A Compact and Portable Deposition Chamber to Study Nanoparticles in Air-Exposed Tissue

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Abstract

Background: Epidemiological studies show that elevated levels of particulate matter in ambient air are highly correlated with respiratory and cardiovascular diseases. Atmospheric particles originate from a large number of sources and have a highly complex and variable composition. An assessment of their potential health risks and the identification of the most toxic particle sources would require a large number of investigations. Due to ethical and economic reasons, it is desirable to reduce the number of in vivo studies and to develop suitable in vitro systems for the investigation of cell–particle interactions.

Methods: We present the design of a new particle deposition chamber in which aerosol particles are deposited onto cell cultures out of a continuous air flow. The chamber allows for a simultaneous exposure of 12 cell cultures.

Results: Physiological conditions within the deposition chamber can be sustained constantly at 36–37°C and 90–95% relative humidity. Particle deposition within the chamber and especially on the cell cultures was determined in detail, showing that during a deposition time of 2 hr 8.4% (24% relative standard deviation) of particles with a mean diameter of 50 nm [mass median diameter of 100 nm (geometric standard deviation 1.7)] are deposited on the cell cultures, which is equal to 24–34% of all charged particles. The average well-to-well variability of particles deposited simultaneously in the 12 cell cultures during an experiment is 15.6% (24.7% relative standard deviation).

Conclusions: This particle deposition chamber is a new in vitro system to investigate realistic cell–particle interactions at physiological conditions, minimizing stress on the cell cultures other than from deposited particles. A detailed knowledge of particle deposition characteristics on the cell cultures allows evaluating reliable dose–response relationships. The compact and portable design of the deposition chamber allows for measurements at any particle sources of interest.

Key words: ultrafine particles, particle–cell interaction, aerosol, particle deposition, in vitro toxicology

Introduction

Aerosol particles present in the ambient atmosphere and engineered nanoparticles used in industrial processes and consumer products are potentially involved in a number of respiratory and cardiovascular diseases. The main route of undesired uptake of submicrometer particles into the body is via inhalation.1) Epidemiological studies provide ample and consistent evidence for a link between adverse health effects and increased concentrations of ambient particles.2–4) There are indications for a specific toxicological role of ultrafine or nanoparticles, i.e., particles with diameters of <100 nm.5–7) Health effects assumed to be associated with aerosol particles are one of the major environmental public health issues in industrialized countries, especially as the environmental burden of nanoparticles is likely to increase in the future.8) Besides atmospheric particles, a large number of engineered nanoparticles are being introduced in industrial...
processes, many of them containing potentially toxic metals. These engineered nanoparticles may become aerosolized at some stage of their life cycle and pose additional health risks for specific workplace settings, but also for the general public.\(^\text{39}\) It should be noted that real-life release mechanisms may strongly alter the size distribution of such engineered nanoparticles, and as a result of this they might not be present in the atmosphere as nanometer-size particles or have altered physical–chemical properties.

Mechanistic in vivo animal and in vitro studies support the findings of epidemiological studies,\(^\text{10–12}\) but the current understanding on a cellular or biochemical level is still incomplete. Thus, a thorough understanding of (nano)particle–lung interactions is urgently needed. In addition, an increasing use of aerosol particles for pharmaceutical drug delivery requires a detailed understanding of particle deposition and drug uptake processes via the lung.

Increasing diversity of engineered nanoparticles, pharmaceutical aerosol drug delivery systems, and also the large number of sources of atmospheric particles call for comprehensive investigations of particle–lung interactions. But an extensive use of animal models is hampered by ethical, technical, and economic reasons, and therefore in vitro techniques are strongly favored. A realistic experimental setup to assess aerosol particle–lung interactions and associated potential toxic effects with in vitro studies should include the following essential parts: (1) nanoparticles need to be deposited onto the cell cultures as real aerosol particles, i.e., from a continuous air flow at realistic particle concentrations, composition, and size; (2) cell cultures used to assess nanoparticle effects need to closely mimic the complex multicellular structure of the human lungs\(^\text{13,14}\), and (3) the exposure and deposition scenario should mimic the corresponding processes in the lungs as accurately as possible.

Current experimental procedures for in vitro particle studies rarely meet these essential requirements, which might result in severe biases when interactions between particles and lung cells are interpreted. Particles are often deposited onto lung cells all at once (e.g., pipetted). This does not reflect the extended period of time during which particles are usually deposited on the lung surface and results in often unrealistically high particle concentrations in in vitro studies. In addition, suspending particles in a solvent can substantially alter their chemical and physical properties compared with conditions where they are freely suspended in air.\(^\text{13,14}\)

We present here a new instrument, which aims to mimic (nano)particle deposition and interaction with lung cells in in vitro experiments as accurately as possible compared with conditions in the lung. A first version of our exposure system introduced the concept of electrostatic deposition of biopolarly charged aerosol particles onto lung cell cultures from a continuous air flow, which is much more efficient than diffusional deposition.\(^\text{13}\) Further, we demonstrated that the associated very weak currents did not induce any cell effects. We also showed that the particles were evenly deposited onto the cell cultures, an essential aspect when toxicological dose–effect relationships are established. Particles were humidified (to \(>80\%\) relative humidity (RH)) and heated to 36–37°C outside of the deposition chamber to mimic incubator-like conditions.

In the significantly improved in vitro system described here, the realistic particle deposition is further refined. Essential parts of the experimental setup, i.e., aerosol humidification, heating, and cell exposure, are combined into one portable instrument, which allows performing experiments outside the cell biology laboratory at any particle source of interest. Besides a detailed description of design and operation of the new deposition chamber, the deposition efficiency of particles is evaluated in detail. This is an essential aspect if biological responses are interpreted in a toxicological context.

**Materials and Methods**

**Design of the deposition chamber**

A schematic of the deposition chamber is shown in Figure 1. The instrument consists of three subunits: a humidifier, a distributor, and a cell chamber unit. Immediately after entering the deposition chamber, the aerosol is humidified by passing through a GORE-TEX\textsuperscript{®} tube (length, 10 cm; inner diameter, 0.6 cm) that is immersed in a heated water bath (1.5 L, 18 MΩ cm water). Thus, the aerosol flow is conditioned to 36–37°C and 90–95% RH. In the distributor, the aerosol is evenly divided into 12 circularly arranged delivery tubes (inner diameter, 1 cm). When exiting the delivery tubes, the aerosol passes over the cell cultures grown on filter inserts [polyester membrane, 0.4 μm pore size; 24-mm Transwell\textsuperscript{®} inserts (Corning, Vitaris, Baar, Switzerland) or 24-mm Falcon\textsuperscript{™} inserts (Becton Dickinson AG, Milan, Geneva, Switzerland)] positioned approximately 2 mm below each tube. The filter inserts are placed in a home-built 12-well plate (for the simultaneous exposure of 12 cell cultures), which is routinely autoclaved before each cell culture exposure experiment. All other parts of the deposition chamber, except the short GORE-TEX tube, can also be sterilized, if required. The air is pumped out of the chamber through a centrally positioned outlet, which is connected to a membrane pump. Total air flow rate is 600 mL min\(^{-1}\), i.e., 50 mL min\(^{-1}\) in each

![FIG. 1. Design of the deposition chamber. The aerosol particle deposition chamber consists of a cylindrical humidifier, a distributor, and the cell chamber containing the 12-well plate (5) and the delivery tubes (4) (see also Fig. 2a). The chamber is heated in the water bath (3) as well as at the bottom (6). The device is insulated by a polystyrene housing (1). The continuous air flow from the aerosol source is humidified by passing through a GORE-TEX tube (2). The flow path of the aerosol is illustrated by the blue arrows.](image-url)
delivery tube. The deposition chamber is heated by a 120 W resistor in the humidifier’s water bath, as well as a 6.2 W silicone rubber thermofoil at the bottom of the chamber. An insulating polystyrene housing (4 cm thickness) around the entire deposition chamber assures temperature homogeneity within the instrument. Each of the 12 delivery tubes is equipped with lead-throughs to allow for installation of optical fibers to detect ciliary beat frequency online during the particle exposure, as described in detail by Savi and coworkers. Potentially, also other optical signals, e.g., from fluorescent markers, could be observed online with these optical fibers. The radial design of the entire instrument assures an even distribution of the particles to the 12 cell culture wells.

Before entering the deposition chamber, trace gases that are potentially present in the aerosol (such as nitrogen oxides, ozone, and volatile organic compounds) are removed by a charcoal denuder to assure that observed biological effects are due only to particles. Five percent CO₂ (by volume) is added to the aerosol flow to maintain the cell culture medium in the physiological pH range. Particles then pass a 85Kr bipolar charger (185 MBq), which assures that the aerosol particles are in charge equilibrium when they enter the deposition chamber. After passing the bipolar charger, particles carry 1–2 elemental charges (positive or negative), which is very similar to biological cells. After passing the bipolar charger, particles carry 1–2 elemental charges (positive or negative), which is very similar to biological cells. Particles then pass a 85Kr bipolar charger (185 MBq), which assures that the aerosol particles are in charge equilibrium when they enter the deposition chamber. After passing the bipolar charger, particles carry 1–2 elemental charges (positive or negative), which is very similar to the particle charge equilibrium in the atmosphere. This charge results in a very low current deposited on the cell culture, typically in the sub-femtoampere range, as discussed in detail by Savi et al. and is not expected to affect the cell cultures. Inside the deposition chamber in the cell chamber unit, the charged particles are deposited evenly (as discussed in detail by Savi et al.) and efficiently onto the cell cultures by applying an alternating square-wave electrical field of 4 kV cm⁻¹ between an electrode below each well (diameter, 1.8 cm) and a fine metal mesh at the outlet of each delivery tube as counter electrode (diameter, 1.2 cm). To avoid particle losses, only stainless steel or conductive rubber tubing is used.

Particle generation

A thorough characterization of the deposition efficiency of particles (i.e., the fraction of particles deposited out of the aerosol flow) is essential to assess how many particles are deposited on the cell cultures during an exposure experiment. Only a precise knowledge of the particle deposition efficiency allows the establishing of particle dose–cell effect relationships, the ultimate goal of many toxicological particle–lung interaction studies.

For the deposition characterization experiments, adipic acid aerosol particles were produced using a conventional recirculation atomizer (model 3076, TSI, Shoreview, MN) connected to a filtered air supply (model 3074, TSI). Adipic acid particles were used for these experiments, because this compound has favorable physical–chemical properties (see below) and the size of the particles is representative of particles found in the ambient atmosphere. The inlet air pressure for the atomizer was set to about 3 bar and the outlet flow was 2.1 L min⁻¹. We kept the adipic acid concentration within the nebulizer constant by using a large volume (1 L) of the solution. The aerosol generator produced a stable aerosol, which was checked by performing repetitive size distribution measurements, before and after every experiment. Three 85Kr sources placed after the atomizer (2 × 74 MBq, 1 × 185 MBq) were used to assure that the atomized particles are in charge equilibrium. An aqueous solution (170 μmol L⁻¹) of adipic acid [hexanedioic acid: HOOC-(CH₂)₆-COOH] was atomized for these measurements, generating particles with a diameter mode of 50 nm (number-weighted particle size distribution). The mass-weighted mode was at about 100 nm. Typical concentrations were 7–8 × 10⁵ particles cm⁻³ with a polydisperse size distribution (sigma 1.7). Adipic acid has a very high deliquescence point (~100% RH), and therefore the RH of 90–95% in the chamber does not affect the size distribution of adipic acid particles. The low vapor pressure of adipic acid (about 9.8 × 10⁻⁶ Pa at 296 K) assures that no adipic acid evaporates after deposition. Particles were collected during 15 min (total aerosol volume collected: 9 L). The filter was positioned upstream of the deposition chamber. The filter was extracted in 2 mL of 0.05 mM NaOH solution, and the extract was then diluted by a factor of 10. This solution was analyzed with ion chromatography coupled to a mass spectrometer (IC/MS) like all other samples. From the measured mass of adipic acid per milliliter, we calculated the mass of adipic acid per cubic meter of air. The comparison of IC/MS and SMPS measurements resulted in an effective density for adipic acid of 0.5 g cm⁻³, which was used to convert the SMPS volume data into mass concentrations.

IC/MS analysis

For a detailed quantification of the particles deposited on the cell culture inserts and to assess the variability of the deposition between the twelve inserts, the following chemical analysis method was developed: The adipic acid mass deposited by the particles in the deposition chamber was quantified with IC/MS. Particles were deposited in the deposition chamber as in normal operation, except that the filter membranes of the inserts (on which the cell cultures are normally grown) were removed. The 12 wells were filled with 3 mL of an aqueous sodium hydroxide (NaOH) solution, and the extract was then diluted by a factor of 10. The solution had a pH of 9–10 (0.05 mM), such that dissolved adipic acid deprotonates and cannot evaporate. The aerosol flow in these deposition experiments was 50 mL min⁻¹ in each particle delivery tube (600 mL min⁻¹ in total), and the exposure time was 120 min. Three different settings were considered, as described in Figure 2b: bypassing the entire deposition chamber (1) and passing the aerosol through the deposition chamber with the electrical field off (2) or on (3).

Particle deposition on the fine metal meshes at the end of each particle delivery tube (i.e., at the counter electrode) was
also quantified. Adipic acid particles were dissolved from the meshes by immersing the ends of the delivery tubes for 5 min in aqueous NaOH solution (2 mL, pH 9–10) filled into each well of a separate 12-well plate. The adipic acid concentration in the NaOH solutions was then quantified with IC/MS using a guard column (AG11-HC, Dionex, Sunnyvale, CA) and an analytical column (AS11-HC, Dionex). The eluent (flow rate, 0.35 mL min⁻¹) consisted of an OH⁻ gradient (0 min, 0.95 mM OH⁻; 5 min, 0.95 mM OH⁻; 15 min, 15 mM OH⁻; 21 min, 60 mM OH⁻; 23 min, 60 mM OH⁻; 23.1 min, 0.5 mM OH⁻; 29 min, 0.5 mM OH⁻). The gradient was generated online by an Eluent Generator (EG40, Dionex) from an EluGen potassium hydroxide cartridge (EGC-KOH, Dionex) and ultrapure water (18 MΩ cm). After the analytical column, OH⁻ anions were suppressed by an anion self-regenerating suppressor (ASRS 300 2 mm, Dionex). A conductivity detector and a quadru-
pole mass spectrometer with unity mass resolution were used as detectors. The IC/MS instrument was calibrated by direct injection of aqueous adipic acid standards (1.103, 10.95, and 109.6 ng mL⁻¹; corresponding to 7.5, 75, and 750 nM), and a first-order least-squares regression was fitted to the data. The retention time of adipic acid was 19.9 min. The reproducibility of this method is 3.6%, which is the standard deviation of three measurements of a 75 nM adipic acid standard solution [i.e., within the concentration range of our analyzed samples (20–200 nM)]. Samples were injected via a 100-μL sample loop without preconcentration step. The comparison of IC/MS and SMPS measurements resulted in an effective density of adipic acid in the particles of 0.5 g cm⁻³. Although the density of solid adipic acid in the bulk phase is 1.36 g cm⁻³, atomization and subsequent drying may lead to fragmented particles with a low effective density.

Results and Discussion

Chamber operation

The 12-well plate containing the cell culture inserts is easily inserted into the deposition chamber by opening the bottom plate of the cell chamber (see Fig. 1). The insulating polystyrene enclosure assures that a homogeneous temperature and humidity are established quickly in the deposition chamber after closing, as shown in Figure 3. The chamber was first brought from room temperature to the set temperature of 37 °C, which takes about 2 hr (Fig. 3, data at times before exposure start). Then the chamber was opened for about 4 min, which is enough to exchange the cell cultures, and closed again. The fast recovery of the desired temperature and humidity prevents condensation of water within the chamber at high RH, which is crucial to avoid particle losses, as well as temperature and humidity–induced stress for the cells. Seventeen minutes after the chamber is closed, the temperature reaches 35 °C and RH exceeds 85% (Fig. 3); after 30 min, the temperature stays constant between 36 °C and 37 °C and RH between 90% and 95%, which allows the

FIG. 2. Experimental setup of particle deposition efficiency measurements. (a) Schematic of experimental setup used to quantify particle deposition in the wells and the particle delivery tubes. Deposited particles were quantified with IC/MS. (b) Three different settings of the aerosol flow were used to quantify the overall particle deposition in the chamber using an SMPS: (1) bypassing the deposition chamber (total aerosol particle number, gray line); (2) passing the aerosol through the deposition chamber without electrical field (dotted line); and (3) passing the aerosol through the deposition chamber with electrical field on (black line).

FIG. 3. Temporal evolution of temperature and RH during exposure. After the deposition chamber is opened to replace a 12-well plate, the temperature and RH recover rapidly, i.e., after 17 min, temperature reaches 35 °C and RH exceeds 85%. After 30 min, temperature is between 36 °C and 37 °C and RH between 90% and 95%, which allows the
performance of cell exposure experiments for several hours. This mimics physiological conditions in the airways very well, and hence stress factors on the cells other than induced by aerosol particles are minimized. After a typical exposure time of a few hours, the 12-well plate is removed from the deposition chamber for further analysis of the cell cultures.

In two recent studies, the deposition chamber was used to expose various lung cell cultures to aerosol particles from diesel exhaust, wood burning, and secondary organic aerosols. These experiments and the observed biological effects are described by Gaschen and co-workers. Particle deposition characteristics

Overall particle losses in the deposition chamber. The fraction of adipic acid particles (generated online with a nebulizer) deposited within the entire deposition chamber, with and without electrical field, was determined using an SMPS. The experimental setup is shown in Figure 2b. Three different experiments were performed: bypassing the deposition chamber (1) and passing the aerosol through the deposition chamber with the electrical field off (2) and on (3). This setup allows the effect of the electrostatic deposition to be determined and the overall diffusional losses in the entire deposition chamber to be quantified.

Figure 4a shows the number-weighted particle size distribution of adipic acid aerosol particles for the three different measurement conditions described in Figure 2b: bypassing the chamber (black) and passing through the deposition chamber with the electrical field off (light gray) and with the electrical field on (gray). A significant fraction of the particles is lost in the deposition chamber even when the electrical field is off, as illustrated by the difference between the black (bypass mode) and light gray (electrical field off) distributions. An increasing fraction of particles is lost without electrical field between 30 nm and 50 nm, which is likely due to turbulence effects within the deposition chamber. Turning on the electrical field leads to a further decrease of the particles passing through the chamber. Figure 4b displays the size dependence of the fraction of particles deposited when the electrical field is on, relative to the particle number passing through the chamber without electrical field. This is calculated from four SMPS scans, and the resulting standard deviations are shown as error bars. Due to low counting statistics at larger diameters, the error increases strongly with increasing diameter. The smallest particles (<20 nm) are deposited with an efficiency of about 15%, 40–120 nm particles with about 40–45% efficiency, and particles larger than 120 nm with about 30% efficiency. Over a wide particle size range, the particle deposition efficiency is only weakly dependent on the particle diameter, which can be explained by the different deposition processes for small and large particles, i.e., diffusional processes for small particles and impaction for large particles as described in detail by Savi and co-workers.

In terms of particle mass concentration, 41% of the aerosol passes the deposition chamber with the electrical field off and 22% with the electrical field on, relative to the bypass mode (black distribution in Fig. 4a). Thus, on a mass basis, 19% of the particles entering the deposition chamber are deposited in the chamber due to the electrical field. However, these measurements do not allow verifying whether all these particles are actually deposited on the cell cultures. This was determined in more detail with the IC/MS method described below.

Particle deposition efficiency on the cell cultures. For a detailed quantification of the particles deposited on the cell cultures, six different experiments were performed (see Table 1). In these experiments, adipic acid particles were quantified using IC/MS. The variability of the deposition was determined for two different plates (experiments 1–4, Table 1). These data were compared with particle deposition without an electric field (experiment 6) and an experiment where adipic acid was quantified in the wells and on the counter electrode, i.e., the mesh of the delivery tubes (experiment 5).

Table 2 lists the results of these experiments. Numbers are given as fractions of the adipic acid mass deposited relative to the adipic acid mass entering the deposition chamber (in percent). Figure 5 shows the average deposition efficiencies of experiments 1–5 for every well of the plates. The left y-axis shows the fraction of deposited particle mass in percent. An
average deposition efficiency of 8.4% with a relative standard deviation of 24% is observed. The average particle deposition efficiency was 7.6% (21% relative standard deviation) onto plate 1 (average of experiments 1, 2, and 5) and 9.8% (19% relative standard deviation; see Table 2) onto plate 2 (average of experiments 3 and 4). The difference is likely due to slight variation in the machining of the two plates where the distance between the end of the delivery tubes and the wells is a critical parameter. The average well-to-well variability of the deposition efficiency is 15.6% (i.e., average of the standard deviations of all wells in experiments 1–5; see Table 2) with a standard deviation of 24.7%.

The right y-axis in Figure 5 shows the fraction of particle mass deposited, considering only the charged particles. The mode of the adipic acid particles is about 50 nm (Fig. 4a), and the corresponding mass median diameter (MMD) is about 100 nm. According to Wiedensohler, about 28.5% of 100-nm particles carry a positive and 28.5% a negative charge at equilibrium. Thus, about 71.5% are neutral or carry a charge with the opposite polarity, and therefore are not deposited on the inserts due to the electrical field. This means that about 24% of all charged particles are deposited in plate 1 and about 34% in plate 2.

A decreased frequency of switching between positive and negative polarity could potentially increase the deposition efficiency (see Fig. 3 in Savi et al.(15)). Therefore, in experiment 5 (Table 2), the frequency was lowered to 1/3 Hz compared with 1 Hz (experiments 1–4). No statistically significant increased deposition could be observed, indicating that for the geometry of the deposition chamber, 1 Hz is an optimal value.

Particle deposition decreased significantly to about 2.7% when no electrical field was applied (experiment 6), showing that the electrical field increases the deposition efficiency by about a factor of 3.1 compared with deposition by diffusion processes (i.e., with the electrical field off) for particles with 100 nm MMD.

Due to the 85Kr bipolar charger, particles are charged positively and negatively, which mimics the charge distribution present in the atmosphere. However, this also results in a limited deposition efficiency on the cell cultures, because only particles of one polarity are attracted to the cell cultures, whereas particles of the other polarity are attracted to the counter electrode, i.e., the metal meshes at the end of the delivery tubes. This can be clearly seen in experiment 5 where the adipic acid concentration was measured in the wells and on the meshes at the end of the delivery tubes. On average, about 35% more particles were deposited on the meshes (12.2±2.0%) compared with the wells (8.8±1.3%). The combined deposition efficiency on the meshes and the wells is 21% (±2%) and includes particles deposited due to the electrical field, but also particles that are deposited due to diffusional/turbulent processes. As described above, about 57% of all 100-nm particles are charged using a bipolar charger. Assuming that about half of the particles are lost in the chamber (see Fig. 6 below) and assuming that these losses affect neutral and charged particles equally, about 74% of the charged particles are charged.
corresponding standard deviation. On average, 8.4% efficiency on all 12 wells, and the dashed lines illustrate the solid line indicates the averaged particle deposition efficiency on all 12 wells, and the dashed lines illustrate the corresponding standard deviation. On average, 8.4±0.8% (mass) of all particles or 30±2.9% (mass) of the charged particles are deposited on the wells.

(21% divided by 28.5%) are deposited in the wells and on the meshes due to the electrical field applied.

Figure 6 summarizes and compares the results from SMPS and IC/MS measurements. The left bar is derived from SMPS mass concentration measurements without electrical field where 41% of the particle mass passes the deposition chamber (light gray fraction) and 59% is deposited in the instrument. From the IC/MS measurements, we know that about 5% of the particles deposit on the meshes and in the wells if no electrical field is applied (experiment 6). This fraction is represented by the white part of the bar, and thus about 54% are lost in other parts of the instrument.

The middle bar shows the SMPS measurements with the electrical field applied. Here only 22% of the particles exit the chamber. As the overall diffusional losses in the deposition chamber are assumed to be the same as for conditions without electrical field, i.e., 59%, about 19% are deposited on the wells and the meshes due to the effect of the electrical field (gray fraction).

The right bar shows the particles deposited as quantified with the IC/MS technique (experiment 5). Twenty-one percent of the particle mass is deposited on the meshes and the wells. Diffusional deposition is assumed to account again for 5%, resulting in 16% of the particles deposited due to the electrical field, which compares well with the 19% determined from the SMPS experiment (middle bar).

Higher total deposition efficiencies could be reached with a unipolar charging of the particles as shown by others, and other systems described in the literature where four or less cell cultures are exposed simultaneously, whereas Sillanpää et al. reached even 95%. Lenz et al. determined a deposition efficiency of 7.2% for a deposition chamber, which was developed to investigate the deposition of nanoparticles in liquids.

Conclusions

We developed and characterized a new instrument to deposit aerosol particles onto cell cultures. Within the deposition chamber, aerosol particles are deposited on cell cultures directly out of a continuous aerosol flow at realistic physiological conditions, i.e., at 36–37°C and 90–95% RH. This tool can be used to investigate the impact of atmospheric, engineered, as well as therapeutic aerosols on cell cultures. Twelve cell cultures can be exposed simultaneously, allowing for a larger number of exposures compared with our previous instrument and other systems described in the literature where four or less cell cultures are exposed simultaneously.

The new deposition chamber is a compact and portable system allowing for particle sampling at any aerosol source, even outside a cell biology or particle laboratory.

A key feature of the deposition chamber is the efficient electrostatic particle deposition on the cell cultures using bipolarly charged aerosol particles, which mimics the charge distribution of particles in the ambient atmosphere. About 24–34% of all charged particles are deposited on the cell cultures, which is about a factor of 3–4 larger than deposition solely due to diffusional processes for the 100-nm particles (MMD) (sigma 1.7) used in this study (which is a typical size for ambient atmospheric aerosol particles). It is expected that this factor is even more favorable for larger particles, where diffusional deposition is less efficient. Deposition of particles of both polarities assures that, on average, no net charge is deposited on the cells. However, bipolar charging has the disadvantage that a fraction of the particles always carries no net charge and is thus not efficiently deposited in our chamber. The fraction of charged particles is mainly a function of particle size and varies between about 20% (one polarity) for 50-nm particles and about 45% for 1,000-nm particles.

The two analytical methods applied in this study to quantify the deposition efficiency of particles within the deposition chamber are in good agreement. The variability of the deposition efficiency between the 12 cell culture wells was assessed to be between 10% and 20%.

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Author Disclosure Statement

All authors declare that no conflicts of interest exist.

References


