Mesomorphism Dependence on Cis-Trans Configuration and Varying Terminal/Lateral Group

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Abstract: A chalconyl trans vinyl ester homologous series of thermotropic liquid crystals RO-C$_6$H$_4$-CH=CH-COO-C$_6$H$_4$-CH=CH-CO-C$_6$H$_3$ (OCH$_3$)$_2$ meta-para is synthesized and studied with a view to understand and establish the relation between molecular structure and mesomorphism or mesomorphic behaviors. Novel homologous series consists of thirteen homologues (C$_1$ to C$_{18}$). First five (C$_1$ to C$_5$) homologues are nonmesogenic. Rest of the homologues (C$_6$ to C$_{18}$) homologues is mesogenic. Monotropically smectogenic character appears from C$_6$ to C$_{10}$ homologue and C$_{12}$ to C$_{18}$ homologues are enantiotropically smectogenic. Nematogenic property is totally absent throughout a series. Textures of smectic phase are of the type smectic A or C. Transition curves of a phase diagram behaved in normal manner. Analytical, spectral and thermal data supports the molecular structures of homologues. Odd-even effect is observed for Sm-I/I-Sm transition curve. Thermal stability for smectic is 93.7$^\circ$C and the mesomorphic phase lengths from enantiotropic homologues for smectic from minimum to maximum is 6.0$^\circ$C to 28.0$^\circ$C at the C$_{12}$and C$_{18}$ homologue respectively. Series is smectogenic without exhibition of nematic property. The group efficiency order on the basis of thermal stability is derived; from comparative study of present series and analogous series.

Introduction

The study of mesomorphism and mesomorphic behaviors of liquid crystalline (LC) compounds [1] is one of the important and interesting aim, object and view to predict and exploited their characteristics in the benefit of mankind and living creatures or matters [2,3,4,5,6,7,8,9,10]. Therefore the object in view to understand and establish the effect of molecular structure [11,12,13,14,15,16] on liquid crystal properties with reference to varying flexibility and cis-trans molecular configuration as a consequence of combined effect of molecular rigidity and flexibility [17,18,19,20,21]. Present investigation is planned to synthesized novel LC compounds through homologous series containing three phenyl rings bond ed through two central bridges -CH=CH-COO- and -CH=CH-CO- and varying -OR terminal with fixed lateral -OCH$_3$ group throughout a novel series of proposed investigations. Study will include the characterization of novel compounds by thermal, analytical and spectral data. Results of the evaluated data will be discussed in terms of molecular rigidity and flexibility. The conclusions will include group efficiency order for smectic and nematic and probable importance and probable possibility of application will be suggested for the researchers working in application fields of present investigation. Number of homologous series have been reported till the date [22,23,24,25,26,27,28,29].

Experimental

Synthesis

4-n-Alk oxybenzaldehydes were synthesized by refluxing 4-hydroxybenzaldehyde (1 equiv.) with corresponding n-alkyl bromides (1 equiv.) in the presence of potassium carbonate (1 equiv.) and acetone as a solvent [30]. The resulting 4-n-alkoxybenzaldehydes were reacted with malonic acid (1.2 equiv.) in the presence of 1-2 drops piperidine as catalyst and pyridine as solvent to yield corresponding trans 4-n-alkoxy cinnamic acids (A) [31]. β-4-Hydroxy phenylα-3’-4’
dimethoxybenzoyl ethylene (B) was prepared by an established method [32] M.P. 148.0 °C, Yield-65.5%. Coupling of compound A and compound B is done by steglich esterification to yield β-4-(4'-n-alkoxy cinnamoyloxy) phenyl α-3”-4”dimethoxy benzoyl ethylene [33].

The synthetic route to the novel homologous series is under mentioned in scheme-1.

**Scheme-1: Synthetic route to the novel series**

- **Acetone, K₂CO₃, 4-5 hrs, Reflux**
- **RO-CHO**
- **Pyridine, 1-2 drops Piperidine**
- **H₂CO₂**
- **RO-CHO**
- **[A]**
- **CH₂Cl₂, DCC, 5 mol% DMAP**
- **RO-CH=CH-COO-[B]**

Where, R=CₙH₂n₊₁, n=1 to 8, 10, 12, 14, 16 and 18

**Characterization**

Some of members of a novel series as the representative member of a series were characterized by elemental analysis (Table-1), Infrared spectroscopy, ¹H NMR spectra and mass spectroscopy. Microanalysis was performed on EuroEA Elemental Analyzer. IR spectra were recorded on Shimadzu FTIR Model-IRAfinity-1S (MIRacle 10), ¹HNMR spectra were recorded on Bruker spectrometer using CDCl₃ as a solvent and mass spectra were recorded on Shimadzu GC-MS Model No.QP-2010.

**Analytical data**

**Table-1: Elemental analysis for (1) Pentyloxy (2) Heptyloxy (3) Dodecyloxy derivatives**

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Molecular formula</th>
<th>Elements %Found</th>
<th>Elements %Calculated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>C</td>
<td>H</td>
</tr>
<tr>
<td>1</td>
<td>C₃₁H₃₂O₆</td>
<td>74.44</td>
<td>6.42</td>
</tr>
<tr>
<td>2</td>
<td>C₃₃H₃₆O₆</td>
<td>75.05</td>
<td>6.83</td>
</tr>
<tr>
<td>3</td>
<td>C₃₈H₄₆O₆</td>
<td>76.28</td>
<td>7.71</td>
</tr>
</tbody>
</table>
Spectral Data

1H NMR in ppm for Methyloxy Derivative

3.87 (s, 3H, -CH₃ of -OCH₃ group of cinnamate ring), 3.94 (s, 6H, two -OCH₃ groups of dimethoxy phenyl ring), 7.38-7.42 (d, 1H, -CH=CH–CO–O–), 6.49-6.53 (d, 1H, -CH=CH–CO–), 7.55-7.57 (d, 1H, -CH=CH–CO–), 7.24-7.26 & 7.84-7.88 (4H, middle phenyl ring), 6.90-6.94 & 7.60-7.63 (4H, phenyl ring with single methoxy group), 7.17-7.18, 7.31-7.33 & 7.77-7.81 (3H, phenyl ring with two methoxy group). The NMR data are reliable with the molecular structure.

1H NMR in ppm for Propyloxy Derivative

1.03-1.08 (t, 3H, -CH₃ of -OCH₃ group), 1.80-1.89 (m, 4H, CH₂-CH₂-CH₂-O–), 3.97-3.98 (t, 2H, -CH₂-O–), 3.95 (s, 6H, two -OCH₃ groups of dimethoxy phenyl ring), 7.42-7.43 (d, 1H, -CH=CH–CO–), 6.49-6.53 (d, 1H, -CH=CH–CO–), 8.08-8.10 (d, 1H, -CH=CH–CO–), 7.54-7.56 (d, 1H, -CH=CH–CO–), 7.26-7.28 & 7.84-7.88 (4H, middle phenyl ring), 6.90-6.95 & 7.62-7.66 (4H, phenyl ring with propyloxy group), 7.17-7.18, 7.32-7.34 & 7.77-7.81 (3H, phenyl ring with two methoxy group). The NMR data are reliable with the molecular structure.

IR in cm⁻¹ for Butyloxy Derivative

3076 (C-H str. of alkene disubstituted), 2960 & 2868 (C-H str. of (-CH₂)ₙ group of -OC₄H₉), 1737 (C=O str. of carbonyl carbon of ester group), 1643 (C=O str. of α,β unsaturated ketone), 1587 (C=C str. of alkene), 1508 & 1421 (C=C str. of aromatic ring), 1357 (C-H bending of alkene disubstituted), 1305 (C-O str. of ether linkage), 1134 (C-O str. of ester group), 972 & 862 (C-H bending of alkene). The IR data are consistent with the molecular structure.

IR in cm⁻¹ for Decyloxy Derivative

3068 (C-H str. of alkene disubstituted), 2918 & 2846 (C-H str. of (-CH₂)ₙ group of -OC₁₀H₂₂), 1732 (C=O str. of carbonyl carbon of ester group), 1651 (C=O str. of α,β unsaturated ketone), 1629 (C=C str. of alkene), 1593, 1510 & 1463 (C=C str. of aromatic ring), 1354 & 1309 (C-H bending of alkene disubstituted), 1259(C-O str. of ether linkage), 1145 (C-O str. of ester group), 983 & 829 (C-H bending of alkene). The IR data are consistent with the molecular structure.

Mass spectra of Ethyloxy Derivative

m/z (rel.int%): 458 (M)⁺, 293, 175, 165, 133, 118, 103, 77

Mass spectra of Heptyloxy Derivative

m/z (rel.int%): 514 (M)⁺, 349, 231, 175, 165, 133, 118, 103, 77

Results and discussion

Transition temperatures of novel homologues are relatively lower as compared to the corresponding dimerised trans n-alkoxy cinnamic acids. Extension of a molecule by linking β-4-hydroxy phenyl α-3’-4’ dimethoxy benzoyl ethylene with trans 4-n-alkoxy cinnamic acid yielded mesomorphic homologues from C₆ to C₁₈ derivatives. C₆ to C₁₀ homologues are monotropicsmectic and the rest of the homologues C₁₂ to C₁₈ homologues are enantiotropicsmectic. None of the homologue is smectogenic in character. C₁ to C₅ homologues are nonliquid crystals. Transition temperatures (Table-2) as determined by an optical polarizing microscopy (POM) were plotted against the number of carbon atoms present in n-alkyl chain ‘R’ of the left n-alkoxy -OR terminal.
Table-2: Transition temperatures in °C

<table>
<thead>
<tr>
<th>Compound No.</th>
<th>n-alkyl chain C_nH_{2n+1} (n)</th>
<th>Sm</th>
<th>N</th>
<th>Isotropic</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>166.0</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>-</td>
<td>-</td>
<td>159.0</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>-</td>
<td>-</td>
<td>154.0</td>
</tr>
<tr>
<td>4</td>
<td>4</td>
<td>-</td>
<td>-</td>
<td>161.0</td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td>144.0</td>
</tr>
<tr>
<td>6</td>
<td>6 (94.0)</td>
<td>-</td>
<td>-</td>
<td>131.0</td>
</tr>
<tr>
<td>7</td>
<td>7 (107.0)</td>
<td>-</td>
<td>-</td>
<td>128.0</td>
</tr>
<tr>
<td>8</td>
<td>8 (104.0)</td>
<td>-</td>
<td>-</td>
<td>133.0</td>
</tr>
<tr>
<td>9</td>
<td>10 (112.0)</td>
<td>-</td>
<td>-</td>
<td>151.0</td>
</tr>
<tr>
<td>10</td>
<td>12</td>
<td>91.0</td>
<td>-</td>
<td>97.0</td>
</tr>
<tr>
<td>11</td>
<td>14</td>
<td>80.0</td>
<td>-</td>
<td>90.0</td>
</tr>
<tr>
<td>12</td>
<td>16</td>
<td>71.0</td>
<td>-</td>
<td>92.0</td>
</tr>
<tr>
<td>13</td>
<td>18</td>
<td>68.0</td>
<td>-</td>
<td>96.0</td>
</tr>
</tbody>
</table>

Sm- Smectic; N- Nematic; ( ) indicate monotropy

Figure-1: Phase Behavior of Series

Transition curves Cr-Sm/I, Sm-I/I-m are drawn through like or related points as depicted in a phase diagram (Figure-1) showing phase behaviors of a series. Cr-Sm/I transition curve adopted normal zigzag path of rising and falling with overall descending tendency. Sm-I/I-Sm transition curves for odd and even homologues initially rises from C_6, then it passes through maxima at C_{10} and finally descended upto C_{14} and then ascended by 4.0°C to 6.0°C for C_{16} and C_{18} homologues. Thus, it behaved almost in normal manner, with exhibition of odd-even effect from C_{6} to C_{10} homologue. Curves for I-Sm for odd-even members merge into each other at the C_{10} homologue and then odd-even effect disappears as series is ascended upto C_{18} homologue. The texture of smectic phase are of the type A or C. Thermal stability for smectic is 93.7°C whose mesophase lengths from 6.0 °C to 28.0 °C at the C_{12} and C_{18} homologues respectively. Mesomorphic properties from homologue to homologue in present series are varied from C_{6} to C_{18} homologue. C_{1} to C_{5} homologues are nonmesogenic. Lowering of transition temperatures and the disappearance of dimerisation of trans n-alkoxycinnamic acids are attributed to the breaking of hydrogen bonding.
between two molecules of carboxylic acids by the process of esterification. The exhibition of smectogenic character is attributed to the suitable magnitudes of intermolecular cohesion by end to end and lateral -OCH₃ groups which causes lamellar packing of molecules commencing from C₆ homologue to C₁₈ homologue to float the molecules on the surface with sliding layered molecular arrangement either in monotropic or enantiotropic condition. Odd-even effect observed for Sm-I or I-Sm transition curve is due to the sequentially added methylene unit at the n-alkyl chain ‘R’ of -OR flexible group. The disappearance of odd-even effect from and beyond C₁₀ homologue is attributed to the coiling bonding or flexing or coupling of longer in alkyl chain or ‘R’ with the major axis of a core structure of a molecule. The non-mesomorphicity observed for C₁ to C₅ is due to their high crystallizing tendency arising from low magnitudes of dispersion forces and low magnitudes of dipole-dipole interactions which abruptly breaks crystal lattices and sharply transforms into isotropic liquid directly from crystalline solid with bypassing of LC state, a high order of molecular disorder occur and molecules are randomly oriented in all possible directions. However, on cooling the isotropic mass for C₆ to C₁₀ homologues below their isotropic temperature, the monotropic smectic mesophase appears in irreversible manner. None of the C₁ to C₅ homologue did exhibit monotropic mesomorphism, but C₁₂ to C₁₈ homologue exhibited smectic mesophase formation enantiotropically in reversible manner due to the presence of highly polar two -OCH₃ groups bonded as terminal and lateral groups; the molecular flexibility contributed in combination with flexibility due to -OR group, facilitated to induce smectogenic character in reversible manner. The absence of nematic property is attributed to the weaker or unsuitable magnitudes of end to end intermolecular cohesion and closeness occurred due to meta substituted -OCH₃ group, which hindered the nematic mesophase formation but favored smectogenic mesophase formation through the suitable magnitudes of molecular polarizability contribution. The variations in mesogenic properties of present novel series-1 are compared with the structurally similar analogous series-X [34] and Y [35] as mentioned below in figure-2.

**Figure-2: Structurally similar series.**

Above novel homologous series-1, X and Y appear identical with respect to three phenyl rings, two central bridges and left n-alkoxy (-OR) group for the same homologue from series to series. But they differ with respect to Trans and cis configuration. Present homologous series-1 is of Trans configuration and the homologous series-X and Y selected for comparison are with cis configuration. Therefore their molecular rigidity differs from series to series keeping the molecular flexibility due to left -OR flexible group unaltered from series to series for the same homologue. Moreover, the right handed groups bonded to third phenyl rig (viz. meta-para dimethoxy (-OCH₃)₂, para -OCH₃ and meta -NO₂) differs in polarities and polarizability affecting molecular flexibility for the same homologue from series to series. Thus, variations in mesomorphic properties and mesomorphic behaviors can vary with the changing features and their magnitudes of molecular rigidity and/or flexibility for the same homologue from series to series. Following table-3 represents some thermometric mesogenic properties and behaviors for the present series-1 and series-X and Y chosen for comparative study.
is related to intermolecular cohesion forces of attractions in case of solarizability as well as transluc nematic i

Therefore mesophase commences from C\text{5} to C\text{6} neighboring homologues. Lowest thermal stability

<table>
<thead>
<tr>
<th>Series ( \rightarrow )</th>
<th>( 1 )</th>
<th>X</th>
<th>Y</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sm-I or I-Sm or Sm-N</td>
<td>93.7 (C\text{12}-C\text{18})</td>
<td>-</td>
<td>154.6 (C\text{5}-C\text{8})</td>
</tr>
<tr>
<td>Commencement of</td>
<td>C\text{6}</td>
<td>-</td>
<td>C\text{5}</td>
</tr>
<tr>
<td>Smectomesophase</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N-I</td>
<td>-</td>
<td>164.6 (C\text{5}-C\text{14})</td>
<td>163.8 (C\text{5}-C\text{16})</td>
</tr>
<tr>
<td>Commencement of</td>
<td>-</td>
<td>C\text{5}</td>
<td>C\text{5}</td>
</tr>
<tr>
<td>nematic phase</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Mesophase length</td>
<td>6.0 (C\text{12})</td>
<td>10.0 (C\text{14})</td>
<td>10.0 (C\text{16})</td>
</tr>
<tr>
<td>range (Sm+N)</td>
<td>to 28.0 (C\text{18})</td>
<td>to 58.0 (C\text{8})</td>
<td>to 46.0 (C\text{6})</td>
</tr>
</tbody>
</table>

Above table-3 represent that

- Present series-1 and X are only smectogenic and only nematogenic respectively, whereas series-Y is smectogenic plus nematogenic.
- Smectogenic mesophase commences from C\text{6} to C\text{5} homologues for series-1 and Y respectively whereas it does not commence till the last homologue of series-X.
- Nematogenic mesophase formation commences identically from C\text{5} homologue for series-X and Y whereas it does not commence till the last member of a present series-1.
- Smectic thermal stability of present series-1 is lower than a series-Y whereas it does not stabilize for series-X.
- Nematic thermal stability of series-X and Y are almost equivalent (164.6 $\approx$ 163.8) whereas it does not stabilize for series-1 of present investigation.
- Upper mesophase length are in increasing order form series-X $>$ Series-Y $>$ series-1 and the lower mesophase lengths equal for series-X and Y and more than series-1, i.e. series-X = series-Y $>$ series-1.

The exhibition of only smectic or nematic or both mesophases smectic plus nematic is attributed to the suitable magnitudes of anisotropic forces of intermolecular end to end attractions and lateral attractions causing statistically parallel orientational order or/and sliding layered molecular organization for definite range of temperature as result of favorable molecular rigidity and flexibility under floating condition depending upon polarities and/or polarizability of terminal or/and lateral group or groups. Homologous series-1 and X contain two and one highly polar methoxy (-OCH\text{3}) groups or group respectively, whose polarity or polarizability as well as trans-cis configuration causes variations in intermolecular cohesion and closeness of magnitudes in more or less proportions. In present case of series-1 which bears one more polar and polarizable lateral -OCH\text{3} group as compared to series-X which strengthens the possibility of lamellar packing of molecules from C\text{6} to C\text{18} homologues in their preoccupied crystal lattices which facilitated appearance of smectic mesophase formation either irreversibly or reversibly in monotropic or enantiotropic condition respectively. But in case of series-X the intermolecular cohesion is undergone reduction with the decreasing number of -OCH\text{3} group of lateral substitution. Therefore end to end attractions predominated to cause the appearance of nematic phase only in case of series-X. The polarity and polarizability of meta substituted -NO\text{2} group being more and the more suitable than the para substituted -OCH\text{3} group which shows the exhibition of smectic plus nematic phase due to the favorable magnitudes of dispersion forces, dipole-dipole interactions, permanent dipole moment across the long molecular axis related to intermolecular cohesion forces of attractions in case of series-Y. The early or late commencement of mesophase or mesophases depends upon the extent of molecular non-coplanarity. The homologous series-X and Y are more noncoplanar (cis configuration) than present series-1 which is a little bit more coplanar or less noncoplanar. Therefore mesophase commences from C\text{5} to C\text{6} neighboring homologues. Lowest thermal stability
(smectic) and the total mesophase lengths of series-1 as compared to series-X and Y are attributed to its symmetrical trans configuration of homologue molecules. Thus, variations in mesogenic behaviors, properties, and the degree of mesomorphism for the same homologue from series to series are attributed to variations in cis-trans configurations, molecular rigidity and flexibilities, and the changing magnitudes of dipolarities and polarizability of molecules.

Conclusions

- Present novel series is only smectogenic with low mesophase lengths, low thermal stability and low ordered melting type.
- The group efficiency order derived on the basis of (a) thermal stability (b) early commencement mesophase and (c) total upper and lower mesophase lengths for smectic and nematic are as under.
  
  (a) Smectic: -NO₂> (-OCH₃)₂ > -OCH₃
  Nematic: -OCH₃ ≈ -NO₂ > (-OCH₃)₂
  
  (b) Smectic: -NO₂> (-OCH₃)₂ > -OCH₃
  Nematic: -NO₂ = -OCH₃ > (-OCH₃)₂
  
  (c) Smectic + Nematic:
  Upper: -OCH₃ ≈ -NO₂ > (-OCH₃)₂
  Lower: -OCH₃ = -NO₂ > (-OCH₃)₂

- Mesomorphism is very sensitive and susceptible to a molecular structure.
- Present investigation may be useful for LC devices to be operated at room temperature through study at binary system.
- Presence of chalconyl group being antimicrobial and antifungal and bioactive with presence of added ester group may be useful for reducing consumption of pesticides and insecticides as well as to produce agricultural products of better healthy quality.
- Present investigation has raised and supported conclusions drawn earlier.

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References


