										P*
Solute	$\delta_0(eq 8),$ (cal/ cm ³) ^{1/2}	$\delta_0(\text{calcd}),$ $(\text{cal/cm}^3)^{1/2}$	V ₀ (exptl), cm ³	Vo(calcd), cm ³	$\epsilon = \frac{V_0(\text{exptl})}{V_0(\text{caled})}$	Imax	Xmax	Mol wt	P*, atm	(previ- ous), atm
Carbowax 4000	10.8	10.7 ± 0.9	520	2800	0.186	5.6×10^{54}	2.4×10^{-4}	3350	162	190
Carbowax 1000	10.9	10.7 ± 0.8	206	865	0.238	1.1×10^{21}	1.5×10^{-3}	1000	91	115
Stearic acid	7.9	8.5	257	329	0.780	1.7×10^{17}	1.6×10^{-3}	284	76	
1-Octadecanol	7.6	8.6	136	328	0.416	$6.7 imes10^8$	1.5×10^{-3}	270		

Table I: Solubility Properties of the Four Solutes in CO2 at 40°

acid with increasing pressure. This effect, consistent with eq 2 and 3, has not been previously observed.

The apparent existence of maxima in the continuous solubility-density curves of Figure 3 which lead to this peculiar phenomenon are, in fact, analogous to the formation of maxima in solubility-polarity plots with liquid mixtures. The explanation is presumably the same. The excess energy of mixing is a minimum when the cohesive energy densities are roughly equal for solute and solvent. Therefore the dense gas solubility parameter at a given maximum should equal the solubility parameter of the particular solute. This is in accord with eq 2 and 3, and, as we shall see in the next subsection, can be broadly confirmed for our specific group of solutes. Since these solutes have different solubility parameters, the maxima in Figure 3 are horizontally displaced from one another.

Although the present experimental pressure range, 270-1900 atm, is rather broad, it covers a relatively incompressible region well above the critical pressure. Thus the density and solubility parameter vary only about 25% in our experiment. While this has a very large effect on solubility, as Figure 3 illustrates, the limited range makes it difficult to show a clear solubility maximum with both descending branches distinctly exhibited. Nonetheless, the existence of a maximum is rather certain from (a) the shape of the curves in the experimental range, and (b) the fact that octadecanol and stearic acid are effectively nonvolatile at low pressures and must therefore have a descending branch on the left to complement the observed one on the right. In fact, previous work has been done with octadecanol on other instrumentation which shows a threshold pressure of 98.7 atm.¹³ This is equivalent to another point (actually a cluster of points) on the plot at (0, 4.8). Such an addition clearly requires a maximum in the solubility curve.

Figure 3 shows that the solubility peaks differ in width. From eq 2 it can be shown that the half-width is

$$\delta_{g} - \delta_{g,\max} = \left[\frac{2.3RT}{V_{0}} \left(\log I_{\max} - \log I\right)\right]^{1/2}$$
(9)

so that the width at any level below the peak maximum is related to solute mole volume as $V_0^{-1/2}$. Thus larger molecules have narrower solubility peaks, as strikingly

confirmed by the Carbowax 4000 and 1000 peaks in Figure 3. This is another instance where liquid and dense gas solubility phenomena follow parallel trends.

Solubility Enhancement. The enhancement in solubility (or volatility) relative to that for the ideal gas reaches very large proportions. At the solubility maxmum it is given by

$$I_{\rm max} = \exp(V_0 \delta_0^2 / RT) \tag{10}$$

Values of I_{max} are tabulated in Table I. For Carbowax 4000 the maximum enhancement is seen to be in excess of 10⁵⁰, reflecting mainly its extremely low vapor pressure under ideal conditions.

Absolute Solubility. The absolute magnitude of the equilibrium solute concentration can be approximated using the calibration procedure mentioned earlier. However, some uncertainty exists because of the need to estimate effective carbon numbers (θ in eq 5). For stearic acid and octadecanol we assumed θ as the number of carbon atoms in the molecule minus one, the excluded one corresponding to a carbon presumably tied up in nondetectable form (CO₂ or CO) in the pyrolysis products. For Carbowax 4000 (mean molecular weight, 3350) we assumed $\theta = 75$, the other half of the carbon atoms again presumably tied up in nondetectable form.

The results for the mole fraction of solute at the solubility peak, X_{max} , calculated from eq 5, are shown in Table I. Solubilities at other δ values may be calculated from eq 6 or obtained from the relative solubility plots in Figure 3.

Threshold Pressures. Of great experimental importance, particularly to gas chromatography, is the threshold pressure, P^* , the applied pressure which is barely sufficient to bring volatility to a measurable level. Since the solubility is an extremely steep function of pressure for large molecules, this parameter is nearly constant with reasonable variations in the sensitivity of the detection device.

For the present system, $R \times E = 1$ serves as the lowest limit of detection. Since none of the measurements was made near the threshold level, the latter was calculated assuming a parabolic dependence of the solubility signal on δ as in eq 6. Conversion from the obtained δ value to density and then to pressure yields the P^* values shown in Table I. Values obtained pre-