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Use of an open-tubular trapping column as phase-switching interface in on-line coupled reversed-phase liquid chromatography-capillary gas chromatography*

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ABSTRACT

The applicability of open-tubular traps for phase switching in coupled RPLC-GC was studied. The phase-switching process involves sorption of the analytes of interest from a methanol-water mobile phase into the stationary phase of an open-tubular column, removal of the aqueous phase by purging the trap with nitrogen and desorption of the analytes with hexane. Water elimination carried out in this manner appears to be highly efficient. In the sorption step the sampling flow-rate and the capacity factors of the analytes in the trap are critical parameters. Using a 2 m x 0.32 mm I.D. trap with a swollen 5-µm stationary phase at flow-rates not exceeding 100 µl/min, polycyclic aromatic hydrocarbons are trapped quantitatively from 300 µl of aqueous phases containing up to 65% (v/v) of methanol. For desorption 70–125 µl of hexane are needed. These volumes are easy to handle in solvent elimination carried out using a PTV injector prior to transfer of the analytes to a GC column.

INTRODUCTION

The main issue in on-line coupled LC-GC currently is the coupling of reversed-phase LC (RPLC) with capillary GC, i.e., the introduction of aqueous eluents into the GC system. The most straightforward solution is direct transfer of the eluent fraction of interest into the GC system as is done in normal-phase LC–GC[1]. This involves refocusing of the analytes by means of a retention gap or by using a PTV injector with a modified liner [2,3]. Both techniques rely on the formation of a solvent film, which in turn presupposes wettability. As water does not form a uniform solvent film on deactivated surfaces, it is not a suitable solvent for the above-mentioned techniques [4]. Moreover, water has a high boiling point, requiring high solvent evaporation temperatures, and it produces a very large volume of vapour per unit volume of liquid, which seriously limits the speed of introduction into the GC system. Also, it is chemically aggressive at higher temperatures, destroying the chemically deactivated surface of precolumns. So far, there is no fully water-resistant deactivation procedure [5]. As a consequence, for direct injections of aqueous samples into the GC system, problems with adsorption of analytes in the precolumn will arise sooner or...
later. Nevertheless, the direct transfer of aqueous solvents has been used in a limited number of studies, i.e., when only small volumes are involved (packed capillary LC-GC) [6,7] and when applying concurrent solvent evaporation [8]. Furthermore, introduction of mobile phases containing water can be successful if water evaporates before the other solvent(s) as in co-solvent trapping [9] and with solvent mixtures containing an amount of water not exceeding the azeotropic composition [10].

Eliminating the aqueous solvent prior to introduction into the GC system enhances the applicability of coupled RPLC-GC. Removal of water can be done by sorption-thermal desorption and phase-switching techniques. Pankow et al. [11] described a sorption-thermal desorption system in which water was sampled through a Tenax tube. The adsorbed analytes were thermally desorbed and introduced into the GC system after refocusing. Schomburg et al. [12] used a similar approach based on a two-oven system. More recently, the use of a PTV injector with packed liners was studied for the same purpose [13], thereby eliminating the need for two ovens. For trace analysis of organics in water, also non-coated [14,15] or coated capillaries [16-18] or entire GC columns [19-21] were used for trapping analytes from water. After thermal desorption and subsequent (cryogenic) refocusing the analytes were analysed.

In phase-switching techniques the analytes are transferred from the aqueous phase to an organic phase which is introduced into the GC system. Inspired by the promising results obtained in the field of automated sample preparation of water samples [22,23], Van Zoonen et al. [24] studied the use of on-line liquid-liquid extraction in a segmented flow system as a means of transferring the analytes from a methanol-water LC phase to an organic phase. Phase switching can also be carried out by the use of small packed trapping columns [25,26]. After trapping the analytes from the aqueous phase, the residual water has to be removed by drying with a flow of nitrogen. Incomplete elimination of water leads to reduced desorption efficiencies as part of the pores of the packing material are blocked by residual water. The need for a drying step can be avoided by using a desorption solvent which is (slightly) miscible with water, e.g., ethyl acetate [27], but then again small amounts of water are transferred to the GC system. Grob and Schilling [28] proposed the use of open-tubular trapping columns as an alternative to the use of packed adsorption tubes. The main advantage of the use of open-tubular traps is that complete removal of the remaining reversed-phase eluent can be obtained by simply purging a short plug of gas through the capillary. Grob and Schilling’s attempts to trap analytes on a 2-m GC column were not very successful. According to the authors this was due to the low diffusion speeds in the liquid and insufficient retention power of the open-tubular trap.

The aim of this paper is to present a thorough study on the applicability of open-tubular traps as phase-switching devices for coupled RPLC–GC. A number of selected test compounds from a volume of methanol-water, which represents an average RPLC eluent, is trapped in a coated capillary. The eluent is then removed by emptying the trap by means of nitrogen prior to desorption of the analytes with hexane, which is next introduced directly into the GC system. Refocusing of the analytes is achieved by using a PTV injector. The phase-switching process was studied in two steps. First the sorption step was studied by determining the breakthrough volumes of polycyclic aromatic hydrocarbons (PAHs) in the LC elution mode. The effects of column dimensions, flow-rate, modifier concentrations and temperature are discussed. Second, starting from 100% sorption conditions, experiments on liquid desorption were carried out. Here the effect of desorption flow-rate and length of the trapping column on the desorption volume are discussed. The applicability of the system is demonstrated by the analysis of a PAH mixture.

**EXPERIMENTAL**

**Instrumentation**

Sorption experiments were carried out using an LC system which consisted of an LC pump (LKB 2 150; Pharmacia, Woerden, Netherlands), a pulse damper (Free University, Amsterdam, Netherlands), a six-port valve with an 11-μl loop, an open-tubular trapping column, a fluorescence detector (LS-4; Perkin-Elmer, Norwalk, CT, USA) and a recorder (BD40; Kipp & Zonen, Delft, Netherlands). The trapping capillaries were cut from new 0.32 mm I.D. GC columns with either a 1.1- or a 5.1-μm film
of CP-Sil-5-CB (Chrompack, Bergen op Zoom, Netherlands). The valve and the trapping capillary were positioned in a thermostatted water-bath. The fluorescence excitation/emission wavelength for naphthalene, phenanthrene and pyrene were 262/330, 243/365 and 236/392 nm, respectively.

The phase-switching device was built around two ten-port valves (Valco, Houston, TX, USA), depicted schematically in Fig. 1. The system consisted of a sampling pump (P1) (syringe pump, type MF-2 from Azumadenki Kogyo, Japan) and a pump used for desorption (P2). This pump was a microprocessor-controlled syringe pump (Digisampler, Gerstel, Mülheim a/d Ruhr, Germany) which allowed the introduction of defined volumes of up to 1 ml with a speed of 1-2000 µl/min. A fraction loop of 300 µl was connected between two ports of valve 1. The open-tubular coated traps were connected between two ports of valve 2. The nitrogen flow applied to valve 1 was controlled by a pressure regulator (adjusted to 1 bar) and a flow controller adjusted to 0.6 ml/min.

A gas chromatograph (Model 5890; Hewlett-Packard, Avondale, PA, USA) with flame ionization detection (FID) and thermal conductivity detection (TCD), and provided with an automated cold temperature-programmed injection system (PTV injector) (KAS 502; Gerstel) was used. For data collection a Nelson integration system (Perkin-Elmer) was used. The PTV injector was equipped with a liner containing a porous glass bed and a modified split vent as described elsewhere [2,3]. With this configuration low outlet pressures could be obtained, allowing high purge gas flow-rates (up to 620 ml/min). By means of a splitter, about 1% of the purge gas is transferred to the TCD instrument for monitoring the solvent elimination.

Operating conditions

Large-volume injections. Large-volume injections were carried out using the microprocessor-controlled syringe pump. A sample of PAHs dissolved in hexane was introduced into the PTV injector (initial temperature between −30 and 30°C) at a predefined rate. The sample was introduced in the solvent split mode, applying a purge gas flow-rate of 620 ml/min. When almost no solvent was left in the liner, as could be seen from the TCD signal, the GC run was started. The start of the GC run involves closing of the split valve (splitless time, 1.0 min), starting the temperature programming of the column and heating the PTV injector to 275°C at 12°C/s. The PTV injector was kept at this temperature for 1 min.

For the GC separations a 12.5 m x 0.32 mm I.D. column coated with 0.12-µm CP-Sil-5-CB was used with helium as the carrier gas at an inlet pressure of 45 kPa. The GC temperature programme was as follows: initial temperature, 40°C for 1 min, then increased at 15°C/min to 250°C, the final temperature being held for 5 min.

On-line phase-switching GC. The position of the valves at the start of a phase-switching cycle is shown in Fig. 1. The LC fraction loop of valve 1 is filled manually with a PAH solution in water-methanol, simulating an LC fraction. When valve 1 is switched the sample is transferred to the open-tubular trap by pump 1 (containing methanol-water) and the trap is flushed with an additional 150 µl of solvent. Then valve 1 is switched back again. The nitrogen flow now slowly pushes the remaining aqueous solvent out of the trap (2 min for the 2-m trap and 4 min for the 5-m trap). Meanwhile pump 2 (containing hexane) is started and the transfer capillary is inserted into the PTV injector. When valve 2 is switched the analytes are desorbed from the trap and directly introduced into the PTV injector as described above for large-volume injections. Two minutes before the next sampling step valve 2 is switched back again to remove hexane remaining in the trap after desorption by the nitrogen flow. Meanwhile the sample loop is filled again, after which the next phase switching can take place.

Desorption experiments. For desorption experiments the trap was sampled using an aqueous phase containing 20% (v/v) of methanol at a low sampling flow-rate (28 µl/min). For this purpose 50-ng/ml PAH solutions were used. After sorption the aqueous solvent was removed as described above. De-

Fig. 1. Phase-switching system. V1 and V2 = valves; P1 and P2 = syringe pumps; L = 300-µl loop.
sorption was then carried out applying a variety of conditions to be discussed below. Desorption profiles were obtained by repeated sampling and subsequent desorption with a stepwise increased volume of hexane. Compounds remaining in the trap were flushed to waste before the next experiment was started.

**THEORETICAL**

The maximum volume of sample that can be passed through an adsorption column is determined by the breakthrough volume of the solute \( V_b \) defined as \( V_r - 3\sigma_v \), where \( V_r \) is the retention volume and \( \sigma_v \) the standard deviation of the Gaussian peak eluting from the trapping column. The breakthrough volume of a component in a trapping column is given by the equation [29]

\[
V_b = V_0 \left( 1 + k \left( 1 - 3\sigma_v \right) \right)
\]

where \( V_0 \) is the void volume of the trapping column, \( k \) the capacity factor of the solute in the trap and \( N \) the plate number of the trapping column (with \( N > 9 \)).

The process of trapping analytes in an open-tubular capillary is very similar to that in open-tubular LC with extremely wide-bore columns. Band broadening in an open-tubular trap can therefore be described by the Golay equation:

\[
H = \frac{2D_m}{u} + \frac{1 + 6k + 11k^2}{96 (k + 1)^2} \cdot \frac{d_s^2}{D_m} \cdot u + \frac{2}{3} \cdot \frac{k}{(k + 1)^2} \cdot \frac{d_t^2}{D_s} \cdot u
\]

where \( D_m \) is the diffusion coefficient of the solute in the mobile phase, \( u \) the linear velocity, \( d_s \) the column diameter, \( d_t \) the stationary phase thickness and \( D_s \) the diffusion coefficient of the solute in the stationary phase. The diffusion coefficients of the PAHs selected as test solutes were calculated using the modified Wilke-Chang equation [30]; the values are about \( 0.40 \cdot 10^{-5} \text{cm}^2/\text{s} [25\% \text{ (v/v)} \text{ methanol in water at } 20^\circ\text{C}] \). The diffusion coefficients of the PAHs in a polysiloxane stationary phase are cu. \( 10^{-7} \text{cm}^2/\text{s} [31] \), i.e., almost two orders of magnitude lower than those in the mobile phase. As the linear velocity is far above the optimum and \( d_t \) is 5 \( \mu \text{m} \) or less, the first and the last terms on the right-hand side of eqn. 2 can be neglected. When the capacity factor is very large, eqn. 2 reduces to

\[
H = \frac{11}{96} \cdot \frac{d_s^2}{D_m} \cdot u
\]

or, expressed in terms of the volumetric flow-rate, \( F \), through the column,

\[
H = \frac{44}{96} \cdot \frac{F}{\pi D_m}
\]

As can be seen from eqn. 4, band broadening at a constant flow-rate depends on \( D_m \) only and is independent of the column diameter. Substitution of eqn. 4 into eqn. 1 yields

\[
V_b = V_0 \left( 1 + k \left( 1 - \sqrt{\frac{396F}{96\pi D_mL}} \right) \right)
\]

After substitution of \( V_0 \) and \( k \), with \( k = K\beta \), where \( K \) is the distribution constant of the analyte and \( \beta \) the phase ratio given by the equation

\[
\beta = \frac{V_m}{V_s} = \frac{(d_c - 2d_t)^2}{4d_t(d_c - d_t)}
\]

eqn. 5 can be rewritten as

\[
V_b = \frac{1}{4} \pi d_s^2 L \left( 1 + \frac{K}{\beta} \right) \left( 1 - \sqrt{\frac{396F}{96\pi D_mL}} \right)
\]

Eqn. 7 shows that in order to obtain \( V_b \) values larger than zero, \( F \) should meet the following requirement:

\[
F < \frac{96\pi}{396} \cdot \frac{D_mL}{D_s}
\]

In the situation described so far, the breakthrough volume was defined as \( V_r - 3\sigma_v \). Under these conditions there is a one-sided loss of only 0.15% of a Gaussian-shaped band when the sampled volume equals the breakthrough volume. For practical applications this can be considered as quantitative trapping. When higher losses are tolerated, the breakthrough volume can be redefined, which will result in higher maximum allowable flow-rates. Table I shows that when losses of 1.0 or 2.5% are accepted, the maximum sampling flow-rates can be increased by a factor 1.7 or 2.3, respectively. For trapping analytes in a 2-m trap and with the earlier \( D_m \) value of 0.40 \( 10^{-5} \text{cm}^2/\text{s} \), the maximum allow-
Maximum allowable sampling flow-rates for different definitions of breakthrough volume

<table>
<thead>
<tr>
<th>Breakthrough volume</th>
<th>Loss (%)</th>
<th>$F_{\text{max}}$ ($\mu l/s$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V_b = V_i \sim 3 \sigma$</td>
<td>0.15</td>
<td>0.762 $D_m L$</td>
</tr>
<tr>
<td>$V_b = V_i \sim 2.326 \sigma$</td>
<td>1.0</td>
<td>1.267 $D_m L$</td>
</tr>
<tr>
<td>$V_b = V_i \sim 1.960 \sigma$</td>
<td>2.5</td>
<td>1.784 $D_m L$</td>
</tr>
<tr>
<td>$V_b = V_i \sim 1.645 \sigma$</td>
<td>5.0</td>
<td>2.533 $D_m L$</td>
</tr>
<tr>
<td>$V_b = V_i \sim 1.28 \sigma$</td>
<td>10.0</td>
<td>4.184 $D_m L$</td>
</tr>
</tbody>
</table>

a One-sided loss for a Gaussian-shaped band.
b Maximum allowable sampling flow-rate assuming $k = \infty$.

able sampling flow-rates corresponding to trapping efficiencies of 99.85, 99.0 and 97.5% are 37, 61 and 88 $\mu l/min$, respectively. Here, the retention power and capacity of the open-tubular trap are assumed to be sufficient. The capacity of the trap is expected to be sufficient for the relatively clean LC fractions. With direct sampling of real water samples however, overloading of the trap by other analytes or matrix components cannot be excluded.

From eqn. 7, one readily sees that larger breakthrough volumes are obtained when either $d_e, L, K$ or $D_m$ is increased or $\beta$ is decreased. Further, reducing the sampling flow-rate is an alternative means of increasing the breakthrough volume. Although it is evident that large trapping column dimensions ($d_e, L$) are favourable for sorption, desorption puts limitations on the dimensions of the trapping column as the volume of organic solvent needed for desorption should not be too large. Too large desorption volumes will lead to a time-consuming introduction of the desorption liquid into the GC system. The effect of the other parameters on the breakthrough volume will be discussed in the next section.

Results and Discussion

Sorption

Maximum allowable sampling flow-rates. The plate number of the open-tubular trap was determined to assess whether the trap meets the requirement of $N > 9$ (see above). Table II shows that the experimental values are 2040% higher than the theoretical values. Apparently the $D_m$ values are higher than calculated. In order to have $N > 9$, flow-rates should not exceed 50 $\mu l/min$ for a 2-m trap with the mobile phase used. Analyte losses due to sampling at much higher flow-rates cannot be calculated by the theory described in the previous section because the compounds then no longer elute as Gaussian-shaped bands (see Fig. 2).

Retention on the open-tubular trap. As can be seen from eqn. 1 and by comparing Fig. 2A and B, an increase in capacity factor causes a distinct increase in the breakthrough volume. The capacity factor needed for quantitative (>99%) trapping depends on the sampling flow-rate and the volume that has to be sampled, i.e., the volume of the LC fraction should not exceed the breakthrough volume of the trap. The LC fraction volume depends on the inside diameter of the LC column used. In this study volumes of 300 $\mu l$ were used, which are typical for 1-2 mm I.D. LC columns. Calculating the breakthrough volume, with $V_0 = 0.15 ml, F = 45 \mu l/min, L = 2 m$ and $D_m = 0.52 \cdot 10^{-5} cm^2/s$ [naphthalene in methanol-water (26:74, v/v) at 25°C], breakthrough volumes exceeding 300 $\mu l$ are already obtained for a $k$ value of 7. This means that naphthalene (Fig. 2A, at 45 $\mu l/min$) will be trapped quantitatively under the conditions mentioned. For the example in Fig. 2B (at 45 $\mu l/min$), a breakthrough volume of 800 $\mu l$ is achieved with $k = 16$.

### TABLE II

<table>
<thead>
<tr>
<th>$F(\mu l/min)$</th>
<th>$u (cm/s)$</th>
<th>Naphthalene ($k = 4$)</th>
<th>Phenanthrene ($k = 15$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>$N_{th}$</td>
<td>$N_{exp}$</td>
</tr>
<tr>
<td>10</td>
<td>0.21</td>
<td>61</td>
<td>-</td>
</tr>
<tr>
<td>25</td>
<td>0.52</td>
<td>24</td>
<td>33</td>
</tr>
<tr>
<td>50</td>
<td>1.04</td>
<td>12</td>
<td>15</td>
</tr>
</tbody>
</table>

a Calculated plate numbers in methanol-water (26:74, v/v) at 25°C. $D_m$ for naphthalene and phenanthrene = 0.52 $10^{-5}$ and $0.44 \cdot 10^{-5} cm^2/s$, respectively.
b Plate numbers determined from elution chromatograms (injection: 11 $\mu l$ of 1-2 ppm PAH dissolved in the mobile phase).
If the capacity factors are too low, the retention power can be increased by working at lower organic modifier percentages. To study the influence of methanol on the sorption process, the capacity factors of the PAHs on the open-tubular trap were determined for several methanol percentages. Fig. 3 shows plots of log $k$ vs. percentage of methanol for the PAHs on 2 m x 0.32 mm I.D. traps coated with a 1- and a 5-$\mu$m film. Although these plots are not really linear over a large range of methanol percentages [32], estimated capacity factors obtained by extrapolation can be safely used to calculate breakthrough volumes. The values obtained with the 1-$\mu$m film trap are low [for naphthalene $k = 5$ at methanol-water (20:80, v/v)]. Therefore, a second trap was examined with a thicker film of stationary phase (5 $\mu$m). The phase ratio of the 5-$\mu$m trap is 14.5, which is almost five times lower than that of the 1-$\mu$m trap. It was therefore expected that the capacity factors would increase by a factor of five. The experimental capacity factors, however, increased only 3.5-fold. With the 5-$\mu$m film trap phenanthrene and pyrene can be trapped from up to 35-40% methanol in water mobile phases ($k > 16$).

A higher retention power can be achieved by using traps with still lower phase ratios, i.e., with very thick films such as are used for trapping analytes from headspace samples [33,34]. With hexane as the desorption liquid, uptake of the desorption liquid into the stationary phase can cause a significant swelling, which results in an increased film thickness. As the hexane remaining in the trap after desorption is not removed until just before the next sorption, a swollen phase will be created in each cycle. A tremendous phase swelling was observed for the 5-$\mu$m film: the internal volume of the trap decreased from 154 to 104 $\mu$l after flushing with hexane. If the retained 50 $\mu$l of hexane form a uniform film in the trap, this means that the film thickness increases from 5 to about 30 $\mu$m, and the phase ratio now is a mere 1.2. The nature of the
stationary phase also changes, and capacity factors were found to increase strongly. In the swollen phase they were 15-20-fold higher than in the original 5-µm film trap. It should now be possible to trap phenanthrene and pyrene from 65% (v/v) and naphthalene from 55% (v/v) methanol in water (k > 16).

As hexane is slightly soluble in methanol-water mixtures, the mobile phase will strip the hexane from the stationary phase. Fig. 4 shows the decrease in retention power of the trap after the passage of various methanol-water mixtures. For methanol-water (60:40, v/v), there is essentially no decrease. For sample volumes not exceeding 1 ml, even 70-75% (v/v) methanol in water mixtures can be used without severe effects on k caused by stripping of the hexane, as the swollen film will be restored prior to the next run, viz., during desorption.

Effect of temperature on sorption. The breakthrough volume is affected by the temperature via the diffusion coefficient of the analytes in the mobile phase and via the capacity factor. Theoretically, $D_m$ values will increase with a factor of about 2.5 on increasing the temperature from 25 to 60°C, thereby allowing for higher sampling flow-rates (eqn. 8). The capacity factors of the PAHs were found to decrease by a factor of 2.2-3.4 (for an example, see the dashed line in Fig. 3). As in most instances retention power is expected to be more critical than sampling flow-rates, trapping of the analytes at room temperature is generally preferable over trapping at elevated temperatures.

Stability of trap. An important practical aspect of the use of open-tubular traps for phase-switching is the stability of the stationary phase on flushing with large volumes of water-methanol and hexane. To this end, capacity factors were again determined after 5 months of continuous usage (ca. 600 ml of methanol-water passed through the trap). The decreases in the capacity factors were found to vary between 20% (naphthalene) and 33% (pyrene) and are no doubt merely due to degradation and dissolution of the stationary phase. From this it can be concluded that the solvent resistance of the trap is sufficient to ensure trouble-free use for many hundreds of phase-switchings.

Elimination of the aqueous phase

The aqueous phase was removed from the trap with a nitrogen flow-rate of 0.6 ml/min. Removal was carried out slowly (during 2 min) to prevent possible breaking up of the aqueous plug. No additional drying with nitrogen was performed, i.e., introduction of hexane was started immediately after completion of the solvent removal. Hexane leaving the trap was transferred directly to the GC system. Elimination of the aqueous phase appears to be satisfactory as there was no indication (irregular solvent peak, peak shape of the solutes) that water entered the GC system. Chromatograms obtained after phase-switching were not different from those obtained on splitless injection of a hexane solution. A more detailed study using (more polar) analytes sensitive to adsorption or decomposition on active surfaces in the PTV liner or column is, however, necessary to evaluate whether water is really completely removed.

Desorption and introduction into the GC system

After sorption and elimination of water the analytes are desorbed with hexane, which is introduced directly into the GC system. As the desorption volume is relatively large, refocusing of the solutes is necessary. This is done by selective solvent elimination in a PTV injector. Both liquid desorption from the trap and solvent elimination in the PTV injector can affect the recovery of the analytes. The occurrence of losses during solvent elimination in the PTV injector was studied by performing large-volume
injections of PAHs dissolved in hexane. Next the combined desorption–solvent elimination process was examined. Low sorption flow-rates and low concentrations of modifier were used to ensure quantitative trapping of the PAHs during sorption.

Large-volume injections using a PTV injector. The PTV injector is equipped with a liner containing a deactivated porous glass bed. The PTV injector can be cooled for retaining volatile components more efficiently. The solvent introduction rate depends on the initial PTV temperature. Data on optimum introduction speeds have been published previously [2,3]. The initial PTV temperature and the volume injected can both affect the recovery of the solutes after solvent elimination. The effects of initial PTV temperature and injected volume on recovery of three PAHs are shown in Fig. 5. With the volatile naphthalene, even on cooling the PTV to -28°C (introduction rate applied at this temperature 25 μl/min) significant losses were observed when more than 50 μl of the hexane solution were introduced. For the other PAHs solvent elimination was not so critical. Phenanthrene is still quantitatively recovered at an initial PTV temperature of 30°C (introduction rate 250 μl/min), provided that the volumes are not too large. An initial temperature of -10°C (introduction rate 50 μl/min) would offer a good compromise with regard to introduction time and recovery for naphthalene.

Desorption of PAHs with hexane. Two aspects are of major interest where desorption is concerned: desorption speed and desorption volume needed for quantitative recovery of the analytes from the trap. These two aspects are interrelated. Desorption of the analytes with hexane is a chromatographic process. The volume needed for desorption depends on the chromatographic band broadening during desorption, which can be described similarly to broadening during sorption. Instead of a breakthrough volume which corresponds to the start of an eluting peak, a desorption volume \( V_d \) corresponding to the end of an eluting peak now has to be used. Here \( V_d = V_t + 3σ_v \) or (cf., eqn. 1),

\[
V_d = V_0 (k + 1) \left( 1 + \frac{3}{\sqrt{N}} \right)
\]  

Because the trap is empty when desorption starts, the volume of liquid leaving the trap will be \( V_d = V_d - V_0 \). Small desorption volumes are obtained if the void volume and the capacity factor are small and the plate number is large. The need for a small void volume in order to obtain small desorption volumes is in contradiction with the demand for a large void volume in order to create efficient sorption. This means that, especially with regard to the column diameter, a compromise has to be made in order to keep the desorption volume at an acceptable level.

The contribution of chromatographic band broadening to \( V_d \) cannot be determined reliably because accurate values for the diffusion coefficients of the analytes in the swollen stationary phase are not available. However, once the values for \( k \) and \( V_d \) for the desorption of an analyte from a trap (length \( L_1 \)) have been experimentally determined (see below), it is possible to calculate the volume needed for desorption of that component from a similar trap of different length \( L_2 \). The following equation for the unknown desorption volume \( V_{d_2} \) can be derived:

\[
V_{d_2} = V_{o_1} (1 + k) \left( \frac{L_2}{L_1} \right) \left( \frac{L_2}{\sqrt{L_1}} \right) + \sqrt{\frac{L_2}{L_1}} \cdot V_{d_1}
\]  

As regards the capacity factors, their values are determined by the distribution constant (K) and the phase ratio. During desorption the analytes partition between two very similar phases, a silicone phase strongly swollen by hexane (stationary phase) and hexane (mobile phase). It can therefore be expected that \( K \) will have a value of ca. 1. With the known phase ratio of 1.2, the expected capacity

![Fig. 5. Effect of initial PTV temperature and volume introduced on recovery for large-volume injections of PAHs in hexane.](image-url)
factor of the PAHs during desorption is then about 0.8. Experimental values of capacity factors can be estimated from the desorption curves in Fig. 6. The desorption volume corresponding to 50% recovery is the adjusted retention volume ($V_i'$) of the PAHs. Introduction of the known void volume yields experimental capacity factors of about 0.45. Apparently the PAHs have a slight preference for the pure hexane phase; $K = 0.54$.

The desorption volumes are dependent on the desorption speed (see above). This was experimentally verified by desorbing the analytes at different flow-rates. The desorption profiles for two traps obtained at different desorption flow-rates are shown in Fig. 6. The higher the desorption flow-rate, the stronger is the band broadening and the larger the volume needed for complete desorption. Using the 2-m trap and applying desorption flow-rates of 10, 50 and 250 µl/min, the desorption volumes ($V_d$) (> 99% recovery) were 70, 80 and 125 µl, respectively. Knowing the capacity factors and the desorption volumes for the PAHs in the 2-m trap, the desorption volumes for longer traps can be calculated. The results are given in Table III. Calculated and measured $V_d$ values for the 5-m trap are seen to match well.

As far as desorption is concerned, the maximum length of trapping column that can be used depends on the application. For volatile analytes short trapping columns are preferable, for two reasons. First, the desorption volume should be small in order to obtain high recoveries in the solvent elimination step (Fig. 5). Second, short trapping columns yield acceptable desorption time. Volatile analytes require a low PTV temperature, which in turn necessitates a low introduction speed. The volume of desorption liquid needed to transfer the analytes from the trap to the PTV is equal to $V_d = V_0 + V_d'$ (the actual volume introduced is $V_d'$).

For example, with a PTV temperature of -30°C corresponding to a desorption speed of 25 µl/min, desorption from a 10-m trap (swollen phase) would take about 32 min. For less volatile analytes, on the other hand, the use of a 10-m trap is fully acceptable. For a high-boiling analyte, such as pyrene, even a 25-m trapping column can be used with quantitative desorption and analyte recovery after solvent elimination within 8.5 min. Such long trapping columns are, of course, very favourable from the sorption point of view.

Quantitative aspects of on-line phase-switching GC

After completion of the sorption experiments in the LC mode and the desorption experiments, the effect of sampling flow-rate and modifier concentration on the overall recovery in on-line phase-switching GC was studied. Here the 2 m x 0.32 mm I.D. trapping column coated with a 5-µm film was used. The volume of aqueous sample with which phase switching was carried out was 300 µl, which is a typical value for LC peak volumes eluting from
1-2 mm I.D. columns. Phase switching of larger volumes should be possible by taking longer trapping capillaries. Sampling was followed by flushing the trap with 150 μl of methanol-water, which may be necessary in future applications to remove buffer salts.

The effect of the sampling flow-rate on the recovery is shown in Table IV. In this example losses can be caused only by too high sampling rates, because the conditions were such that breakthrough caused by insufficient analyte retention and losses during desorption and solvent elimination in the PTV injector were excluded. In accordance with the predictions of Lökvist et al.[35], the experimental recoveries were higher than those expected assuming Gaussian elution profiles. In other words, with a 2 m x 0.32 mm I.D. trap at room temperature, a flow-rate of 100 μl/min can be used without loss of analytes due to incomplete trapping. This flow-rate results in an acceptable sorption time of 4.5 min for the sample described.

The effect of the methanol percentage on recovery (applying a relatively high sorption flow-rate of 111 μl/min) is given in Table V. Under the applied PTV and desorption conditions naphthalene is partially lost during solvent elimination in the PTV. At a methanol concentration of 80% (v/v), losses due to insufficient retention power of the trap are observed for all analytes. The repeatability of the total phase-switching-solvent elimination process was found to be very good. The relative standard deviation (R.S.D.) was 2.5% (n = 3) or less both when the analyte recovery was quantitative and when losses occurred.

As an example, Fig. 7 shows separation of

---

**TABLE IV**

RECOVERY OF PYRENE AT DIFFERENT SORPTION FLOW-RATES

<table>
<thead>
<tr>
<th>F (μl/min)</th>
<th>Recovery (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Theoretical\textsuperscript{a}</td>
</tr>
<tr>
<td>14</td>
<td>100</td>
</tr>
<tr>
<td>28</td>
<td>100</td>
</tr>
<tr>
<td>42</td>
<td>100</td>
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<td>111</td>
<td>95</td>
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<td>333</td>
<td>83</td>
</tr>
<tr>
<td>1000</td>
<td>71</td>
</tr>
<tr>
<td>2000</td>
<td>65</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Based on Gaussian elution profile; \(D_m\) of pyrene = 0.37 \(10^{-7}\) cm\(^2\)/s.

\textsuperscript{b} Conditions: trap, 2 m x 0.32 mm I.D., 5-μm CP-Sil-5-CB; mobile phase, methanol-water (20:80, v/v); temperature, 20°C. Sample, 300 μl of 37 ppb pyrene. Desorption with 80 μl of hexane at 50 μl/min; PTV initial temperature, –10°C.

**TABLE V**

EFFECT OF METHANOL CONCENTRATION IN WATER ON RECOVERY IN ON-LINE PHASE-SWITCHING GC

<table>
<thead>
<tr>
<th>Methanol in water (%, v/v)</th>
<th>Recovery (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Naphthalene</td>
</tr>
<tr>
<td>20</td>
<td>69</td>
</tr>
<tr>
<td>50</td>
<td>66</td>
</tr>
<tr>
<td>65</td>
<td>61</td>
</tr>
<tr>
<td>80</td>
<td>34</td>
</tr>
</tbody>
</table>

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Fig. 7. On-line phase-switching GC of sixteen PAHs and a system blank (methanol-water). Sample: 300 μl of a solution of sixteen PAHs (100:50, v/v); concentrations: 200–400 ppb (1-3) and 20–40 ppb (4-16). Sampling flow-rate, 111 μl/min. Trap: 2 m x 0.32 mm I.D., 5-μm film (swollen with hexane). Desorption, 80 μl of hexane at 25 μl/min. PTV initial temperature, –30°C. GC temperature programme: 40°C (2 min) –10°C/min to 275°C. Recoveries: (1) naphthalene, 80%; (2) acenaphthylene, 98%; (3) acenaphthene, 97%; (4) thiophene, 99%; (5) phenanthrene, 100%; (6) anthracene, 97%; (7) fluoranthene, 102%; (8) pyrene, 99%; (9,10) benz[a]anthracene + chrysene (97%); (11,12) benz[b]fluoranthene + benz[k]fluoranthene, 93%; (13) benz[a]pyrene, 92%; (14) indeno[1,2,3-cd]-pyrene, 88%; (15) dibenz[a,h]anthracene, 85%; (16) benz[e]perylene, 86%.
sixteen PAHs after phase switching from methanol-water (50:50, v/v) to hexane. Data on recoveries (SO-102%) are given in the caption.

CONCLUSIONS

Theory allows the influence of various operating conditions, such as flow-rate, column dimensions and diffusion coefficients, on the phase-switching process to be described. In order to trap analytes quantitatively the sampling flow-rates should not exceed a certain maximum value which depends only on the diffusion coefficient of the analyte in the aqueous phase and the length of the trapping column. For a 2 m x 0.32 mm I.D. trap, the maximum sampling flow-rate was found to be ca. 100 μl/min. For trapping analytes from conventional LC eluents, i.e., eluents containing high percentages of modifier, the phase ratio of the trap should be very low, i.e., thick films have to be used. Stationary phase swelling due to the uptake of large amounts of desorption fluid (hexane) into the stationary phase leads to a substantial increase in the amount of desorption fluid (hexane) into the stationary phase. The retention power in a thick-film trap with a film thickness and hence the retentive strength of the trap. The retention power in a thick-film trap with a film thickness and hence the retentive strength of the trap.

It can be concluded that open-tubular traps are highly promising for phase-switching in coupled RPLC-GC. The system described here also holds promise for automated pretreatment of aqueous samples.

ACKNOWLEDGEMENT

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REFERENCES
