

Evaluation of the surface properties of 4-(Decyloxy) benzoic acid liquid crystal and its use in structural isomer separation

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Abstract: The selectivity of 4-(Decyloxy) benzoic acid (DBA) liquid crystal in surface adsorption region (303.2–328.2 K) and thermodynamic region (423.2 – 433.2 K) was investigated by inverse gas chromatography at infinite dilution (IGC-ID). The selectivity parameters of the structural isomer series named butyl acetate, butyl alcohol, and amyl alcohol series were calculated for the DBA using IGC-ID technique. Additionally, the surface properties including dispersive surface energy (γ_s^D), free energy (ΔG_A^S), enthalpy (ΔH_A^S), and acidity-basicity constants were calculated with net retention volumes obtained from IGC-ID experiment results. When the ΔH_A^S and ΔG_A^S are constants, DBA surface was found to be an acidic character ($K_D/K_A \cong 0.89$).

Key words: Inverse gas chromatography, liquid crystal, selectivity, surface properties

1. Introduction

The liquid crystal (LC) state, which is different from the known forms, was discovered in the 1880s during the studies of Reinitzer [1] and Lehmann [2] on some cholesterol esters. LC state, also called as the meso-phase, is known as a physical state located between the crystalline solid and the isotropic liquid phase. Although it is considered as a separate phase between the solid and liquid phase, it can have at least one property of the solid and liquid phase. In the LC structure, the spontaneous orientation of the molecules in a certain direction provides a geometric selectivity to the stationary phase [3–6]. The orientation of the molecules in the LC structure can also differentiate the properties and usage fields of the LC. LCs are widely used in sensor technology, technological devices, such as televisions, computers, tablets, and biological fields [7–10].

In the conventional gas chromatography (GC), certain stationary phases suitable for the chromatographic column studied are used. The probes in volatile form to be analysed in the GC are separated from each other depending on their polarity on conventional stationary phases. It is not possible to analyse high molecular weight and nonvolatile materials, such as LCs, polymers, composites, etc. in conventional GC [11–15]. Therefore, IGC-ID is a simple, low cost, high efficiency, and high accuracy technique developed to analyse such substances. This technique is based on filling the substances to be analysed into the chromatographic column as a stationary phase and retaining the probes passed over them in vapor form at different times [16–18].

Separation of the isomer series is crucial industrially. When LCs are used as stationary phase for separation of isomer series, generally better, more efficient results can be obtained compared to conventional stationary phases. By using the IGC-ID technique, faster and more accurate results can be obtained compared to conventional separation methods [19–23].

Surface properties are closely related to important physicochemical phenomena such as colloidal stability, stickiness, and wettability. Besides, the surface properties, especially the surface energy, is an extremely important parameters in understanding the interaction between the surface of the material and various probes [24–26]. Surface energy arises from unbalanced molecular forces on the surface of the materials. The surface energy of the materials can be analysed using liquid adsorption, flow microcalorimetry, and contact angle measurements. Since the application of these techniques is difficult and limited, IGC-ID has become a preferred technique by researchers [27–31].

In the scope of this study, DBA's ability to separate isomer series including butyl acetate series (n-butyl acetate (nBAC), iso-butyl acetate (iBAC) and tert-butyl acetate (tBAC)), butyl alcohol series (n-butyl alcohol (nBAL), iso-butyl alcohol (iBAL)

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and tert-butyl alcohol (tBAL)) and amyl alcohol series (n-amyl alcohol (nAAL), iso-amyl alcohol (iAAL) and tert-amyl alcohol (tAAL)), and surface properties were investigated by IGC-ID technique. The selectivity of DBA was investigated in surface adsorption (303.2–328.2 K) and thermodynamic region (423.2–433.2 K). Additionally, the IGC-ID experiments were carried out to investigate the surface properties of DBA in relation to polar and nonpolar probes in surface adsorption region (303.2–328.2 K). Using the retention data obtained from IGC-ID experiments, the parameters used to determine the selectivity parameters and the surface properties were calculated.

2. Theory of inverse gas chromatography at infinite dilution (IGC-ID)

2.1. The selectivity coefficient

To determine selectivity of materials, the net retention volumes (V_N) in surface adsorption region and thermodynamic region should be calculated as main data. For volatile polar and nonpolar solvents used in the analysis, the V_N is closely related to the interaction of these solvents with the materials [32–36]. V_N is calculated as follows:

$$V_N = QJ(t_R - t_A)T/T_f \quad (1)$$

Here, t_R and t_A are retention times of volatile probes and air, respectively; Q is the volumetric flow rate; T and T_f are the column and ambient temperature, respectively; J is James-Martin pressure correction factor.

The selectivity of the stationary phase contained in the chromatographic column can be calculated from the proportioning of the numerical difference between the retention times obtained from the IGC-ID experiments. Besides, selectivity coefficient can also be calculated from the ratio of V_N calculated according to Eq. (1). The selectivity of stationary phase is determined depending on the size of the selectivity coefficient (α). This value is calculated as follows [37,38]:

$$\alpha = (t_{R1} - t_A) / (t_{R2} - t_A) = V_{N1} / V_{N2} \quad (2)$$

Here, t_{R1} and t_{R2} are the retention time of the first and second isomer from the isomer pairs, respectively; t_A is the retention time of air; V_{N1} and V_{N2} are the net retention volume of the first and second isomer, respectively.

2.2. Surface properties

In recent years, IGC-ID is commonly used for examining the surface properties of the materials. The standard free energy (ΔG_A°) value for the adsorption of volatile probes on the stationary phase is calculated with the help of the V_N resulting from the interaction between probe and stationary phase [39–41]. ΔG_A° is calculated as follows:

$$\Delta G_A^\circ = -RT \ln(V_N) + K \quad (3)$$

Surface energy is an extremely important parameter in explaining the interaction between stationary phase and volatile probes. The greater the surface energy, the more interactions between molecules. On the contrary, when this energy is low, the interaction decreases. Surface energy of the stationary phase (γ_s) can be calculated as a sum of dispersive energy (γ_s^D) generated by weak interactions on surface and specific energy (γ_s^S) generated by strong interactions on surface [42–44]:

$$\gamma_s = \gamma_s^S + \gamma_s^D \quad (4)$$

γ_s^D of stationary phase is determined when non-polar probes are injected at Henry's law region. This energy is due to dispersive interactions between molecules on the surface of the material and non-polar probe molecules [45]. γ_s^D values can be calculated in the surface adsorption region according to the method proposed by Dorris–Gray [46] as follows:

$$\gamma_s^D = (\Delta G_{[CH_2]})^2 / 4(N_A)^2(a_{[CH_2]})^2\gamma_{[CH_2]} \quad (5)$$

Here, γ_s^D is the dispersive energy of the surface (mj/m²), $\Delta G_{[CH_2]}$ is the adsorption free energy of a methylene group, which is determined the slope of the plot between the number of alkanes versus $RT \ln V_N$ values, N_A is the Avogadro's number, $a_{[CH_2]}$ is the molecular area of a methylene group (0.06 nm²) and $\gamma_{[CH_2]}$ is the surface energy of a methylene group. $\gamma_{[CH_2]}$ values are calculated at any temperature (t °C) as follows [47]:

$$\gamma_{[CH_2]} = 35.6 - 0.058t \quad (6)$$

Additionally, the method proposed by Schultz is widely used to calculate dispersive energy of surface [48]. This energy is calculated as follows:

$$-RT \ln(V_N) = 2N_A a (\gamma_s^D)^{0.5} (\gamma_L^D)^{0.5} + K \quad (7)$$

Here, a is the cross-sectional area of the probes, N_A is the Avogadro's number, γ_L^D is the dispersive energy of the probes. The a and γ_L^D values were taken from the literature, and were listed in Table 1. γ_s^D values of stationary phase can be calculated from the slope of plot between $RT \ln V_N$ versus $a(\gamma_L^D)^{0.5}$ of non-polar probes.

$\Delta G_{[CH_2]}$ values are calculated as follows [49]:

$$\Delta G_{[CH_2]} = -RT \ln(V_{N,n} / V_{N,n+1}) \quad (8)$$

Here, R is the universal gas constant; $V_{N,n}$ and $V_{N,n+1}$ are the net retention volumes of two n-alkanes having n and $n+1$ carbon atoms, respectively.

ΔG_A^S for the polar probes are calculated as follows:

$$\Delta G_A^S = -RT \ln(V_N / V_{N(ref)}) \quad (9)$$

Table 1. The values of a and γ_L^D for non-polar and polar probes.

Probes	$a(\times 10^{-10} \text{ m}^2)$	$\gamma_L^D \text{ (mj/m}^2\text{)}$
n-Hexane (Hx)	51.5	18.4
n-Heptane (Hp)	57.0	20.3
n-Octane (O)	62.8	21.3
n-Nonane (N)	69.0	22.7
n-Decane (D)	75.0	23.4
Dichloromethane (DCM)	31.5	27.6
Chloroform (TCM)	44.0	25.9
Tetrahydrofuran (THF)	45.0	22.5
Ethyl acetate (EA)	48.0	19.6
Acetone (Ace)	42.5	16.5

When the studies are carried out at different temperatures, ΔH_A^S and ΔS_A^S values can be calculated as follows [50]:

$$\Delta G_A^S = \Delta H_A^S - T\Delta S_A^S \quad (10)$$

The value of ΔH_A^S is linked with K_A (donor or acidity group) and K_D (acceptor or basicity group) parameters. This situation is due to the interactions that occur between probes and surfaces that do not have dispersive and entropic interactions. These values are calculated as follows [51,52]:

$$-\Delta H_A^S = K_A(\Delta N) + K_D(AN^*) \quad (11)$$

Here, DN is an electron donor or acidity number and AN^* is an electron acceptor or basicity number determined by Gutmann [53]. By calculating the value of ΔH_A^S for polar probes, a linear plot is drawn between $-\Delta H_A^S/AN^*$ and $\Delta N/AN^*$. The values of K_A and K_D of solid materials can be obtained from the slope and intercept of the line, respectively. If $K_D/K_A > 1$, the surface is considered to be a basic; whereas, if $K_D/K_A < 1$, the surface is considered to be an acidic.

3. Materials and methods

All the properties of the chemicals used in this study are given in Table 2.

All measurements in IGC-ID studies were carried out using an Agilent Technologies HP-6890N device combined with thermal conductivity detector (TCD) (Hewlett-Packard, Palo Alto, CA, USA). The stainless-steel column (1/8" o.d., 2.10 mm i.d. 10 m) was purchased from Alltech Associates, Inc. (Chicago, IL, USA). Chromosorb W (AW-DMCS-treated, 80/100 mesh) was used as the support material and obtained from Sigma Aldrich. The DBA liquid crystal was dissolved in the Chloroform, and Chromosorb W was added slowly. A homogeneous mixture was obtained by continuous stirring in heating controlling water bath, and the LC was coated on support. Silane-treated glass wool used to plug the ends of the column was obtained from Alltech Associates Inc (Deerfield, IL, USA). The ends of the column were loosely plugged with silanized glass wool. After the column was cut to a size of 1 m and cleaned thoroughly, approximately 1.21 g of the prepared column interior material was filled. The total loading of DBA liquid crystal on the support was determined as 10.38% by weighing. Helium (He), which kept at a constant flow rate of 3.6 mL/min, was used as the mobile phase during the experiments. Probes and air were injected into the column with 1 mL and 10 mL Hamilton syringes, respectively. For infinite dilution, the probe (0.1 mL) was taken into the syringe and flushed into the air. Then, the retention times for probe and air were determined. At least four consecutive injections were made for each probe and air at each set of measurements.

4. Results and discussion

The main data (V_N) obtained from IGC-ID studies were calculated for all probes injected surface adsorption region (303.2–328.2 K) and thermodynamic region (423.2–433.2 K) according to Eq. (1). The retention diagrams of DBA used as separator stationary phase in two regions were given in Figure 1 and 2, respectively. The " α " values for nBac/iBac, nBac/tBac, nBal/iBal, nBal/tBal, nAal/iAal, and nAal/tAal were obtained using their V_N in two regions. " α " values calculated according to Eq. (2) determined the separation ability of DBA. The higher the values of the separation factor calculated according to Eq. (2), the better the selectivity for isomers. Table 3 and 4 shows the calculated values for the isomer pairs in two regions. Considering these values, it is seen that isomers are separated. Besides, it was observed that structural isomers were better separated in the surface adsorption region than in the thermodynamic region.

Table 2. Source, assay, and CAS registry numbers of the chemicals.

Chemicals	Source	CAS No	Assay
DBA	Sigma Aldrich	5519-23-3	0.980
nBAc	Supelco	123-86-4	≥ 0.995
iBAc	Sigma Aldrich	110-19-0	≥ 0.980
tBAc	Sigma Aldrich	540-88-5	≥ 0.990
nBAL	Sigma Aldrich	71-36-3	≥ 0.994
iBAL	Supelco	78-83-1	≥ 0.990
tBAL	Sigma Aldrich	75-65-0	≥ 0.990
nAAl	Sigma Aldrich	71-41-0	≥ 0.990
iAAl	Sigma Aldrich	123-51-3	≥ 0.990
tAAl	Supelco	75-85-4	≥ 0.990
Hx	Supelco	110-54-3	≥ 0.997
Hp	Supelco	142-82-5	≥ 0.990
O	Sigma Aldrich	111-65-9	≥ 0.990
N	Sigma Aldrich	111-84-2	≥ 0.990
D	Sigma-Aldrich	124-18-5	≥ 0.940
EA	Supelco	141-78-6	≥ 0.998
Ace	Supelco	67-64-1	≥ 0.998
DCM	Supelco	75-09-2	≥ 0.998
THF	Supelco	109-99-9	≥ 0.998
TCM	Supelco	67-66-3	≥ 0.998

The main data (V_N) obtained from IGC-ID studies was calculated for all probes between 303.2 and 328.2 K according to Eq. (1). Retention diagrams of non-polar and polar probes were given in Figure 3 and 4, respectively.

The surface energy of a solid materials depends on the chemical structure, physical properties, and composition. Interactions between molecules on a solid surface and polar or nonpolar probe molecules are due to long- and short-range interactions known as weak interactions (London dispersive forces) and strong interactions (acid-base interactions). Dispersive surface energy occurs as a result of nonspecific interactions caused by the London dispersive forces known as weak or long-range interactions [54]. γ_s^D can be calculated using IGC-ID technique based on well-known approaches for data analysis, such as Dorris–Gray (Eq. (5)) and Schultz (Eq. (7)) methods. In these α_s^D calculations, homologous alkane vapor series are used in infinite dilution, resulting in a single numerical α_s^D value. ΔG_A^S for the all probes were calculated from the Schultz method using Eq. (7) in the surface adsorption region (303.2–328.2 K). A plot of $RT \ln V_N$ versus $a(\gamma_L^D)^{0.5}$ for all probes was plotted at 303.2 K in Figure 5. From Eqs. (5) and (7), γ_s^D of Δ BA was calculated using Schultz and Dorris-Gray methods. The results obtained from studies were listed in Table 5.

It is showed that the value determined for γ_s^D of DBA have different ranges from 47.51–44.06 (Schultz method) to 47.74–46.19 mj/m^2 (Dorris-Gray method). Besides, it is observed that the γ_s^D values calculated by Dorris-Gray method are higher than those obtained from the Schultz method. The γ_s^D values obtained from the Schultz method decrease faster than the γ_s^D values obtained by the Dorris–Gray method with increasing temperature. The results obtained for the Schultz and Dorris–Gray method at surface adsorption region are close to each other, showing that these two methods are compatible and feasible. There is no study on DBA in the literature. A rough comparison can be made with reported LCs. In the literature, γ_s^D values for LCs were ranged from 30 to 42 mj/m^2 in agreement with this study [50, 55].

The values of $-\Delta G_A^S$ were calculated by the numerical difference between the calculated value of $RT \ln V_N$ and that which was obtained from Eq. (7) of the linear plot of the nonpolar reference line. The variation of ΔG_A^S between Δ BA and the polar probes for the studied temperatures is given in Table 6. Regarding the Table 6, it was seen that the temperature did not change the ΔG_A^S values much. ΔH_A^S values were calculated for polar probes and the results were given in Table 7. The ΔH_A^S values were calculated as the degree of interaction between the Δ BA molecule surface and the polar probe

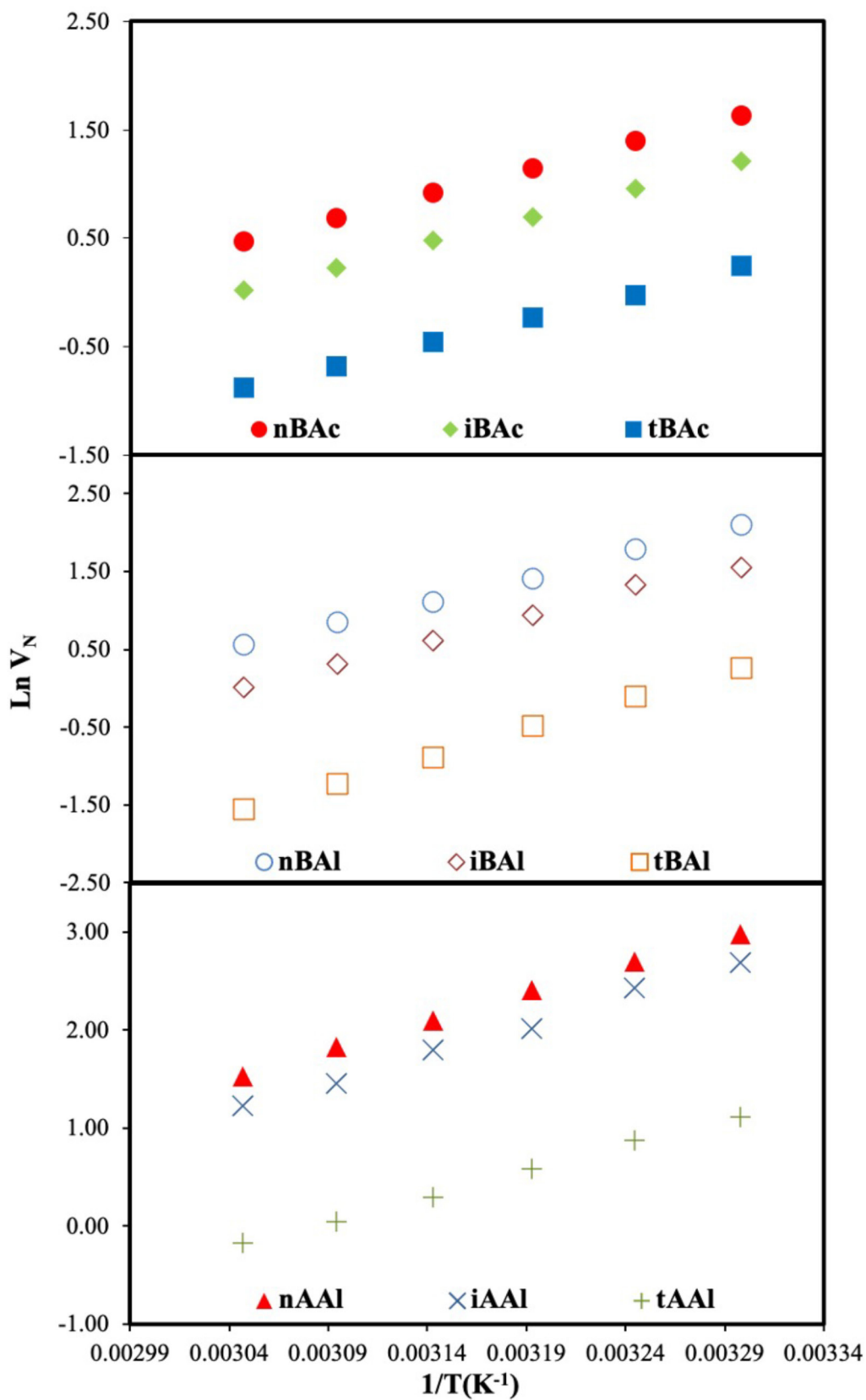


Figure 1. Net retention volumes (V_N) of isomer series on DBA (Surface adsorption region).

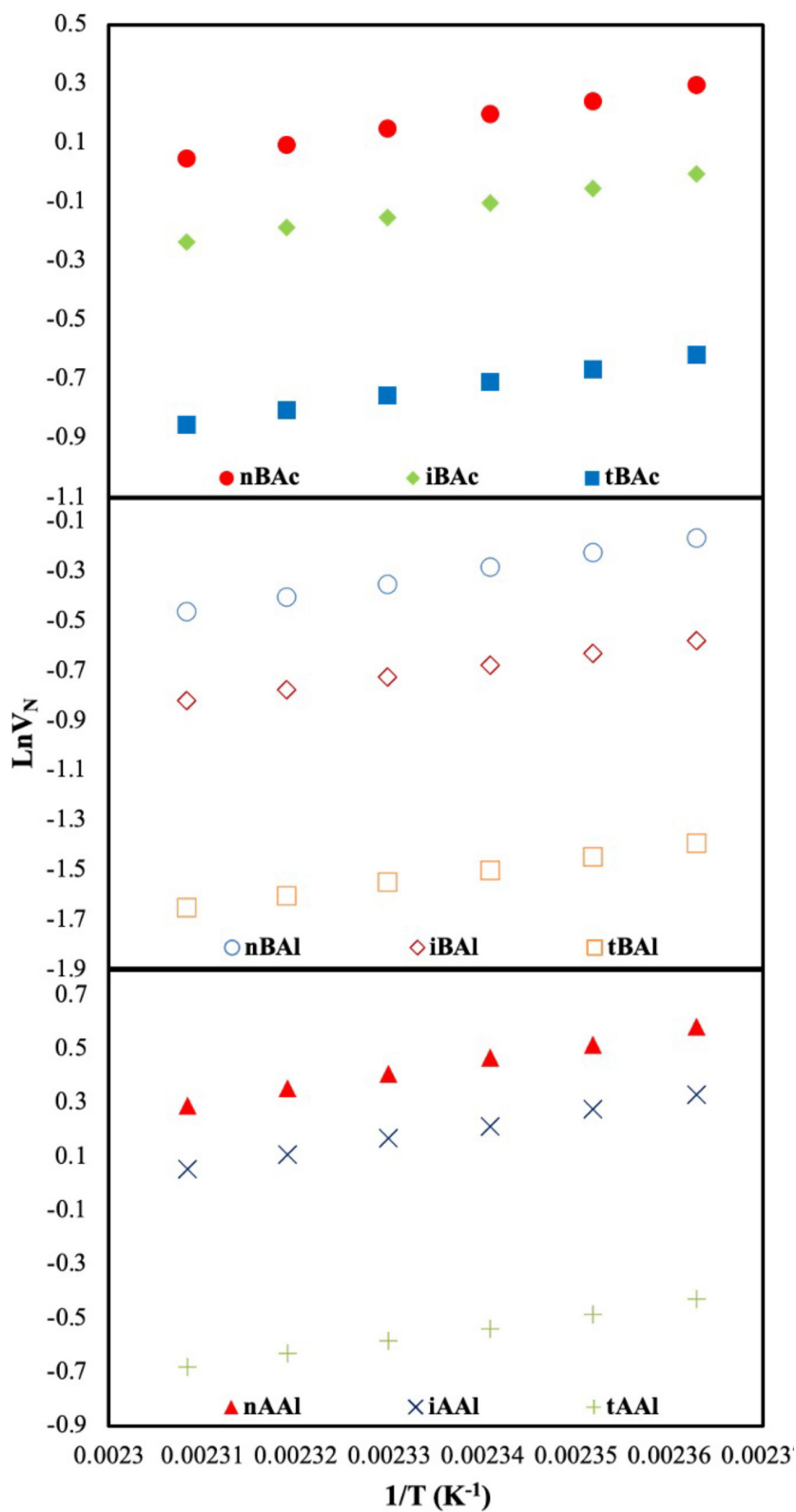


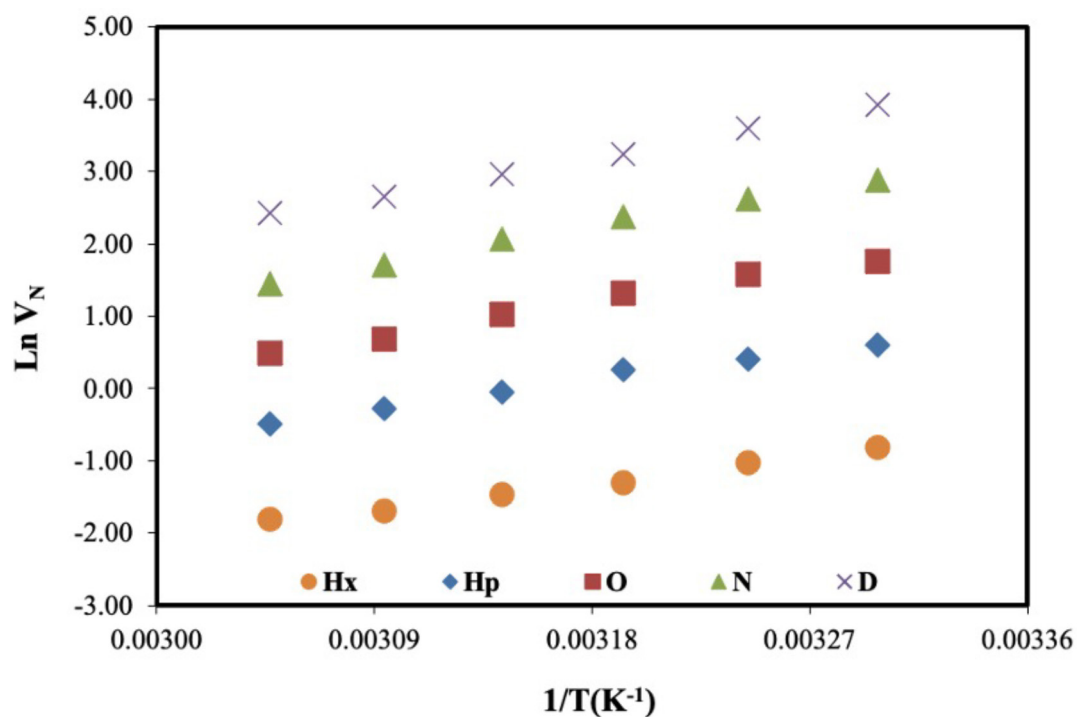
Figure 2. Net retention volumes (V_N) of isomer series on DBA (Thermodynamic region).

Table 3. The separation factor of DBA for the isomer pairs: nBac/iBac, nBac/tBac, nBal/iBal, nBal/tBal, nAAl/iAAl, and nAAl/tAAl (303.2–328.2 K).

$\alpha = V_{N1} / V_{N2}$						
T (K)	V_{NnBac} / V_{NiBac}	V_{NnBac} / V_{NtBac}	V_{NnBal} / V_{NiBal}	V_{NnBal} / V_{NtBal}	V_{NnAAl} / V_{NiAAl}	V_{NnAAl} / V_{NtAAl}
303.2	1.52	3.99	1.72	6.26	1.34	6.49
308.2	1.55	4.15	1.58	6.69	1.31	6.19
313.2	1.57	3.98	1.60	6.66	1.48	6.20
318.2	1.57	4.00	1.65	7.41	1.35	6.03
323.2	1.58	3.90	1.70	7.89	1.46	5.97
328.2	1.57	3.88	1.74	8.34	1.35	5.46

Table 4. The separation factor of DBA for the isomer pairs: nBac/iBac, nBac/tBac, nBal/iBal, nBal/tBal, nAAl/iAAl, and nAAl/tAAl (423.2–433.2 K).

$\alpha = V_{N1} / V_{N2}$						
T (K)	V_{NnBac} / V_{NiBac}	V_{NnBac} / V_{NtBac}	V_{NnBal} / V_{NiBal}	V_{NnBal} / V_{NtBal}	V_{NnAAl} / V_{NiAAl}	V_{NnAAl} / V_{NtAAl}
423.2	1.35	2.50	1.52	3.40	1.29	2.75
425.2	1.35	2.51	1.51	3.33	1.27	2.71
427.2	1.36	2.49	1.49	3.45	1.29	2.76
429.2	1.35	2.47	1.45	3.38	1.28	2.70
431.2	1.33	2.44	1.46	3.35	1.28	2.70
433.2	1.33	2.44	1.43	3.32	1.26	2.63

**Figure 3.** Net retention volumes (V_N) of nonpolar probes on DBA.

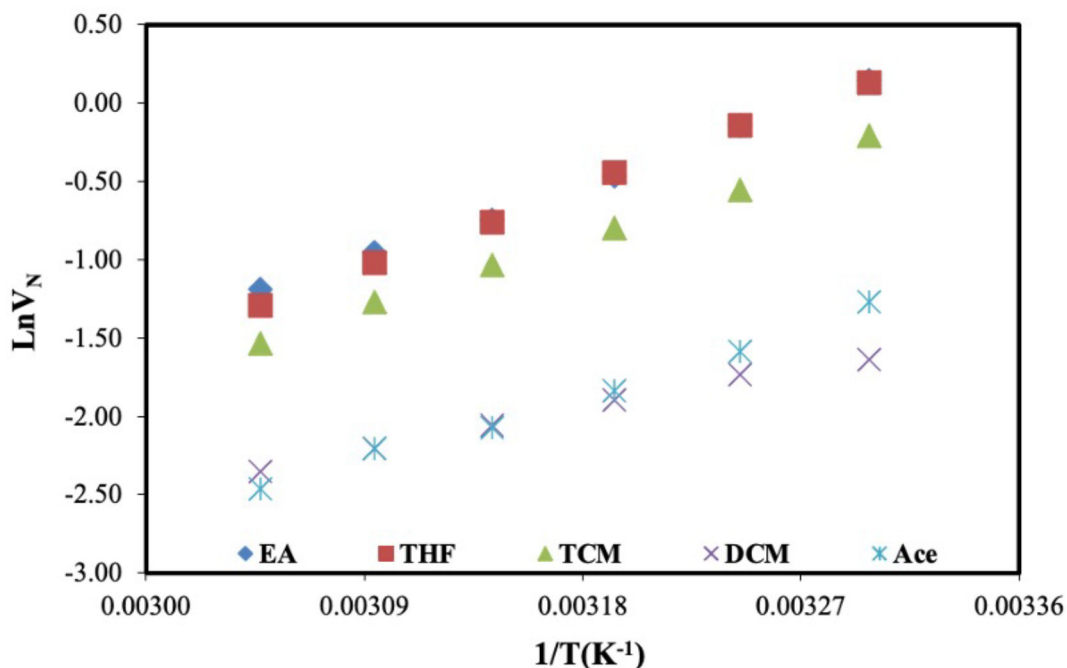


Figure 4. Net retention volumes (V_N) of polar probes on DBA.

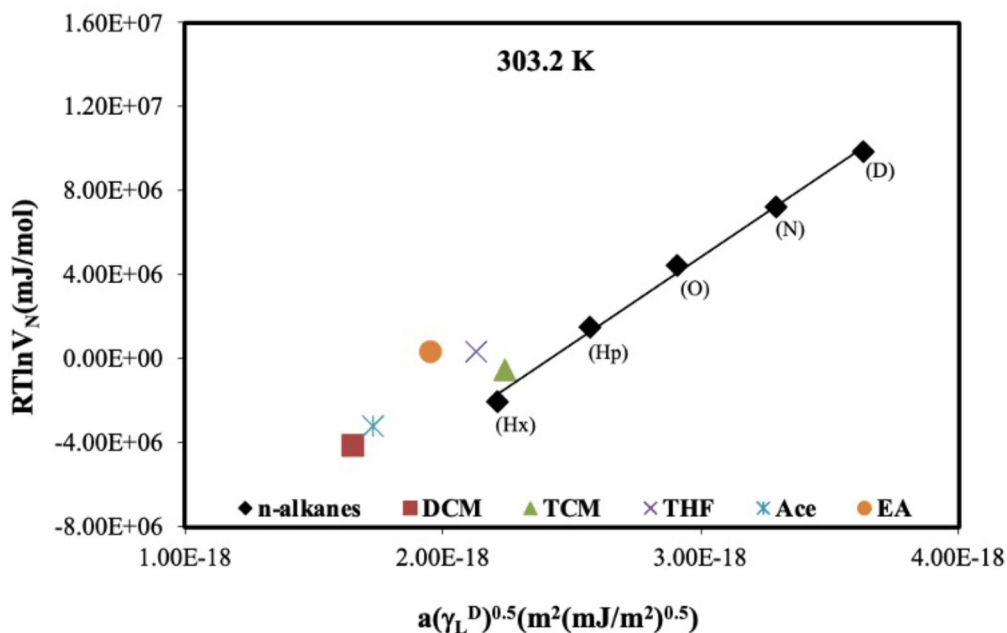


Figure 5. A linear plot of $RT \ln V_N$ vs $a(\gamma_L^D)^{0.5}$ for all probes on DBA at 303.2 K.

molecules. These values were followed the order THF>EA>Ace>TCM>DCM. The DCM probe molecule ($DN = 0.0$, $AN = 16.4$) showed the lowest $-\Delta H_A^S$ value which is to be expected taking into account the acidic properties of this molecule and the acidic properties of LC surface given by the K_A value. THF is a basic probe molecule ($DN = 84.4$, $AN = 2.1$), it may be expected to interact strongly with acid surfaces [56,57]. Considering the values of ΔH_A^S and ΔG_A^S for each polar probe, adsorption occurs exothermically and spontaneously for all studied temperatures. The specific intermolecular interactions are derived from the interaction between the polar probe and the Lewis acidic-basic sites on surface [58–60].

Table 5. Dispersive surface energy (γ_s^D) of DBA.

T (K)	γ_s^D $mj.m^{-2}$	
	Schultz	Dorris-Gray
303.2	47.51	47.74
308.2	47.06	47.69
313.2	46.38	47.36
318.2	45.89	47.27
323.2	44.71	46.45
328.2	44.06	46.19

Table 6. The variation of free energy of specific interactions, $-\Delta G_A^s$ (kJ/mol), between DBA and the polar probes for the studied temperatures.

T (K)	EA	Ace	DCM	TCM	THF
303.2	4.21	2.48	2.21	0.91	2.69
308.2	3.99	2.10	2.39	0.54	2.50
313.2	3.68	1.92	2.42	0.45	2.28
318.2	3.54	1.83	2.53	0.42	2.03
323.2	3.63	2.03	2.67	0.44	2.00
328.2	2.95	1.24	2.17	-0.34	1.21

Table 7. Values of enthalpy (ΔH_A^s) of adsorption on DBA for the polar probes.

Polar probes	$-\Delta G_A^s$ (kJ/mol)
Ace	13.68
EA	17.20
THF	18.39
DCM	0.87
TCM	12.12

A plot of $-\Delta H_A^s/AN^*$ versus DN/AN^* was plotted by K_A as the slope and K_D as the intercept using Eq. (11), and it is shown in Figure 6. The character of DBA surface was determined by the ratio of K_D/K_A . The obtained K_A and K_D values were listed in Table 8. Due to the K_D/K_A value is lower than 1, DBA surface is an acidic character.

5. Conclusion

IGC-ID technique was used to investigate the separation of isomer series in surface adsorption (303.2–328.2 K) and thermodynamic region (423.2–433.2 K) and the surface properties of DBA in surface adsorption region. Considering the separation factors, it was determined that the DBA in IGC-ID technique can be used to separate the isomer series in surface adsorption and thermodynamic region. The values of γ_s^D for DBA were determined to be 47.51–44.06 mj/m^2 using the Schultz method and 47.74–46.19 mj/m^2 using the Dorris–Gray method. γ_s^D values from both calculation methods decrease linearly with the increase in temperature in the range from 303.2 to 328.2 K. The values of K_A and K_D were found to be 0.2134 and 0.1907, respectively. As shown that, the K_D value is lower than the K_A . In this case, it can be said that

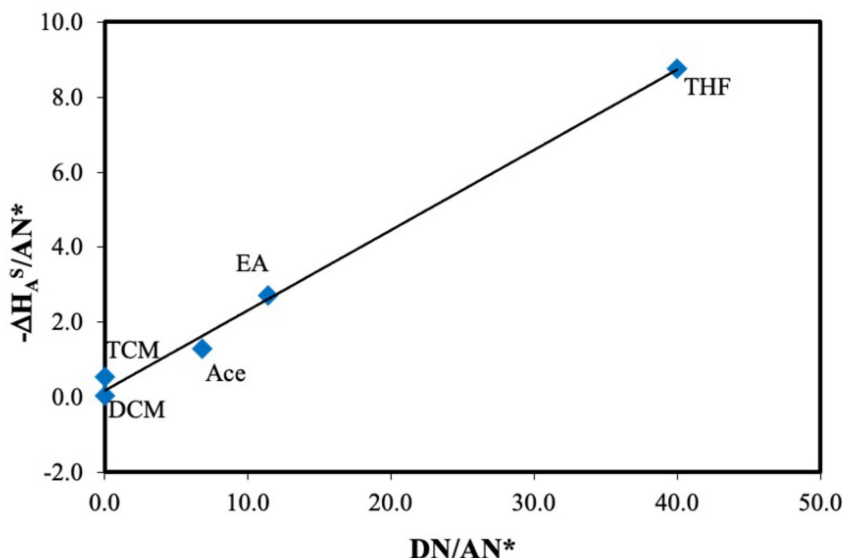


Figure 6. The plot of $-\Delta H_A^S/AN^*$ vs DN/AN^* .

Table 8. Lewis acid-base parameters, K_A and K_D , of DBA.

Liquid crystal	K_A	K_D	K_D/K_A
DBA	0.2134 ± 0.012	0.1907 ± 0.009	0.89

the DBA surface is an acidic character. The IGC-ID technique is very important in improving the quality of products for industrial fields, since isomers can be separated effectively and the surface energy of samples can be easily determined.

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