Molecular Flexibility Operated Mesomorphism

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Keywords: Mesomorphism; Nematic; enantiotropy; Liquid Crystal; Smectic

ABSTRACT. Novel homologous series: RO-C₆H₄-CH=CH-COO-CH₂-C₆H₄-Br (p) synthesized and studied with a view to understand and establish the effects of molecular structure on liquid crystal (LC) behavior of a series. Series consists of eleven homologues. C₁ to C₇ members of a novel series are non-liquid crystals. Mesomorphism commences from Octyloxy (C₈) homologue and continued upto hexadecyloxy homologue (C₁₆) as enantiotropic nematic without exhibition of smectic property. The textures of a nematic phase are threaded or Schlieren. Transition temperatures (table-3) and textures are determined by an optical polarizing microscopy equipped with a heating stage. Cr-N/I and N-I transition curve behaved in normal manner in phase diagram, showing their phase behavior (figure-2). Odd-even effect is absent for N-I transition curve. Analytical and spectral data confirmed the molecular structures of a series. The LC properties are compared with the structurally similar series. The transition temperatures are relatively lower than the corresponding n-alkoxy benzoic acid. Thus, present novel series is partly nematogenic with absence of smectic property whose mesogenic phase length is low and of low ordered melting type. Thermal stability for nematic is 93.0°C and the mesophase length ranges from 8 to 21°C.

1. INTRODUCTION

Liquid crystalline (LC) state [1] of substance is an intermediate state of existence between crystalline solid and isotropic liquid, which is partly crystalline and partly liquidous, flowing as liquid on the surface but possess an optical property as crystal. Exploiting such abnormal behaviours, it has number of applications in the benefit of mankind as to manufacture LC devices and proven its ability in the biological systems [2-8]. Present investigation is planned with a view to understand and establish the effects of molecular structure on LC properties [9-12] with references to the molecular flexibility, keeping molecular rigidity unaltered for all homologues of a homologous series. Therefore, novel LC substances will be generated as the alternative source of LC materials, useful to the group of scientific research community specifically working in the application part and it will enhance the knowledge of constructing LC materials workable as per our desire and need. Number of homologous series have been reported till the date [13-21] with ester, central or terminal group. Thus, novel ester series with three phenyl rings and two central bridges –COO- and –CH=CH-COO-CH₂- will be synthesized, characterized and its thermometric evaluated data will be interpreted and compared with structurally similar series on the basis of molecular rigidity and flexibility [22-25]. Comparative study will help to understand and establish the effects of molecular structure on mesomorphic properties.

2. EXPERIMENTAL

4-Hydroxy benzoic acid, Alkyl halides, Methanol, KOH, 4-bromo benzy1 bromide or alcohol, N,N-Dimethyl Formamide, NaHCO₃, HCl, Dry Pyridine, Thionyl chloride, EtOH etc required for the synthesis were used as received, except solvents which were dried and purified prior to synthesis.
2.1 General method for the preparation of n-alkoxy benzoic acid and 4-Hydroxy-4-bromobenzyl cinnamate.

4-Hydroxy benzoic acid was alkylated by suitable alkylating agent R-X to form n-alkoxy benzoic acid by modified method of Dave and Vora [26] as main component of a series. 4-Hydroxy-4'-bromobenzyl cinnamate (m.p:95-96°C) was prepared from 4-bromo benzyl bromide or alcohol and 4-Hydroxy Cinnamic acid by applying the method of European patent and the modified method of Doshi, Patel and Marathe [27].

2.2 General method for the preparation of final esterified products.

The n-alkoxy benzoic acid through their corresponding acid chlorides and 4-Hydroxy-4'-bromobenzyl cinnamate were condensed in dry cold Pyridine to form final products by usual established method [28]. Final esterified products were individually decomposed, filtered, washed, dried and purified till their constant transition temperatures obtained. Synthetic route to the series is mentioned below as scheme-1.

Scheme-1: Synthetic route to the novel series
3. CHARACTERIZATION

The selected members of a novel homologues series were characterized and analyzed by elemental analysis and the structure elucidation by Infra-red spectra, \(^1\)HNMR spectra and mass spectra. Textures and Transition temperatures of homologues as well as of related materials were determined by an optical polarizing microscope, equipped with a heating stage. Elemental analysis was performed on Perkin-Elemer PE 2400 C,H,N analyzer. The percentage halide content is determined by usual analytical method. IR spectra were recorded on Shimadzu FTIR Model-IRAfinity-1S (MIRacle 10). \(^1\)HNMR spectra were determined on Bruker spectrometer using CDCl3 solvent and mass spectra were recorded on Shimadzu GC-MS Model No.QP-2010. Textures of nematic mesophase of some homologues were recognized either directly from the microscopic observations or by miscibility method.

4. ANALYTICAL DATA

Table-1: Elemental analysis for (1) Ethoxy (2) Propyloxy and (3) Hexyloxy derivative

<table>
<thead>
<tr>
<th>Sr.No.</th>
<th>Molecular formula</th>
<th>% Elements calculated (Experimental %)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>C</td>
</tr>
<tr>
<td>1</td>
<td>C(<em>{25})H(</em>{21})BrO(_{5})</td>
<td>62.37 (65.22)</td>
</tr>
<tr>
<td>2</td>
<td>C(<em>{26})H(</em>{24})BrO(_{5})</td>
<td>62.90 (63.20)</td>
</tr>
<tr>
<td>3</td>
<td>C(<em>{29})H(</em>{29})BrO(_{5})</td>
<td>64.80 (64.80)</td>
</tr>
</tbody>
</table>

Spectral Data:

\(^1\)HNMR data for series

\(^1\)HNMR in ppm for the Methoxy Derivative: 1.07-1.19 (-CH\(_3\) groups), 2.66 (–CH\(_3\)), 5.13-5.27 (–CH=CH–), 6.90-6.92 (p-substituted phenyl ring), 7.85-7.91 (p-substituted benzene).

\(^1\)HNMR in ppm for the Dodecylxylo Derivative: 0.86-88 (–CH\(_3\) of C\(_{12}\)H\(_{25}\)), 1.24-1.77 (–CH\(_2\)- of –C\(_{12}\)H\(_{25}\)), 3.97-3.99 (–O–CH\(_2\)), 5.16-5.30 (–CH=CH–), 6.90-6.93 (p-substituted phenyl ring), 7.30-7.40 (p-substituted benzene).

NMR data confirm the structure.

IR in cm\(^{-1}\) for Butyloxy Derivative

2954, 2870 (C-H str.), 1465 (str. of (–CH\(_2\)-) group of –OC\(_4\)H\(_9\)), 1675 (–COO- ester group), 1604 (C-H stretching of –CH=CH–), 1512 (C=C str. of alkene), 1396 (C-H bending of alkene disubstituted), 1257 (C-O str. of ether linkage), 1165 (C-O str. of ester group), 1016 (Aromatic linkage of -Br), 848 (para substituted phenyl ring), 771 (Polymethylene group), 640-695 (-CH\(_2\)- Stretch). The IR data are consistent with the molecular structure.

IR in cm\(^{-1}\) for Pentyloxy Derivative

2931, 2870 (C-H str.), 1489 (str. of (–CH\(_2\)-) group of –OC\(_4\)H\(_{11}\)), 1597 (–COO- ester group), 1712 (C-H stretching of –CH=CH–), 1512 (C=C str. of alkene), 1396 (C-H bending of alkene), 1257 (C-O str. of ether linkage), 1165 (C-O str. of ester group), 1010 (Aromatic linkage of -Br), 848 (para substituted phenyl ring), 756 (Polymethylene group), 648-689 (-CH\(_2\)- Stretch). The IR data are consistent with the molecular structure.

Mass spectra of Tetradecyloxy Derivative

Theoretical Mass = 649.0
Experimental Mass = 649.80

5. RESULT AND DISCUSSION

LC homologues or non-liquid crystal homologues excluding Octyloxy and onward (C\(_8\) to C\(_{16}\)) novel homologues are the derivatives of dimeric 4-n-alkoxy benzoic acids. The transition temperatures of novel homologues are lower than corresponding to 4-n-alkoxy benzoic acids.
Enantiotropic nematic phase commences late from C\textsubscript{8} homologue of a series, with absence of smectogenic character. Numbers of carbon atoms of n-alkyl chain are plotted versus the transition temperatures of left n-alkoxy polar end group. Like or related points are linked to form Cr-N/I and N-I transition curves, showing phase behaviours of series in a phase diagram (figure-2). Odd-even effect is absent for N-I transition curve. The analytical and spectral data confirmed the molecular structures of homologues. The changing Thermometric behaviours from homologue to homologue in a series varied keeping right sided tail end group intact throughout the novel series. Negligible abnormal behaviour is attributed to more polarizable –Br end group and the effect of unexpected status of longer n-alkyl chain linked with first phenyl ring through oxygen atom. Thus, present series is partly nematogenic without exhibition of smectic property whose mesophase length is relatively shorter, low thermal stability (93.0°C) and lower ordered melting type.

The disappearance of dimerization of 4-n-alkoxy benzoic acids and the lowering of transition temperatures of present homologues are attributed to the breaking of hydrogen bonding between two molecules of aromatic acids by esterification process. The exhibition of only nematic phase from C\textsubscript{8} homologue attributed to the commencement of suitable magnitudes of anisotropic forces of end to end intermolecular attractions, which causes an environmental situation of floating molecules to float on the surface with statistically parallel orientational order with restricted magnitudes of entropy (\(\Delta S=\Delta H/T\)), within definite range of temperature. However, the intermolecular attractions are failed to build up lamellar packing of molecules to cause sliding layered arrangement to induce smectic mesophase formation. Thus, absence of smectogenic character and exhibition of only nematic property is facilitated under the influence of exposed thermal vibration late from C\textsubscript{8} homologue. The early homologues, prior to commencement of nematogenic mesophase are abruptly breaking their crystal lattices and sharply transform directly from solid crystal lattices to isotropic liquid without passing through LC state due to their high crystallizing tendency which arises from their inability to resist exposed thermal vibrations as a consequence of unfavourable magnitudes of combined effect of molecular rigidity and flexibility induced by low dipole-dipole interactions and the low magnitudes of dispersion forces through interactions between instantaneous dipole produced by the spontaneous oscillations of electron clouds of the molecules. All the molecules of mesomorphic homologues on heating above the isotropic temperatures randomly orient in all possible directions with high order of disorder or with high order of unrestricted entropy. But, on cooling the same, a reversible nematic mesophase reappears from and below isotropic temperature till it solidify to crystalline state. The absence of odd-even effect in N-I transition curve is due the unsuitable magnitudes of anisotropic forces of intermolecular end to end attractions which produced by the homologues of shorter n-alkyl chain of left n-alkoxy terminal end group. Negligible abnormality in N-I transition curve for C\textsubscript{14} homologue is attributed to its unusual status of longer n-alkyl chain. The changing trends in LC properties from homologue to homologue in the present novel series is attributed to the changing number of methylene unit or units of n-alkyl chain, keeping the rest of the molecular part unchanged throughout the novel series under discussion. Some LC properties of present series-1 are compared with the structurally similar homologous series-X [29] and Y [30] as under in figure-1.

Figure-1: Structurally similar series.
Homologous series-1, X and Y are identical with respect to three phenyl rings and two central bridges –COO- and –CH=CH-COO-CH2- which contributes to the total molecular rigidity. They are also identical with respect to n-alkoxy (-OR) terminal end groups for the same homologue from series to series. However, series-1, X and Y differ with respect to tail end groups –Br, -OCH₃ and –Cl which contributed to the total molecular flexibility from homologue to homologue in the same series and for the same homologue from series to series. Thus, the variations in mesomorphic (LC) properties and the degree of mesomorphism will depend upon the differing features and their magnitudes of combined effects of total molecular rigidity and flexibility. Following table-2 represent some LC properties of series-1, X and Y under comparative study.

<table>
<thead>
<tr>
<th></th>
<th>Series→ 1(-Br)</th>
<th>X(-OCH₃)</th>
<th>Y(-Cl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nematic – Isotropic Or Smectic-Nematic</td>
<td>-</td>
<td>-</td>
<td>92.78 (C₁ - C₁₆)</td>
</tr>
<tr>
<td>Commencement of Smectic phase</td>
<td>C₈</td>
<td>C₁</td>
<td>C₈</td>
</tr>
<tr>
<td>Nematic – Isotropic</td>
<td>93.0</td>
<td>211.54</td>
<td>100.8 (C₈ – C₁₆)</td>
</tr>
<tr>
<td>Commencement of Nematic phase</td>
<td>(C₈ - C₁₆)</td>
<td>(C₁ – C₁₆)</td>
<td>C₈</td>
</tr>
<tr>
<td>Total Mesophase length range in °C from Ti to Tj</td>
<td>8.0 to 21.0</td>
<td>17.0 to 72.0</td>
<td>21.7-72.2</td>
</tr>
<tr>
<td>Ci</td>
<td>C₁₄</td>
<td>C₁₄</td>
<td>C₆</td>
</tr>
<tr>
<td>Cj</td>
<td>C₈</td>
<td>C₃</td>
<td>C₈</td>
</tr>
<tr>
<td></td>
<td>C₁₆</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table-2 indicates that,
- Homologous series-1 and X are only nematogenic whereas series-Y is smectogenic in addition to nematogenic.
- Smectic mesophase commences from very first member of a series-Y, but it does not commence till the last member of the series-1 and X.
- Nematic mesophase commences from very first member of a series-X, but it commences late from C₈ membered homologue for series-1 and Y.
- Nematic thermal stabilities are in increasing order from series-1 to series-Y to series-X.
- Smectic thermal stability of series-Y is 92.78 but smectogenic mesophase did not stabilize for series-1 and X.
- Total mesophase length ranges are varied in decreasing order from series-X to series-Y and series-1.

Variations in the mesogenic behaviours of the series under comparison are attributed to the magnitudes of molecular flexibility arising from changing tail end groups –Br, -OCH₃ and –Cl, whose individual group polarities group atomicity, inductive effect, effects due to molecular length to breadth ratio, ratio of the polarity to polarizability, permanent dipole moment across the long molecular axes, dipole-dipole and electronic interactions. C-Xi (where Xi= terminal tail group)
bond polarity, dispersion forces, Vander Waals forces, the suitable magnitudes of anisotropic forces of intermolecular attractions as a consequence of varied molecular flexibility for the same homologue from series to series and from homologue to homologue in the same series etc; show variations in the mesogenic behaviours from series-1 to series-X to series-Y. Mono atomic –Cl tail end group of series-Y is more efficient and capable to build up lamellar packing of molecules in the crystal lattices as compared to –OCH3 and –Br terminal ends of greater size, which maintained sliding layered molecular arrangement for series-Y but fails to maintain the same for –OCH3 and –Br of series-1 and X to facilitate smectic mesophase formation. The formation of nematic mesophase of low thermal stability by –Br and –Cl terminal end groups is attributed to their low dipolarity of C-Xi bond but, -OCH3 terminal end group being highly polar and polarizable as compared to –Br and –Cl enhances nematic thermal stability more than halogen C-Xi substituent. Thus, resistivity towards exposed thermal vibrations being comparatively more for the series involving –OCH3 tail group stabilizes and facilitates nematic mesophase formation at the cost smectic phase with highest degree of mesomorphism.

6. CONCLUSIONS

- Presently investigated novel series is partly nematogenic without exhibition of smectogenic character of low melting and short degree of mesomorphism.
- Group efficiency order derived for smectic and nematic on the basis of (i) thermal stability (ii) early commencement of mesophase (iii) the total mesophase length.

(i) Smectic

\[
\begin{align*}
-\text{CL} & \gg -\text{Br} = -\text{OCH}_3 \\
\text{Nematic} & \\
-\text{OCH}_3 & \gg -\text{CL} \gg -\text{Br}
\end{align*}
\]

(ii) Smectic

\[
\begin{align*}
-\text{CL} & \gg -\text{Br} = -\text{OCH}_3 \\
\text{Nematic} & \\
-\text{OCH}_3 & \gg -\text{CL} = -\text{Br}
\end{align*}
\]

(iii) (Sm+N) Total Mesophase Length

\[
\begin{align*}
-\text{OCH}_3 & \gg -\text{CL} \gg -\text{Br}
\end{align*}
\]

- Present investigation may be useful for the study of binary system for LC devices workable between 57°C and 90°C.
- Present novel compounds may be useful as fire resistance material.
- Thus, present investigation supports the conclusions drawn earlier and raises the credibility to the existing literature on LC.
### Table-3: Transition temperatures in °C

<table>
<thead>
<tr>
<th>Compound No.</th>
<th>n-alkyl chain C&lt;sub&gt;n&lt;/sub&gt;H&lt;sub&gt;2n+1&lt;/sub&gt; (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>94</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>-</td>
<td>-</td>
<td>103</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>-</td>
<td>-</td>
<td>92</td>
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<tr>
<td>4</td>
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<td>89</td>
</tr>
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<td>5</td>
<td>-</td>
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<td>97</td>
</tr>
<tr>
<td>12</td>
<td>16</td>
<td>-</td>
<td>78</td>
<td>86</td>
</tr>
</tbody>
</table>

Sm- Smectic; N- Nematic

**Figure-2: Phase Behavior of Series**

**Acknowledgements**

Authors acknowledge thanks to the Green Circle Inc. Laboratory for providing research facilities services as and when needed. Authors acknowledge thanks to the Mr. Baldev Prajapati of Rajesh Pharma Ankleshwar, for providing the free sample of 4-Bromobenzyl bromide. Authors are also thankful to Dr. Vipul Patel and Dr. M.L. Chauhan, P.T. Arts and Science College, Godhara, for their valuable helping hand and microscopic facility. Also thanks are due to the Sophisticated Analytical Instrumentation Facility, Punjab University, (Chandigarh) for extending their help for analytical services.
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