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THE CHARACTERIZATION OF SOLUTES AND SOLVENT PHASES(II)
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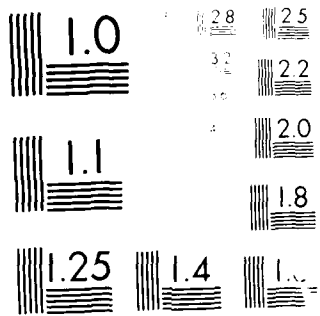
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THE CHARACTERIZATION OF SOLUTES & SOLVENT PHASES

PROGRESS REPORT

14 OCTOBER 1987

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Summary

The characterisation of five gas-liquid chromatographic stationary phases, using retention data obtained by Laffort and co-workers for 240 solutes, has been carried out by the method of multiple linear regression analysis. Relative gas-liquid partition coefficients for as many solutes as possible were correlated against various combinations of parameters. The best general equation was found to be one containing the exploratory variables π_2^* , δ , α_m , β_m , and $\log L^{16}$. Attempts to replace π_2^* by the dipole moment (μ^2) were not very successful, and neither were attempts to use refractive index functions or molar refractions in combination with μ^2 . However, replacement of π_2^* by the dipole moment itself was more successful, and led to quite good equations in μ , δ , α_m , β_m , and $\log L^{16}$.

A number of new $\log L^{16}$ values have been determined experimentally, using the gas-chromatographic method.

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Introduction

Over the past few years, Abraham, Doherty, Kamlet, Taft and co-workers^{1,2} have constructed equations for the correlation and prediction of a very large number of physicochemical and biochemical phenomena. These equations are based on a cavity theory of solution, in which the process of dissolution of a solute in a solvent may be broken down into a number of hypothetical steps: (i) the endoergic formation of a cavity in the bulk solvent, (ii) rearrangement of solvent molecules round the cavity, and (iii) the exoergic interaction of the solute with the surrounding solvent molecules after the solute has been inserted into the cavity. If the Gibbs energy change in step (ii) is zero, or very nearly zero as is usually assumed, only steps (i) and (iii) need to be modelled. The energy of formation of a cavity can be taken as proportional to the solvent cohesive energy density, $(\delta_H^2)_1$, where δ_H is the Hildebrand solubility parameter, and to some function of the solute size, or volume, V_2 ,⁺ leading to a term $(\delta_H^2)_1 \cdot V_2$ with the units of energy. Various solvent-solute interactions can take place in step (iii). If both solvent and solute are dipolar, a term in $\pi_1^* \cdot \pi_2^*$ will arise, where π^* is the solvent or solute dipolarity. Hydrogen-bond interactions will also be set up, either between a solvent acting as a hydrogen-bond base and a solute acting as a hydrogen-bond acid, $\beta_1 \cdot \alpha_2$, or between a solvent acting as the acid and the solute as the base, $\alpha_1 \cdot \beta_2$. In these two terms, α_1 and β_1 refer to the solvent hydrogen-bond acidity and basicity, and α_2 and β_2 to the solute hydrogen-bond acidity and basicity. The full equation for the correlation of some solubility related property, SP, is then given by the multiple linear regression equation,

⁺ We denote, as usual, solvent properties by the subscript 1 and solute properties by the subscript 2.

$$SP = A + B\pi_1^* \pi_2^* + C\beta_1 \alpha_2 + D\alpha_1 \beta_2 + E(\delta_H^2)_1 V_2 \quad (1)$$

Now for a process involving a series of solutes in a given solvent, all the solvent parameters in equation (1) are constant, leading to equation (2). For solutes that are aromatic or polyhalogenated, a polarisability

$$SP = C + s.\pi_2^* + a.\alpha_2 + b.\beta_2 + M.V_2 \quad (2)$$

correction term is needed, which takes the form $\delta_2 = 1$ for aromatic solutes, 0.5 for polyhalogenated solutes, and zero for all other solutes. This leads to the final equation, used extensively in the correlation of a wide variety of phenomena in condensed phases, equation (3). An early application of

$$SP = C + s.\pi_2^* + d.\delta_2 + a.\alpha_1 + b.\beta_2 + m.V_2 \quad (3)$$

equation (3) to a process involving a gaseous phase, namely the solubility of gases and vapours in polymers,³ revealed a possible deficiency in that equation (3) contains no term that corresponds to solute-solvent dispersion, or van der Waals, interaction. An alternative equation was therefore put forward, with a new solute parameter, $\log L^{16}$, replacing the volume term V_2 . This new parameter was defined as the logarithm of the solute Ostwald solubility coefficient, L , on n-hexadecane at 298K.⁴

$$L = \frac{\text{concentration of solute in solution}}{\text{concentration of solute in the gas phase}} \quad (4)$$

Two possible equations for the correlation of the solubility of a series of gases and vapours in a given condensed phase are therefore equations (3) and (5), and we set out to investigate the use of these two equations. There are several ways of describing the solubility of gases in liquids, but in view

of the use of the Ostwald solubility coefficient in equation (5) shall define gas solubility through equation (4). It should be noted that L is actually the same as the gas-liquid partition coefficient, K, used in the description of gas-liquid chromatography (GLC). The coefficient L or K is related to the specific retention volume at the column temperature, V_G , through equation (6) where ρ_1 is the stationary phase density.

$$SP = C + s.\pi_2^* + d.\delta_2 + a.\alpha_2 + b.\beta_2 + 1.\log L^{16} \quad (5)$$

$$L \text{ (or } K) = \beta_1 \cdot V_G \quad (6)$$

In order to apply equations (3) and (5), values of SP (i.e. $\log L$ or $\log V_G$) should be available for a wide selection of solutes on the same liquid phase. By far the most convenient and accurate method of obtaining such a series of SP values is by GLC itself, and there are numerous compilations of such data. Probably the most extensive, carefully measured, values are those recorded by Laffort and co-workers⁵ for 240 solutes on 5 stationary phases, and this is the data we have chosen first to analyse. Laffort and co-workers published their data in the form of Kovat's retention indices, defined for isothermal GLC through equation (7).

$$I(x) = 100 \frac{\log V_G(x) - \log V_G(P_n)}{\log V_G(P_{n+1}) - \log V_G(P_n)} + 100n \quad (7)$$

Here, $I(x)$ is the retention index of solute x , V_G is the specific retention volume, and (P_n) and (P_{n+1}) represent n -alkanes of carbon number n and $n + 1$. The equation used to calculate their retention indices is,

$$I = 100 \frac{\log \tau(x) - \log \tau(P_{10})}{b} + 1000 \quad (8)$$

where $\tau(x)$ and $\tau(P_{10})$ are the corrected retention times for compound x and n -decane, and b is the slope for n -alkanes. From equation (6) and equation (8) it follows that

$$\log L(x) = \frac{I-1000}{100} b + \log L(P_{10}) \quad (9)$$

and hence values of $\log L(x)$ may be calculated for the 240 solutes relative to $\log L$ for n -decane. These relative values of $\log L(x) - \log L(P_{10})$ can be used in equations (3) and (5) to yield exactly the same coefficients etc. as would be found with the absolute values of $\log L(x)$.⁺ The constant quantity $\log L(P_{10})$ will be subsumed into the value of C .

The five phases studied by Laffort and co-workers are listed in Table 1, and the 240 solutes are given in Table 2. In Table 3 are given those listed in our database, i.e. those for which we had some, or most, of the required solute parameters and in Table 4 are the b -values used in equation (9). The parameters that were chosen to be used in regressions are as follows:

- π_2^* These values are those that have been extensively used by Kamlet and co-workers.^{1,2,7-10}
- δ_2 This is a trivially-calculated parameter, taken as 1.0 for aromatic solutes, 0.5 for polyhalogenated solutes, and zero for all others.
- $\alpha_2(\alpha_m)$ These values were taken from recent papers of Kamlet and co-workers.⁷⁻¹⁰

⁺ This is not the case if only Kovat's retention indices are known. It is therefore fortunate that Laffort and co-workers⁵ had the foresight to record values of b for each of the five stationary phases studied. Note that slightly different b -values are given in Laffort's table III. We are indebted to Professor Laffort⁶ for suggesting that those in the first two would be the most suitable to use in our equation (9).

$\alpha_2(\alpha_2^H)$	This is a new hydrogen-bond acidity parameter recently developed by Abraham and co-workers using log K values for hydrogen-bond complexation. ¹¹
$\beta_2(\beta_m)$	These were, again, taken from papers of Kamlet and co-workers. ⁷⁻¹⁰
$\beta_2(\beta_2^H)$	This is a new hydrogen-bond basicity parameter, ¹² obtained by the same procedure as α_2^H .
$V_2(V_x)$	In all our calculations we used the trivially calculatable characteristic volume, as detailed by Abraham and McGowan. ¹³
$\log L^{16}$	Many of values were taken from the experimental paper of Abraham, Grellier, and McGill. ⁴ Other values have been obtained in this work, see experimental section and the section on log L ¹⁶ values.
μ_2	Dipole moments were taken from standard literature sources. ^{14,15}
$f(n^2)$	This refractive index function, defined as $f(n^2) = (n^2-1)/(n^2+2)$ was calculated from literature values of the refractive index of the solute liquid at 293 K and the sodium-D line. ¹⁶
MR	The molar refraction was obtained from the usual definition, $MR = f(n^2).M/P$, where M and P are the solute molecular weight and density; MR is the same as the electron polarisation, P_E .
MR_x	To eliminate the need for a new parameter (the density), a modified molar refraction was calculated as $MR_x = f(n^2).V_x$.

In the present report, we set out equations, based on the α_m and β_m values of Kamlet and co-workers,⁷⁻¹⁰ and in a subsequent report we shall investigate the use of the new parameters α_2^H and β_2^H . To some extent equations (3) and (5) are straightforward in that, apart from α_m and β_m taken as standard, the other parameters V_x and $\log L^{16}$ are well-defined solute parameters. However π_2^* is partly derived from the solvent parameter π_1^* , and

partly obtained through a dipole moment correlation. One aim of the present work was therefore to see if π_2^* could be replaced either by the dipole moment (μ or μ^2) or some combination of dipole moment with $f(n^2)$ or MR. Of course, another aim is to obtain the best regression equation that could be used to predict new values of the chromatographic parameter.

Results and discussion

Regressions were run for the relative values of $\log L(x)$ on the five phases listed in Table 1. The coefficients of the parameters are listed in a series of Tables, together with the number of data points (n), the multiple correlation constant (r), and the standard deviation (s.d.). Also given are the per cent confidence levels for the coefficients (correlation %). The Tables are as follows:

Table 5:	$\delta, \pi_2^*, \alpha_m, \beta_m, Vx$	All phases
	$f(n^2), \mu^2, \alpha_m, \beta_m, Vx$	
Table 6:	$\delta, \pi_2^*, \alpha_m, \beta_m, \log L^{16}$	All phases
	$f(n^2), \mu^2, \alpha_m, \beta_m, \log L^{16}$	
Table 7:	$\delta, \mu^2, \alpha_m, \beta_m, Vx$	Polyph ether only
	$\delta, \mu^2, \alpha_m, \beta_m, \log L^{16}$	
	$MR_x, \mu^2, \alpha_m, \beta_m, Vx$	
	$MR_x, \mu^2, \alpha_m, \beta_m, \log L^{16}$	

Since the various regressions in μ^2 were not very good, an additional series was run that included only solutes with a single dominant dipole moment - i.e. solutes such as acids, esters, nitro compounds and so forth were removed. Further regressions were carried out as follows:

Table 8:	$f(n^2), \mu^2, \alpha_m, \beta_m, Vx$	TCEP only
	$f(n^2), \mu^2, \alpha_m, \beta_m, \log L^{16}$	
	$MR_x, \mu^2, \alpha_m, \beta_m, Vx$	
	$MR_x, \mu^2, \alpha_m, \beta_m, \log L^{16}$	

Although μ^2 is theoretically a better parameter to use than μ itself, two sets of regressions were run using μ as a dipolar parameter. In each set, regressions were carried out for all solutes for which parameters were available, and for solutes with $\delta = 0$:

Table 9:	$f(n^2), \mu, \alpha_m, \beta_m, \log L^{16}$	All phases
Table 10:	$\delta, \mu, \alpha_m, \beta_m, \log L^{16}$	All phases
	$\mu, \alpha_m, \beta_m, \log L^{16}$	

A direct comparison of Tables 5 and 6 shows quite clearly that regressions in $\log L^{16}$ are always markedly superior to those in Vx , and hence we shall consider only the $\log L^{16}$ correlations henceforth. Results in Table 6 for the general equation (5) are quite good, with correlation constants ranging from 0.9943 to 0.9805 for the five phases with about 90 solutes. The constants in equation (5) make general chemical sense: all five phases are hydrogen-bond bases and have no hydrogen-bond acidity, and hence the coefficient in the term $b.\beta_m$ should be statistically not significant. This is true for all the phases except Zonyl E7. This phase is supposed to be a

fluorinated ester of "pyromellitic acid and a trihydrofluoro alcohol". The former is 1,2,4,5-benzene tetra-carboxylic acid, and it is possible that the commercial product contains either unesterified carboxylic acid or hydroxyl groups.

One difficulty over a physicochemical interpretation of equation (5) is that polarisability effects are contained in the $s.\pi_2^*$ term as well as on the $d.\delta$ term. It would be particularly useful if the dipolarity and polarisability effects, both contained in $s.\pi_2^*$, could be separately counted. To this end, we have investigated the effect of replacing the δ/π_2^* terms by various combinations of the dipole movement, as μ^2 , and polarisability functions such as $f(n^2)$ or MR. In Table 6 are results of a direct replacement in equation (5) of π_2^* and δ by $f(n^2)$ and μ^2 . The overall correlation constants are not as good as those in the original equation (5), but are not too bad. However, most surprisingly, the $b.\beta_m$ terms are statistically significant for all five phases, thus making the entire regression equations rather suspect from a chemical point of view.

In Table 7 are results of replacing π_2^* and δ by either μ^2 and δ or by μ^2 and MR_x , for the polyphenyl ether phase as an example. There is an excellent correlation with δ , μ^2 , α_m , β_m , and $\log L^{16}$ with $r = 0.9922$ and $sd = 0.093$, which must be close to an exhaustive fit. But once again, the $b.\beta_m$ term is highly significant. Regressions with $f(n^2)$ and μ^2 or MR_x and μ^2 for TCNE using a restricted set of solutes that contain either no dipole or else a single dominant dipole, are in Table 8. The only chemically reasonable regressions are those of the original form in δ/π_2^* .

Our conclusion as a result of the regressions set out in Tables 5-8 is that replacement of the δ/π_2^* symbolism by μ^2 in combination with a polarisability term leads (i) to regressions that are not as good, and (ii) to regressions that contain an unacceptable $b.\beta_m$ term.

In terms of chemical theory, correlations of an energy-related quantity such as $\log V_G$ or $\log L$ with dipole moment should certainly involve μ^2 and not μ . However, we thought it useful on an empirical level to investigate the use of μ as a solvent parameter. Table 9 gives details of regressions where $f(n^2)$ and μ replace δ and π_2^* . Once again, the $b.\beta_m$ term is highly significant for all five phases. Finally, in Table 10, are results of simply making a direct replacement of π_2^* by μ in equation(5), to give:

$$SP = C + s.\mu + d.\delta_2 + a.\alpha_2 + b.\beta_2 + l.\log L^{16} \quad (10)$$

The regressions are all very good, and the only difficulty is that the $b.\beta_m$ term is still too significant for the phases TCEP, Polyphenyl ether, and DEGS. We carried out another set of regressions using equation (10) for a restricted set of solutes for which $\delta = 0$ (about 60-65 such solutes). The regression coefficients and sd values are the best we have obtained (compare Table 10 with Table 6), and now the $b.\beta_m$ terms are statistically not significant, as required (except for zonyl E7 !). We are much encouraged by the results in Table 10, and intend to pursue this line of regression analyses. If, indeed, π_2^* can be replaced by μ , not only would interpretation be much easier, but it would be possible to predict the dipole parameter rather easier from a knowledge of molecular structure.

Work in Progress

Further work is in hand on the refinement of equation (10) for the correlation and prediction of gas-liquid and gas-polymer partition coefficients. We hope that on the next report we shall be able to set out a modified equation (10) that will deal with the Laffort data set.

We also have in hand the analysis of a large number of $\log L$ values (or the equivalent $\log V_G$ values) on the nonpolar phases Apiezon and squalane. It

will be possible to extract from this data a rather extended list of secondary $\log L^{16}$ values that will considerably extend our data base.

Our next projected experimentation will be the acquisition of $\log L$ values for a range of solutes on some simple organic solvents, so that a direct comparison can be made with GLC stationary phases and with polymer phases.

Experimental

In order to increase the number of Laffort solutes for which we had all the parameters, additional $\log L^{16}$ values were determined, at the standard temperature of 298.15 K. A short column was used of length 50 cm and internal diameter 2 mm, containing 8.34% w/w of n-hexadecane on chromosorb B, mesh size 45/60. The standard used was n-octane of $\log L^{16}$ value 3.677, and $\log L^{16}$ values for other solutes were obtained relative to n-octane, using a flame ionisation detector, as described before.⁴ Results are given in Table 11. Attempts were also made to obtain $\log L^{16}$ values for proprionic acid and higher carboxylic acids, but without success, but further attempts will be made to determine these quantities, either directly or indirectly.

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Table 1. The five stationary phases studied by Laffort and co-workers.

Zonyl	Zonyl E7 (Dupont). This is a fluoro-ester.
CW 1540	Carbowax 1540 (Applied Science).
TCEP	Tricyanoethoxypropane
Polyph. ether	Polyphenyl ether, six rings.
DEGS	Diethyleneglycol succinate.

Table 2

Kovats retention indexes in GLC of 240 substances on five stationary phases. [corresponds to extrapolated values. The numbers of Handbook followed by x or an additional digit are not reported in the 50th edition but correspond to the same logic of classification.

Substances	Handbook	Zonyl	C-max	ICLP	Polyph.	DEGS
		1540			ether	
ALCOHOLS						
1 METHANOL	M0349	457	916	1228	522	1062
2 ETHANOL	E0334	704	933	1238	587	1083
3 1-PROPANOL	P1387	823	1045	1333	703	1188
4 ISOPROPANOL	P1388	741	916	1261	613	1037
6 ALLYL ALCOHOL	F1815	832	1112	1432	713	1284
6 1-BUTANOL	B2044	944	1164	1473	820	1311
7 ISOBUTANOL	F1433*	897	1104	1394	772	1242
8 2-BUTANOL	B2044	867	1023	1312	737	1144
9 TER-BUTANOL	F1433	816	1009	1198	654	1034
10 1-PENTANOL	P0313	1053	1271	1502	927	1427
11 ANYL ALCOHOL (ACT C)	B2099	1010	1220	1514	884	1361
12 ISOBANYL ALCOHOL	B2095	1025	1229	1529	883	1375
13 TER-ANYL ALCOHOL	B2096	917	1014	1298	771	1162
14 CYCLOPENTANOL	C0092	1083	1327	1477	994	1518
15 1-HEXANOL	H0470	1160	1383	1641	1032	1644
16 2-HEXANOL (TRANS)	H0372*	1167	1416	1737	1044	1681
17 2-HEXANOL	H0400	1090	1241	1531	950	1384
18 3-HEXANOL	H0403	1063	1212	1497	941	1337
19 2-METHYL-2-PENTANOL	P0334	1022	1110	1393	848	1233
20 3-METHYL-3-PENTANOL	P0335	1038	1132	1427	909	1289
21 1-HEPTANOL	H0144	1242	1489	1794	1138	1648
22 2-HEPTANOL (TRANS)	H0209*	1278	1538	1806	1163	1737
23 1-OCTANOL	O0106	1370	1590	1982	1241	1764
24 2-OCTANOL (TRANS)	O0214	1383	1639	1979	1262	1832
25 CP. CR. 2,4-DIURETIC CYCLOHEXANOL	C07311	1235	1423	1714	1168	1583
26 CP. TR. 2,4-DIURETIC CYCLOHEXANOL	C07312	1247	1492	1824	1219	1687
27 TR. TR. 2,4-DIURETIC CYCLOHEXANOL	C07313	1272	1440	1737	1173	1614
28 2-METHYL-3-HEPTANOL	H0179	1222	1310	1582	1063	1435
29 3-METHYL-3-HEPTANOL	H0180	1220	1307	1584	1078	1456
30 1-NONANOL	N0343	1470	1712	2012	1350	1855
31 1-DECANOL	D0037	1371	1704	2182	1439	1959
32 A-TERPINEOL (DL)	T0049	1326	1709	2100	1426	1948
33 1-UNDECANOL	U0031	1702	1919	2234	1549	2068
34 1-DODECANOL	D0311	1813	2028	2343	1670	2208
35 EXO-STYRYL FENEBOL	F0013*	1340	1478	2010	1480	1890
ALDEHYDES						
36 ACETALDEHYDE	A0010	738	742	1073	541	914
37 PROPIONAL	P1083	830	823	1170	654	980
38 PROPENAL	P1707	842	864	1224	659	1033
39 BUTYRALDEHYDE	B2471	933	909	1243	753	1078
40 ISOBUTYRALDEHYDE	F1099*	887	854	1173	704	992
41 2-BUTENAL (TRANS)	B2933	1004	1004	1517	870	1305
42 ISOVALERALDEHYDE	B2481	1002	948	1290	813	1103
43 FURFURAL	F0277	1358	1493	2037	1133	1816
44 BENZAL	B0316	1149	1114	1473	664	1209
45 2-HEXENAL (TRANS)	H0546	1313	1274	1712	1077	1502
46 HEPTANAL	H0000	1249	1224	1573	1064	1398
47 2-HEPTENAL (TRANS)	H0197*	1400	1343	1793	1172	1600
48 OCTANAL	O0110	1343	1311	1660	1167	1504
49 2-OCTENAL (TRANS)	O0208*	1287	1447	1729	1154	1602
50 BENZALDEHYDE	B0035	1432	1364	2076	1272	1863
51 SALICYLALDEHYDE	S0122	1321	1483	2194	1356	2030
KETONES						
52 ACETONE	F1649	904	809	1239	632	1034
53 2-BUTANONE	B2912	993	932	1329	764	1121
54 BIACETYL	B2432	1001	999	1498	872	1220
55 2-PENTANONE	P0204	1092	1030	1424	857	1230
56 CYCLOPENTANONE	C0094	1270	1250	1748	1050	1380
57 2-HEXANONE	H0317	1193	1123	1512	966	1302
58 3-HEXANONE	H0318	1149	1004	1434	943	1239
59 CYCLOHEXANONE	C0785	1384	1343	1863	1160	1623
60 2-HEPTANONE	H0104	1293	1223	1612	1061	1411
61 CYCLOHEPTANONE	C0417	1491	1479	1990	1284	1740
62 2-OCTANONE	O0197	1400	1332	1718	1163	1524
63 CYCLOOCTANONE	C0037	1600	1590	2111	1404	1899
64 ACETOPHENONE	A0045	1612	1609	2243	1387	2013
65 2-NONANONE	N0332	1608	1427	1810	1244	1622
66 CYCLONONANONE	C0850	1710	1702	2232	1521	2000
67 2-DECANONE	D0042	1597	1526	1915	1346	1733
68 CYCLODECANONE	C0449	1820	1812	2333	1639	2142
69 CARBONE	C0212	1732	1747	2154	1566	2063
70 2-UNDECANONE	U0039	1696	1619	2012	1469	1810
71 CYCLOUNDECANONE	C06103	1931	1923	2474	1737	2249
72 2-DODECANONE	D0313	1799	1717	2182	1571	1916
73 CYCLODODECANONE	C04004	2040	2033	2593	1873	2394
74 CYCLOTRIDECANONE	C0424*	2151	2143	2714	1994	2522
75 CYCLOTETRADECANONE	L0030*	2261	2236	2837	2111	2640
ETHERS						
76 ETHYL ETHER	E0077	658	627	750	574	760
77 BUTYL ETHER	B0471	800	774	1000	634	1034
78 FURAN	F0103	693	603	1040	620	964
79 1,4-DIOXANE	D0198	1104	1100	1328	924	1347
80 2-ACETYL-3-METHYL FURAN	F0194*	1379	1310	2106	1303	1801
81 2-ACETYL-3-METHYL FURAN	F0194*	1612	1641	2211	1319	1831
82 2-PROPYL-3-METHYL FURAN	F0194*	1479	1364	1987	1300	1801
83 2,4,5-TRIMETHYL OXAZINE	O0204*	1731	1747	2303	1044	1443
84 THIOFURAN	T0033	1291	1133	1783	1130	1614
85 THIOPHENE	T0044	1302	1070	1574	1167	1711

Table 2 (Continued)

Substances	Handbook	Zonyl	C. max 1540	TCFP	Polyph. ether	DEGS
NITROGEN COMPOUNDS						
87 1-NITROETHANE	M0204	1010	1149	1439	766	1402
88 1-NITROETHANE	E0235	1101	1184	1443	873	1423
89 1-NITROPROPANE	F1210	1184	1248	1722	944	1493
90 2-NITRYL-2-NITROPROPANE	F1201*	1201	1138	1379	933	1334
91 3-NITROTOLUENE	T0335	1743	1826	2386	1521	2204
92 ACETONITRILE	A0241	942	1048	1508	781	1242
93 BUTYRONITRILE	B2680	1132	1144	1489	866	1332
94 VALERONITRILE	F0229	1251	1234	1493	987	1451
95 BENZONITRILE	B1291	1330	1443	2172	1298	1942
96 TRIMETHYLAMINE	A0939	700	584	973	474	1036
97 ALLYLAMINE	F1714	784	877	1227	474	1327
98 2-AMINOBUTANE	B2489	828	870	1200	698	1318
99 PYRAZOLE	F2126	1123	1516	1941	1001	1742
100 PYRIDINE	F1909	1153	1242	1660	994	1387
101 2,3,6-TRIMETHYL PYRIDINE	F2032	1343	1423	1811	1232	1683
102 TRIMETHYL PYRAZINE	F19043	1383	1449	1854	1235	1793
103 2-METHYL-3-ETHYL PYRAZINE	F19042	1354	1438	1825	1236	1685
104 2-METHOXY-3-ISOBUTYL PYRAZINE	F19041	1421	1350	1821	1338	1727
CARBOXYLIC ACIDS						
105 ACETIC ACID	A0058	1035	1423	1823	734	1480
106 PROPIONIC ACID	F1302	1150	1344	1914	884	1744
107 BUTYRIC ACID	B2451	1235	1618	1994	983	1840
108 ISOBUTYRIC ACID	F1310	1206	1372	1908	943	1773
109 3-BUTENOIC ACID	B3016	1248	1738	2140	983	1991
110 4-METHYLBUTYRIC ACID	B2797	1343	1674	2012	1094	1891
111 VALERIC ACID	F0216	1371	1749	2099	1108	1954
112 ISO-VALERIC ACID	B2802	1382	1664	2016	1042	1877
113 HEXANOIC ACID	B0427	1483	1858	2191	1233	1963
114 ISOHXANOIC ACID	F0291	1417	1779	2128	1148	1877
115 HEPTANOIC ACID	B0144	1593	1964	2283	1338	2059
116 OCTANOIC ACID	O0161	1707	2074	2375	1483	2147
117 NONANOIC ACID	B0334	1819	2182	2467	1608	2236
118 DECAHOIC ACID	D0042	1931	2298	2559	1733	2333
119 UNDECANOIC ACID	B0022	2043	2398	2651	1858	2426
120 DODECAHOIC ACID	D0293	2155	2506	2744	1982	2519
ESTERS						
121 METHYL ACETATE	A0221	884	849	1164	646	1024
122 ETHYL ACETATE	A0189	960	913	1221	731	1078
123 METHYL PROPIONATE	F1324	953	929	1241	770	1099
124 PROPYL FORMATE	F0144	950	932	1260	742	1112
125 PROPYL ACETATE	A0239	1034	984	1298	844	1152
126 BUTYL ACETATE	A0178	1170	1110	1422	938	1271
127 PROPYL BUTYRATE	B2686	1218	1143	1444	1024	1297
128 AMYL ACETATE	A0223	1269	1209	1513	1058	1349
129 ISOAMYL ACETATE	A0223	1231	1147	1444	1010	1304
130 ISOBUTYL ISOBUTYRATE	F1530	1226	1119	1371	1012	1238
131 METHYL SALICYLATE	B1671	1237	1497	1807	969	1766
132 ISOAMYL ISOVALERATE	B2807*	1432	1325	1592	1217	1444
133 BENZYL ACETATE	A0173	1451	1749	2344	1378	2082
134 METHYL BENZOATE	B1286	1394	1924	2382	1442	2290
135 2-ETHOXY ETHYL ACETATE	A0444	1358	1333	1776	1112	1573
HALOGEN COMPOUNDS						
136 1-FLUOROOCTANE	O0129	1090	1050	1237	972	1139
137 1,18-FLUOROTETRACHLOROCTANE	E0223	890	889	1013	784	976
138 1,23-FLUOROTETRACHLOROCTANE	E0224	899	898	1029	796	991
139 CHLOROFORM	M0304	773	1029	1240	773	1157
140 CARBON TETRACHLORIDE	M0294	780	987	1047	794	1013
141 ETHYLENE CHLORIDE	E0210	863	1081	1398	834	1233
142 TRICHLOROETHYLENE	E0423	862	1023	1210	859	1134
143 1,1,2,2-TETRACHLOROETHANE	E0207	1151	1493	1859	1160	1741
144 HEXACHLOROCYCLOHEPTANE	B2453	1343	1322	1733	1376	1713
145 1,1-DICHLOROBENZENE	T0391	1333	1342	1978	1273	1807
146 1,2-DICHLOROBENZENE	B0330	1008	1043	1266	968	1171
147 1,2-DICHLOROBENZENE	B0333	1317	1323	1862	1279	1749
148 1,1,2-TRICHLOROETHANE	E0498	1347	1312	1941	1273	1764
149 2-CHLOROPHENOL	F0613	1367	1033	2274	1270	2120
150 ETHYL BROMIDE	E0195	477	793	1012	645	914
151 1-BROMOPENTANE	F0090	934	1081	1300	973	1199
152 2-BROMOOCTANE	O0117	1242	1320	1523	1214	1418
153 METHYL IODIDE	M0282	643	844	1036	698	967
154 1-IODOBUTANE	B2368	954	1111	1340	989	1234
155 2-IODOBUTANE	B2369	926	1054	1275	946	1178
SULFUR COMPOUNDS						
156 1-ETHANETHIOL	E0328	425	733	970	435	804
157 1,2-ETHANEDITHIOL	E0312	1044	1341	1771	1099	1583
158 1-PROPANETHIOL	F1297	730	863	1079	743	979
159 2-PROPANETHIOL	F1294	693	780	975	677	882
160 ALLYL MERCAPTAN	F1739	740	909	1146	737	1033
161 1-BUTANETHIOL	B2443	849	963	1193	856	1084
162 ISOBUTANETHIOL	F1293	817	920	1123	812	1020
163 TERT-BUTANETHIOL	F1294	717	780	937	693	878
164 TETRAHYDROTHIOPHENE	T0224	1038	1179	1509	1033	1343

Table 2 (Continued)

Substances	Handbook	Zonyl	C.wax		TCEP	Polyph. ether	DEGS
			1540				
163 TRIOPHENE	T0187	841	1456	1353	871	1236	
164 1-PENTANETHIOL	P0209	951	1473	1293	936	1193	
167 ISOPENTANETHIOL	B2648	817	069	1018	803	983	
168 DITHIAPENTANE	P0138*	1132	1767	1739	1144	1574	
169 2-METHYL THIOPHENE	T0215	996	1154	1441	970	1314	
170 HEXANETHIOL	H0420	1042	1189	1407	1070	1297	
171 2,3-DIMETHYL THIOPHENE	T0200	1094	1224	1508	1052	1377	
172 BENZENETHIOL	B0354	1374	1632	2183	1345	1918	
173 1-HEPTANETHIOL	H0142	1164	1282	1508	1166	1407	
174 BENZYL MERCAPTAN	T3543	1379	1658	2112	1372	1920	
175 1-OCTANETHIOL	O0158	1249	1386	1615	1271	1514	
176 1-NONANETHIOL	N0333*	1374	1490	1822	1377	1621	
177 1-DECANETHIOL	D0041	1479	1594	1729	1482	1728	
178 METHYL SULFIDE	S0313	694	776	1015	654	912	
179 ETHYL SULFIDE	S0303	864	930	1171	831	1057	
180 PROPYL SULFIDE	S0332	1049	1112	1339	1021	1228	
181 PROPYLENE SULFIDE	S0350*	821	965	1238	818	1110	
182 ALLYL SULFIDE	S0292	1063	1168	1483	1034	1333	
183 ISOAMYL SULFIDE	S0298	1374	1387	1596	1303	1469	
184 DIMETHYL DISULFIDE	D0242	935	1138	1425	947	1286	
185 DIETHYL DISULFIDE	D0258	1113	1292	1563	1112	1423	
186 DIBUTYL SULFIDE	S0295	1255	1322	1525	1217	1480	
187 METHYL ETHYL SULFIDE	S0341	626	664	799	611	803	
188 METHYL PROPYL SULFIDE	S0340	885	961	1196	833	1076	
189 METHYL THIOCYANATE	T0744*	1165	1456	1779	1230	1646	
190 METHYL THIOCYANATE	T0175	1160	1325	1821	985	1583	
191 METHANETHIOL ACETATE	M03451	1029	1093	1438	899	1286	
192 ETHYL ISOTHIOCYANATE	E0253	1077	1265	1651	1019	1461	
193 METHANETHIOL PROPARGATE	M03454	1109	1179	1507	995	1357	
194 ALLYL ISOTHIOCYANATE	I0216	1144	1383	1765	1101	1591	
195 METHANETHIOL BUTYRATE	M03452	1188	1255	1580	1079	1428	
196 METHANETHIOL ISOVALERATE	M03453	1239	1284	1586	1117	1444	
197 PHENYL ISOTHIOCYANATE	I0236	1486	1731	2104	1437	1985	
HYDROCARBONS							
198 ETHYLENE	E0401	222	308	363	270	465	
199 PROPYLENE	P1713	324	395	450	360	533	
200 1-BUTENE	B2940	422	483	536	430	601	
201 PENTENE	P0375	522	581	632	532	689	
202 1-HEXENE	H03461	598	634	679	614	712	
203 1-HEPTENE	H0197	722	744	794	723	805	
204 1-OCTENE	O0208	822	843	890	819	893	
205 2-OCTENE(C19)	O0209	848	879	939	849	931	
206 2-ETHYL HEXENE	H0564	837	854	922	824	894	
207 2-BUTYNE	B3071	595	798	890	587	838	
208 1-OCTYNE	O0221	937	1038	1187	907	1118	
209 2-OCTYNE	O0222	975	1075	1229	979	1167	
210 BENZENE	B0202	871	979	1257	833	1141	
211 TOLUENE	T0273	994	1078	1363	958	1242	
212 ETHYL BENZENE	B0758	1084	1162	1439	1035	1333	
213 STYRENE	S0159	1150	1280	1620	1094	1510	
214 ETHYNYL BENZENE	B0778	1144	1386	1698	1096	1567	
215 O-XYLENE	O0460	1112	1182	1475	1046	1360	
216 M-XYLENE	M0461	1115	1188	1479	1050	1362	
217 P-XYLENE	P0462	1113	1180	1473	1041	1350	
218 MESITYLENE	M0975	1233	1311	1587	1155	1481	
219 A-PINENE	P0923	1007	1064	1143	1015	1131	
<i>Alkanes</i>							
220 ETHANE	E0162	200	200	200	200	200	
221 PROPANE	P1107	300	300	300	300	300	
222 BUTANE	B2485	400	400	400	400	400	
223 ISOBUTANE	I1290*	376	364	359	358	359	
224 PENTANE	P0854	500	500	500	500	500	
225 HEXANE	H0319	600	600	600	600	600	
226 CYCLOHEXANE	C0437	675	734	814	735	821	
227 HEPTANE	H0493	700	700	700	700	700	
228 2-METHYL HEPTANE	H0126	780	761	769	760	763	
229 3-METHYL HEPTANE	H0127	793	778	789	774	784	
230 2,4-DIMETHYL PENTANE	P0133	614	594	544	608	664	
231 1-OCTANE	O3112	800	800	800	800	800	
232 1-NONANE	N0519	900	900	900	900	900	
233 2,2,3-TRIMETHYL HEXANE	H0370	813	757	759	759	757	
234 DECANE	D0020	1000	1000	1000	1000	1000	
235 DECALIN	D0006	1184	1284	1408	1241	1376	
236 HYDRINDANE	H0713	1057	1136	1273	1117	1239	
237 UNDECANE	N0919	1100	1100	1100	1100	1100	
238 DODECANE	D0204	1200	1200	1200	1200	1200	
239 TRIDECANE	T0716	1300	1300	1300	1300	1300	
240 TETRADECANE	T0077	1400	1400	1400	1400	1400	

1351 ethanol	0.000	0.221	2.850	2.791	0.400	0.320	0.450	0.500	2.435	1.449
1354 1-propanol	0.000	0.234	2.877	2.781	0.400	0.320	0.450	0.400	2.097	0.590
1355 2-propanol	0.000	0.230	2.756	2.757	0.400	0.320	0.510	0.400	2.021	0.596
1411 CH ₂ =CH-CH ₂ COH	0.000	0.250	2.561	2.569	0.450	0.370	0.450	0.370	2.096	0.547
1353 1-butanol	0.000	0.242	2.755	2.757	0.400	0.320	0.450	0.400	2.601	0.731
1358 2-Me-propan-1-ol	0.000	0.240	2.679	2.762	0.400	0.320	0.450	0.400	2.299	0.731
1357 2-butanol	0.000	0.241	2.721	2.762	0.400	0.320	0.510	0.500	2.326	0.731
1359 1-butanol	0.000	0.236	2.657	2.725	0.400	0.320	0.570	0.400	2.119	0.731
1360 1-pentanol	0.000	0.248	2.756	2.763	0.400	0.320	0.450	0.500	2.126	0.872
1361 2-Me-butanol-1-ol	0.000	0.248	-	2.760	0.400	0.320	0.450	0.500	-	0.872
1362 3-Me-butanol-1-ol	0.000	0.245	2.569	2.736	0.400	0.320	0.450	0.500	-	0.872
1367 2-Me-butanol-2-ol	0.000	0.245	2.956	2.736	0.400	0.320	0.570	0.500	-	0.872
1440 cyclopentanol	0.000	0.270	2.890	2.900	0.400	0.320	0.510	0.510	-	0.742
1368 1-hexanol	0.000	0.252	2.890	2.553	0.400	0.320	0.450	0.500	2.610	1.013
1369 2-hexanol	0.000	0.250	2.890	2.530	0.400	0.320	0.510	0.500	2.340	1.013
1370 3-hexanol	0.000	0.251	-	2.543	0.400	0.320	0.510	0.500	2.440	1.017
1372 2-Me-pentanol-2-ol	0.000	0.248	-	2.512	0.400	0.320	0.570	0.500	2.119	1.013
1374 3-Me-pentanol-2-ol	0.000	0.252	-	2.553	0.400	0.320	0.570	0.500	2.277	1.013
1389 1-heptanol	0.000	0.256	2.924	2.954	0.400	0.320	0.450	0.500	4.115	1.154
1405 1-octanol	0.000	0.258	2.958	3.241	0.400	0.320	0.450	0.500	4.619	1.295
1416 1-nonanol	0.000	0.263	2.956	2.731	0.400	0.320	0.450	0.500	5.124	1.435
1426 1-decanol	0.000	0.262	2.592	4.129	0.400	0.320	0.450	0.500	5.628	1.576
1551 acetaldehyde	0.000	0.205	7.236	0.830	0.670	0.100	0.420	0.237	1.270	0.406
1552 propionaldehyde	0.000	0.223	8.350	1.200	0.650	0.000	0.410	0.261	1.815	0.547
1553 butyraldehyde	0.000	0.234	7.792	1.510	0.630	0.100	0.410	0.46	2.270	0.688
1554 iso-butyraldehyde	0.000	0.228	-	1.569	0.620	0.100	0.410	0.479	-	0.688
1570 trans-MeCH=CHCHO	0.000	0.262	12.530	1.890	0.750	0.000	0.35	0.445	-	0.645
1555 pentanal	0.000	0.239	8.760	1.981	0.607	0.100	0.41	0.571	2.770	0.829
1590 benzaldehyde	1.000	0.317	7.563	2.767	0.920	0.100	0.440	0.606	3.925	0.873
1651 2-propanone	0.000	0.220	8.294	1.207	0.710	0.100	0.430	0.280	1.760	0.547
1652 2-butanone	0.000	0.231	7.616	1.539	0.670	0.100	0.430	0.477	2.237	0.698
1657 2-pentanone	0.000	0.237	7.290	1.955	0.670	0.100	0.467	0.574	2.755	0.829
1705 cyclopentanone	0.000	0.262	10.890	1.886	0.750	0.100	0.521	0.500	2.120	0.720
1659 2-hexanone	0.000	0.243	7.020	2.257	0.65	0.100	0.480	0.67	2.262	0.970
1660 3-hexanone	0.000	0.243	7.023^e	2.257	0.650	0.100	0.430	0.668	2.310	0.970
1708 cyclohexanone	0.000	0.269	9.000	2.216	0.750	0.000	0.520	0.571	2.616	0.861
1664 2-heptanone	0.000	0.247	8.310	2.744	0.630	0.100	0.480	0.767	3.760	1.111
1675 2-octanone	0.000	0.250	7.400	3.128	0.610	0.100	0.430	0.850	4.257	1.251
1730 acetophenone	1.000	0.312	9.000	3.164	0.900	0.000	0.490	0.690	4.483	1.014
1685 2-nonanone	0.000	0.254	7.300	3.536	0.610	0.100	0.480	0.966	4.755	1.392
1690 2-decanone	0.000	0.256	7.200^c	2.924	0.610	0.100	0.430	1.051	5.260	1.533
1752 Et ₂ O	0.000	0.217	1.220	1.586	0.270	0.000	0.470	0.505	2.061	0.731
1353 n-Bu ₂ O	0.000	0.242	1.359	2.134	0.240	0.000	0.450	0.397	4.001	1.295
1414 furan	1.000	0.254	0.456	1.361	-	0.600	-	0.270	-	0.536
1421 dioxan	0.000	0.254	0.000	1.730	0.550	0.000	0.740	0.491	2.777	0.681
1450 PhOMe	1.000	0.303	1.904	2.775	0.730	0.000	0.320	0.630	3.926	0.916
1550 methyl acetate	0.000	0.220	2.956	1.337	0.600	0.000	0.420	0.424	1.960	0.606
1561 ethyl acetate	0.000	0.227	3.168	1.696	0.550	0.000	0.450	0.521	2.276	0.747
1581 methyl propionate	0.000	0.230	2.890	1.718	0.500	0.000	0.410	0.504	2.459	0.747
1582 n-propyl acetate	0.000	0.234	3.420	2.078	0.520	0.100	0.450	0.620	2.378	0.888
1584 n-butyl acetate	0.000	0.239	3.240	2.457	0.500	0.100	0.450	0.716	3.279	1.028
1587 pentyl acetate	0.000	0.244	3.067	2.852	0.450	0.100	0.450	0.721	3.881	1.169
1873 benzyl acetate	1.000	0.306	3.240	3.715	0.530	0.000	0.540	0.879	-	1.214
1921 methyl benzoate	1.000	0.302	3.240	3.240	0.770	0.100	0.740	0.776	4.634	1.075
1937 MeCOOH	0.000	0.227	2.025	1.826	0.450	0.110	0.54	0.27	2.29	0.465
1957 EtCOOH	0.000	0.235	2.067	-	0.450	0.270	0.530	0.419	-	0.606
1954 n-PrCOOH	0.000	0.241	1.510	-	0.450	0.500	0.540	0.510	-	0.747
1955 i-PrCOOH	0.000	0.239	1.190	-	0.45	0.600	0.54	0.516	-	0.747
1956 n-BuCOOH	0.000	0.247	1.590	-	0.450	0.600	0.540	0.507	-	0.698
1957 i-BuCOOH	0.000	0.244	1.740	-	0.500	0.600	0.54	0.516	-	0.698
1958 n-pentCOOH	0.000	0.251	1.220	-	0.500	0.600	0.540	0.708	-	1.028
1954 i-hexCOOH	0.000	0.251	1.30^e	-	0.450	0.600	0.540	0.804	-	1.169
1959 n-heptCOOH	0.000	0.258	1.30	-	0.480	0.670	0.54	0.90	-	1.310
1979 n-octCOOH	0.000	0.259	1.30^e	-	0.450	0.670	0.54	0.90	-	1.592
1957 PhCO ₂ Me	0.500	0.267	1.320	1.647	0.580	0.000	0.11	0.427	2.48	0.617
1957 PhCO ₂ Et	0.500	0.267	1.320	1.647	0.580	0.000	0.11	0.427	2.48	0.617

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174 n-tridecane
175 n-tetradecane

δ	$H(n^2)$	u	(n^2/V_x)	π_2^*	α_2	B_2	V_2	$\log 4u$	V_x
0.1	0.255	0.001	4.466	0.001	1.000	0.001	1.122	6.289	1.941
0.1	0.255	0.001	5.267	0.001	1.000	0.001	1.322	6.289	1.941

Table 4. Values of b used in equation (8) and (9).

Stationary phase	b
Zonyl	0.203
CWAX 1540	0.214
TCEP	0.178
Polyph ether	0.262
DEGS	0.190

Table 5. PARAMETER COEFFICIENTS OBTAINED FROM REGRESSIONS USING V_x

ZONYL E7										CORRELATION %				
d	π_2^*	α_m	β_m	V_x	C	n	r	s.d.	d	π_2^*	α_m	β_m	V_x	C
0.144	1.781	0.602	0.354	1.511	-2.313	104	0.9779	0.133	99.7	100	99.9	99.8	100	100
f(n^2)	μ^2	α_m	β_m	V_x	C	n	r	s.d.	f(n^2)	μ^2	α_m	β_m	V_x	C
9.266	0.063	0.281	1.205	1.158	-4.0	100	0.9579	0.187	100	100	92.4	100	100	100
CARBOWAX														
d	π_2^*	α_m	β_m	V_x	C	n	r	s.d.	d	π_2^*	α_m	β_m	V_x	C
0.236	2.17	2.343	-0.443	1.446	-2.206	103	0.9714	0.172	99.9	100	100	99.7	100	100
f(n^2)	μ^2	α_m	β_m	V_x	C	n	r	s.d.	f(n^2)	μ^2	α_m	β_m	V_x	C
13.051	0.068	1.809	0.842	0.968	-4.648	98	0.9548	0.215	100	100	100	100	100	100
TRICYANOETHOXYPROPANE														
d	π_2^*	α_m	β_m	V_x	C	n	r	s.d.	d	π_2^*	α_m	β_m	V_x	C
0.231	2.552	1.873	0.065	1.189	-1.787	108	0.9639	0.213	99.6	100	100	30.4	100	100
f(n^2)	μ^2	α_m	β_m	V_x	C	n	r	s.d.	f(n^2)	μ^2	α_m	β_m	V_x	C
13.92	0.088	1.445	1.388	0.651	-4.368	98	0.9494	0.248	100	100	100	100	100	100

Table 6. Parameter Coefficients obtained from regressions using Log L16

ZONYL E7										CORRELATION %				
δ	π_2^*	α_m	β_m	Log L16	C	\bar{N}	r	S.d.	δ	π_2^*	α_m	β_m	Log L16	C
-0.038	1.029	-0.065	0.708	0.422	-1.978	90	0.9910	0.09	68.6	100	58.9	100	100	100
$f(\bar{N}^2)$	μ^2	α_m	β_m	Log L16	C	\bar{N}	r	S.d.	$f(\bar{N}^2)$	μ^2	α_m	β_m	Log L16	C
3.389	0.052	-0.094	1.126	0.395	-2.611	87	0.9826	0.124	100	100	60.9	100	100	100
CARBOWAX														
δ	π_2^*	α_m	β_m	Log L16	C	\bar{N}	r	S.d.	δ	π_2^*	α_m	β_m	Log L16	C
0.034	1.549	1.743	-0.141	0.416	-1.93	94	0.9847	0.128	50.7	100	100	82.1	100	100
$f(\bar{N}^2)$	μ^2	α_m	β_m	Log L16	C	\bar{N}	r	S.d.	$f(\bar{N}^2)$	μ^2	α_m	β_m	Log L16	C
7.559	0.065	1.514	0.67	0.339	-3.376	88	0.9735	0.166	100	100	100	100	100	100
TRICYANOETHOXYPROPANE														
δ	π_2^*	α_m	β_m	Log L16	C	\bar{N}	r	S.d.	δ	π_2^*	α_m	β_m	Log L16	C
0.012	2.178	1.572	0.112	0.335	-1.63	94	0.9838	0.149	16.3	100	100	60	100	100
$f(\bar{N}^2)$	μ^2	α_m	β_m	Log L16	C	\bar{N}	r	S.d.	$f(\bar{N}^2)$	μ^2	α_m	β_m	Log L16	C
10.903	0.086	1.247	1.402	0.25	-3.749	89	0.9566	0.246	100	100	100	100	100	100

Table 9.

										CORRELATION %				
										μ	α_m	β_m	$\text{Log}L^6$	C
ZONYL E7														
$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C	n	r	s.d.	$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C
2.528	0.191	-0.293	0.997	0.404	-2.472	85	0.9815	0.128	99.9	100	99.1	100	100	100
CARBOWAX														
$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C	n	r	s.d.	$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C
6.933	0.254	1.44	0.30	0.352	-3.302	85	0.9753	0.162	100	100	100	96.2	100	100
TRICYANOETHOXYPROPANE														
$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C	n	r	s.d.	$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C
9.300	0.292	0.967	1.194	0.232	-3.482	85	0.9666	0.212	100	100	99.9	100	100	100
POLYPHENYL ETHER														
$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C	n	r	s.d.	$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C
5.166	0.172	0.261	0.231	0.470	-3.423	85	0.9886	0.106	100	100	98.5	98.6	100	100
DIETHYLENE GLYCOL SUCCINATE														
$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C	n	r	s.d.	$f(n^2)$	μ	α_m	β_m	$\text{Log}L^6$	C
9.468	0.221	0.995	0.976	0.268	-3.513	85	0.9666	0.191	100	100	100	100	100	100

Table 10.

ZONYL E7										CORRELATION %				
δ	μ	α_m	β_m	$\text{Log } L^6$	C	N	r	S.d.	δ	μ	α_m	β_m	$\text{Log } L^6$	C
0.345	0.193	-0.188	1.014	0.415	-1.940	85	0.9916	0.091	100	100	97	100	100	100
	μ	α_m	β_m	$\text{Log } L^6$	C	N	r	S.d.		μ	α_m	β_m	$\text{Log } L^6$	C
	0.252	-0.053	0.735	0.415	-1.935	62	0.9951	0.071		100	50.2	100	100	100
CARBOWAX														
δ	μ	α_m	β_m	$\text{Log } L^6$	C	N	r	S.d.	δ	μ	α_m	β_m	$\text{Log } L^6$	C
0.552	0.315	1.616	0.075	0.404	-1.851	88	0.9837	0.135	100	100	100	48.8	100	100
	μ	α_m	β_m	$\text{Log } L^6$	C	N	r	S.d.		μ	α_m	β_m	$\text{Log } L^6$	C
	0.369	1.850	-0.115	0.410	-1.902	65	0.9884	0.120		100	100	65.6	100	100
TRICYANETHOXYPROPANE														
δ	μ	α_m	β_m	$\text{Log } L^6$	C	N	r	S.d.	δ	μ	α_m	β_m	$\text{Log } L^6$	C
0.812	0.395	1.262	0.833	0.336	-1.535	83	0.9865	0.136	100	100	100	100	100	100
	μ	α_m	β_m	$\text{Log } L^6$	C	N	r	S.d.		μ	α_m	β_m	$\text{Log } L^6$	C
	0.530	1.810	0.075	0.343	-1.589	63	0.9872	0.138		100	100	34.5	100	100

Table 11. Further values of $\log L^{16}$ obtained at 298.15 K relative to n-octane standard.

Solute	$\log L^{16}$
n-Octane	3.677
Benzonitrile	3.994
Benzaldehyde	3.985
2-Methyl-3-pentanol	3.183
3-Methyl-3-pentanol	3.227
3-Hexanol	3.440
Ethyleneglycol diacetate	4.083
Pyrrrole	2.866
2-Chlorophenol	4.937
N-Methylaniline	4.494
2-Chlorotoluene	4.160
Methyl benzoate	4.634
Mesitylene	4.399

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