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## Graphical tools for the planning and interpretation of polyurethane foam based passive air sampling campaigns†

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Due to low cost and easy handling during sampling and extraction, passive air samplers (PASs) using polyurethane foam (PUF) as a sorbent have become the most commonly deployed PASs for semivolatile organic compounds (SVOCs). However, depending on the scenario, PUF-PAS may not always be operating in the linear uptake phase, which implies the need to consider how temperature, wind speed, deployment length and chemical properties interact to determine the amount of a target chemical taken up and the fraction of a depuration compound (DC) being lost during deployment. Guidance is, therefore, necessary to quantitatively interpret curvi-linear uptake in the PUF-PAS and avoid selection of DCs unsuited to the deployment conditions. In this study, the PAS-SIM model is used to generate graphical tools that aid in addressing important questions frequently arising during the use of PUF-PASs. Specifically, we generated five charts that display (i) the inherent sampling rate as a function of wind speed and a chemical's molecular diffusivity, (ii) the length of the linear uptake period as a function of chemical properties, temperature and the acceptable deviation from linearity, (iii) the time to 95% equilibrium as influenced by chemical properties, temperature and wind speed, (iv) the dependence of the fractional loss of DCs on chemical properties, temperature, wind speed and deployment length, and (v) the influence of chemical properties, temperature and the total suspended particle concentration on the extent of sorption to atmospheric particles. The charts also facilitate the assessment of the influence of parameter uncertainty. It is hoped that these charts assist with planning and interpreting sampling campaigns based on a mechanistic and quantitative understanding of uptake in PUF-based PASs.

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#### **Environmental significance**

Even though polyurethane foam based passive air samplers (PUF-PAS) are widely used for sampling semi-volatile organic chemicals, interpretation of the results is not always based on a comprehensive understanding of how these samplers work. While a number of numerical simulation models of the uptake in the PUF-PAS exist and could provide valuable guidance, they have not been widely adopted. In order to lower the thresholds for their adoption, this study makes simulation results of one such model accessible without the need to actually run it. This is accomplished through the construction of graphs, that allow variables such as the length of the linear uptake period, the time to equilibrium, and the fractional loss of depuration compounds to be obtained from the length, temperature, and wind speed of deployment and the chemical partitioning properties.

#### 1. Introduction

Polyurethane foam (PUF) is made by the reaction of diisocyanates and polyols, and depending on the production method yields PUF-ether or PUF-ester. Due to the reasonably high and consistent sorptive capacity PUF-ether is a widely used sorbent in the sampling of semi-volatile organic compounds (SVOCs) from the atmosphere.1-6 Because of wide availability, easy handling and low cost of PUF compared to other sorbents,

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passive air samplers based on PUF (PUF-PASs) are commonly utilized for collecting SVOCs around the world.7-11

The PUF-PAS was originally designed to be a linear uptake sampler, which implies that evaporation of target chemicals from the PUF is assumed to be negligible. If that assumption is valid, a volumetric air concentration averaged over the time of deployment in units of ng m<sup>-3</sup> is obtained by simply dividing the amount of chemical sorbed to the PUF ( $m_S$  in ng, possibly corrected for the amount in field blanks) by the product of the deployment period  $\Delta t$  in days and an inherent sampling rate SR<sub>inh</sub> in m<sup>3</sup> day<sup>-1</sup>. This inherent sampling rate varies between chemicals, because their molecular diffusivity is somewhat dependent on molecular size. However, in many sampling scenarios, it is not valid to assume that the evaporation of chemical from the PUF is negligible. In that case, the sampler is said to leave the linear uptake period and enter the curvi-linear section of the generic uptake curve displayed in Fig. 1. When this transition occurs is chemical-specific, with more volatile chemicals having much shorter linear uptake periods than less volatile ones.

The real sampling rate of a chemical in the curvi-linear uptake period is lower than the inherent one. Because uptake curves in the curvi-linear uptake period can still appear to be linear (Fig. 1), these sampling rate are also referred to as "apparent" sampling rates SR<sub>app</sub>. An alternative nomenclature refers to an effective sampling volume  $V_{\rm eff}$ , which is equivalent to the product of SR<sub>app</sub> and deployment time. The volumetric air concentration during the deployment  $C_G$  can then be calculated by the simple equation:

$$C_{\rm G} = m_{\rm S}/({\rm SR}_{\rm app}\Delta t) = m_{\rm S}/V_{\rm eff} \tag{1}$$

But not only is eqn (1) no longer valid if  $C_G$  is variable during the sampling period,  $^{12}$  SR<sub>app</sub> or  $V_{\rm eff}$  are also difficult to establish with high confidence. Li and Wania<sup>13</sup> have shown that the uncertainty in the equilibrium partition ratio between PUF and the gas phase  $(K_{PUF/G})$  alone causes uncertainty in estimated  $V_{eff}$ or  $SR_{app}$  of at least  $\sim 50\%$  and the uncertainty will be even higher when accounting for the influence of ambient conditions during the sampling, especially temperature and wind speed.14 While it is therefore often desirable to avoid sampling outside of the linear uptake range, use of the PUF-PAS in the curvi-linear regime is generally the rule rather than the exception. Sampling in the linear uptake range is more easily achieved by using a high uptake capacity sorbent such as styrenedivinylbenzene co-polymeric resin (commercial name XAD).

 $SR_{app}$  or  $V_{eff}$  are functions of the sampled chemical, namely its molecular diffusivity and  $K_{PUF/G}$ , the deployment conditions, most notably wind speed and temperature, and the deployment length. Theoretical methods for estimating  $SR_{app}$  or  $V_{eff}$  have been presented.15 Alternatively, they can also be estimated from the measured fraction of a depuration compound (DC) lost from the PUF during the deployment of a PAS.16,17 Isotopically labelled SVOCs and chemicals that are not present in the sampled environment are often used as DCs to distinguish them from the SVOCs in the field. Importantly, using DCs does not prevent the uncertainty in  $K_{PUF/G}$  from being carried forward to the uncertainty in  $SR_{app}$  or  $V_{eff}$ , because  $K_{PUF/G}$  of the DCs at the temperature of deployment need to be known.<sup>13</sup>

Alternatively, one could also envisage to use the PUF-PAS as an equilibrium sampler for volatile substances. If a chemical reaches equilibrium between PUF and atmospheric gas phase, the air concentration (in ng m<sup>-3</sup>) can be obtained by dividing the concentration of the chemical in the PUF (in ng m<sup>-3</sup>) by the  $K_{\text{PUF/G}}$  (m<sup>3</sup> air m<sup>-3</sup> PUF) (Fig. 1).

When planning to use a PUF-PAS or after having used one, a number of questions often arise, such as:

- For how long can I deploy a PUF-PAS under specific meteorological conditions for my target compound(s) to remain in the linear uptake phase?
- Having deployed a sampler for a period of time under a set of conditions, can I use the SR<sub>inh</sub> for my target compound(s) or do I need to estimate an  $SR_{app}$  or  $V_{eff}$ ?
- How long would a deployment under a set of conditions need to be if I wanted my target compound(s) to reach equilibrium with the PUF?
- What deployment length can provide the best balance between the ability to detect a compound and being able to interpret the sequestered amount quantitatively?
- What DC(s) should I spike onto my PUF to make sure the fraction lost during a deployment of a particular length and under a set of conditions is suitable for deriving a SR<sub>app</sub> reliably?
- Will the target chemical be sampled from the atmospheric gas phase or do I have to consider the possibility that some of it is being sampled with particles and therefore experiences different uptake kinetics?

As indicated above, several methods for estimating SR<sub>app</sub> or V<sub>eff</sub> have been presented and could serve to answer such questions. Zhang and Wania<sup>18</sup> developed a model to illustrate how

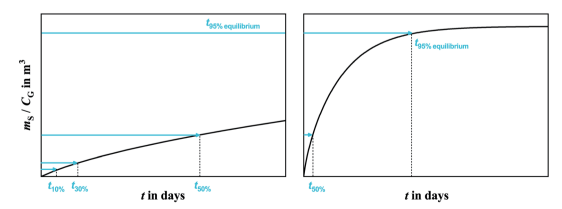


Fig. 1 Generic uptake curve of a chemical in a passive air sampler calculated with the PAS-SIM model, displaying the change in the effective sampling volume ( $V_{\text{eff}} = m_5/C_G$ ) during the linear uptake period (left panel) and the curvi-linear and equilibrium uptake periods (right panel). Shown are the lengths of the linear uptake periods, if a 10%, 30% or 50% deviation from linearity is tolerable  $(t_{10\%}, t_{30\%}, t_{50\%}, respectively)$ , and the time to 95% equilibrium (t95% equilibrium). The air concentration and meteorological conditions are assumed to be constant and no degradation of the chemical was assumed to occur.

the mass transfer processes in general and the sampling rate in particular is affected by various parameters, but it is not a model suitable for routine applications. Harner<sup>15</sup> published a Microsoft Excel template to calculate the  $V_{\rm eff}$  based on the physicochemical parameters of target compounds, the deployment length and ambient conditions such as temperature and particle concentration. However, the template uses fixed constants, e.g.,  $SR_{\rm inh}$  and particle concentration, and is only suitable for specific SVOCs. To solve this problem, Herkert  $et~al.^{19}$  developed a MATLAB-based model to predict the  $V_{\rm eff}$  accounting for the influence of external wind speed and temperature. However, the user interface is not straightforward for new users. In addition, the model ignores any resistance to mass transfer within the PUF, the potential for degradation of chemicals sorbed to the sampler and the influence of aerosol.

The PAS-SIM model by Armitage et al.20 is a user-friendly Microsoft Excel-based model developed to quantitively simulate the transport of chemicals from the atmospheric gas phase to the sorbent of a PUF- or XAD-based PAS. PAS-SIM considers the potential existence of a sampler-side resistance to uptake and provides daily outputs for the amount sorbed to the PUF or XAD and the SR<sub>app</sub>. In addition, the influence of different parameters is embedded in PAS-SIM, such as chemical properties, temperature, wind speed, deployment length, and atmospheric particle concentration. The performance of PAS-SIM model for XAD-PAS was evaluated using data for polychlorinated biphenyls, polycyclic aromatic hydrocarbons, and pesticides.20,21 Overall, these studies show that PAS-SIM model is capable of accounting for the influence of temperature and wind speed on passive sampling rates and modeled PAS uptake curves show a good agreement with the empirical data (generally within a factor of two).20,21

Even though these models have been developed, none of them are widely used. PUF-PAS derived data continue to be interpreted with generic sampling rates without consideration of the specific uptake conditions applicable to a particular set of compounds and deployments.22,23 Multiple DCs are often used without consideration of the specific deployment conditions, even though they are expensive and often do not yield the desired information.20 Possible reasons for this situation are a general reluctance to adopt numerical models and a lack of visualization of model outputs. The main objective of this study therefore is to use the PAS-SIM model, supplemented with theoretical calculations, to generate charts to provide graphical guidance on how to use PUF-PASs and to interpret data obtained from PUF-PASs. These charts allow anyone to benefit from the information provided by PAS-SIM without the need to actually run it. It is hoped that these charts facilitate the planning and interpretation of sampling campaigns based on a mechanistic and quantitative understanding of how the PUF-PAS functions.

## 2. Method

#### 2.1 Characteristics of the PUF-based passive air sampler

In this study, the height (h), radius (r) and porosity ( $\theta$ ) of the PUF in the PUF-PAS are 0.0135 m, 0.07 m and 0.97, respectively. The total surface area of the PUF  $A_{\rm S}$  of 0.037 m<sup>2</sup> is calculated using:

$$A_{\rm s} = 2 \times \pi \times r \times (r+h) \tag{2}$$

The total volume of the PUF  $V_s$  of 0.0002 m<sup>3</sup> is obtained using:

$$V_{\rm s} = \pi \times r^2 \times h \tag{3}$$

In this study, the sampler is divided into 50 layers (n = 50) to calculate the mass transfer coefficient in the sampler.

#### 2.2 Inherent sampling rate

The first chart displays the inherent sampling rate for the PUF-PAS ( $SR_{inh}$  in  $m^3$  day<sup>-1</sup>) as a function of a chemical's molecular diffusivity in the gas phase (B in  $m^2$   $h^{-1}$ ) and external wind speed (WS in m s<sup>-1</sup>). Following the approach taken in PAS-SIM,  $SR_{inh}$  was calculated using:

$$SR_{inh} = k_{SA}A_S + 1.32 \times WS \tag{4}$$

where  $A_{\rm s}$  is the surface area of the sampler available for mass transfer (m²) and  $k_{\rm SA}$  is the overall mass transfer coefficient between gas phase and PUF sorbent (m h⁻¹). The first term in eqn (4) describes the SR<sub>inh</sub> under wind-still conditions, whereas the second term is based on an empirical relationship between WS and SR in the range 0 to 8 m s⁻¹ presented in Thomas *et al.*²⁴ For WS greater than 8 m s⁻¹, the SR<sub>inh</sub> is assumed to be constant (*i.e.*,  $k_{\rm SA}A_{\rm S}$  + 10.6). This approach is consistent with empirical data published by Tuduri *et al.*²⁵ for experiments representative of average wind speeds typically encountered in the field (*i.e.*, ≤8 m s⁻¹) but may underestimate SR at higher WS where there is greater variability and uncertainty in the experimental SR.

According to two-film theory and assuming diffusion through the PUF material to make a negligible contribution to mass transfer,  $^{21}$   $k_{\rm SA}$  can be estimated as:

$$k_{\rm SA} = \left(\frac{1}{k_{\rm A}} + \frac{1}{k_{\rm s-p}}\right)^{-1} \tag{5}$$

where  $k_A$  and  $k_{s-p}$  (m day<sup>-1</sup>) are the mass transfer coefficients across the air boundary layer and through the air-filled pores of the PUF, respectively.  $k_A$  can be obtained by dividing the molecular diffusivity in air  $(B, m^2 h^{-1})$  by the air side boundary thickness  $(\delta_A, m)$ .

$$k_{\rm A} = \frac{B}{\delta_{\rm A}} \tag{6}$$

Here,  $\delta_A$  under wind-still conditions is assumed to be 0.0075 m.<sup>21</sup>  $k_{s-p}$ can be obtained using:

$$k_{\rm s-p} = \frac{B \times \theta^{3/2}}{\delta_{\rm s}} \tag{7}$$

where  $\theta$  is the porosity of the sampling medium, and  $\delta_s$  is the diffusion path length within the PUF, which in PAS-SIM is estimated as half of the layer thickness:

$$\delta_{\rm s} = \frac{h}{2n} \tag{8}$$

If the height of PUF is larger than the radius, PAS-SIM calculates the layer thickness as the ratio of radius and the number of layers. In this study, the height is less than the radius; thus, eqn (8) was used to calculate the diffusion path length on the sampler side.

B is a chemical-specific parameter that is influenced by ambient temperature. In PAS-SIM, B is estimated using the semi-empirical Fuller-Schettler-Giddings equation (eqn (9)):26

$$B = 10^{-3} \times T^{1.75} \times U \times \frac{\left(\frac{1}{\text{MW}_{A}} + \frac{1}{\text{MW}}\right)^{0.5}}{p \times \left(\text{MV}_{A}^{\frac{1}{3}} + \text{MV}^{\frac{1}{3}}\right)^{2}}$$
(9)

$$U = 60 \times 60/10000 \tag{10}$$

where T is air temperature (K), MW<sub>A</sub> is the molecular weight of air (28.96 g mol<sup>-1</sup>), MW is the molecular weight of the chemical  $(g \text{ mol}^{-1}), p \text{ is atmospheric pressure (atm), } MV_A \text{ is the molar}$ volume of air (20.1 cm<sup>3</sup> mol<sup>-1</sup>), MV is the molar volume of the chemical, and U is used to convert the units from cm<sup>2</sup> s<sup>-1</sup> to m<sup>2</sup> h<sup>-1</sup>. In this study, due to lack of information on the molecular density for most SVOCs, the density is assumed to be a constant of 1.6 g mL<sup>-1</sup>. The MV is the ratio of MW and the molecular

When generating the data displayed in the chart, the molecular weight of SVOCs was set to range between 100 to 1000 g mol<sup>-1</sup>, and T from -40 to 50 °C. Atmospheric pressure was set to 1 atm.

#### 2.3 Length of the linear uptake period

The second chart displays the length of the linear uptake period as a function of chemical properties, temperature, deployment length and the deviation from linearity a user is willing to accept. The data displayed in this chart were directly generated with PAS-SIM using molecular weights (MW) of 100, 300 or 1000 g mol<sup>-1</sup> and  $\log (K_{PUF/G}/m^3 m^{-3})$  values ranging from 3 to 10. No aerosol is assumed to be present, wind speed is constant at 5 m s<sup>-1</sup>, and no degradation of target compound is assumed to occur during sampling.

Specifically, the percentage of the deviation from linearity was calculated using:

Deviation(%) = 
$$\frac{(SR_{inh} \times i \times C_G - m_{Si})}{SR_{inh} \times i \times C_G} \times 100\%$$
 (11)

where  $m_{Si}$  (ng) is mass of a compound in the PUF on the ith day as calculated by PAS-SIM,  $C_G$  (ng m<sup>-3</sup>) is the concentration of a chemical in the air. The length of the linear uptake period was then determined as the first days on which the deviation exceeds 10, 30 or 50%.

#### Time to equilibrium

The third chart displaying the time to 95% equilibrium as influenced by chemical properties, temperature and wind speed was again based on data generated with PAS-SIM. A common

molecular weight of 300 g mol<sup>-1</sup>, a range of log  $K_{PUF/G}$  between 4 to 7 and three different wind speeds (2, 5 and 8 m s<sup>-1</sup>) were used. The time to 95% equilibrium is determined as the first day when  $m_{Si}$ , as calculated by PAS-SIM for different log  $K_{PUF/G}$ , exceeds 0.95 times the mass at equilibrium.

### 2.5 Fractional loss of depuration compounds

The fourth chart, displaying the fractional loss of a DC in its dependence on chemical properties, temperature, wind speed and deployment length, is based on data generated with a version of PAS-SIM (PUF-Depurator) specially developed for simulating the loss of DCs. A common molecular weight of 300 g mol<sup>-1</sup>, a range of  $\log K_{PUF/G}$  between 4 to 10, three different wind speeds (0, 4 and 8 m s<sup>-1</sup>) and deployment lengths between 20 and 300 days were used. The percent loss of a depuration compound (R) is estimated as:

$$R = \left(1 - \frac{m_{\mathrm{D}i}}{m_{\mathrm{D}0}}\right) \times 100\% \tag{12}$$

where  $m_{\mathrm{D0}}$  (ng) is the amount of a DC spiked onto the PUF, and  $m_{\rm Di}$  (ng) is the amount of the DC on the PUF after the ith days of deployment as calculated by PAS-SIM.

#### **Particle sorption**

The fifth chart displays the influence of the chemical's equilibrium partitioning ratio between particles and the gas phase  $(K_{O/G})$  and the total suspended particle (TSP) concentration  $(C_{TSP} \text{ in g particles m}^{-3} \text{ air})$  on the extent of sorption of a compound to atmospheric particles, expressed as the percentage of a compound in the gas phase  $\varphi_G$ :

$$\varphi_{\rm G} = \frac{1}{1 + K_{\rm Q/G} \times \frac{C_{\rm TSP}}{\rho_{\rm Q}}} \times 100\%$$
(13)

where  $\rho_{\rm Q}$  is the density of TSP, assumed to be 1.7  $\times$  10<sup>6</sup> g particles m<sup>-3</sup> particles.<sup>27</sup> Relying on eqn (13), the chart displays combinations of  $C_{\text{TSP}}$  and  $K_{\text{Q/G}}$  that yield a  $\varphi_{\text{G}}$  of 1%, 10%, 50%, 90% and 99%.

#### 2.7 Estimating $K_{PUF/G}$ and $K_{O/G}$

In most of the charts in this study, a parameter is plotted against the logarithm of an equilibrium partitioning ratio  $(K_{PUF/G})$  and  $K_{O/G}$ . To apply these charts to a specific chemical therefore requires knowledge of the partitioning ratio, either through measurement or by estimation. Because  $K_{PUF/G}$  and  $K_{O/G}$  are functions of temperature, the partitioning ratio at the temperature of the sampler's deployment is required. To facilitate the placement of a number of commonly sampled SVOCs on the x-axes, we provide estimates of  $K_{PUF/G}$  and  $K_{O/G}$  at different temperatures at the bottom of each of these plots. The  $K_{PUF/G}$  was estimated using the following poly-parameter linear free energy relationship (ppLFER):6

$$\log K_{\rm PUF/G15^{\circ}C} = 1.63S + 3.39A + 0.71L + 0.27V - 0.02 + \log \rho_{\rm PUF} \tag{14} \label{eq:Kpuf}$$

where A and B are the H-bond acidity and basicity, S is the polarizability/bipolarity, V is the McGowan molar volume and L is the logarithm of the partition ratio between hexadecane and air and  $\rho_{PUF}$  is the density of the PUF (0.02 g mL<sup>-1</sup>). The original ppLFER provides a  $K_{PUF/G}$  with units of mL g<sup>-1</sup>. The last term in eqn (14) serves to convert the K into units of  $m^3$   $m^{-3}$ . Partition ratios at temperatures T other than 15 °C were obtained using:29,30

$$\ln K_{\text{PUF/G}} = \ln K_{\text{PUF/G15}^{\circ}\text{C}} - \frac{\Delta U_{\text{PUF/G}}}{R} \left( \frac{1}{T} - \frac{1}{288.15} \right) \quad (15)$$

where  $K_{PUF/G15^{\circ}C}$  is the value obtained with eqn (14) and R is the ideal gas constant (8.314 J K<sup>-1</sup> mol<sup>-1</sup>). The internal energy of phase transfer between PUF and the gas phase  $\Delta U_{\mathrm{PUF/G}}$  was obtained from another ppLFER:6

$$\Delta U_{\text{PUF/G}} \text{ (J mol}^{-1}) = (-17.6S - 46.6A - 12.8V - 4.3L + 2.7) \times 1000 + RT$$
 (16)

The  $K_{O/G}$ , required for the fifth charts, was similarly obtained with ppLFERs.28 Details are provided in the ESI.†

The solute descriptors for commonly targetted SVOCs and 16 commonly used DCs17,19,31-34 were taken from the UFZ LSER database,35 and are listed in Tables S1 and S2,† respectively. The calculation results and other data used in the making of the charts are compiled in Tables S3-S5.†

## Results

Fig. 2 illustrates the inherent sampling rate's dependence on wind speed and a chemical's molecular diffusivity. The bottom panel in Fig. 2 shows the relationship between molecular weight MW and molecular diffusivity B. B is size-dependent, with small chemicals having a higher B compared to large chemicals. B is also a much stronger function of MW for smaller than for larger compounds, i.e. the rate at which B changes decreases as MW increases. The five lines in the bottom panel demonstrate that B increases with temperature, but this temperature dependence is not very dependent on MW as the five lines are generally equidistant. The top panel in Fig. 2 shows that SR<sub>inh</sub> increases linearly with both *B* and wind speed WS.

We can use Fig. 2 to estimate the SR<sub>inh</sub> range we might expect for typical SVOCs under commonly encountered meteorological conditions (e.g., 5 °C and 4 m s<sup>-1</sup>). The light blue shaded area in Fig. 2 yields a SR<sub>inh</sub> around 7 and 7.8 m<sup>3</sup> day<sup>-1</sup> for commonly sampled SVOCs ranging in MW from 285 to 360 g mol<sup>-1</sup> (e.g. HCB to PCB-153). We can expect this SR<sub>inh</sub> to vary only a little between chemicals and with temperature: the range of MW from 100 to 1000 g mol<sup>-1</sup>, which comprises all SVOCs conceivably being sampled with the PUF-PAS, translates into a change in SR<sub>inh</sub> of only  $\sim$ 30% at -40 °C (e.g. from 6.1 to 8 m<sup>3</sup>  $day^{-1}$  when WS is 4 m s<sup>-1</sup>) or  $\sim$ 50% at 50 °C. Similarly, the temperature range of -40 to 50 °C, comprising all conceivable deployments, only results in a change in SR<sub>inh</sub> of 23% and 11% for chemicals with a MW of 100 and 1000 g mol<sup>-1</sup>, respectively. Clearly, wind speed is the dominant factor controlling SR<sub>inh</sub>.

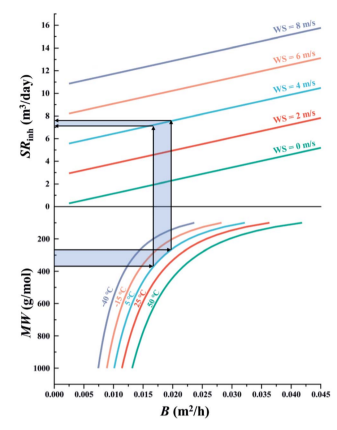


Fig. 2 Chart displaying the inherent sampling rate (SR<sub>inh</sub>) as a function of wind speed (WS) and a chemical's molecular diffusivity (B), which in turn is a function of deployment temperature and the chemical's molecular weight (MW).

Fig. 3 displays the length of the linear uptake period as a function of a chemical's  $K_{PUF/G}$ . The relationship between this length and  $\log K_{PUF/G}$  is not linear: as  $\log K_{PUF/G}$  increases, the linear uptake period increases first very slowly, but eventually very rapidly. When this transition to a rapid increase occurs depends strongly on the acceptable deviation from linearity (comparing differently coloured lines in Fig. 3). For example, during a PUF-PAS deployment of one month, chemicals with a log  $K_{PUF/G}$  above 6.5, 7.2 or 8.2 can be assumed to experience linear uptake, if deviations of 10, 30, or 50% are tolerated. During a seasonal deployment (90 days), these thresholds shift to slightly higher values above 6.8, 7.5 and 8.5. Because many of the SVOCs sampled with the PUF-PAS have a  $\log K_{PUF/G}$  in the range between 6.5 and 8.5 at temperatures typical of ambient deployment, the length of the linear uptake period is very strongly dependent on the applied linearity criterion. In fact, applying the strictest criterion (10%) very few SVOCs would remain in the linear uptake phase during typical deployment lengths. Different line styles in Fig. 3 indicate that the influence of the chemical's molecular weight on the length of the linear uptake period is minor.

The  $K_{PUF/G}$  of selected chemicals is displayed in the bottom of Fig. 3-5 over the temperature range from -40 to 50 °C, with bars shaded in a color gradient from blue to red indicating different temperatures. The SVOCs cover a very wide range in

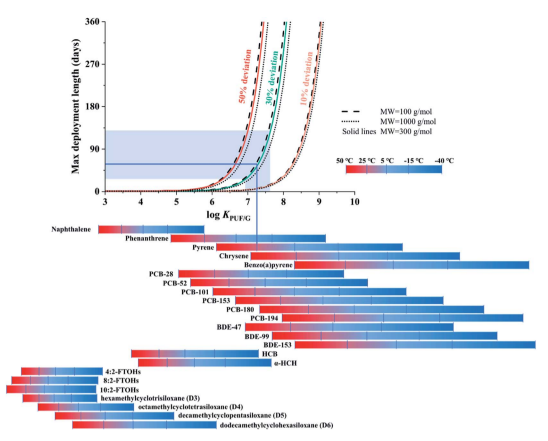


Fig. 3 Graph displaying the maximum deployment length of a PUF-PAS as a function of a chemical's  $K_{PUF/G}$  and molecular weight and the acceptable deviation from linearity in uptake (wind speed fixed at 5 m s $^{-1}$ ). To use this plot choose a target chemical and deployment temperature to get log  $K_{PUF/G}$  on the x-axis. Deduce  $t_{max}$  on the y-axis based on % deviation, e.g.  $t_{max}$  of 2 months for pyrene at 25 °C, if 30% deviation from linearity is tolerable.

 $\log K_{PUF/G}$  over this very large, but environmentally relevant, temperature range (see also Table S3 $\dagger$  in the ESI). The  $K_{PUF/G}$  for the same chemical can vary over many orders of magnitude within this temperature range. Fig. 3 conveys that the PUF-PAS cannot be used as a linear uptake sampler for more volatile SVOCs with a log  $K_{PUF/G}$  below  $\sim$ 6, such as the fluorotelomer alcohols (FTOHs), the cyclic volatile methylsiloxanes (cVMS), or smaller polycyclic aromatic hydrocarbons (PAHs) such as naphthalene. Even with a high tolerance for non-linearity and at low temperatures, the linear uptake periods for such compounds are shorter than a few days. At warm temperatures, this is also the case for the less chlorinated PCBs and three-ring PAHs. On the other hand, the PUF-PAS can generally be assumed to function as a linear uptake sampler for compounds with a log  $K_{PUF/G}$  above 8.5, as long as deployment periods do not exceed a few months. However, such SVOCs will often partition to atmospheric particles (see Fig. 6 below) and the sampling rate for particle-bound substances is uncertain and likely dependent on the size of the particles.

The chart in Fig. 3 can be used to estimate the maximum deployment length of a PUF-PAS if the goal is to sample a specific chemical within the linear uptake phase under a specific set of deployment conditions. For example, if pyrene (MW is  $202.25 \text{ g mol}^{-1}$  (ref. 36)) is to be sampled in summer (25)

°C) we can read from the respective bar below the diagram that its  $\log K_{\rm PUF/G}$  is  $\sim$ 7.25. Moving from this value on the x-axis up vertically until the line intersects with the 30% deviation curve, we can move left to obtain a maximum deployment length of  $\sim$ 60 days on the y-axis. Much shorter and longer values would be obtained with a different tolerance for deviation from linearity. We can further explore graphically, how the uncertainty in  $K_{\text{PUF/G}}$  propagates to the uncertainty in the length of linear uptake period. The uncertainty in  $\log K_{PUF/G}$  of pyrene at 25 °C is approximately  $\pm 0.36$ , which is represented by the area shaded in light blue in Fig. 3. The maximum deployment lengths of 60 days thus has an uncertainty range from  $\sim$ 30 to  $\sim$ 130 days.

Fig. 4 displays the time to 95% equilibrium as a function of the  $\log K_{PUF/G}$ , with three differently coloured lines indicating different wind speed conditions. Again the increase in  $t_{95\%}$  $_{\text{equilibrium}}$  is not linear with increasing log  $K_{\text{PUF/G}}$ , but slow for more volatile SVOCs and faster for less volatile ones. While a higher WS shortens  $t_{95\%\text{equilibrium}}$ , the change in  $t_{95\%\text{equilibrium}}$ is not proportional to the change in WS, because the three curves are not equidistant. For example, as WS increases from 5 to 8 m s<sup>-1</sup>,  $t_{95\%\text{equilibrium}}$  increases less than if it increases from 2 to 5 m  $s^{-1}$ .

The transition from a  $t_{95\% equilibrium}$  of a few days to half a year occurs over a fairly small range in  $\log K_{\text{PUF/G}}$  fom 4.5 to 6.

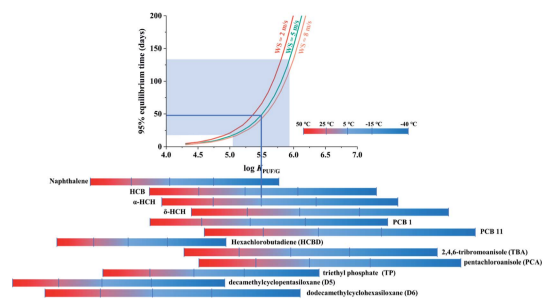


Fig. 4 Chart displaying the time to 95% equilibrium as influenced by chemical properties, temperature and wind speed. To use this plot, choose a target chemical and deployment temperature to get log  $K_{PUF/Gas}$  on x-axis. Then deduce  $t_{95\%}$  on y-axis based on windspeed (e.g.  $t_{95\%}$  of 50 days for  $\alpha$ -HCH at 5 °C and 5 m s<sup>-1</sup>).

The log  $K_{PUF/G}$  of the most volatile SVOCs, such as 4:2-FTOHs or D3, are not included in Fig. 4, as they will reach 95% equilibrium very quickly within a few days, even at the coldest temperature (-40 °C). The plot does not displays the less volatile SVOCs either, such as chrysene, benzo(a)pyrene, and some highly halogenated PCBs and PBDEs, which have  $\log K_{\text{PUF/G}}$ above 6.5, even at 50 °C, and thus always have  $t_{95\%\text{equilibrium}}$  in excess of 200 days. For the SVOCs that are displayed, it is readily apparent that the time to equilibrium is very strongly dependent on the deployment temperature. The chart suggests that the PUF-PAS could function as an equilibrium sampler for compounds such as naphthalene, hexachlorobutadiene (HCBD), phosphate (TP) triethyl decamethylcyclopentasiloxane (D5), with  $t_{95\%\text{equilibrium}}$  in the range of typically used PUF-PAS deployment periods.

This chart can be used to estimate the time a specific chemical requires to approach equilibrium with the PUF of a PAS during a particular set of deployment conditions: find the log  $K_{\rm PUF/G}$  value of the target substance at the temperature of the planned deployment on the x-axis, move up vertically to the intersection with the curve representing the WS of the planned deployment, and then read the  $t_{95\%\rm equilibrium}$  from the y-axis. For instance,  $\alpha$ -HCH sampled at 5 °C and an average WS of 5 m s<sup>-1</sup> will reach 95% equilibrium within  $\sim$ 50 days (blue lines in Fig. 4), whereas that time is only  $\sim$ 10 days if the temperature is 25 °C. The uncertainty of  $K_{\rm PUF/G}$  of  $\alpha$ -HCH at 5 °C is  $\pm$ 0.44, <sup>13</sup> leading to a range in  $t_{95\%\rm equilibrium}$  from  $\sim$ 20 to  $\sim$ 130 days (shaded area in Fig. 4). <sup>13</sup>

More reliable SR<sub>app</sub>s can be obtained if the percentage loss of a DC is approximately between 20% and 80%.<sup>37</sup> Because chemical properties, meterological conditions and deployment length jointly influence the extent of loss of DCs, choosing a DC that is lost to the appropriate extent is not trivial. Fig. 5 displays

the dependence of the fractional loss of a DC on  $\log K_{\rm PUF/G}$ , wind speed and deployment length. The three panels in the figure represent different assumed average wind speeds (0, 4, and 8 m s<sup>-1</sup>) and differently coloured lines in each panel are used to distinguish different deployment lengths. The grey shading represents loss between 20% to 80%. Bars at the bottom of the chart indicating the  $\log K_{\rm PUF/G}$  at different temperatures of a number of SVOCs commonly used as DCs allow for the selection of the appropriate value along the x-axis.

The loss of a DC decreases with an increase in  $\log K_{PUF/G}$ , the function resembling a sigmoidal curve, i.e. the loss is a particularly strong function of  $\log K_{PUF/G}$  in a transitional  $\log K_{PUF/G}$ range. Below that range, almost all of the DC is lost in a short period of time. Above that range, almost no loss occurs, even after longer deployments. Fig. 5 can aid in the identification of the DCs that have a  $\log K_{PUF/G}$  in the desirable steep transition area under a given set of conditions. This is done as follows: estimate the wind speed during deployment (choose one of the three panels) and decide on a desired deployment length (choose one of the lines in that panel). This indicates the  $\log K_{PUF/G}$  range that will result in a loss between 20 and 80%. Finally choose a DC that has a  $\log K_{PUF/G}$  in that range at the temperature of the planned deployment. The bottom part of Fig. 5 indicates that three groups of PCBs, namely congeners 101, 111 and 118, congeners 28, 31 and 32, and congeners 107 and 108 have very similar  $K_{PUF/G}$ . It would not be necessary to pick more than one DC from each of these groups, as their fractional loss would be expected to be similar.

Fig. 5 can also be used to estimate the expected fractional loss of a particular DC under a specific set of deployment conditions. For example, the blue lines indicate that half of the PCB-54 added to a PUF-PAS can be expected to be lost after a deployment of 50 days at 25 °C and a wind speed around 4 to

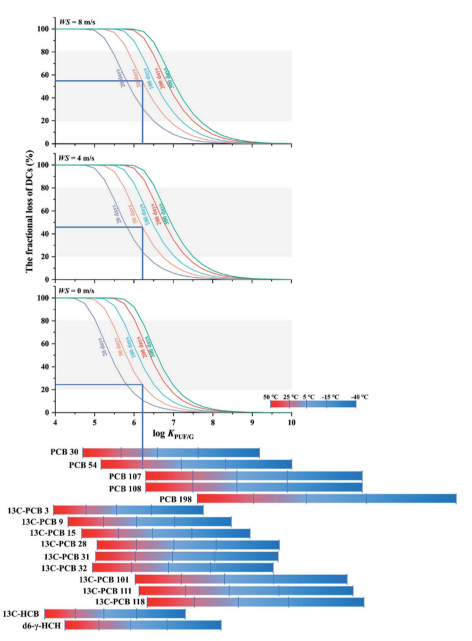


Fig. 5 Chart displaying the fractional loss of a DC in its dependence on chemical properties, temperature, wind speed and deployment length. To use this plot choose a DC and deployment temperature to get log  $K_{PUF/G}$  on the x-axis. Deduce  $\%_{loss}$  on the y-axis based on deployment length and wind speed (e.g. around half of PCB 54 is lost after 50 days at 25 °C with wind speed of 4 m s<sup>-1</sup>).

8 m s<sup>-1</sup>. Under windstill conditions, only about a quarter would be lost.

There is clear evidence that the PUF-PAS collects SVOCs in both the gas and particle phase.38-40 However, there is no consensus on whether the SR of particle-bound substances is similar to that of gaseous compounds. For example, Harner et al. 39 reported that both gas- and particle-phase polycyclic aromatic hydrocarbons (PAHs) were sampled with a similar SR, whereas Klánová et al. found that the SR of particle-bound PAHs was 10 times lower than that of gaseous compounds.38 The inconsistency of the findings is likely because the uptake of particle-bound substances may be influenced by the atmospheric particles (concentration, composition, size

distribution), the meteorological conditions (wind speed and its variability, angle of incidence), and the density of the PUF. 25,40,41 Even though the chamber design was the same in these two studies38,39 the design of the PAS housing can also be a factor.42 The PAS-SIM model describes the uptake of gas phase compounds only and does not attempt to simulate the uptake of particle-bound substances. Modelling the particle uptake processes within the PUF-PAS would be extremely complex and require deployment-specific input information, such as data on the local wind conditions, the particles in the sampled atmosphere and the particle size distribution of the target compounds, that is typically not available.

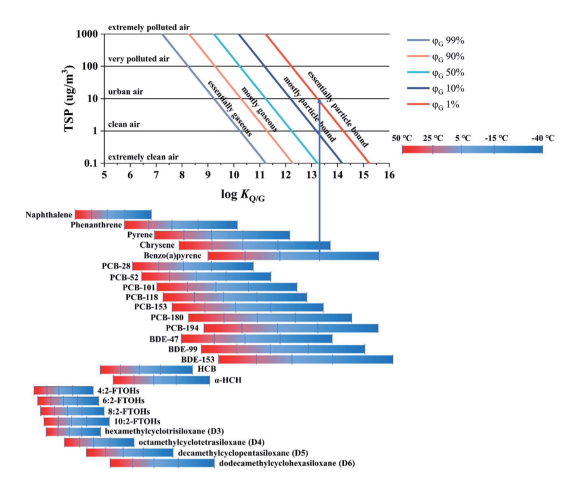


Fig. 6 Chart displaying the influence of chemical properties, temperature and the total suspended particle concentration on the extent of sorption to atmospheric particles. To use the plot, choose a target chemical and deployment temperature to get log  $K_{O/G}$  on the x-axis. Then deduce  $\phi_G$  based on TSP during deployment (e.g. benzo(a)pyrene is essentially particle bound,  $\phi_G < 1\%$ , in urban air in winter -15 °C).

The only purpose a model such as PAS-SIM can serve in this regard is therefore to estimate what fraction of the total atmospheric load of a target compound can be expected to be in the gas and particle phases under a specific set of deployment conditions. This information is provided in Fig. 6, which shows the extent of sorption to particles, expressed as the percentage of a compound in the gas phase,  $\varphi_G$ , as a function of  $\log K_{Q/G}$ and the total suspended particle (TSP) concentration. The charts in Fig. 2–5 can then be assumed to apply to the fraction in the gas phase. A user may also decide to refrain from a quantitative interpretation of data obtained for substances that are estimated to be mostly particle bound during a particular deployment.

In the top panel of Fig. 6, five air pollution levels are defined based on the TSP concentration. Increasing either the TSP concentration or the  $K_{Q/G}$  value will increase the extent of sorption to atmospheric particles. At the bottom of the plot, the  $K_{O/G}$  of a number of commonly sampled SVOCs is displayed, again within the temperature range -40 to 50 °C (numerical values of  $K_{Q/G}$  and  $\Delta U_{Q/G}$  are reported in Table S5† in the ESI). One way to use this chart is as follows: find the  $\log K_{O/G}$  value of the target compound at the temperature of the planned deployment. Move up vertically from there until you intersect

with the TSP concentration prevailing in the sampled atmosphere, then read the extent of sorption to particles from the coloured line (e.g., benzo(a)pyrene is essentially particle bound,  $\varphi_{\rm G}$  < 1%, in urban air in winter -15 °C).

### Discussion

For PUF-PAS users, there are two primary ways to use these charts to guide sampling and data interpretation:

Chemical centred: if a user has target compounds or a group of target compounds and a region of potential deployment (i.e., a specific range of temperature and wind speed), these charts can help to establish how the PUF-PAS can be applied to measure these compounds: as a linear or an equilibrium sampler? For what lengths of deployment? Is the use of a DC necessary, e.g. because it is inevitable that sampling occurs in the curvi-linear region?

Deployment centred: if a user has deployed, or plans to deploy, samplers for a certain length of time under a particular set of meteorological condition, these charts help to establish what chemicals can be expected to be interpretable using linear uptake assumptions, equilibrium assumptions, or require the use of DCs within the curvi-linear range. If there are some Paper

unexpected data, these charts may help to explain whether these data are reasonable or not and what causes this situation.

In the chemical centred approach, the process of using the charts starts with the colored bars representing different chemicals at the bottom of Fig. 3–6 and proceeds to the corresponding upper panel. In the deployment centred approach, one starts with the upper panel and goes down to the bars.

In an application resembling a model sensitivity analysis, the charts may facilitate the identification of the parameters that will have the largest influence on the uptake of a particular set of target compounds during a particular set of deployment conditions. Attention can then be paid to the study design by assuring values for these parameters are appropriately chosen and well characterised.

Here we used  $K_{\rm PUF/G}$  values predicted for a selected group of SVOCs with ppLFERs to generate the bars at the bottom of Fig. 3–5. If a user is interested in other compounds not currently displayed, it is of course possible to predict  $K_{\rm PUF/G}$  at the desired temperature using eqn (14)–(16). Solute descriptors required for the ppLFER predictions of eqn (14) and (16) can be obtained for thousands of compounds from the UFZ-LSER database, <sup>35</sup> or – if no such data are available – can be predicted from molecular structure using the quantitative structure property relationships implemented in the UFZ-LSER website. <sup>35</sup> ppLFER estimates are uncertain and sometimes diverge from available experimental  $K_{\rm PUF/G}$  data. <sup>13</sup> It is of course possible to use the charts with  $K_{\rm PUF/G}$  obtained by other means, *e.g.* experimentally determined values or values predicted with other techniques. Attention needs to be paid to using a  $K_{\rm PUF/G}$  in the appropriate units.

While we have tried to vary influential parameters within the entire plausible range and to depict the most commonly encountered scenarios when constructing the charts, they are inevitably based on only a limited set of data combinations. In particular, all of the charts assume invariant air concentrations, temperatures and wind speeds. Obviously, the PAS-SIM model itself is not constrained in the same way and is capable of simulating a much wider array of possible deployment scenarios, including those that allow for changes in many parameters over time. It is hoped that using the charts displayed here provides a gateway that helps PUF-PAS users to become familiar with the capabilities of the PAS-SIM models and to overcome the barriers to using the actual software.

Comparing Fig. 3 and 6 highlights that the range of  $\log K_{\rm PUF/G}$  in which the PUF-PAS can function as a true linear uptake sampler, where sorbed amounts can be interpreted with the help of inherent sampling rates, is very narrow and only includes a small number of commonly sampled SVOCs under a limited set of temperature conditions. In other words, when applied to a group of compounds or a range of temperature, the PUF-PAS is almost inevitably operating in the curvi-linear uptake regime, with all of the additional complexity and uncertainty this entails. The same applies to PASs using other sorbents with limited uptake capacity, such as polyethylene. High capacity sorbents such as XAD resin can widen the range of both SVOCs and temperatures for which true linear uptake can be achieved. This can reduce errors propagating from

uncertain sorption coefficients and the need for, and the expense of,  $DCs.^{12}$ 

The charts presented here provide only very limited guidance on the uptake of particle-bound substances in the PUF-PAS. This sampler takes up particle-bound SVOCs, because the housing design allows for wind to impinge on the sorbent (the foam), i.e., the double bowl design is a poor wind shelter. Large variations in the sampling rate for particle-bound SVOCs when using such a housing design are inevitable.42 These uncertainties are tolerated by PUF-PAS users, because this housing design allows for relatively high sampling rates and maybe also because they are interested in sampling particle-bound substances. Rather than interpreting data for compounds that are particle-bound during a PUF-PAS's deployment without consideration of the uncertainties this entails, we recommend refraining from such quantitative interpretation altogether. Alternatively, a user may consider relying on a sampler with a housing that acts as an effective wind shelter, i.e., largely prevents particle impaction on the sorbent. This facilitates quantitative interpretation and has the added benefit of a much weaker dependence of the sampling rate for gaseous compounds on wind speed.43 However, it comes at the cost of a lower sampling rate and the inability to record the presence of particle-bound substances.

## Conflicts of interest

There are no conflicts to declare.

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