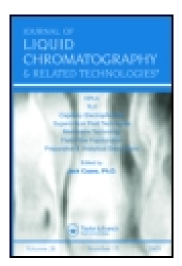
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## Comparison of Eluents in Supercritical Fluid Chromatography

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### COMPARISON OF ELUENTS IN SUPER-CRITICAL FLUID CHROMATOGRAPHY

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#### Summary

The chromatographic behavior of a number of low boiling eluents has been compared in supercritical fluid chromatography (SFC) with packed columns. Unmodified silica gel has been used as the stationary phase and a test mixture of polycyclic aromatic hydrocarbons (PAH's) as the substrate. Carbon dioxide, nitrous oxide, trifluoromethane, chlorotrifluoromethane, n-pentane, n-butane, i-butane, propane, ethane, diethylether, and dimethylether have been investigated as mobile phases.

Of the mobile phases  ${\rm CO_2}$ ,  ${\rm N_2O}$ ,  ${\rm C_2H_6}$ ,  ${\rm CHF_3}$ , and  ${\rm CClF_3}$ , which all possess low critical temperatures, comparisons were made at the same pressure and at similar reduced pressure. Poor transport properties were observed for  ${\rm CClF_3}$ , exemplified by extremely high capacity ratios for the PAH's. Of the remaining four mobile phases, at the same pressure, the highest capacity ratios were found for  ${\rm C_2H_6}$ , followed by  ${\rm CO_2}$ ,  ${\rm N_2O}$  and  ${\rm CHF_3}$ . The highest resolution was obtained with  ${\rm C_2H_6}$ , followed by  ${\rm CHF_3}$ ,  ${\rm N_2O}$ , and  ${\rm CO_2}$ . At similar reduced pressure,  ${\rm C_2H_6}$  showed again the highest capacity ratios, while  ${\rm CHF_3}$ ,  ${\rm CO_2}$ , and  ${\rm N_2O}$  nad lower values. The resolution showed a similar order.

For a given mobile phase, capacity ratio and resolution were correlated. To a first approximation, a higher capacity ratio leads to higher resolution. This correlation differs, however, for different eluents. For instance, among all mobile phases, the highest resolutions at a given k' were found for CHF3.

#### Introduction

For HPLC the choice of a suitable mobile phase is of greatest importance. Because this may be expected for supercritical fluid chromatography (SFC), a comparative study has been undertaken for a series of mobile phases

which are suitable for SFC. Some of these phases, like  ${\rm CO}_2$  and pentane, have frequently been used in SFC, while others, like ethane and  ${\rm CHF}_3$ , have not or only rarely been used. The choice of a mobile phase depends in large measure on the solubility power of the mobile phase for the substrate to be analysed. The common rule that "like dissolves like" is valid also for supercritical fluids if not only the molecular structure, but also the density of the supercritical fluid is considered. Therefore, the solubility parameter,  $\delta$ , of Hildebrandt and Scott <1>, as extended by Giddings to supercritical fluids <2,3>, considers not only the critical pressure,  $\rho_{\rm C}$ , as a molecular constant, but also the reduced density,  $\rho_{\rm r}$ , as a variable of state

1) 
$$\delta \text{ (gas)} = (3 \text{ p}_c)^{0.5} \cdot \frac{\rho \text{ (gas)}}{\rho c} = (3 \text{ p}_c)^{0.5} \cdot \rho_c$$

According to this simple equation, fluids of high  $p_c$  and  $\rho_r$  should lead to a high  $\delta$ . With a higher  $\delta$  the solubility power for more polar substrates usually increases. However, the solubility parameter concept does not account for specific interactions between fluid and substrate, e.g. for specific hydrogen bonding. Modification of Eq. 1 has been suggested <2,3> and the  $\delta$  have also been calculated considering a thermodynamic equation of state <4,5>. Again, specific interactions have not been considered.

Besides solubility, the selection of a supercritical mobile phase clearly depends on the critical temperature,  $T_{\rm C}$ . The  $T_{\rm C}$  should not be higher than the thermal stability of the substrate permits. On the other hand, it is convenient to choose an eluent whose  $T_{\rm C}$  is above ambient temperature. In this case the mobile phase can be pumped and metered as a liquid which, at the same pressure, possesses a lower compressibility than the supercritical gas. Moreover, the  $\delta$  and the solubility power tend to decrease with decreasing  $T_{\rm C}$ .

Mobile phases having a  $T_{c}$  not far above ambient possess the advantage of allowing low operating temperatures for SFC. They also allow admixing of a second component of higher T to the mobile phase, without raising the T of the resulting eluent to excessive levels. The second component is usually added for its higher solubility power.  $CO_2$  with  $T_C = 31.3$  °C has often been used because of its low price, noninflammability and nontoxicity. Its good dissolution properties for nonpolar to moderately polar substrates is indicated by a relatively high  $p_C$  of 78.5 bar.  $N_{\gamma}O$  possesses similar critical data, but due to its small permanent dipole moment it may be capable of specific interactions. Fluorocarbons having a T shortly above room temperature are  $CHF_3$  and  $CClF_3$ . Both are noninflammable and of low toxicity, but are more expensive in chromatographic purity and may have harmful effects on

the ozone layer of the upper atmosphere. In the homologous series of the alkanes, it is only  ${\rm C_2H_6}$  which has a low  ${\rm T_c}$  above room temperature.

The results obtained with the five eluents of low  $T_C$  are presented in this work, together with results on mobile phases of higher  $T_C$  studied earlier, i.e. alkanes <6,7> and linear ethers <8>. The earlier work on alkanes and ethers has given capacity ratios, k', and resolutions, R, in dependence of temperature, T, and column outlet pressure  $p_e$  <6-9>. As an outstanding feature, the k' and R exhibited temperature dependent maxima above  $T_C$ . It is of interest now whether the five mobile phases of low  $T_C$  will show a similar behavior.

#### Experimental

The apparatus and the column have been described previously <8>. All chromatographic runs were performed on columns (25 x 0.46 cm) packed with unmodified silica gel and a test mixture of the polycyclic aromatic hydrocarbons naphthalene, anthracene, pyrene and chrysene (PAH's) as the substrate. Purities and sources of mobile phases and substrates are given in Table I. For all runs the flow rate was 1 ml/min, measured at the pumps in the liquid state and checked volumetrically at the column outlet.

Table I: Mobile phases and substrates

pentane	technical product	<pre>dried over sodium, distilled</pre>
butane	Linde AG, Höllriegels- kreuth, F.R.G.	99.5 %
i-butane	Linde AG	99.5 %
propane	Linde AG .	99.5 %
ethane	Messer-Griesheim, Hermülheim, F.R.G.	99.5 %
diethyl ether	technical product	dried over sodium distilled
dimethyl ether	Merck AG, Darmstadt F.R.G	99 %, 0.3 % methanol
co <sub>2</sub>	Messer-Griesheim	99.995 %
N <sub>2</sub> O	Messer-Griesheim	99.8 %
CHF <sub>3</sub>	Messer-Griesheim	99.8 %
CC1F <sub>3</sub>	Messer-Griesheim	99.8 %
heptane	technical product	<pre>dried over sodium, distilled</pre>
naphthalene, anthracene, pyrene, chrysene		recry- stallized

Capacity ratios,  $k^*$ , and resolutions, R, were calculated as described earlier <10>:

$$k' = \frac{t_r - t_o}{t_o}$$

3) 
$$R_{ij} = \frac{f_{ij}}{g_{ij}} + \frac{d_{ij}}{w'_{i} + w'_{j}} \cdot \sqrt{\ln 4}$$

$$R_{m} = \frac{\sum_{i=1}^{n} R_{ij}}{n}$$

 $t_r$  = retention time

t = dead time

 $R_{ij}$  = resolution between neighboring peaks i and j

 $f_{ij}$  = depth of valley between neighboring peaks, as measured by starting from the average peak height

 $g_{ij}$  = average peak height

d<sub>ij</sub> = baseline distance between tangent on peaks

w' = peak width at half height

 $R_m = mean resolution$ 

Results and discussion

The critical data and the boiling points of all mobile phases are collected in Table II. The low boiling temperatures of  ${\rm CO_2}$ ,  ${\rm N_2O}$ ,  ${\rm C_2H_6}$ ,  ${\rm CHF_3}$ , and  ${\rm CClF_3}$  and the prepressurization of the mobile phase by helium in the storage tank require considerable working pressure to prevent gas formation at ambient temperature in the apparatus, e.g. in the detector. This is particularly true for  ${\rm C_2H_6}$  and  ${\rm CClF_3}$  which require a rather high minimum column outlet pressure  ${\rm p_e}$ . The temperature and the pressure ranges for the chromatographic runs are listed in Table III.

Table II: Critical data and boiling points of eluents

eluent	critical temper- ature T <sub>C</sub> /°C	critical pressure p <sub>C</sub> /bar	critical density $\rho/g \text{ cm}^{-3}$	boiling temper- ature T <sub>b</sub> /°C
n-pentane	196.5	33.7	0.237	36.1
n-butane	152.0	38.0	0.228	- 0.5
i-butane	135.0	36.5	0.221	-11.7
propane	96.7	42.5	0.217	-42.1
ethane	32.3	48.8	0.203	-88.7
diethyl ether	192.6	35 <b>.6</b>	0.265	34.6
dimethyl ether	126.9	52.6	0.271	-24.8
co <sub>2</sub>	31.3	73.8	0.468	-78.5 (s)*
N <sub>2</sub> O	36.4	72.7	0.435	-88.5
CHF <sub>3</sub>	25.6	48.4	0.516	-82.1
cclF <sub>3</sub>	28.9	39.2	0.579	-81.5

(s)\* = sublimation

Comparison between different mobile phases may be conducted in different ways. Generally applicable is a comparison at the same reduced free volume <11> since this comparison is independent of the molecular weight of the eluents. Related is a comparison at equal reduced density, because the reduced density and free volume are both relative densities. Using simply the same density is less informative, an exception being homologous series, because

Table III: Experimental temperature and pressure ranges

eluent	temperature/	pressure/bar
	°C	
pentane	20 ~ 280	6, 20, 36, 70
butane	20 - 280	39, 70
i-butane	20 - 280	37, 70
propane	20 - 280	43, 70
ethane	20 - 250	180, 210, 250
diethyl ether	20 - 280	6, 38, 70
dimethyl ether	20 - 280	56, 70
co <sub>2</sub>	20 - 250	120, 150, 180, 210, 250
N <sub>2</sub> O	20 - 230	120, 180, 250
CHF <sub>3</sub>	20 - 200	150, 200
cclr <sub>3</sub>	100 - 200	250, 300

then the same density represents approximately the same free volume <12>. For practical reasons, a comparison at equal pressure and temperature or at the same reduced pressure and temperature is useful. Comparison at the same capacity ratio, k', has also been used in SFC <13> and leads to a comparison by way of a chromatographic parameter.

The behavior of k' versus temperature at  $p_e$  = const. for the two fluorinated hydrocarbons CHF $_3$  and CClF $_3$  is presented in Figs. 1 and 2, respectively. The CHF $_3$  shows

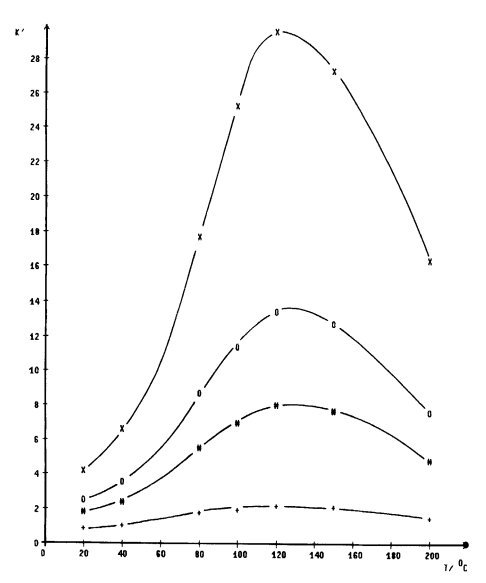


Fig. 1 Dependence of capacity ratios, k', on column temperature, T. Eluent: CHF<sub>3</sub> (Freon 23).

Column end pressure, p<sub>e</sub>: 150 bar. Substrates: (+) naphthalene, (#) anthracene, (O) pyrene, (X) chrysene (PAH's).

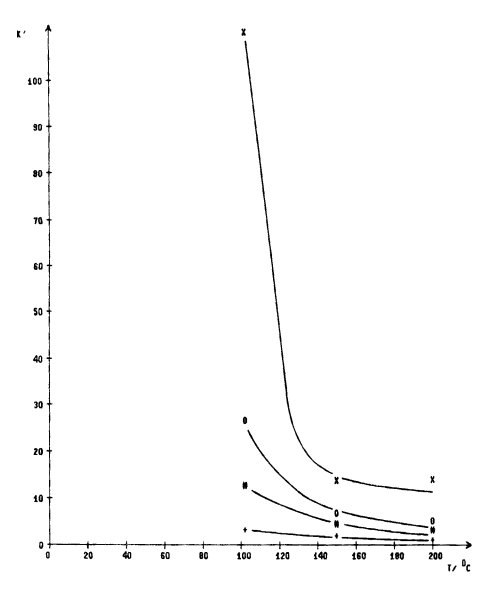


Fig. 2 Dependence of capacity ratios, k', on column temperature, T. Eluent:  ${\rm CClF}_3$  (Freon 13).  ${\rm p}_e\colon$  250 bar. Substrates as in Fig. 1.

a strong maximum of k' for the four PAH's in analogy to the maxima found for eluents of higher  $T_{\rm C}$ , i. e. alkanes <6,7> and ethers <8>. For  ${\rm CClF}_3$  very high k' were indicated at temperatures between  $T_{\rm C}$  and 100 °C, values which could not be determined because of very slow elution. The very high k' are found despite of an increase in pressure from  $p_{\rm e}$  = 150 bar for  ${\rm CHF}_3$  to  $p_{\rm e}$  = 250 bar for  ${\rm CClF}_3$ . Therefore, the elution power and probably also the dissolution ability of  ${\rm CClF}_3$  is much smaller than for  ${\rm CHF}_3$  and too small to yield measurable k'. As a consequence, the  ${\rm CClF}_3$  is not included further in the comparisons between mobile phases.

The  $T_{\rm C}$  of  ${\rm CO_2}$ ,  ${\rm N_2O}$ ,  ${\rm C_2H_6}$ , and  ${\rm CHF_3}$  are all between 25 and 37 °C. Nevertheless, there exist large differences in k' at the same  ${\rm p_e}$ . For chrysene as the substrate, the k'(c) (c = chrysene) versus T curves of different  ${\rm p_e}$  are shown in Figs. 3 - 5. The highest k' are found for  ${\rm C_2H_6}$ , followed by  ${\rm CO_2}$ ,  ${\rm N_2O}$ , and  ${\rm CHF_3}$ . The outstanding feature of the isobaric curves in Figs. 3 - 5 is a more or less well developed maximum. The temperature at which the maximum occurs and the pressure dependence of this temperature is different for different eluents. For instance, the k' maximum for  ${\rm C_2H_6}$  is located at about 100 °C at a pressure of 180 bar (Fig. 3). This is significantly below the temperature of the maximum for  ${\rm N_2O}$  or  ${\rm CO_2}$  which both lie at approximately 120 °C. At

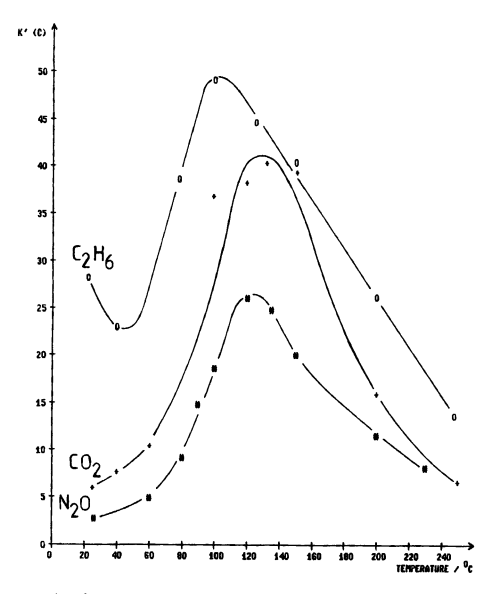


Fig. 3 Dependence of the capacity ratios of chrysene, k'(C), on column temperature, T. Eluents: (+)  $CO_2$ , (#)  $N_2O$ , (O)  $C_2H_6$ .  $P_e$ : 180 bar.

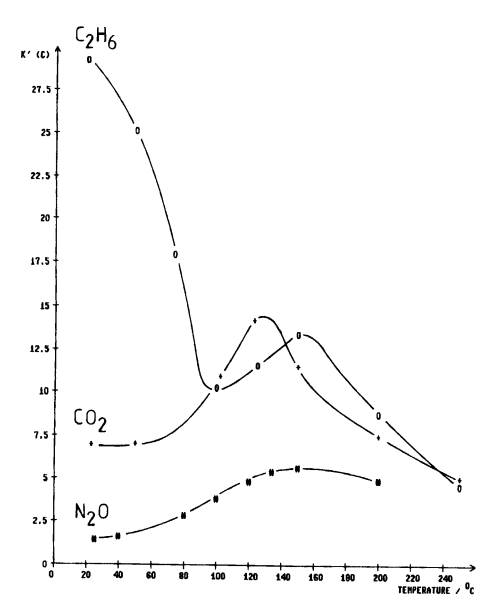


Fig. 4 Dependence of the capacity ratios of chrysene, k'(C), on column temperature, T. Eluents: (+)  $CO_2$ , (#)  $N_2O$ , (O)  $C_2H_6$ .  $p_e$ : 250 bar.

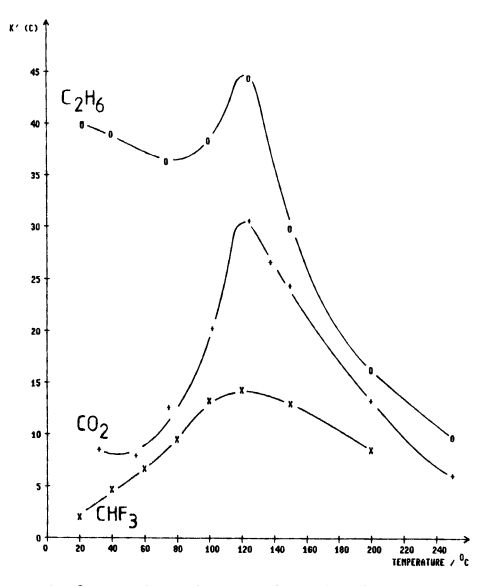


Fig. 5 Dependence of the capacity ratios of chrysene, k'(C), on column temperature, T. Eluents: (+)  $\rm CO_2$ , (O)  $\rm C_2H_6$ , (X)  $\rm CHF_3$ .  $\rm p_e$ : 210 bar for  $\rm CO_2$  and  $\rm C_2H_6$ , 200 bar for  $\rm CHF_3$ .

250 bar, however, the  $\mathrm{C_2H_6}$  and  $\mathrm{N_2O}$  maxima are situated both at about 160 °C, while that for  $\mathrm{CO_2}$  has hardly moved (Fig. 4). At an intermediate pressure of 210 bar, the maxima of  $\mathrm{C_2H_6}$  and  $\mathrm{CO_2}$  are seen at the same temperature of 120 °C. Besides the dependence of location of the maximum on pressure, there is a dependence of arcal and height of the maximum on this variable. A higher pressure reduces both area and height (cf. Figs 3 and 4). This type of pressure dependence of area and height of the k' maxima has also been found before <6,7>.

The mean resolution  $R_m$ , obtained for the four mobile phases, is given in Figs. 6 - 8. The chromatographic conditions are the same as for Figs. 3 - 5. Accordingly,  $C_2H_6$  yields not only the highest k', but also the highest  $R_{m}$ . In Table IV the eluents have been put in approximate order of increasing  $k^{\,\prime}$  and  $R_{_{\boldsymbol{m}}}.$  The order at equal  $p_e$  is for k' and  $R_m$  not the same since  $CO_2$  and  $\mathrm{CHF}_3$  have changed places. Therefore, a high  $\mathrm{R}_\mathrm{m}$  at p\_=const. is not necessarily obtained at the expense of a higher k', i. e. higher analysis time. The temperature of the  $R_{m}$  maximum is found close to that of the  $k^{\,\prime}$  maximum, the largest differences between the two being found for  $C_2H_6$  in Figs. 3 - 8. Like the k' maxima, the maxima for  $R_{m}$  move to higher temperatures and become smaller in area and height with rising pressure. Also noteworthy is, that  $C_2H_6$  exhibits very high k' and  $R_m$ , both in the

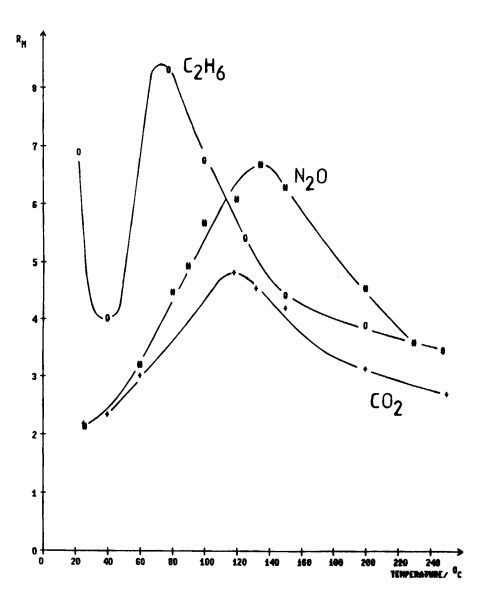


Fig. 6 Dependence of the mean resolution between the four PAH's,  $R_{\rm m}$ , on column temperature, T. Eluents: (+)  ${\rm CO_2}$ , (#)  ${\rm N_2O}$ , (O)  ${\rm C_2H_6}$ .  ${\rm p_e}$ : 180 bar.

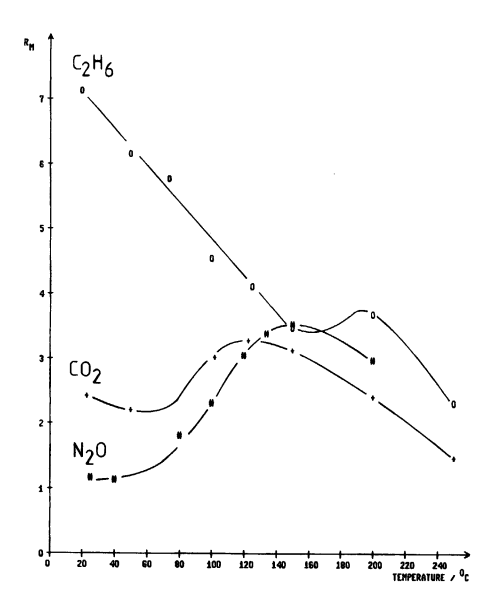


Fig. 7 Dependence of the mean resolution between the four PAH's,  $R_{\rm m}$ , on column temperature, T. Eluents: (+)  ${\rm CO_2}$ , (#)  ${\rm N_2O}$ , (O)  ${\rm C_2H_6}$ .  ${\rm p_e}$ : 250 bar.

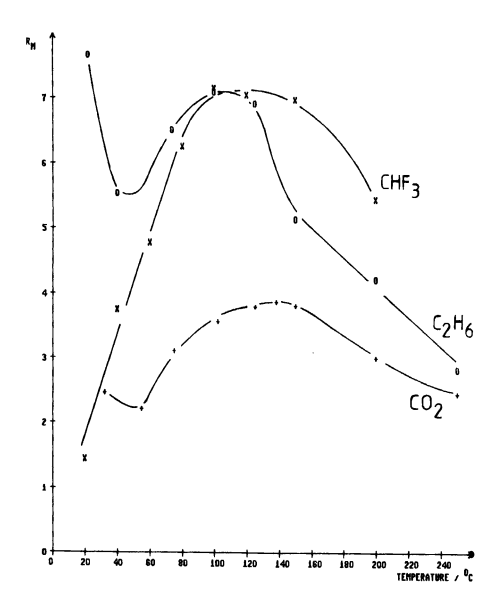


Fig. 8 Dependence of the mean resolution between the four PAH's,  $R_{\rm m}$ , on column temperature, T. Eluents: (+)  ${\rm CO_2}$ , (0)  ${\rm C_2H_6}$ , (X)  ${\rm CHF_3}$ ,  ${\rm P_e}$ : 210 bar.

Table IV: Comparison of k' and R for eluents of low  $T_c$ . Comparisons carried out at equal pressure,  $p_e$ , at equal reduced pressure  $p_r$ , or at equal capacity ratio, k'.

supercritical and in the liquid region. At higher pressures, the k' and R $_{\rm m}$  observed for C $_2$ H $_6$  near T $_{\rm c}$  are even larger than those at the maximum in the supercritical region.

Conducting the comparison between different mobile phases at equal reduced pressures,  $\mathbf{p_r}$ , is expected to lead to results which are different from those at the same pressure,  $\mathbf{p_e}$ . The critical pressures of  $\mathrm{CO_2}$  and  $\mathrm{N_2O}$  are much larger than for  $\mathrm{C_2H_6}$  and  $\mathrm{CHF_3}$ . Therefore, a comparison at equal  $\mathbf{p_r}$  for  $\mathrm{CO_2}$  and  $\mathrm{N_2O}$  on one hand, and  $\mathrm{C_2H_6}$  and  $\mathrm{CHF_3}$  on the other, is equivalent to a comparison at a higher  $\mathbf{p_e}$  for the first two eluents to a lower  $\mathbf{p_e}$  for the last two eluents. In Figs. 9 and 10 such

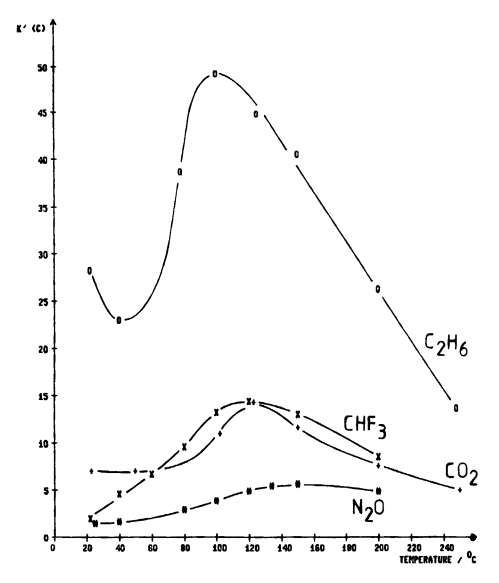


Fig. 9 Dependence of the capacity ratios of chrysene, k'(C), on column temperature, T at similar reduced pressures, p<sub>r</sub>. Eluents and pressures:

- (+)  $CO_2$ ,  $p_e = 250$  bar,  $p_r = 3.39$ ;
- (#)  $N_2O$ ,  $r_1$  = 250 bar,  $p_r$  = 3.44;
- (0)  $C_2H_6$ ,  $p_e = 180$  bar,  $p_r = 3.71$ ;
- (x) CHF<sub>3</sub>,  $p_e = 200$  bar,  $p_r = 4.13$ .

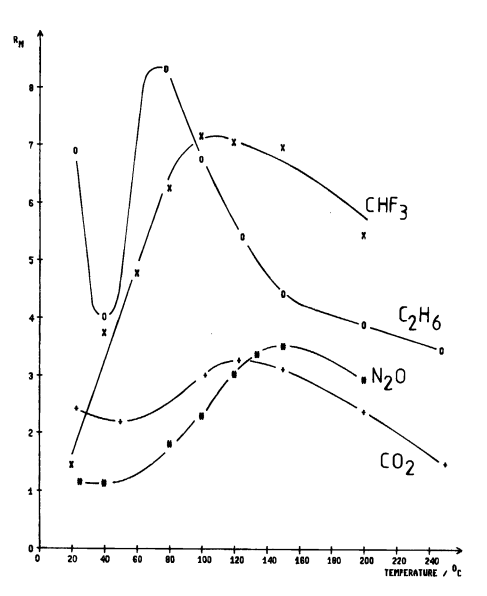


Fig. 10 Dependence of the mean resolution between the four PAH's,  $R_{\rm m}$ , on column temperature, T, at similar reduced pressures,  $p_{\rm r}$ . Eluents and pressures:(+)  ${\rm CO_2}$ ,  $p_{\rm e}$  = 250 bar,  $p_{\rm r}$  = 3.39; (#)  ${\rm N_2O}$ ,  $p_{\rm e}$  = 250 bar,  $p_{\rm r}$  = 3.44; (O)  ${\rm C_2H_6}$ ,  $p_{\rm e}$  = 180 bar,  $p_{\rm r}$  = 3.71; (X)  ${\rm CHF_3}$ ,  $p_{\rm e}$  = 200 bar,  $p_{\rm r}$  = 4.13.

a comparison is seen for k' and  $R_m$ .  $CO_2$  and  $N_2O$  $(p_r (CO_2) = 3.39; p_r (N_2O) = 3.44)$  show about the same area under the  ${\rm R}_{\rm m}\text{-curve}$  and the same height of the maxima (Fig. 10), but these features are smaller for the k'-curves of  $N_2^0$  versus  $CO_2$  (Fig. 9).  $C_2^H_6$  and  $CHF_3 (p_r (C_2H_6) = 3.71; p_r (CHF_3) = 4.13)$ exhibit also about the same area and height for  $R_{\mathrm{m}}$ , but  $\mathrm{C_{2}H_{6}}$  has a decidedly higher k'. Because the  $\mathrm{t_{o}}$  are similar for  $C_2H_6$  and  $CHF_3$ , analysis time is in favor of CHF, at comparable resolution. Comparing now  ${\rm CO}_2$  and  $N_2O$  on one hand to  $C_2H_6$  and  $CHF_3$  on the other, it is to be admitted that the pr for the latter are significantly higher. However, the differences between the two pairs of eluents are so large as to make the smaller k' and  $R_m$  for  $CO_2$  and  $N_2O$  certain. The k' and  $R_m$  will become even smaller when the  $p_{_{m{r}}}$  would be increased to meet the  $p_1$  of  $C_2H_6$  and  $CHF_3$ . Also, according to Eq. 1, the eluents  ${
m CO}_2$  and  ${
m N}_2{
m O}$  give higher solubility parameters than  $C_2H_6$  and  $CHF_3$  at conditions of comparable  $\mathbf{p_r}$  because the  $\mathbf{p_c}$  of the former two phases is higher than those of the latter.

A comparison between mobile phases at the same k' is per definition a comparison at the same relative elution time, i. e. at the same ratio of the times the substrate spends in the stationary and in the mobile phase. Equal k' area, however, not necessarily mean equal retention times, tr,

because the same capacity ratio for different mobile phases is obtained by different t<sub>o</sub>. This arises because for different p - T pairs the liquid volume feed rate was kept constant. Nevertheless, k' gives also an indication of analysis time.

Prior to comparison between different mobile phases at the same k', it is important to know whether there exists a correlation between R<sub>m</sub> and k' already for a single mobile phase. The data for CO<sub>2</sub> at different p<sub>e</sub> - T pairs are given for Fig. 11 as an example, whereby only runs at supercritical conditions are included. A coarse but definite correlation becomes apparent between the resolution of naphthalene and anthracene, R(NA), and the capacity ratio of anthracene, k'(A). The band, defining the range of scatter is at lower k' relatively narrow and of high slope whereas at higher k' wider and of lower slope. In the k' range between 0 and 10, which is the range of greatest practical importance, a small increase in k' will be accompanied by a large increase in R.

Because a correlation might exist between R and k' also for other single mobile phases, the R(NA) versus k'(A) data are plotted in Fig. 12 for the five alkanes n-pentane, n-butane, i-butane, propane, and ethane. Although the scatter for a given alkane is considerable and the scattering ranges overlap each other, correlations of the

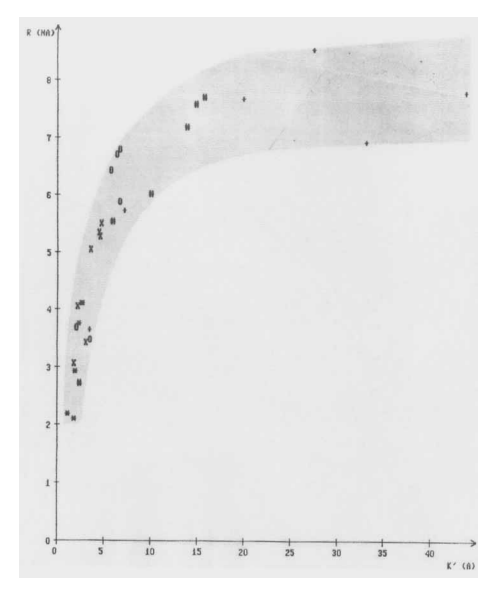


Fig. 11 Dependence of the resolution between naphthalene and anthracene, R(NA), on the capacity ratio of anthracene, k'(A). Eluent: CO<sub>2</sub>. p<sub>e</sub>: (+) 120 bar, (#) 150 bar, (0) 180 bar, (X) 210 bar, (\*) 250 bar (p<sub>r</sub> = 1.63 - 3.39). Different temperatures as shown in Table III.

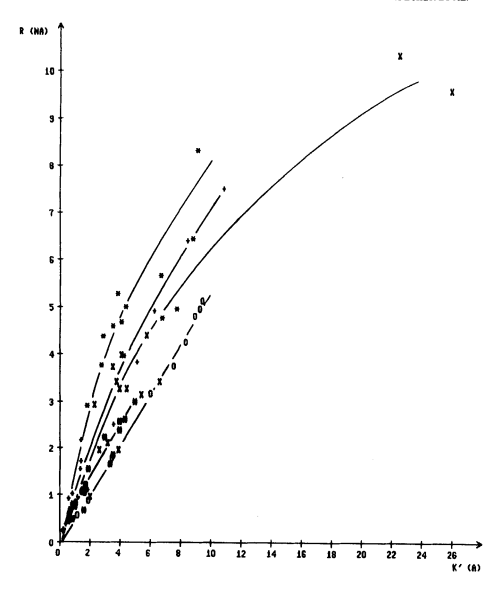


Fig. 12 Dependence of the resolution between naphthalene and anthracene, R(NA), on the capacity ratio of anthracene, k'(A). Eluents: (+) pentane, (#) butane, (O) i-butane, (X) propane, (\*) ethane. Different p<sub>e</sub> and T.

individual alkanes apparently not only exist but the correlations also seem to differ from each other to a significant extent. In Fig. 13 this type of comparison is extended to mobile phases which do not belong to the same homologous series or the same class of compounds. Here, the differences between the curves are larger than in Fig. 12, particularly at higher k'. Larger differences in the performance of the eluents in terms of resolution and analysis times is therefore observed. Thus the CHF $_3$  combines high resolution with high analysis speed, particularly at higher k', while this is less the case for  $N_2$ O,  $CO_2$ , pentane, or diethylether.

In Fig. 14 the resolution of all peak pairs, R<sub>m</sub>, is plotted versus k' of the most retained solute, chrysene, instead of only the resolution between naphthalene and anthracene, R(NA). In this way additional experimental data at higher k' are included in the comparison between different mobiles phases. Obviously, the differences of the mobile phases in Fig. 14 are basically the same as in Fig. 13.

The present study has been carried out to provide practical guide lines for the selection of mobile phases in SFC.

Because the study has been carried out solely with unmodified silica gel and a test mixture of polycyclic aromatic hydrocarbons, the comparisons described should be extended to other stationary phases and other substrates.

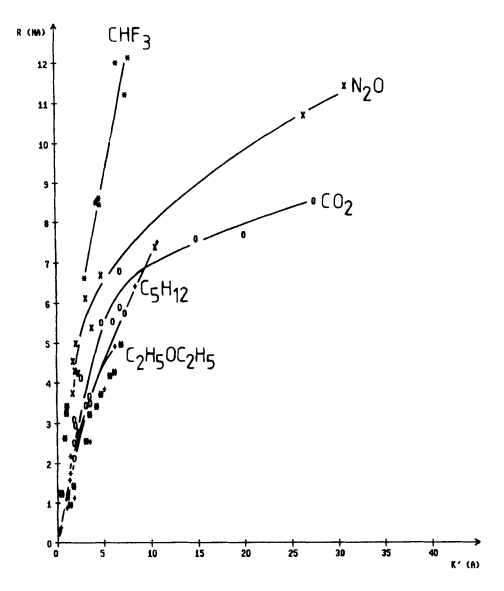


Fig. 13 Dependence of the resolution between naphthalene and anthracene, R(NA), on the capacity ratio of anthracene, k'(A). Eluents: (+) pentane, (#) diethylether, (0) CO<sub>2</sub>, (X) N<sub>2</sub>O, (\*) CHF<sub>3</sub>. Different p<sub>e</sub> and T.

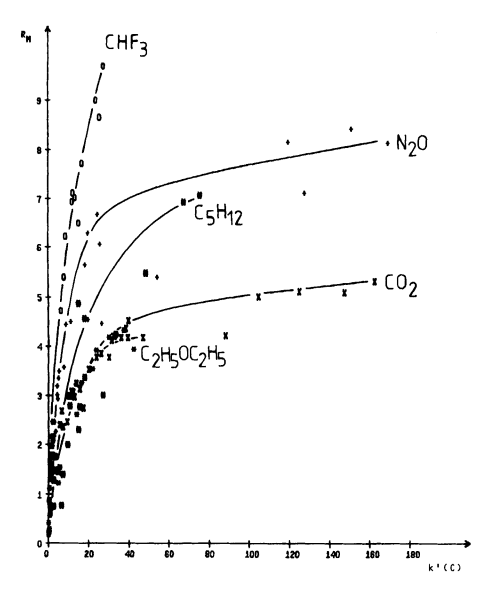


Fig. 14 Dependence of the resolution between all PAH's,  $R_{\rm m}$ , on the capacity ratio of chrysene k'(C). Eluents: (#) pentane, (\*) diethylether, (X)  ${\rm CO_2}$ , (+)  ${\rm N_2O}$ , (O)  ${\rm CHF_3}$ . Different  ${\rm p_e}$  and  ${\rm T}$ .

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