Performance evaluation of a Semivolatile Aerosol Dichotomous Sampler (SADS) for Exposure Assessment: impact of design issues.

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Abstract. Aerosols of semivolatile organic compounds (SVOCs) pose significant health risks to workers in various occupational settings. Measuring human exposure to these aerosols requires a separate assessment of the contribution of particles and gases, which is not resolved by existing sampling techniques. Here, we investigate experimentally the performance of the Semivolatile Aerosol Dichotomous Sampler (SADS), proposed in previous studies, for sampling monodisperse liquid particles with aerodynamic diameters between 0.15 and 4.5 µm, corresponding to workplace aerosols. The measured sampling performances are compared to their theoretical counterparts computed by computational fluid dynamics. The effects of leakage rate, repeatability of the assembly, imprecision of the actually machined nozzle diameters and SADS parts misalignment are examined. The SADS assembly is found easily leaky, but consequences on sampling can be overcome when a prior leak test with leakage rate below 4 Pa.s⁻¹ is passed. Variation of nozzle diameters in the range (−4.5%, +3.7%) with respect to nominal values affects marginally (<3%) aerosol transmission efficiency, but sampling performance is little reproducible during successive SADS assemblies (CV=22.1% for wall losses). Theoretically unpredicted large (40–46%) wall losses are measured for particles larger than 2 µm, located mostly (80%) on the external walls of the collection nozzle. Assembly repeatability issues and simulations of SADS parts misalignment effect by CFD suggest that these undesirable particle deposits are due to the mechanical backlashes of the assembly. Thus, the current design does not guarantee a nozzle misalignment of less than 5% of the acceleration nozzle diameter, and other important geometric parameters are not further constrained. The promising theoretical sampling performance of the SADS for SVOCs aerosol larger than 1 µm thus falls short of expectations due to mechanical design issues that can be improved before possible field use.

1 Introduction

Semivolatile organic compounds (SVOCs) represent a significant subgroup of volatile organic compounds (VOCs), and their presence in the environment raises concerns due to their association with carcinogenic, mutagenic, and reprotoxic effects (Raffy et al., 2018). One of the defining features of SVOCs is their ability to exist simultaneously in both vapour and particle phases, making their sampling and analysis a complex task. There are varying definitions of SVOCs, with the U.S. Environmental Protection Agency (Technical Overview of Volatile Organic Compounds, 2020) proposing a classification based on boiling points (240-380 °C at atmospheric conditions).
The measurement of both vapour and particulate phases can often suffer from the issue of evaporation of the particulate phase during sampling. The collection nozzle, located 1.2 mm from the acceleration nozzle, has a diameter of 1.1 mm, and filters and adsorbent beds are placed at each outlet (major and minor flows). The SADS operates at a total sampling flow rate of 2 L.min⁻¹, split into 1.8 L.min⁻¹ at the major flow outlet and 0.3 L.min⁻¹ at the minor flow outlet (split ratio of 0.143).

The SADS was further optimized numerically and tested both in the laboratory and in the field by its designers (Kim and Raynor, 2010a, b; Kim et al., 2014). The optimized version is characterized essentially by a revised split ratio of 0.1 with a total sampling flow rate of 2 L.min⁻¹ and with a length of the separation space reduced to 0.48 mm instead of 1.2 mm. The angle of the acceleration nozzle was also changed from 19° to 45° between the 2009 and 2010 versions. Despite these modifications, the overall mechanical design of the SADS has not changed.

As of now, no applicable model exists to theoretically calculate the evaporation of a semivolatile aerosol during workplace air sampling, which hampers the use of these techniques. Other techniques that do not instantaneously separate particles and vapour also face the problem of evaporation during sampling (Raynor and Leith, 1999; Leith et al., 2010; Lilienberg et al., 2008; Wlaschitz and Höflinger, 2007; Sutter et al., 2010; Kim and Raynor, 2010a). An alternative approach is the Virtual Impactor (VI) principle, inspired by classical inertial impactors with collection plates (Loo and Cork, 1988; Marple and Chien, 1980). The VI is employed for size classification of particles based on their aerodynamic diameter. In 2009, the Semivolatile Aerosol Dichotomous Sampler (SADS), a novel variant of the VI dedicated to SVOCs, was proposed by Kim and Raynor (2009) and raised great hopes for this application.

In its original version, the SADS features an inverted flow configuration between the major and minor flows, resulting in 86 % of the total air being directed into the collection nozzle, while the remaining 14 % is suctioned perpendicular to the acceleration nozzle axis (Figure 1). The aerosol is sampled through a 4 mm inlet orifice and accelerated through a convergent shape called the “acceleration nozzle”, which narrows to a 0.8 mm orifice diameter. In the separation space, inertial particles are directed into the collection nozzle, while non-inertial particles and vapours follow both the major and minor flows. The collection nozzle, located 1.2 mm from the acceleration nozzle, has a diameter of 1.1 mm, and filters and adsorbent beds are placed at each outlet (major and minor flow). The SADS operates at a total sampling flow rate of 2.1 L.min⁻¹, split into 1.8 L.min⁻¹ at the major flow outlet and 0.3 L.min⁻¹ at the minor flow outlet (split ratio of 0.143).

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between these two versions (Figure 3). The overall shape of the device is that of a 37 mm cassette, and it is made
up of two parts that fit together via a cylindrical bearing surface. The parts are held together by 2 screws. Sealing
is ensured by an O-ring between the two parts, pressed together by the two screws.
Thus, the work of Kim et al. led to the creation of the SADS concept and revealed its interest in the sampling of
semi-volatile aerosols. However, many questions remain before SADS can be considered sufficiently mature for
widespread use as a portable sampling device for SVOC aerosols.
Firstly, the sampling performance of the device was not evaluated in detail for particles with aerodynamic
diameters greater than 1 µm, as the initial device was not designed for this. However, for field use, the evaluation
of the performance of the SADS for particles above 1 µm is especially important because workplace SVOCs
aerosols showed a presence of particles with diameters up to 10 µm (Cooper et al., 1996; Park et al., 2009). Since
the metric of SVOC exposure is mass concentration, and the mass carried by particles increases with the cube of
their diameter, sampling errors on the most inertial particles generate biases in exposure measurements that are far
more problematic than sampling errors on sub-micron particles. Optimization work by Kim et al. focused on
reducing the cut-off diameter of the device around an aerodynamic diameter of 0.7 µm, but the impact on super-
micron particle sampling was not assessed. Subsequent tests on real aerosols revealed significant deposits in the
device that had not been anticipated by the theoretical study, and the exact origin of these deposits is still unknown
(26.5 % of wall losses for an aerosol with MMAD of 2.17 µm in Kim and Raynor (2010b) and separate evaluations
by NIOH, Norway and Fraunhofer ITEM showed similarly high deposition ratios (Olsen et al., 2013).
Secondly, it’s important to emphasize the absence of published documentation or feedback regarding the
mechanical realization and the necessary operating procedures for obtaining measurements in line with theoretical
performance for the SADS. It is well known that the details of mechanical design and manufacture have as much
to do with impactor performance as the theoretical design: sealing, nozzle alignment (Loo and Cork, 1988),
geometric assembly tolerances - these are all necessary qualities which are the consequence of a suitable
mechanical design. So far, the SADS design proposed by Kim et al. has not been studied from these aspects, and
it is possible that a more definitive version of the SADS will require a review of its overall mechanical design,
without modification of the interior volume, which is perhaps optimal. Summarily, the authors who worked on the
SADS have neither published documentation related to these aspects nor investigated them in previously published
articles. In particular, it is doubtful whether the device as shown in Kim and Raynor (2010b) is leak-tight, with
only two diametrically opposed clamping points. Also, in both versions, the proposed design does not seem to
guarantee a precise control of nozzle spacing and alignment during assembly (limited guides and ground seats).
Finally, it is not certain that the optimized version proposed in 2010 is really optimal for sampling semi-volatile
aerosols encountered at workplaces, for various reasons. Firstly, from a methodological point of view, the
optimization carried out is based on a Computational Fluid Dynamics (CFD) model, which does not appear to be
sufficiently accomplished to provide enough accurate results for the optimization approach to have been
conclusive, as exposed in Belut et al. (2022). This is notably illustrated by significant differences between CFD
predictions and measurements, for both the airflow and the aerosol phase (modelled pressure drop on the major
flow side is more than twice the measured value for Kim and Raynor (2009), measured and simulated particles
separation efficiencies depart by as much as 30 % in Kim and Raynor (2009) and by as much as 53 % in Kim and
Raynor (2010b), simulations report almost no particle deposition but experimental evaluations found important
wall losses: 26.5 % for an aerosol with MMAD of 2.17 µm in Kim and Raynor (2010a) and separate evaluations
by NIOH, Norway and Fraunhofer ITEM showed similarly high deposition ratios (Olsen et al., 2013). Secondly, the 2010 version has a much steeper inlet convergent angle (acceleration nozzle) than the 2009 version, which increases the probability of undesirable wall loss for the most inertial aerosols (Belut et al., 2022).

In this context, the present article proposes to:

- Investigate experimentally the size-resolved sampling performances of the SADS on liquid SVOCs particles in the range of aerodynamic diameter 0.15-4.5 µm, i.e. beyond previous studies;
- Identify practical issues related to the design, manufacturing and operation of the SADS such as proposed in Kim and Raynor (2009, 2010b);
- Investigate the effect of small variations of SADS nozzle diameters linked to inevitable random manufacturing uncertainties;
- Detail the localization of wall losses in the device evoked in prior researches and identify their cause.

In doing so, our aim is to suggest improvement targets for future realization of the SADS, a small step to obtain a device suitable for the dichotomic measurement of particles and vapours composing SVOCs aerosols at workplaces. We shall base our study on the 2009 version of the SADS, because of the smaller cut-off diameter and also considering that the issues related to the overall design of the SADS are common to both versions. To reach our objectives, five SADS prototypes were constructed and their sampling behaviour was characterized using monodisperse liquid aerosols of various sizes. After evaluating the leakage resistance of the assembly, and its consequences on wall deposition, the actual sampling performances are compared to their theoretical counterparts computed by Belut et al. (2022). Origin of discrepancies are examined in terms of influence of the actually machined nozzle diameter and of the repeatability of the SADS assembly. An analysis of the distribution of deposits within the SADS is then used to estimate the likely cause of deposits in the device. The results are then discussed to propose improvement targets for the realization of the SADS, in terms of design and assembly.

Where necessary, CFD simulation results are used to support the observations. The approach of Belut et al. (2022) is then used for this purpose, including systematic calculation verification steps.

![Figure 1: Picture and schematic representation of a SADS prototype.](https://doi.org/10.5194/ar-2024-1)
2 Definitions and principle of particle-vapour dichotomous sampling

For a given aerodynamic diameter of particles $d_a$, the particles transmission efficiency $\eta_p(d_a)$ to the particle major flow outlet is defined as the ratio of particle mass collected at the major flow outlet to the total particulate sampled mass of particles with the same diameter (Eq. 1):

$$\eta_p(d_a) = \frac{m_{\text{major}}(d_a)}{m_{\text{inlet}}(d_a)}$$  \hspace{1cm} (1)

Similarly, $\eta_v(d_a)$ is the particles transmission ratio to the particle minor flow outlet, defined by the ratio of the particle mass collected at the minor flow outlet to the total particulate sampled mass (Eq. 2), for a given particle size.

$$\eta_v(d_a) = \frac{m_{\text{minor}}(d_a)}{m_{\text{inlet}}(d_a)}$$ \hspace{1cm} (2)

Finally, we defined a particles deposition ratio $\eta_d(d_a)$ that correspond to the ratio of the mass deposited on the inner wall of the SADS to the total particulate sampled mass, for a given particle size (Eq. 3):

$$\eta_d(d_a) = \frac{m_{\text{dep}}(d_a)}{m_{\text{inlet}}(d_a)}$$ \hspace{1cm} (3)

In ideal working conditions of the SADS as a gas-particle separator, we expect $\eta_d$ to be zero while $\eta_p=1$. If this is verified, the particle and vapour concentration of SVOC in the sampled air can be easily obtained by collecting the total mass (vapour+particles) of SVOC at each outlet (minor and major) and by knowing that the minor outlet sample corresponds to vapours only (it is particle-free). The major outlet sample corresponds to particles and vapours, but the vapour contribution, known from the minor flow, can be subtracted to yield the particle concentration. However, this principle is valid only if $\eta_d=0$ and $\eta_p=1$ for the sampled aerosol size range, whence the necessity of finding the conditions under which this is valid in actual realization of the SADS.

3 Material and methods

3.1 Aerosol generation

To assess the performance of the Semivolatile Aerosol Dichotomous Sampler (SADS), aerosols were generated using a specialized equipment called the Bench for Organic Aerosol (BOA), as depicted in Figure 2. The BOA is a vertical wind tunnel designed to operate with controlled airflow velocities ranging from 0.1 to 0.5 m.s\(^{-1}\) and humidity levels between 10 to 90 %RH. Aerosols were introduced countercurrent at the head of the tunnel to ensure thorough mixing with the airflow. To achieve spatially homogeneous mixtures at the sampling zone, the air/aerosol mixture passed through a series of diaphragms with different meshes. The SADS prototypes, along with an isokinetic nozzle connected to online instrument measurements, were placed on a crown support in the sampling zone.
To maintain consistent experimental conditions, room temperature was set to 21 °C, atmospheric pressure was measured at 1018 ± 10 hPa, relative humidity was regulated at 20 %RH, and airflow velocity was fixed at 0.2 m.s\(^{-1}\). The airflow inside the tunnel was considered isothermal, incompressible, and turbulent, with a Reynolds number based on the tunnel size of approximately 4×10\(^3\).

The BOA was carefully calibrated to meet the requirements of the standard NF EN 13205-2:2014. Spatial homogeneity of velocities in the sampling section was confirmed, with the standard deviation below 1 % over the entire sampling zone. Similarly, the spatial homogeneity of particle concentration demonstrated standard deviation values below 10 %.

### 3.2 Aerosol generator

To produce aerosols for testing the SADS prototypes, a specialized Condensation Monodisperse Aerosol Generator (CMAG - TSI 3475) based on Sinclair-La Mer was employed. Originally designed for use with diethyhexyl sebacate (DEHS) and NaCl, the CMAG was modified to accommodate the use of glycerol and fluorescein. This modification was necessary as DEHS is not water-soluble, making the analysis of particles collected on filters or internal walls of the sampler challenging and less sensitive. By replacing DEHS with glycerol and NaCl with fluorescein, water-soluble droplets were generated, and fluorescein could be quantified at a very low concentration (i.e., LoQ < 1 ng.L\(^{-1}\) within water extract).

The aerosol production process within the CMAG initiates with the nebulization of a water solution, specifically composed of 0.8 g.L\(^{-1}\) fluorescein and 5 g.L\(^{-1}\) sodium hydroxide in pure water, within an atomizer. This step is succeeded by the drying of the droplets in a diffusion dryer. Following the diffusion dryer, small nuclei, constituted of a blend of fluorescein and sodium hydroxide, were generated. These nuclei, serving as condensation nuclei,
exhibited sizes ranging from 10 to 100 nm. These nuclei were then exposed to a saturated vapour of glycerol downstream of the saturator. The resulting mixture of glycerol vapour and nuclei was directed to a re-heater and subsequently cooled down in a condensation chimney to produce the monodispersed aerosol. It is important to note that the size of the generated particles could be adjusted by modifying the temperature of the saturator or the number concentration of nuclei. For this study, aerosols with mass median aerodynamic diameters (MMAD) of circa 0.15, 2, 3, and 4.5 µm were produced and used for the experiments.

3.3 Aerosol characterization

Characteristics of the generated aerosols were measured continuously during the generation process. Aerodynamic particle sizes and geometric standard deviations (GSD) were measured using a TSI Aerodynamic Particle Sizer (APS 3321) associated with an aerosol diluter (TSI 3302 A) for particles ranging from 0.5 to 20 µm. For particles ranging from 0.056 to 0.560 µm, a TSI Fast Mobility Spectrometer (FMPS–3091) was used. The FMPS apparatus measures a mobility diameter that was converted in this study in an aerodynamic diameter using the following equation:

\[ d_{ae} = \frac{d_{ev}}{\left(\frac{C_u(d_{ae})\rho_p}{C_u(d_{ev})}\right)^{1/2}.} \]  

(4)

Where \(d_{m}\) the particle mobility diameter, \(d_{ae}\) the aerodynamic diameter, \(d_{ev}\) called the equivalent volume diameter: \(d_{ev} = \sqrt[3]{\frac{6V_p}{\pi}}\) with \(V_p\) the particles volume, \(C_u(d_{m})\) the Cunningham slip correction factor based on electrical mobility diameter, \(C_u(d_{ae})\) the Cunningham slip correction factor based on aerodynamic diameter, \(\rho_p\) particle density, \(\rho_0\approx 1\) reference density, and \(x\) the shape factor (taken equal to 1 for the considered spherical particles).

The particle density exhibits variability between nuclei and condensed glycerol particles. Based on the initial composition of the fluorescein solution utilized for generating nuclei, the density of the nuclei was determined to be 1.72 after total desiccation. In contrast, the density of the condensed particles is approximated to the density of pure glycerol, given the negligible mass of the nuclei compared to the mass of glycerol that condenses on them. Consequently, particles with diameters of 2 µm and above are considered to possess a density of approximately 1.26.

To further enhance the relevance of this study, the physical diameter of the particles is approximated by the measured aerodynamic diameter, considering the spherical nature of the particles. This approximation facilitates the conversion of the number-based particle size distribution into a mass-based particle size distribution, a parameter of greater significance for our research objectives. Following the conversion from a number-based to a mass-based particle size distribution, we proceeded to calculate the mass median aerodynamic diameter (MMAD). This parameter serves as a valuable metric, providing a comprehensive characterization of the aerosol particles in our investigation.

The measurement of aerosol characteristic by APS and FMPS apparatus allow modulating the particle diameter produced by the CMAG and verifying the stability of the aerosol concentration during the experiment. Averages are shown in Table 1.

Table 1: Averaged particle size distributions of the test aerosols (N=3 ± SD).
Aerosol reference diameter (µm)  

<table>
<thead>
<tr>
<th>Minimum Diameter</th>
<th>MMAD ± SD (µm)</th>
<th>GSD* ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.15</td>
<td>0.16 ± 0.05</td>
<td>1.56 ± 0.02</td>
</tr>
<tr>
<td>2</td>
<td>2.04 ± 0.15</td>
<td>1.16 ± 0.03</td>
</tr>
<tr>
<td>3</td>
<td>3.17 ± 0.21</td>
<td>1.14 ± 0.01</td>
</tr>
<tr>
<td>4.5</td>
<td>4.70 ± 0.12</td>
<td>1.10 ± 0.02</td>
</tr>
</tbody>
</table>

*GSD: geometrical standard deviation.

The stability of aerosol concentration in the sampling zone was verified, with a mass concentration deviation below 6% across experiments.

### 3.4 SADS prototypes

The STAMI, Norway, had five titanium SADS Kim & Raynor (2009) prototypes manufactured by a precision mechanics workshop. Each prototype consisted of two main pieces: the acceleration part and the collection part, as shown in Figure 3. For all SADS prototypes, the actually machined nozzle diameters were measured using scanning electron microscopy (SEM). One of the manufactured SADS presented nozzle diameters exactly equal to those of the 2009 version of the SADS proposed by Kim & Raynor (2009): it was denoted SADS R and considered as the reference case for comparisons. The other SADS prototypes (SADS 1, SADS 2, SADS 3, and SADS 4) exhibited slight deviations in their nozzle diameters with respect to the SADS R, due to the inherent tolerance of the manufacturing process. The measured diameters of the acceleration and collection nozzles for each SADS prototype are presented in Table 2. The maximum deviation of nozzle diameter with respect to the reference dimensions of SADS R (Kim and Raynor (2009) dimensions) was +0.03 mm (+3.7%) for the acceleration nozzle and -0.05 mm (-4.5%) for the collection nozzle.
Figure 3: Photographs of the SADS prototype, consisting of two main components - the acceleration nozzle (A') and the collection nozzle (B'). The upper left photo (A) shows the inlet side of the acceleration nozzle, while the lower left photo (B) displays the outlet side of the collection nozzle. On the outlet side of the collection nozzle (B), a 37 mm cassette is easily connected to the SADS sampler.

Table 2: Nozzles diameters measured by SEM.

<table>
<thead>
<tr>
<th>SADS name</th>
<th>Acceleration nozzle [mm] (absolute difference with reference / relative difference with reference)</th>
<th>Collection nozzle [mm] (absolute difference with reference / relative difference with reference)</th>
<th>Ratio Collection nozzle / Acceleration nozzle</th>
</tr>
</thead>
<tbody>
<tr>
<td>SADS 1</td>
<td>0.83 (+0.03 / 3.7 %)</td>
<td>1.05 (-0.05 / -4.3 %)</td>
<td>1.27</td>
</tr>
<tr>
<td>SADS 2</td>
<td>0.82 (+0.02 / 2.5 %)</td>
<td>1.06 (-0.04 / -3.6 %)</td>
<td>1.29</td>
</tr>
<tr>
<td>SADS 3</td>
<td>0.798 (-0.002 / 0.3 %)</td>
<td>1.11 (+0.01 / +0.9 %)</td>
<td>1.39</td>
</tr>
<tr>
<td>SADS 4</td>
<td>0.818 (+0.018 / 2.2 %)</td>
<td>1.11 (+0.01 / +0.9 %)</td>
<td>1.36</td>
</tr>
<tr>
<td>SADS R</td>
<td>0.8 (+0.00 / 0 %)</td>
<td>1.10 (+0.00 / 0 %)</td>
<td>1.38</td>
</tr>
</tbody>
</table>

3.5 Sampling procedure

The performance evaluation of the five SADS prototypes was conducted simultaneously in the sampling zone of the Bench for Organic Aerosol (BOA) (Figure 2). Prior to testing, each prototype was meticulously cleaned using ethanol and pure water. The samplers were equipped with Whatman Quartz Microfiber Filters (QMA) placed into...
37 mm and 25 mm Open Face Cassettes (OFC) and connected at the major outlet and the minor outlet (Figure 1), respectively. The flow rates at the major flow outlet (1.8 L.min⁻¹) and the minor flow outlet (0.3 L.min⁻¹), corresponding to a total inlet flow rate of 2.1 L.min⁻¹, were precisely controlled using flow meters (Gilian Gilibrator-2).

### 3.6 Fluorescence analysis

After each generation test, the sampling procedure for fluorescence analysis was carried out. The Whatman Quartz Microfiber Filters (QMA) contained in the 37 mm and 25 mm Open Face Cassettes (OFC) were extracted separately and analysed for fluorescence content. Each filter in the CFC and OFC was inserted into independent vials. A volume of 2 to 8 mL of the extraction solution, consisting of ultrapure water with a concentration of 5 g.L⁻¹ of NaOH, was added to the vial to dissolve the collected droplets of glycerol and their fluorescein/sodium hydroxide nuclei. The walls of the CFC were also washed with the extraction solution (pure water basified with 5 g.L⁻¹ of NaOH), and the resulting volume was combined with the one in the vial containing the CFC filter. After 20 minutes of mechanical shaking, the extracts were filtered through a PTFE syringe filter with a pore size of about 0.2 µm to prevent any disruption of the fluorescence measurement.

Wall deposition inside the SADS was determined by using 2 mL of the extraction solution to wash each wall of the SADS separately. Three different extracts were obtained: one from the acceleration nozzle wall (carried particle mass \( m_{\text{depa}} \)), one from the collection nozzle wall (carried particle mass \( m_{\text{depn}} \)), and one from the plenum wall (carried particle mass \( m_{\text{deppl}} \)). (Figure 1).

The extracts were then analysed for fluorescence using a portable ESElog Fluorescence Detector (Qiagen, Germany), with an excitation wavelength of 485 nm and an emission wavelength of 520 nm. The linear range of the ESElog Fluorescence Detector defined the lower (LLOQ) and upper (ULOQ) limits of quantification, which covered the concentrations encountered in this work (LLOQ = 0.33 ng.L⁻¹, ULOQ = 4×10⁴ ng.L⁻¹).

For each tested aerosol aerodynamic diameter \( d_a \), the total sampled mass \( m_{\text{inlet}} \) is evaluated as the sum of sampled masses:

\[
m_{\text{inlet}} = m_{\text{depa}} + m_{\text{depn}} + m_{\text{deppl}} + m_{\text{major}} + m_{\text{minor}}
\]

and the fractional deposition ratio \( \eta_d(d_a) \) is computed as:

\[
\eta_d(d_a) = \frac{m_{\text{depa}} + m_{\text{depn}} + m_{\text{deppl}}}{m_{\text{inlet}}}
\]

and local deposition ratios at the acceleration nozzle (\( \eta_{d_a} \)), collection nozzle (\( \eta_{d_c} \)) and in the plenum (\( \eta_{d_p} \)) are respectively computed from:

\[
\eta_{d_a} = m_{\text{depa}}/m_{\text{inlet}}, \quad \eta_{d_c} = m_{\text{depn}}/m_{\text{inlet}}, \quad \eta_{d_p} = m_{\text{deppl}}/m_{\text{inlet}}
\]

so that \( \eta_d = \eta_{d_a} + \eta_{d_c} + \eta_{d_p} \).

Because monodispersed aerosols are used, the masses evoked in this paragraph are all linearly related to the amount of fluorescent dye that they carry. Hence, deposition ratios are directly computed from the measured masses of fluorescent dye.

### 3.7 Mass balance verification

A mass balance verification step was adopted to verify that the protocol allowed recovering all particles sampled by the SADS. The SADS prototypes and 37-mm Closed Face Cassettes (CFCs) were arranged alternately on the...
crown support (Figure 4). The 37-mm CFCs served as reference samplers, enabling the determination of the total mass of particles collected, in comparison to the particle mass measured by the SADS prototypes.

All SADS and 37-mm CFCs presented a standardized 4 mm aerosol-sampling orifice and operated at the same sampling flow rate of 2.1 L.min\(^{-1}\). This standardization ensured that the total mass collected inside the 37-mm CFCs corresponded to \(m_{\text{inlet}}\), the total mass sampled by the SADS, which includes the contribution of both the Open Face Cassettes (OFC) placed at the minor and major flow outlets and the wall deposits. By comparing \(m_{\text{inlet}}\), the total particle mass measured by the SADS, with the mean reference mass measured by the 37-mm CFCs \(m_{\text{CFC}}\), we introduce the mass balance ratio of SADS \(i\) as:

\[
M_{b,SADS} = \frac{m_{\text{inlet}}}{m_{\text{CFC}}} 
\]

This mass balance makes it possible to assess the overall efficiency of the protocol used to measure the distribution of particles collected by SADS, between deposits and major and minor outlets.

Additionally, the use of the 37-mm CFCs allows assessing the spatial homogeneity of the aerosol distribution within the sampling zone. No significant spatial variation was observed across different positions of the CFCs (CV < 5 %). With this assurance of spatial homogeneity, the individual SADS mass balance \(M_{b,SADS}\) was calculated using equation 5 for each of the SADS prototypes.

### Figure 4: Schematic representation of the positions of the samplers on the crown support.

#### 3.8 Leak evaluation

Any form of leakage is known to compromise the sampling performances of aerosol samplers such as SADS, by disrupting the airflow and path lines within the nozzles and separation zone. Experimental tests were carried out to examine the leakage resistance of the proposed SADS assembly, to observe the effects of leaks and to determine an acceptable leakage limit for the SADS.

As the SADS operates under depression, a leakage test was performed using a digital pressure calibrator (DPC - FSM AG) set to a depression of -4000 Pa, equivalent to the operating pressure of the system. Following a stabilization period, the DPC’s internal pump was deactivated, and the pressure was continuously measured to determine the leakage rate (LR) in Pa.s\(^{-1}\) (Eq. 5). Three levels of leakage rates were defined: low, medium, and high, corresponding to LR values of LR \(\geq 13\) Pa.s\(^{-1}\), \(4\) Pa.s\(^{-1}\) \(\leq LR < 13\) Pa.s\(^{-1}\), and \(LR < 4\) Pa.s\(^{-1}\), respectively.

\[
LR = \frac{|P_0 - P_f|}{\Delta t} \quad (6)
\]

Where \(P_0\) and \(P_f\) represent the pressures at \(t = 0\) s and at the final time, respectively, and \(\Delta t\) is the duration of the leak test.
3.9 Supporting CFD Model

The CFD modelling approach employed in this study to support observations is documented in detail in Belut et al. (2022). Simulations are conducted using ANSYS FLUENT V.19.3 software. After due examination of the most influential modelling and physical factors affecting the significance of results, a 2D axisymmetric reduction of the inner volume of the SADS is used to perform simulations. A low-Reynolds realizable k-ε turbulence model is used to model the incompressible airflow, with a free-inlet boundary condition at the entrance of the SADS, following the guidelines of Belut et al. (2022). Aerosol particle fates are computed through a Lagrangian tracking of their centre of mass, taking into account turbulent dispersion and using a free-inlet boundary condition at the inlet. External forces acting on particles are reduced to drag force, including rarefaction effect. Impaction and interception phenomena are taken into account for wall losses, particles being assumed trapped when hitting a wall (consistent with the liquid nature of present aerosols). An extensive verification of computations with respect to grid size, numerical resolution tolerances and number of used aerosol trajectories was performed, exactly as exposed in Belut et al. (2022). For further insights into the model’s design and its applicability to the SADS, interested readers are encouraged to refer to the aforementioned study.

4 Results and discussion

4.1 Leaks effects

The airtightness tests conducted revealed significant leakages, primarily occurring at the O-ring seal between parts A and B (Figure 1) of the SADS prototypes. Additional leaks were also identified at the connection points between the Open Face Cassette (OFC) and the major and minor outlets of the SADS. Notably, SADS 1, 2, 3, and 4 displayed varying levels of airtightness during the tests, with the exception of SADS R, which consistently exhibited high airtightness across all tests. Fig. 6 illustrates the evolution of mass balance and mass distributions for the five SADS prototypes with varying levels of air tightness.
The aerosol generated in the three tests was monodisperse with a size distribution centred on a MMAD of 3.11 ± 0.21 µm, with a GSD of 1.14 ± 0.03 and a particle number concentration of 10124 ± 320 pcl.cm⁻³. Tests on the leaks effects on the performance of the SADS were also conducted with aerosols having MMAD of 2 and 4.5 µm. The results and conclusions were consistent with the distributions presented, indicating that the outcomes converge towards those presented in the following section.

Figure 5: Mass balance in the SADS prototypes in function of the airtightness level. Error bars represent the standard deviation calculated on five replicates for each condition.

Low airtightness led to a substantial decrease in the mass balance of SADS 1, 2, 3, and 4 (Figure 5), with mass balances reaching 66 ± 6.2 %, 74 ± 6.8 %, 70 ± 6.58 %, and 55 ± 7.59 %, respectively. Only when a high level of airtightness was achieved could a mass balance of 90 % or higher be attained for all prototypes. Moreover, low airtightness resulted in an undesirable increase in the mass fraction collected at the minor flow outlet. Presumably, leaks disturbed the airflow in the separation space, leading to the deviation of larger particles to the minor flow outlet than theoretically expected.

Globally, leaks not only influenced the total amount of particles collected within the SADS but also affected the particles transmission ratio to the major and minor outlets, which make results from leaky SADS unreliable. A systematic leak test is then mandatory before using the SADS for sampling purposes. To ensure reliable and accurate results, the SADS should only be considered suitable for sampling when the leakage rate (LR) is below 4 Pa.s⁻¹. Keeping the leakage rate within this acceptable limit will help maintain the integrity of the SADS and improve the reliability of the data collected during sampling operations. For further development, we also recommend revising the design of future realization of the SADS to guarantee its airtightness.

Due to difficulties in maintaining a constant level of airtightness for SADS 3 and 4 throughout all experimental tests, further experimentation with these two prototypes was not conducted.
4.2 Theoretical effect of actually manufactured nozzle diameters

Before comparing theoretical and actual performances of manufactured SADs, the effect of the lack of precision on the actually manufactured nozzle diameters is examined from a theoretical point of view, using the CFD model with the measured nozzle diameters of SADS 1, 2, 3, 4 and R. The numerical model computed the $\eta_p$ curves for each SADS prototype across a range of aerodynamic particle diameters from 0.1 to 20 µm, and the results are presented in Figure 6. Error bars correspond to one standard deviation of values arising from turbulent dispersion.

![Figure 6: Evolution of CFD mass transmission of the SADS prototypes, accounting for variation in nozzles diameters.](https://doi.org/10.5194/ar-2024-1)

Theoretical $\eta_p$ curves calculated by the numerical model are similar for all SADS prototypes (Figure 6). SADS 3 exhibits the maximum difference compared to the reference SADS R transmission efficiency curve, but this difference remains below 3% for all diameters. Overall, the variations observed in the nozzle diameters actually machined are not expected to lead to radically different sampling performance between the different prototypes.

These results are consistent with findings from a previous study on a VI by Marple & Chien, 1980a, who observed that increased ratios between nozzle diameters led to increased wall deposition, above the recommended value of 1.33 (and thus to decreased transmission ratios). From Table 2, we indeed see that SADS 3 exhibits the largest nozzle diameters ratio (1.39) of the prototypes. All ratios are, however, in the 1.33-1.49 range recommended by Marple & Chien, 1980a.

4.3 Actual vs. theoretical particle transmission efficiency

Figure 7 compares the theoretical and actually measured $\eta_p$ curves of the 3 airtight SADS.
For aerosols with a reference diameter of 0.15 µm, numerical predictions are in accordance with experimental tests, with transmission efficiencies to the major and minor flow outlets close to the ratio between the major and the minor flow ($\eta_p = 86 \pm 0.58\%$ for the model and $\eta_p = 81.7 \pm 6\%$ experimentally). This corresponds to the expected behaviour of non-inertial particles that are not separated by the SADS. We shall see, however, in the following section that a substantial fraction of these particles is actually deposited experimentally, in contradiction with theoretical results.

For the inertial particles tested with nominal diameters 2, 3 and 4.5 µm, $\eta_p$ is measured as always less than about 60\%, whereas 100\% is theoretically expected for the SADS in free-sampling situation. The origin of this difference is examined first by considering the particles deposition ratio in the next sections.

### 4.4 Fate of inertial particles and repeatability issues

To illustrate the origin of the unexpectedly low transmission efficiency of inertial particles in the device, the distribution of all measured $\eta_p, \eta_v$ and $\eta_d$ for 3 repetitions of the experiment and for the 3 airtight prototypes is shown on Figure 8 and Table 3. Only results for the 4.5 µm particles are shown here for brevity. For these particles, we observe that the low transmission efficiency $\eta_p$ is attributable to large (46.6 ± 5.4\%) wall losses ($\eta_d$), and not to the misdirection of particles to the minor outlet. These deposits are not theoretically explained, even if we take into account the lack of precision of machined nozzle diameters (Figure 7), and we can note that they apparently vary randomly across repetitions with a large coefficient of variation for $\eta_d$ (22.1\%). These variations are then likely to be attributable to the assembly process of the SADS, since other influencing parameters were monitored and controlled (flow rates, aerosol particle sizes, homogeneity of concentrations in the BAO, SADS leakage rate similarity of sampled masses).
Figure 8: Distribution of the fate of inertial particles with a reference diameter of 4.5 µm in multiple repetitions, for SADS 1, SADS 2, and SADS R at High Airtightness Level. Error bars represent the measuring uncertainty associated with the data points.

Table 3: Transmission ratio to the major and minor outlets and deposition ratio for SADS 1, 2 and R.

<table>
<thead>
<tr>
<th>SADS name</th>
<th>Tests</th>
<th>( \eta_p ) [%]</th>
<th>( \eta_v ) [%]</th>
<th>( \eta_d ) [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>SADS 1</td>
<td>Test 1</td>
<td>55.9</td>
<td>2.08</td>
<td>42.1</td>
</tr>
<tr>
<td></td>
<td>Test 2</td>
<td>50.3</td>
<td>2.04</td>
<td>47.6</td>
</tr>
<tr>
<td></td>
<td>Test 3</td>
<td>38.0</td>
<td>1.26</td>
<td>61.4</td>
</tr>
<tr>
<td></td>
<td>Average</td>
<td>48.1</td>
<td>1.79</td>
<td>50.4</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>9.2</td>
<td>0.46</td>
<td>10.0</td>
</tr>
<tr>
<td>SADS 2</td>
<td>Test 1</td>
<td>35.0</td>
<td>0.07</td>
<td>64.9</td>
</tr>
<tr>
<td></td>
<td>Test 2</td>
<td>59.9</td>
<td>1.26</td>
<td>38.9</td>
</tr>
<tr>
<td></td>
<td>Test 3</td>
<td>56.0</td>
<td>0.64</td>
<td>43.4</td>
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<tr>
<td></td>
<td>Average</td>
<td>50.3</td>
<td>0.66</td>
<td>49.0</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>13.4</td>
<td>0.60</td>
<td>13.9</td>
</tr>
<tr>
<td>SADS R</td>
<td>Test 1</td>
<td>63.6</td>
<td>1.73</td>
<td>34.6</td>
</tr>
<tr>
<td></td>
<td>Test 2</td>
<td>59.3</td>
<td>0.48</td>
<td>40.2</td>
</tr>
<tr>
<td></td>
<td>Test 3</td>
<td>53.0</td>
<td>0.75</td>
<td>46.3</td>
</tr>
<tr>
<td></td>
<td>Average</td>
<td>58.6</td>
<td>0.99</td>
<td>40.4</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>5.4</td>
<td>0.66</td>
<td>5.8</td>
</tr>
<tr>
<td>Total</td>
<td>Average</td>
<td>52.3</td>
<td>1.15</td>
<td>46.6</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>5.6</td>
<td>0.58</td>
<td>5.4</td>
</tr>
</tbody>
</table>

4.5 Detailed particles fate measured for SADS-R

Figure 9(a) and Figure 9(b) present images depicting a typical deposition that occurs inside the SADS after the sampling process. Notably, a significant amount of liquid particles can be seen on the external walls of the nozzles. Deposits can also be found on the internal walls of the nozzles, but they are difficult to capture in photographs.
Additionally, in certain tests, projections of macroscopic droplets from the nozzles to the walls of the plenum were also observed.

![Picture of deposition](https://example.com/deposition.png)

**Figure 9:** Pictures of particles deposition outside the nozzle.

To better understand the localization and underlying reasons for particle deposition in the SADS, independently of variations between prototypes, the detailed transmission ratios and position-resolved deposition ratios for the reference SADS-R are given on Figure 10, for all tested particle sizes. Values are averaged over 3 repetitions.

For particles with reference diameter 0.15 µm, wall deposition is primarily located on the collection nozzle, with 6 ± 5.8 % of sampled mass, though it should be almost zero theoretically (0.44 ± 0.58 %).

For particles with reference diameters 2, 3 and 4.5 µm, the experimental results show substantial particle depositions (43 % of sampled mass in average), unpredicted by theory either (0.2 ± 0.1 %). These experimental wall deposits seem almost independent of particle diameter for these inertial particles. The distribution of deposits across zones reveals that the majority of particles deposit on the collection nozzle (34.5 ± 3.4 %), followed by the acceleration nozzle (7.6 ± 3.4 %), with minimal deposition in the plenum (0.6 ± 3.4 %).

![Mass distribution graph](https://example.com/mass_distribution.png)

**Figure 10:** Mass distribution in SADS R exposed to four different particle size distributions: experimental and numerical study for precise wall deposition localization and transmission efficiency analysis. (Three repetitions, error bars represent measuring uncertainty). It is essential to note that most of the deposits observed can contaminate the vapour phase measurements at the minor outlet if particles evaporate after deposition. Indeed, wall deposition on the...
acceleration nozzle is located before separation, and most of the deposit of the collection nozzle is actually located on its outside walls (fraction of the deposit which is visible to the unaided eye).

Having ruled out the effects of leakage and machining inaccuracies in nozzle diameter, we can envisage several reasons for these deposits, which are not predicted by the numerical model. Firstly, the simulated geometry may not correspond to the real geometry for aspects other than nozzle diameter. In particular, the variations in deposits between the tests (Figure 8 and Figure 10) suggest variability in the assembly of the 2 parts of the SADS in relation to each other, and therefore a geometry of the interior domain of the SADS that is not only variable but also different from what is simulated. These variations may correspond in particular to a misalignment of the nozzles with respect to each other, which can easily explain the impaction of inertial particles outside the collection nozzle (Loo & Cork, 1988). In the following section, the sensitivity of SADS performances with respect to nozzle misalignment is thus illustrated theoretically.

### 4.6 Theoretical effect of nozzle misalignment

Study by Loo & Cork (1988) emphasized the importance of maintaining axial misalignment between the acceleration and collection nozzle of a VI. In their case, which is very different from the SADS in terms of dimensions and air flow rates, they recommend avoiding an offset of more than 1.6% of nozzle diameter and observe that each 1.6% increase in misalignment leads to a 1% increase in nozzle wall loss. Meeting this criterion in the case of the SADS would mean avoiding a misalignment of more than 0.013 mm, which is challenging from a mechanical design point of view. Experimentally, measuring the misalignment offset of the mounted SADS was not feasible. However, a sensitivity analysis can be performed by means of parametric CFD computations to explore the impact of this parameter.

Simulations were hence carried out with relative displacements of the collection nozzle with respect to the acceleration nozzle (Figure 1) in the Z-axis direction, with likely values of the axial backlash between the two parts of the SADS. These parts are assembled by manually fitting together a shaft and a 37 mm diameter hole. Following ISO system of limits and fits, this corresponds to a H7/h6 clearance fit (location fit), whence a possible axial backlash in the range 0 to 0.041 mm is deduced. Parametric computations were then performed for axial backlashes of 0, 0.025 and 0.041 mm respectively (corresponding to 0, 3.1% and 5.1% of the nozzle diameter respectively). A simulation with an extreme backlash of 0.075 mm (9.4% of nozzle diameter) was also performed for information, which could correspond to a more tolerant H8/f7 ISO clearance fit (close-running fit).

Figure 11 displays the corresponding numerical simulation results for the transmission efficiency and wall deposition, compared with experimental measurements of SADS 1, 2, and R, illustrating the possible effect of axial misalignment on $\eta_p$. 

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**References**

Loo & Cork (1988)
Results show that likely values of the axial mechanical backlash between the two parts lead to a severe decrease of $\eta_p$ due to dramatically increasing wall losses $\eta_d$ on the collection nozzle external walls, especially for the most inertial particles. This finding is compatible with experimental measurements. Axial misalignment of the device is therefore a possible cause of the differences in performance between the ideal version and the mechanical realization of the SADS, for the most inertial particles tested. Of course, present simulations can only qualitatively reproduce the tendency of the experiment, since they were not performed with the actual axial backlash which is unknown and which varies between each SADS assembly. Also, the effects of other existing mechanical backlashes were not numerically tested and necessarily contribute to sampling performances (tolerance on the separation length between nozzles, existing angle between the axes of the two parts of the SADS, etc.). Based on the analysis of the results, we can conclude that the maximum allowable misalignment during the assembly of the SADS, between the collection nozzle and the acceleration nozzle, is established at 0.025 mm, as evidenced by the violet curve. Notably, this curve consistently aligns with that of the SADS R-CFD, representing a curve with perfect alignment.

5 Conclusion and discussion

This study experimentally evaluated the dichotomous sampling performance (gas and particles) of 5 SADS prototypes of identical design, and for an aerosol of liquid particles with aerodynamic diameters of 0.15, 2, 3 and 4.5 µm. Lab tests were carried out in a dedicated controlled generation environment, the BAO, in which monodisperse aerosols marked with a fluorescent dye were emitted, thanks to the modification of the CMAG generator to accommodate the use of glycerol and fluorescein as condensing vapour and nuclei respectively. SADS sampling performance in terms of total mass sampled, particle fraction transmitted to the major outlet and particle losses at the walls were put into perspective with the details of mechanical construction and with the operating conditions of the prototypes: leak rate, repeatability between successive assemblies, imprecision of machined nozzles diameters, and axial misalignment of the assembly.
The measured sampling performances were compared to their theoretical counterparts computed by CFD in Belut et al. (2022), and CFD was also used to study the theoretical effect of variations in the geometry of machined SADS relative to the plans, regarding the nozzles diameters and nozzle misalignments. With the originally proposed design, the SADS assembly was found easily leaky, but consequences on sampling could be overcome when a prior leak test with leakage rate LR below 4 Pa.s⁻¹ was passed. Sampling performances were found little reproducible during successive SADS assemblies (between tests, CV=22.1 % for wall losses). Theoretically unpredicted large (40-46 % of sampled mass) wall losses were measured for particles larger than 2 µm, located mostly (80 %) on the external walls of the collection nozzle. Assembly repeatability issues and simulations of SADS parts misalignment effect by CFD suggest that these undesirable particle deposits are due to the mechanical backlashes of the assembly, and not to the imprecision of actually machined nozzles diameters. Indeed, the measured variation of nozzle diameters in the range (-4.5 %, +3.7 %) with respect to nominal values were found to theoretically affect marginally (<3 %) aerosol transmission efficiencies.

Present results suggest that although the dichotomous sampling performances of the SADS are theoretically interesting for workplace exposure assessment to SVOC aerosols, its actual realization fail in reaching theoretical performances for micron-sized particles, due to mechanical design issues. Airtightness, nozzle alignment and repeatability of assembly are not sufficiently guaranteed by its initial design and future development should focus on improving these aspects to obtain a sampler suitable for field studies.

However, several biases may have affected the findings of the study. For example, rare macroscopic particles are sometimes emitted by the CMAG and may have been sampled by the SADS, biasing the separation performance measurements and especially the deposition measurements. However, we believe that this possibility is largely controlled by the real-time monitoring of particle sizes in the test rig, and by the simultaneous use of several SADS in the test rig. Regarding the plausibility of the simulation results, it is, of course, limited by the physical phenomena actually taken into account. Calculation errors are limited by the verification procedure used (Belut et al., 2022), which guarantees a numerical error of less than 0.5 % on the particulate fractions deposited and transmitted. However, actual variations in the geometry of assembled SADS compared with the drawings (due to machining inaccuracies other than nozzle diameters) are not taken into account, nor is wall roughness, despite its acknowledged effect on deposits. The roughness of the machined acceleration nozzles could therefore help to explain the deposition of particles in this nozzle, which are not predicted by calculations that assume a perfectly smooth nozzle. Similarly, the more or less pronounced sharpness of the sampling orifice actually machined can have a significant influence on the inlet particle velocity and concentration profile, and therefore on the actual performance of the SADS (Belut et al., 2022).

Nevertheless, the lack of repeatability of SADS performance between successive assemblies, its low and variable airtightness level and its radial clearance large enough to cause a significant misalignment of the nozzles (typically 5 %) have sufficiently important effects for these possible limitations of the study not to call into question its conclusions.

By addressing the identified challenges and incorporating further refinements in the SADS design and operation, researchers can enhance its reliability, accuracy, and applicability in various aerosol sampling applications, contributing to advancements in aerosol science and related fields.
7 Table of Symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\eta_p$</td>
<td>Particles transmission efficiency to the particle outlet (major flow)</td>
</tr>
<tr>
<td>$\eta_v$</td>
<td>Particles transmission efficiency to the vapour outlet (minor flow)</td>
</tr>
<tr>
<td>$\eta_d$</td>
<td>Particles deposition ratio</td>
</tr>
<tr>
<td>$\eta_{d_a}\cdot\eta_{d_c}\cdot\eta_{d_p}$</td>
<td>Particles deposition ratio in the acceleration nozzle, collection nozzle and in the plenum respectively</td>
</tr>
<tr>
<td>$m_{\text{major}}$</td>
<td>Mass of particles collected at the major flow outlet (ng)</td>
</tr>
<tr>
<td>$m_{\text{minor}}$</td>
<td>Mass of particles collected at the minor flow outlet (ng)</td>
</tr>
<tr>
<td>$m_{\text{inlet}}$</td>
<td>Mass of sampled particles at the inlet (ng)</td>
</tr>
<tr>
<td>$m_{\text{dep}}$</td>
<td>Mass of deposited particles (ng)</td>
</tr>
<tr>
<td>MMAD</td>
<td>Mass median aerodynamic diameter</td>
</tr>
<tr>
<td>GSD</td>
<td>Geometric standard deviation</td>
</tr>
<tr>
<td>$L_R$</td>
<td>Leakage rate (Pa.s$^{-1}$)</td>
</tr>
<tr>
<td>T</td>
<td>Temperature ($^\circ$C)</td>
</tr>
<tr>
<td>P</td>
<td>Pressure (Pa)</td>
</tr>
<tr>
<td>$M_{\text{SADS}}$</td>
<td>Mass balance of SADS</td>
</tr>
<tr>
<td>$m_{\text{CFC}}$</td>
<td>Mass measured by the closed face cassettes (ng)</td>
</tr>
<tr>
<td>LLOQ</td>
<td>Lower limit of quantification ng.L$^{-1}$</td>
</tr>
<tr>
<td>ULOQ</td>
<td>Upper limit of quantification ng.L$^{-1}$</td>
</tr>
<tr>
<td>WD</td>
<td>Wall depositions</td>
</tr>
<tr>
<td>$m_{\text{WL}}$</td>
<td>Quantity of fluorescein measured in the wall ng</td>
</tr>
<tr>
<td>$m_{\text{tot}}$</td>
<td>Total quantity of fluorescein sampled by the SADS</td>
</tr>
<tr>
<td>$U_1$</td>
<td>Air velocity m.s$^{-1}$</td>
</tr>
<tr>
<td>$\rho$</td>
<td>Air density Kg.m$^{-3}$</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Air viscosity Pa.s$^{-1}$</td>
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8. Author contribution

BS, EB, EG and RO conceptualized the research project. NR, BS and EB developed the methodology and NR was responsible of the investigation. BS and EB validated the results. EG was responsible for the supervision of the project. NR wrote the original draft preparation and BS, EB and EG reviewed and edited the manuscript.

9. Competing interests

The authors declare that they have no conflict of interest.
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8 Bibliography


