 1548, 1448, 1256, 1048, 840 cm⁻¹.

624 **HRMS (EI):** m/z calcd for $C_{29}H_{51}N_2O_5Si_3$ [M + H⁺] 591.3096, found: 591.3097.

625 $[\alpha]_D^{20}$: +9.74 (c 0.029, CHCl₃).

The synthesis of DR257 was carried out through the scheme shown in FIG. 18, further details

627 of each stage are provided below.



628



FIG. 18. General scheme for the preparation of DR257

630 Synthesis and characterization of the intermediates are given below. Structure of the631 compound 5 is given in Fig. 19 and details follow below:

$$Br \longrightarrow O-(CH_2)_6-C_4F_9$$

632 633

FIG. 19. The Chemical structure of compound 5

Na₂S₂O₄ (0.040 g, 0.230 mmol) and Na₂HPO₄ (0.039 g, 0.276 mmol) were added to a 634 635 mixture of 1-iodoperfluorobutane (0.955 g, 2.76 mmol), the alkene 1 (0.590 g, 2.30 mmol) in H₂O/CH₃CN (12 mL 1:3) [42]. The mixture was stirred overnight at room temperature. Et₂O 636 637 and H₂O were added to the mixture and the phases were separed. The organic phase was 638 extracted with Et₂O (2 X 10 mL) and the combined organic phase was dried with magnesium 639 sulphate. The solvent was removed at reduce pressure to led a yellow oil. This iodinated 640 intermediate was used without further purification. Bu₃SnH (0.802 g, 2.76 mmol) and a catalytic amount of AIBN (12 mg) were added to the oil. The reaction mixture was heated to 641

642 110° C for 24 h. The solvent was removed and the crude of reaction was purified by column 643 chromatography on silica gel (hexane:ethyl acetate = 9:1) to give **5** (0.208 g, 19% yield).

644 ¹H NMR (400 MHz, CDCl₃) δ : 8.51 (s, 2H), 4.33 (t, J = 6.5, 2H), 2.17 – 1.95 (m, 2H), 1.90

645 – 1.75 (m, 2H), 1.63 (m, 2H), 1.55 – 1.39 (m, 4H).

646 ¹³C NMR (101 MHz, CDCl₃) δ: 164.11 (C), 159.79 (2CH), 111.89 (C), 68.32 (CH₂), 31.15

647 (CH₂), 30.70 (t, J = 22.3, CH₂), 29.00 (CH₂), 28.69 (CH₂), 25.83 (CH₂), 20.29 (t, J = 3.7,

648 CH₂).

649 ¹⁹F NMR (**376 MHz, CDCl**₃) δ: -81.07, -114.61, -124.51, -126.06.

650 **HRMS (EI):** molecular weight calculated for $C_{14}H_{15}BrF_9N_2O [M + H^+] 477.0224 g$; found as 651 477.0208 g.

The chemical structure of compound 6 is given in Fig. 20 and other details follow:

654

653

FIG. 20. Chemical structure of compound 6

. .

655 4-hydroxyphenylboronic acid (0.070 g, 0.501 mmol) and 5 (0.220 g, 0.462 mmol) were 656 sealed in a schlenk under argon. Toluene (12 mL) was added and the solution degassed for 5 minutes. Degassed methanol (4 mL) and potassium carbonate dissolved in degassed water (1 657 658 mL) were added to the schlenk. Tetrakis(triphenylphosphine)palladium(0) (0.029 g, 0.025 659 mmol) was added as a solid and the solution was refluxed overnight. The crude was 660 concentrated and water (10 mL) was added before extraction with DCM (3 x 10 mL). The 661 organic phase was dried with magnesium sulphate and concentrated in vacuo. The product was purified by column chromatography (ethyl acetate:hexane, 3:7, where $R_f = 0.35$ in ethyl 662 acetate: hexane, 3:7) to yield a white solid (0.169 g, 0.345 mmol, 75%). 663

¹H NMR (400 MHz, CDCl₃) δ: 8.65 (s, 2H), 7.39 (d, J = 8.6, 2H), 6.96 (d, J = 8.5, 2H), 5.58

 $665 \qquad (s,\,1H),\,4.39\ (m,\,2H),\,2.07\ (m,\,2H),\,1.86\ (m,\,2H),\,1.65\ (m,\,2H),\,1.57-1.42\ (m,\,4H).$

666 ¹³C NMR (101 MHz, CDCl₃) δ: 164.56 (C), 157.09 (2CH), 156.32 (C), 128.25 (C), 128.09

667 (2CH), 127.09 (C), 116.51 (2CH), 67.79 (CH₂), 30.94 (t, *J* = 22.3, CH₂), 29.05 (CH₂), 28.86

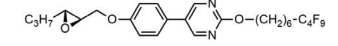
668 (CH₂), 25.90 (CH₂), 20.28 (t, J = 3.7, CH₂).

669 ¹⁹F NMR (**376 MHz, CDCl**₃) δ: -81.07, -114.62, -124.51, -126.06.

670 **HRMS (EI):** The molecular weight for $C_{20}H_{20}F_9N_2O_2$ [M + H⁺] calculated as 491.1381 g, but

671 was found as 491.1389 g.

The chemical structure of compound DR257 is given in Fig. 21 and the other details follow:



674

FIG. 21. Chemical structure of compound DR257

675 **6** (0.220 g, 0.449 mmol), **4** (0.048 g, 1.03 mmol) and triphenylphosphine (0.129 g, 0.490 676 mmol) were dissolved in dry THF (10 mL) under argon. Diethyl azodicarboxylate (0.085 g, 677 0.490 mmol) was dissolved in dry THF (4 mL) under argon before being added dropwise to 678 the reaction flask. The flask was stirred overnight and then concentrated. The product was 679 purified by column chromatography (ethyl acetate: hexane, 1.5:9, where $R_f = 0.60$ in ethyl 680 acetate: hexane, 3:7) to yield a white powder (0.82 g, 0.140 mmol, 34%).

681 ¹H NMR (400 MHz, CDCl₃) δ: 8.65 (s, 2H), 7.42 (m, 2H), 7.01 (m, 2H), 4.39 (t, J = 6.5,

682 2H), 4.23 (dd, J = 11.1, 3.3, 1H), 4.01 (dd, J = 11.1, 5.5, 1H), 3.12 (m, 1H), 2.97 (m, 1H),

683 2.06 (m, 2H), 1.90 - 1.80 (m, 2H), 1.69 - 1.40 (m, 10H), 0.98 (t, J = 7.3, 3H).

- ¹³C NMR (101 MHz, CDCl₃) δ: 164.63 (C), 158.94 (C), 157.09 (2CH), 128.08 (C), 127.87
- 685 (2CH), 127.56 (C), 115.70 (2CH), 68.81 (CH₂), 67.72 (CH₂), 56.61 (CH), 56.20 (CH), 33.81
- 686 (CH₂), 30.91 (t, J = 22.3, CH₂), 29.02 (CH₂), 28.84 (CH₂), 25.88 (CH₂), 20.25 (t, J = 3.7,
- 687 CH₂), 19.41 (CH₂), 14.06 (CH₃).
- ¹⁹F NMR (**376** MHz, CDCl₃) δ: -81.08, -114.63, -124.52, -126.08.
- 689 **HRMS (EI):** m/z calcd for $C_{26}H_{30}F_9N_2O_3$ [M + H⁺] 589.2097, found 589.2104.
- 690 $[\alpha]_D^{20}$: +8.868 (c 0.0106, CHCl₃).

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