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## Molecular Crystals and Liquid Crystals

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### Synthesis and Mesomorphic Properties of the Homologous Series of 4-Alkyl or Alkoxy-4'-Bromo or Cyanotolanes

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# Synthesis and Mesomorphic Properties of the Homologous Series of 4-Alkyl or Alkoxy-4'-Bromo or Cyanotolanes†

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A homologous series of 4-alkyl or alkoxy-4'-bromo or cyanotolanes in which the alkyl group is  $C_4 \rightarrow C_{10}$  and the alkoxy group is  $C_1 \rightarrow C_{12}$ , has been prepared. Interphase transitions between solid, mesomorphic and isotropic phases were studied by hot stage microscopy and differential scanning calorimeter. The bromo compounds have a smectic polymorphism. Three pure products of cyano derivatives with the chains  $C_9H_{19}O$ ,  $C_{10}H_{21}O$  and  $C_{10}H_{21}$  present an enantiotropic or monotropic reentrant nematic phase at atmospheric pressure. The reentrant phenomenon itself will be discussed as well as the absence of correlation between  $T_{NA}/T_{NI}$  McMillan parameter and the heat of transition. A plot of the nematic isotropic transition temperatures against the number of carbon atoms in the alkoxy chain shows the usual odd even effect.

#### INTRODUCTION

Some reentrant nematic phases have been described in binary mixtures at atmospheric pressure<sup>1</sup> or in pure materials but at high pressure.<sup>2</sup> After the first observations of enantiotropic reentrant nematic phases at atmospheric pressure in pure compounds of two series: 4-n-alkoxybenzoyloxy-4'-cyano stilbenes<sup>3-5</sup> and then, the 4-n-alkoxybenzoyloxy-4'-cyanobiphenyls,<sup>6</sup> we have carried on the systematic study of such series with three phenyl rings

<sup>†</sup> Presented at the third Liquid Crystal Conference of Socialist countries, Budapest, Hungary, August 1979.

in the rigid core and a cyano end group. All these compounds present the same general formula.

$$R - COO - CN$$

where X is:  $-CH = CH = ,^{3-5}$  single bound,  $^6 - CH = N - ,^7 - COO - ,^8 - N = N - ,^9$  we present here the results obtained with  $X = -C \equiv C -$  for which preliminary results have been published.  $^{10}$ 

#### RESULTS AND DISCUSSION

Compounds of these series are prepared according to the scheme below:

HO—CH=CH—CH—CH—Br

$$CH_{3}-COO$$
—CH=CH—CH—Br

$$\downarrow Br_{2}$$

$$\downarrow KOH$$

$$C_{2}H_{3}OH$$

$$CH_{3}-COO$$
—CHBr—CHBr—Br

$$\downarrow Br$$

$$\downarrow R$$

$$\downarrow R$$

$$\downarrow R$$

$$\downarrow COO$$
—Br
$$\downarrow R$$

$$\downarrow COO$$
—CEC—Br
$$\downarrow R$$

$$\downarrow COO$$
—CEC—CN
$$\downarrow R$$

$$\downarrow R$$

$$\downarrow COO$$
—COO—CEC—CN
$$\downarrow R$$

The 4-bromo-4'-hydroxystilbene was prepared according to Ref. [11]. The mesophases of the prepared compounds were observed by means of a polarizing microscope equipped with a heating and cooling stage (Mettler

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TABLE 1

Transition temperatures and enthalpies of fusion of compounds with formula

R         K         S <sub>B</sub> S <sub>A</sub> N         1         Kcal/m           C <sub>n</sub> H <sub>2n+1</sub> O         n         164         —         —         275         3.94           2         1466         —         —         271         8.6           3         158         —         271         8.6           4         158.5         —         271         8.6           5         1113         —         259         747           6         1111         —         220         211         4.10           8         122         —         163         225         8.12           9         117         —         182         211         7.07           10         110         —         182         210         6.8           11         114         —         —         189         202         10.8           12         106         —         —         189         196         10.4           12         107         —         189         196         10.4           11         11         —         110.4         10.4         10.4           11 <th></th> <th></th> <th></th> <th><b>22</b></th> <th></th> <th>-coo(</th> <th></th> <th>)=c≡c</th> <th><math>  \bigcirc  </math></th> <th>) →Br</th> <th></th> <th></th> <th></th> <th></th>				<b>22</b>		-coo(		)=c≡c	$  \bigcirc  $	) →Br				
O n 1 164 — — — — — — — — — — — — — — — — — — —	R		K		S		$S_{\mathbf{B}}$		$S_{A}$		Z		-	ΔHf Kcal/mole
1       164       —       —       275         2       146       —       —       271         3       158       —       —       271         4       158.5       —       —       273         5       123       —       —       239         6       111       —       —       192       211         9       117       —       —       182       225         10       110       —       —       182       221         10       110       —       —       182       221         11       114       —       —       188       202         11       114       —       —       189       202         12       106       —       —       189       202         12       106       —       —       189       202         13       —       —       —       189       202         13       —       —       —       189       202         13       —       —       —       189       204         5       107       —       —       189	2n+10	u										i		
2 146		_		164	1		1		l			275		3.94
3 158		7		146	I		1		1			271		9.8
4       158.5       —       —       259         5       123       —       —       239         6       111       —       —       192       211         7       128       —       —       163       225         8       122       —       —       182       211         10       110       —       —       182       211         11       114       —       —       189       202         11       114       —       —       189       202         12       106       —       —       189       196         13       —       —       —       189       196         1       128       —       —       189       196         5       107       —       —       119.5       224.6         6       97.6       (78)       (101)       167       205         7       103.4       —       163       195         8       8       —       101       167       195		3		158	1		1		ł			262.5		3.2
5       123       —       —       239         6       111       —       —       192       211         7       128       —       —       163       225         8       122       —       —       182       211         9       117       —       —       182       211         10       110       —       —       189       202         11       114       —       —       189       202         12       106       —       —       189       196         13       —       —       —       189       196         13       —       —       —       189       196         13       —       —       —       189       196         5       107       —       —       (118.5)       224.6         6       97.6       (78)       (101)       167       205         7       103.4       (79.6)       (101)       167       205         8       8       —       (101)       167       195		4		158.5	1		1		l			259		7.47
6 1111		5		123	1	•	1		1			239		3.70
7       128       —       —       163       225       .         8       122       —       —       182       .       221       .         9       117       —       —       182       .       221       .         10       110       —       —       .       188       .       .       .         11       114       —       —       .		9	٠	111	I	•	1		l	192		211		4.15
8 122		7	٠	128	ł		1		l	163	,	225		8.12
9 117		œ	٠	122	1	•			ı	182		221		4.10
10		6		117	1	•	1			182		211		7.07
11		10		110	I		1			188		210		8.9
12       106       —       —       189       196       .         n       131       —       —       237       .         4       128       —       —       (118.5)       224.6         5       107       —       (79)       119.5       218.4         6       97.6       (78)       (84.3)       141       205         7       103.4       (79.6)       (101)       167       206         8       85       —       (84.1)       163       195		=		114	I	•	1		,	189		202		8.01
n       3       131       —       —       237         4       128       —       —       (118.5)       224.6         5       107       —       —       (79)       119.5       218.4         6       97.6       (78)       (84.3)       141       205         7       103.4       (79.6)       (101)       167       206         8       85       —       (84.1)       163       195		12		106	i	•	ļ		•	189		196	٠	10.4
3       131       —       —       237         4       128       —       —       224.6         5       107       —       (79)       119.5       218.4         6       97.6       (78)       (84.3)       141       205         7       103.4       (79.6)       (101)       167       206         8       85       —       (84.1)       163       195		п												
. (79) . (118.5) . 224.6	- -	m		131	I	,	ı		Į			237		5.05
. (78) . (19.5 . 218.4		4	-	128	I	,	1			(118.5)		224.6	٠	5.87
(78) (84.3) 141 205 (79.6) (101) 167 206 (84.1) 163 195 (		5	٠	107	1			(62)		119.5		218.4		6.3
(79.6) (101) 167 206		9		9.76		(78)		(84.3)		141		205		5.5
. (84.1) . 163 . 195 .		7		103.4		(9.67)		101)		167		506		6.5
		∞	٠	82	1			(84.1)	٠	163		195		5.2

The meanings of the signs used in this table and in the following are: K: crystalline phase; S: smectic phase (s); SA. SB.... smectic phases A, B ...; N: nematic phase; I: isotropic phase; .. the phase exists; - ; the phase does not exist. The temperatures are given in Celsius degrees. Metastable transitions are indicated between bracked.

TABLE II
Transition temperatures and enthalpies of fusion of compounds with formula

	f nole			νο ·ν
	ΔHf Kcal/mole	5.8 9.3 8.6 7.6	12 9.8 7 9.9	4.9 4.1 5.8 5.95 5.8 7.25
		276 268 256 256	239 233.5 227 223	258 242 234 238 227 217
Z	z			
			183 208 213 216	188
	S <sub>A</sub>			11111
C			141	(62)
	z	1111		1111
-000 <del>-</del>		(105) (107) 108 96	(75.7)	(78.5) (63) (53) (44.5)
	S,			1 1
₩.		107 113 102 86	90 84 85 92.5	115 100 87 78 71 65
	×			
		E 2 0 1 8	9 10 11 12	n 5 7 7 8 9
	×	C,H2,+10		$C_nH_{2n+1}$

The notations are that of Table I.

FP 5). The transition temperatures and enthalpies of fusion were determined by means of a differential scanning calorimeter (Dupont 990). The results are listed in Tables I and II.

The structures of the smectic phases were identified using the contact method by their isomorphy with a known reference compound.<sup>12</sup>

#### 1 p-bromo substituted compounds

When  $R = C_n H_{2n+1}O$ , the first five compounds  $(n = 1 \rightarrow 5)$  are only nematic. From the hexyloxy, the compounds exhibit a smectic A phase in addition to the nematic phase. We point out that the stability range of the smectic A phase for even n is about 82°C and for odd n about 65°C. The nematic existence range becomes smaller while the smectic A range becomes larger. The odd even effect on the clearing point is rather well followed, except for the compound with n = 6 (Figure 1). At first, we thought it was a problem of purity of the sample but repeated careful purifications lead to the same transition temperatures. In fact, we have to point out that this compound is the first in this homologous series where the  $S_A$  phase is observed. The same results have been obtained on our series with  $X = CH_2$ — $CH_2$ .

When  $R = C_n H_{2n+1}$ , the compound with n = 3 is only nematic, the butyl derivative has a monotropic  $S_A$  phase. The hexyl and heptyl compounds exhibit in addition, two monotropic smectic phases S and  $S_B$ . The textures of the  $S_A$  and  $S_B$  phases are either focal conics or homeotropic, the textures of the S phase are striated focal conics.

#### 2 p-cyano substituted compounds

These are colourless, stable materials having wide thermal range of mesomorphism (Table II). When  $R = C_n H_{2n+1} O$ , the two derivatives (n = 5, 6) exhibit an enantiotropic nematic phase and a monotropic  $S_A$  phase. The enantiotropic  $S_A$  phase appears from n = 7. The nonyloxy derivative presents a stable reentrant nematic phase; below this nematic phase and from  $75.7^{\circ}C$ , another smectic A phase is observed. It is the third compound having a tetramorphism  $N S_A H S_A$ . The two others are the octyloxy and the nonyloxy of the series.

$$R - COO - CH = CH - CN$$

Another compound with such tetramorphism was then reported. The decyloxytolane derivative presents three enantiotropic phases with increasing temperature N,  $S_A$ , N. From the undecyloxy derivative the

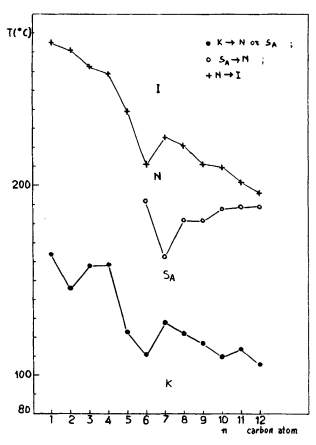
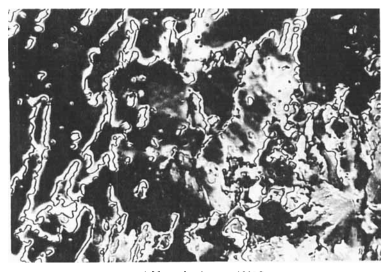


FIGURE 1 Plot of transition temperatures against n, the number of carbon atoms in the alkoxy chain of

$$C_nH_{2n+1}O$$
— $COO$ — $C$  $\equiv$  $C$ — $Br$ 

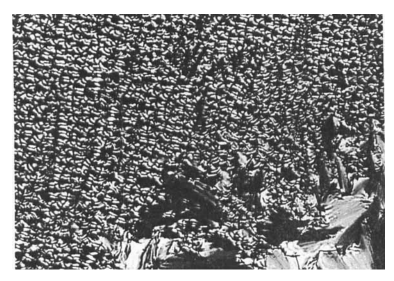
reentrant nematic phase disappears. On cooling the isotropic liquid of the nonyloxy derivative one can observe the nematic phase with a classical schlieren textures (Figure 2a). Below this nematic phase, the smectic A phase with a focal conic or homeotrope texture appears (Figure 2b). On further cooling another schlieren is observed (Figure 2c) followed by a focal conic or homeotropic texture (Figure 2d). The identification of the tetramorphism N, S<sub>A</sub>, N, S<sub>A</sub>, of the nonyloxy derivative has been made by the miscibility method with the four corresponding phases of the 4-cyano-4'-octyloxybenzoyloxystilbene (Figure 3). The odd even effect on the clearing point is rather well followed (Figure 4, Table II).



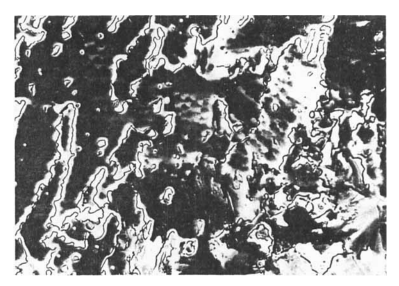
a) Nematic phase at 189°C;

FIGURE 2 Textures of the compound

$$C_9H_{19}O$$
— $COO$ — $C\equiv C$ — $CN$ 



b) Smectic A phase at 160°C:



c) Reentrant nematic phase at 130°C;



d) Reentrant smectic phase A at 74°C.

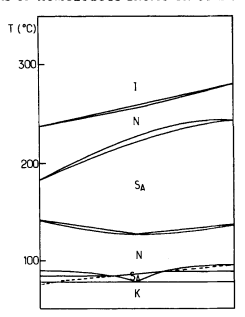


FIGURE 3 Diagram of isobaric state for the mixture of

$$C_9H_{19}O$$
—COO—C $\equiv$ C—CN (on left)

with

$$C_8H_{17}O$$
—COO—CH=CH—CH—CN (on right)

When  $R = C_n H_{2n+1}$ , the pentyl compound is only nematic. From the hexyl to the nonyl, these compounds have, in addition, a monotropic  $S_A$  phase. The decyl derivative exhibit a monotropic reentrant nematic phase. This enantiotropic  $S_A$  phase and this monotropic reentrant nematic phase have been checked by miscibility with the corresponding phases of the decyloxy derivative (Figure 5). This last alkyl derivative exhibits a wide  $S_A$  range (123°C) and has the lowest melting point (65°C). This phenomenon with an alkyl chain is recently observed by Madhusudana *et al.*<sup>14</sup> in the series:

$$C_nH_{2n+1}$$
 COO COO COO COO

n = 11, 12

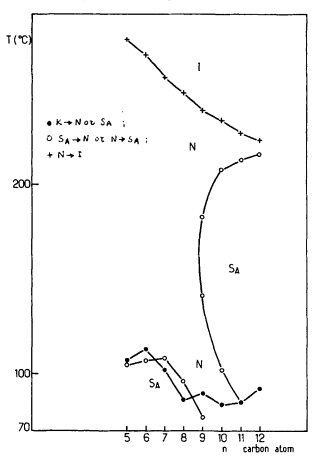


FIGURE 4 Plot of transition temperatures against n, the number of carbon atoms in the alkoxy chain of

$$C_nH_{2n+1}O$$
— $COO$ — $C\equiv C$ — $CN$ 

In the series of the cyanotolanes, another remarkable fact is the evolution of the  $T_{NA}/T_{NI}$  ratio calculated for each compound in Table III.

 $T_{\rm NA}$ ,  $T_{\rm NI}$  are respectively the temperature in Kelvin of the smectic A-nematic transition at the highest temperature and the nematic-isotropic transition. As in the series of 4-alkoxy-benzoyloxy-4'-cyanostilbenes recently discussed,<sup>5</sup> this parameter is almost constant and largely inferior to the McMillan's number <sup>15</sup> ( $\simeq$  0.87) up to octyloxy and nonyl and changes abruptly for nonyloxy and decyl. From the nonyloxy the evolution of this parameter with the

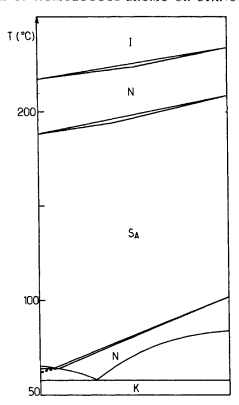


FIGURE 5 Diagram of isobaric state for the mixture of

$$C_{10}H_{21}$$
  $COO$   $C\equiv C$   $CN$  (on left)

with

$$C_{10}H_{21}O - \bigcirc -COO - \bigcirc -CN (on right)$$

highest  $T_{NA}$  seems to agree with the McMillan's theory. It seems that there is a close relation between this parameter and the existence of the reentrant nematic and smectic A phases at low temperature.

We can also compare compounds having the same general formula:

$$C_9H_{19}O$$
— $COO$ — $COO$ — $COO$ 

with various linkage X = -CH = CH -, -C = C -, single bond, -COO -, -CH = N -, -N = N -. All these compounds exhibit a

 $TABLE\ III$   $T_{NA}/T_{NI}\ ratio\ of\ compounds\ with\ formula$ 

TABLE IV

Transition temperatures of compounds with formula

$$C_9H_{19}O$$
— $COO$ — $X$ — $CN$ 

-x-	K		$\mathbf{S}_{\mathbf{A}}$		N		$S_{\mathbf{A}}$		N		Ĭ
-сн=сн-		97		(63)		(93.7)		261		275.3	_
single bond		96		, ,		(71)		217		232	
–č≡c–	,	90		(75.7)		141		183		239	
-coo-		121				(116)		198		228.5	
-CH=N-		96				(92)		228		251	
-N=N-		90	•	(72)		118		214		253	

reentrant nematic phase (Table IV), have a highly polar cyano group attached to one end of the molecule which results in strong antiparallel correlations between neighbouring molecules.<sup>16</sup> This leads to a formation of a bilayer structure with interdigitated molecules in each bilayer.<sup>17</sup> As the temperature is varied the structure is slightly modified and this change is favourable for the occurrence of the reentrant nematic phase.<sup>27</sup>

#### CONCLUSION

We show that 4-alkyl or alkoxy benzoyloxy-4'-cyanotolanes exhibit another example of pure substances with a stable nematic reentrant phase at atmospheric pressure. Our results show that these new materials have the same general feature than the previous ones.

Now, we must point out several new interesting facts:

- the alkoxy series is the first one which presents two stable reentrant nematic phases. One of them exhibit a tetramorphism N, S<sub>A</sub>, N, S<sub>A</sub>
  - the alkyl series present also a monotropic reentrant nematic phase
- the bromo derivatives do not present a reentrant nematic phase but give a polymorphism smectic.

#### **EXPERIMENTAL**

The infrared spectra have been recorded using a Perkin Elmer 225 spectrophotometer and the NMR ones with a Bruker 270 MHz.

*p-Alkoxybenzoic acids* They were prepared from the *p*-hydroxybenzoic-acid and the selected alkyl bromide following the method of Gray *et al.*<sup>18</sup>

p-Alkylbenzoic acid chlorides They were prepared from the alkylbenzene and the oxalyl chloride following the method of Reynolds et al.<sup>19</sup>

4-Acetyloxy-4'-bromostilbene The acetyl chloride (8.6 g, 0.11 mole) was added dropwise to a mixture of the 4-bromo-4'-hydroxystilbene<sup>11</sup> (27.5 g, 0.1 mole) in anhydrous pyridine (200 ml). The mixture stirred magnetically at 0°C for five hours and at room temperature for 20 hours. The reaction mixture was poured onto a mixture of crushed ice (400 g) and concentrated hydrochloric acid (200 ml) and stirred for half hour. It was filtered off, washed with water, 10% aqueous sodium hydroxide solution, water. The product was recrystallized from a mixture of solvents CHCH<sub>3</sub> +  $C_2H_5$ —OH (1:2) into white needles: yield, 80%, m.p. 159°C.

4-Acetyloxy-4'-bromostilbene dibromide Bromine (29 g, 0.18 mole) was added to a solution of 4-acetyloxy-4'-bromostilbene (25.3 g, 0.08 mole) in chloroform (600 ml). This solution was stirred for 3 hours. The excess of bromine was destroyed by the addition of an aqueous solution of sodium metabisulfite. The product was isolated by filtration. It was recrystallized from ethanol. Yield: 76%, m.p. 220°C (dec.)

4-Bromo-4'-hydroxytolane The stilbene dibromide (29.5 g) was added to a hot solution of potassium hydroxyde (84 g) in absolute ethanol (180 ml).<sup>20</sup> The reaction mixture was stirred, heated in oil bath at 140°C for 24 hours and allowed to cool. It was poured onto mixture of crushed ice (180 g) and concentrated hydrochloric acid (150 ml) and stirred for half hour. It was filtered

off, washed with water and recrystallized from a mixture of solvents:  $CH_2Cl_2$  + hexane (30:70). This procedure gave 4.4 g (26%) of the required product m.p. 182°C.

4-Bromo or cyano-4'-alkyl or alkoxybenzoyloxytolane All the alkyl and alkoxy have been prepared following the same general procedure. We give only the example of the preparation of the octyloxy derivatives.

4-Bromo-4'-octyloxy benzoyloxy tolane A mixture of 4-octyloxy-benzoic acid (0.55 g, 21 mmol) and thionyl chloride (2 ml) was refluxed for 2 hours and the excess thionyl chloride was removed under reduced pressure. The acid chloride was added to a solution of 4-bromo-4'-hydroxytolane (0.5 g, 1.8 mmol) in anhydrous pyridine (4 ml). The mixture was stirred magnetically at room temperature for 24 hours and poured onto a stirred mixture of crushed ice and concentrated hydrochloric acid. The product was filtered off, washed with water and air dried. It was chromatographed on silica gel and eluted with the mixture benzene-hexane (1:1) and the finally recrystallized from ethanol. Yield 0.6 g (65%), K 117 SA 199 N 231 I

I.R. (Nujol = 1730 cm<sup>-1</sup> (COO stretching) 840 cm<sup>-1</sup> (
$$-$$
 )  
pmr (CDCL<sub>3</sub>):  $\delta$ : 0.905 ( $t$ -3, CH<sub>3</sub> of  $-$  C<sub>8</sub>H<sub>17</sub>),1.3-2.06 (m-12, six  $-$  CH<sub>2</sub>),  
4.05 ( $t$ -2 CH<sub>2</sub>O), 6.96-8.16 (six  $d$ -12 aromatic)  
Calculated (C<sub>29</sub>H<sub>29</sub>O<sub>3</sub>Br): C% 68.91; H%, 5.74; Br%, 15.84  
Found : C%, 69.08; H%, 5.85; Br%, 16.09

4-Cyano-4'-octyloxybenzoyloxytolane Cuprous cyanide (0.27 g, 3 mmole) was added to a solution of 4-bromo-4'-octyloxybenzoyloxytolane (0.51 g, 1 mmole) in N-methyl-2-pyrolidone (1 ml). This mixture was stirred and heated in an oil bath at 200°C for 2 hours. The cooled reaction mixture was poured onto a solution of ferric chloride (0.8 g) in water (15 ml and concentrated hydrochloric acid (0.4 ml) and heated at 60°C for 30 min. The organic product was extracted into ether. The etheral solution was washed with water, dried (anhydrous sodium sulfate) and then evaporated to dryness. It was chromatographed on silica gel and eluted with benzene and finally recrystallized from ethanol. Yield 0.25 g (53%) K 86 S<sub>A</sub> 96 N 248 I

pmr (CDCL<sub>3</sub>) $\delta$ (ppm) 0.905 (t . 3 CH<sub>3</sub> of C<sub>8</sub>H<sub>17</sub>), 1.308–2.005 (m-12, six CH<sub>2</sub>)

4.056 (t-2, CH<sub>2</sub>O), 6.97-8.16 (six d-12, aromatic)

Calculated: (C<sub>30</sub>H<sub>29</sub>O<sub>3</sub>N): C%, 79.82; H%, 6.43; N%, 3.10

Found : C%, 79.68; H%, 6.50; N%, 3.15

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