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Determination of acrolein in ambient air and in the atmosphere of environmental test chambers†

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Acrolein (2-propenal) is a reactive substance undergoing multiple reaction pathways and an airborne pollutant with known corrosive, toxic and hazardous effects to the environment and to human health. So far, investigating the occurrence of acrolein in indoor air has been challenging due to analytical limitations. The classic DNPH-method has proven to be error-prone, even though it is still recommended in specific testing protocols. Thus, different approaches for an accurate determination of ambient acrolein have been introduced. In this work, an overview of already published data regarding emission sources and air concentrations is provided. In addition, a new method for the quantitative determination of acrolein in environmental test chambers and in indoor air is presented. Analysis is carried out using thermal desorption and coupled gas chromatography/mass spectrometry (TD-GC/MS) after sampling on the graphitized carbon black (GCB) Carbograph™ 5TD. All analytical steps have been carefully validated and compared with derivatization techniques (DNPH and DNSH) as well as online detection using PTR-QMS. The sampling time is short due to the low air collection volume of 4 L. Although derivatization is not applied, a detection limit of $0.1 \mu\text{g m}^{-3}$ can be achieved. By increasing the sampling volume to 6 L, the limit of detection can be lowered to $0.08 \mu\text{g m}^{-3}$. No breakthrough during sampling or analyte loss during storage of the acrolein laden sampling tubes was found. Therefore, the presented method is robust, easy-to-handle and also very suitable for routine analyses and surveys.

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Environmental significance

Acrolein is considered a priority indoor air pollutant. It is formed and released by several chemical processes and is known to have a negative impact on human health. Consequently, the accurate determination of acrolein in the indoor environment is of considerable importance. This paper introduces a new analytical method based on solid sorbent sampling and subsequent analysis by TD-GC/MS. It is easy-to-handle, fast and robust with a low limit of detection (LOD) at small sampling volumes. Validation was performed using recovery and breakthrough tests as well as chamber experiments. The reliability of the method makes it particularly suitable for routine analyses and surveys.

1. Introduction

Acrolein (2-propenal) is the simplest monounsaturated aldehyde and has two theoretically possible conformers.¹ The substance, which is liquid at room temperature, is classified as corrosive, toxic and hazardous to the environment. The International Agency for Research on Cancer (IARC) lists acrolein as a group 3 carcinogen. The toxic effect is mainly based on the formation of protein and DNA adducts.² Irritant effects,³ decreased respiratory function⁴ and cardiovascular diseases^{5,6} were also reported. The various exposure paths (inhalation, dermal, and oral) are discussed by Faroon *et al.*⁷ The Office of Environmental Health Hazard Assessment (OEHHHA) gives the

following Reference Exposure Limits (REL) for the inhalation path: acute REL = $2.5 \mu\text{g m}^{-3}$; 8 h REL = $0.7 \mu\text{g m}^{-3}$; chronic REL = $0.35 \mu\text{g m}^{-3}$.⁸ Trantallidi and co-workers⁹ derived a short-term health-based Critical Exposure Limit (CEL) of $21 \mu\text{g m}^{-3}$ from the results of a chamber study. Health Canada has released Residential Indoor Air Quality Guidelines (RIAQG) for contaminants which impact indoor air. For acrolein, a long-term exposure limit (24 hours) of $0.44 \mu\text{g m}^{-3}$ and a short-term exposure limit (1 hour) of $38 \mu\text{g m}^{-3}$ are recommended.¹⁰

In industrial use, acrolein is an intermediate in the production of acrylic acid and methionine. Acrolein synthesis routes are the aldol condensation of formaldehyde and acetaldehyde, the dehydrogenation of glycerol and, practically most important, the catalyzed oxidation of propene.^{11,12} First, Esterbauer *et al.*¹³ and later, Stevens and Maier¹⁴ reported the exogenous and endogenous pathways of acrolein formation (see Fig. 1).

Acrolein is ubiquitous in the environment and an intermediate in atmospheric chemistry.^{15,16} Although various sources

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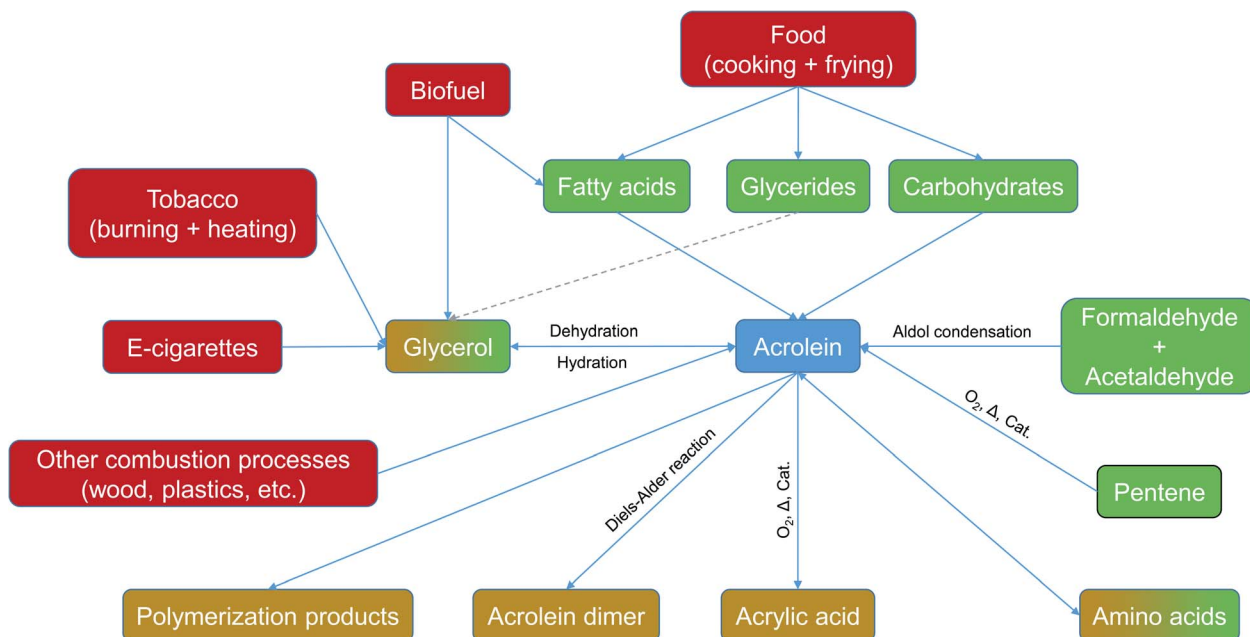


Fig. 1 Exogenous and endogenous pathways of acrolein formation and reaction.

have been identified, human indoor exposure to acrolein has rarely been studied so far, even though Logue *et al.*¹⁷ identified acrolein as one of nine priority hazards in their review on chemical air contaminants in residences. The comparatively small number of publications on acrolein in indoor air may be due to the fact that VVOCs (very volatile organic compounds) have only rarely been included in systematic surveys.¹⁸ Moreover, as an α,β -unsaturated carbonyl compound, acrolein is highly reactive and therefore poses a particular challenge from an analytical point of view. Thus, various methods for determining acrolein in ambient air have been developed.

1.1 Study objectives

With regard to the importance of acrolein in the environment, the current paper discusses published data and already applied analytical techniques. Furthermore, facing the integration of acrolein into the routine analysis of VVOCs¹⁹ and surveys, we are introducing a new method for its quantitative determination in environmental test chambers and in indoor air. By sampling on a solid adsorber without the need of derivatization, only small sampling volumes of air are required to reach sufficiently low detection limits. The analysis is carried out using thermal desorption followed by coupled gas chromatography/mass spectrometry (TD-GC/MS). The method has been tested and validated within the framework of various chamber experiments. During these test series, currently available and published methods based on DNPH- and DNSH-derivatization, respectively, were included for comparison. In addition, sampling on Tenax® TA according to ISO 16000-6²⁰ was considered as it can be applied during sampling protocols if no additional sampling on low-molecular carbonyls is foreseen, even though it is not recommended for the reliable determination of acrolein in air.

2. Acrolein emission sources and published levels in ambient air

2.1 Indoor emission sources

In indoors, acrolein is known to be mainly formed through thermal processes, which include the decomposition of wood,^{21,22} fuel,^{23–25} and plastics,^{26,27} incense burning,²⁸ cooking and deep-frying,^{29–34} smoking,³⁵ tobacco heating³⁶ and vaping.^{37,38} The cause of the formation of acrolein during food preparation are the decomposition processes of fatty acids,³⁹ glycerides and carbohydrates contained in food. Acrolein was identified to be emitted from solid wood and wood-based products in low concentrations, including insulation materials and flooring, as shown in Table 1.^{40,41} Several studies have focused on the release of acrolein due to cooking processes. Depending on the oil type and the temperature, elevated concentrations can be reached.^{39,42,43} Studies performed in hotel kitchens and restaurants revealed indoor levels in the range of $10 \mu\text{g m}^{-3}$ to $590 \mu\text{g m}^{-3}$.^{29,31} The sometimes high acrolein concentrations in the smoke and steam of tobacco products and e-cigarettes can be explained by the addition of glycerol and triacetin.⁴⁴ Heat-not-burn cigarettes have been found to be minor emission sources compared to conventional cigarettes.³⁷ A comprehensive study targeting gaseous emissions from burning scented and unscented candles measured unit specific emission rates between $8 \mu\text{g unit}^{-1} \text{ h}^{-1}$ and $20 \mu\text{g unit}^{-1} \text{ h}^{-1}$ and highlighted the fact that acrolein is released as a by-product which cannot be avoided.⁴⁵

2.2 Acrolein levels in outdoor and indoor air

Table 2 summarizes published acrolein data measured in outdoor and indoor air. In the Oakland-San Francisco Bay area,



Table 1 Acrolein emission rates determined in emission test chambers. See references for experimental details

Category	Value	Unit	Reference
Wood-based products			
Softwood			
Douglas fir	2	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
	8.1 ± 0.90	ng g^{-1}	Seaman <i>et al.</i> ⁴¹
Pine	8	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
	5.9 ± 0.55	ng g^{-1}	Seaman <i>et al.</i> ⁴¹
Spruce	2	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
	2	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
	2.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Hardwood			
Oak	<0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Red oak	1.0 ± 0.14	ng g^{-1}	Seaman <i>et al.</i> ⁴¹
Larch	2	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
	15	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Beech	<0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Yellow poplar	1.0 ± 0.14	ng g^{-1}	Seaman <i>et al.</i> ⁴¹
Redwood lumber	1.3 ± 0.41	ng g^{-1}	Seaman <i>et al.</i> ⁴¹
Laminated veneer lumber			
Hardwood/formaldehyde-resin	<0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Hardwood/polyvinyl acetate-resin	<0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Hardwood, dried without adhesive	<0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Hardwood, cooked	1	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Solid structural timber			
Softwood/melamine-resin	2	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Softwood/polyurethane	1	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Laminated timber			
Softwood/melamine-resin	1.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Softwood/isocyanate	1.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Plywood panel			
Three-layer panels	2.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Plywood	3	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Wooden composite board			
Oriented strand board (OSB)	<0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
	2	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Particle board	<0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
	1.0 ± 0.045	ng g^{-1}	Seaman <i>et al.</i> ⁴¹
	6.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Acoustic panels	2.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Insulation			
Wood fibre insulation	1	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
	6.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Wooden fibreboard	1	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Rigid foam	0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Flooring			
Parquet flooring	<1.25	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Laminate flooring	<1.25	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Schieweck ⁴⁰
Paint, construction materials			
Latex paint	0.35 ± 0.08	ng g^{-1}	Seaman <i>et al.</i> ⁴¹
Plaster board	3	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Plastics			
Foam plastic building materials applied on plywood, burning (max. 600–700 °C)	max. 400	ppm	Morikawa and Yanai ¹⁰²



Table 1 (Contd.)

Category	Value	Unit	Reference
PE-foil	<0.5	$\mu\text{g m}^{-2} \text{ h}^{-1}$	Fraunhofer WKI, unpublished results
Cooking			
Canola oil, heated to 180 °C	52.6 ± 2.5	$\text{mg h}^{-1} \text{ L}^{-1}$	Fullana <i>et al.</i> ⁴²
(Extra-virgin) olive oil, heated to 180 °C	9.3 ± 1.2 – 9.6 ± 0.9	$\text{mg h}^{-1} \text{ L}^{-1}$	Fullana <i>et al.</i> ⁴²
Soybean oil, heated to 180 °C	11.8 ± 67.4	$\text{mg h}^{-1} \text{ L}^{-1}$	da Silva and Pereira ⁴³
Palm oil, heated to 180 °C	2.58 ± 41.5	$\text{mg h}^{-1} \text{ L}^{-1}$	da Silva and Pereira ⁴³
Oil cooking events, 5 min post-cooking events	10.68–27.04	$\text{mg h}^{-1} \text{ kg}^{-1}$	Seaman <i>et al.</i> ³⁰
Different oil types, heated to 150–220 °C	max. 400–4160 at 220 °C	ppb	Endo <i>et al.</i> ³⁹
Stir-fry cooking in a test house	0.56	mg per person meal	Arata <i>et al.</i> ³⁴
Smoking			
Conventional cigarettes	6.5 ± 4.5	$\mu\text{g unit}^{-1} \text{ min}^{-1}$	Ruprecht <i>et al.</i> ³⁷
Heat-not-burn cigarettes (iQOS)	0.09 ± 0.03	$\mu\text{g unit}^{-1} \text{ min}^{-1}$	Ruprecht <i>et al.</i> ³⁷
Candles			
Different fuels, no fragrance	<8	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵
Palm, fruit fragrance	12	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵
Paraffin, fresh fragrance	13	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵
Paraffin, fruit fragrance	12	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵
Paraffin, spice & edibles fragrance	20	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵
Soy, fresh fragrance	10	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵
Soy, fruit fragrance	8	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵
Soy, spice & edibles fragrance	14	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵
Stearin, fruit fragrance	10	$\mu\text{g unit}^{-1} \text{ h}^{-1}$	Salthammer <i>et al.</i> ⁴⁵

acrolein concentrations between $0.032 \mu\text{g m}^{-3}$ and $0.100 \mu\text{g m}^{-3}$ were measured.⁴⁶ In further measurements in California, the concentrations in urban areas were significantly higher than in remote and coastal areas.⁴⁷ As part of “The Relationship of Indoor, Outdoor and Personal Air (RIOPA)” study, the indoor/outdoor ratio in 234 U.S. non-smoking homes, located in California, New Jersey and Texas, was found to be ≈ 1.3 with median values of $0.59 \mu\text{g m}^{-3}$ and $0.46 \mu\text{g m}^{-3}$ for indoor and outdoor air, respectively.⁴⁸ Acrolein concentrations in the range 0.1 ppb to 1.0 ppb (0.2 – $2.7 \mu\text{g m}^{-3}$, at $P = 1013 \text{ mbar}$ and $T = 298 \text{ K}$) were detected in the rural air of Portugal.⁴⁹ Seaman and co-workers⁴¹ investigated acrolein in the air of California apartments and found concentrations of up to $12.2 \mu\text{g m}^{-3}$. The values indoors were always significantly higher than outdoors (3–40 times). For the apartments examined, they calculated emission rates between 0.31 mg h^{-1} and 1.46 mg h^{-1} . Recently, indoor exposure to a large range of volatile organic compounds (VOCs) was investigated with high time resolution both during summer and winter in two Californian residences.⁵⁰ Due to continuous indoor sources (e.g. lumber), the main daily occupant exposure was $\sim 4.6 \text{ ppb-h d}^{-1}$ (mean value). Concentration peaks were observed in dependence of cooking events as specific emission sources. Both oven cooking and stovetop cooking were identified to produce the largest increase in indoor acrolein exposure. Enhancements in acrolein levels caused by an electric oven were higher than those associated with the use of a natural gas stovetop. The results confirmed the fact that heated cooking oil is a major acrolein source indoors.⁵⁰

In Canadian homes, measured acrolein concentrations were between $0.1 \mu\text{g m}^{-3}$ and $4.9 \mu\text{g m}^{-3}$.⁵¹ During the “House Observations of Microbial and Environmental Chemistry (HOMEChem)” study,⁵² the emissions of scripted everyday activities were measured in a test house in Texas. The amount of released acrolein during stir-fry cooking was $0.56 \text{ mg per person meal}$. As measurements were performed by PTR-TOF-MS, the recorded ion $\text{C}_3\text{H}_5\text{O}^+$ can be assigned both to acrolein and to a propionic acid fragment. The formation of acrolein is attributed to high-temperature degradation of cooking oil and heated vegetables, respectively.³⁴ In a Chinese temple, the acrolein level was up to 3 times higher during incense burning than on the outside.²⁸ Williams *et al.*⁵³ determined an average acrolein concentration of 5 – $6 \mu\text{g m}^{-3}$ in London smoking rooms exceeding those measured in comparison rooms. A comprehensive study targeting carbonyl emissions in the exhaust of commercial kitchens in China was performed by Ho *et al.*⁵⁴ revealing substantial amounts of acrolein in dependence of different cooking ingredients. Schieweck⁴⁰ published acrolein data obtained by indoor measurements in wooden prefabricated houses in Germany. Depending on the ventilation type and the construction stage, the concentrations varied between $5.1 \mu\text{g m}^{-3}$ and $18.3 \mu\text{g m}^{-3}$.

3. Analytical approaches

Table 3 provides an overview of analytical methods and procedures for the detection of ambient acrolein. As a rule, the substance is derivatized directly on the collection phase. The



classic DNPH method according to ISO 16000-3⁵⁵ and ASTM D5197⁵⁶ is applied in different testing protocols, but has been criticized to be unsuitable.^{57–59} Alternative techniques have therefore been developed including derivatization with dansylhydrazine (DNSH),^{60–62} sampling on solid sorbents,⁶³ thermal desorption gas chromatography,⁶⁴ online monitoring using proton-transfer-reaction mass spectrometry (PTR-MS),^{50,65,66} solid phase microextraction⁶⁷ and derivatization with *o*-(2,3,4,5,6-pentafluorobenzyl)hydroxylamine (PFBHA).⁶⁸

3.1 DNPH-coated sorbents

The most applied method is a solid phase extraction technique and corresponds to ISO 16000-3⁵⁵ and ASTM D5197.⁵⁶ Sorbent cartridges are filled with silica gel as the substrate which is impregnated with 2,4-dinitrophenylhydrazine (DNPH) as a derivatization agent. Rarely, DNPH was used as a solution filled in impinger bottles according to the US EPA Method TO-5.⁶⁹ After extraction with acetonitrile (ACN), the analysis of the formed hydrazone was performed by HPLC-separation and subsequent detection by UV. Williams *et al.*⁵³ performed comparative measurements of both approaches, but without identifying acrolein after sampling on DNPH-cartridges even though it was highlighted that this method is less expensive and time-consuming with lower material consumption than the wet chemical technique. Several publications have discussed the limitations and uncertainties associated with the determination of airborne unsaturated carbonyls when using DNPH-coated solid sorbent cartridges.^{57–59} Airborne oxidants, such as ozone (O₃), nitrogen oxide (NO) and nitrogen dioxide (NO₂) can interfere with unsaturated carbonyls.^{70–72} The reaction of acrolein and DNPH forming acrolein-2,4-DNP-hydrazone leads also to several dimerization and trimerization products with similar chromatographic properties and overlapping peaks which might lead to an over-estimation of acrolein concentrations.⁵⁸ Using fluorescence detection techniques, different absorption maxima of these side products were observed.^{57,73,74} The quantification is more complicated due to varying formation rates of the dimers and trimers in dependence of environmental factors during sampling and analytical conditions and also due to varying response factors under single-wavelength detection.^{59,73} An additional drawback is the fact that only acrolein-2,4-DNPH is commercially available as external standard solution and is therefore commonly used for calibration. However, this neither allows comparative measurements nor calibration of all other reaction products formed. Also, due to reaction processes and a resulting variation of formed peaks, the acrolein-DNP-hydrazone may not be correctly identified in real air samples. A summation of the acrolein-DNP-hydrazone and associated adduct peak areas might provide an estimation of the detected acrolein concentration, but seems to be limited to a storage period of approximately 5 h.^{58,59} However, an advantage of this established method is that it allows the detection of a broad range of carbonyls in one analytical step. The determination of acrolein can therefore be easily integrated into daily routine.

3.2 DNSH-coated sorbents

Due to the limitations of the DNPH-method, a silica-based-bonded C18 sorbent coated with dansylhydrazine (DNSH) was introduced as a reagent for derivatizing unsaturated carbonyls such as acrolein.^{57,60–62,75} Herrington and co-workers⁶¹ analyzed the formed DNSH-derivatives after ACN-extraction with HPLC-fluorescence techniques; the peaks should be identified and quantified on the basis of individual standard substances.^{61,62} Corresponding to the acrolein-DNSH derivatives formed, three chromatographic peaks can be observed.^{61,76} Two peaks are attributed to *E*- and *Z*-isomers of mono-derivatized acrolein with a molecular weight of 303.38 g mol^{−1}. The third peak is described as di-derivatized acrolein (dimer) with a molecular mass of 568.71 g mol^{−1} and is recommended to be used for acrolein analysis to obtain stable results.⁶¹ The DNSH-based technique is just recommended for acrolein with a low limit of detection (LOD) of 0.24 µg m^{−3} at low sampling volumes (7.5 L) and short sampling times (30 min), as the classic DNPH-method works well for discontinuous sampling of other low-weight carbonyls. The DNSH-method was further modified to increase the sensitivity by using a di-derivatized acrolein-hydrazone for quantification and to extend the analytical sensitivity to crotonaldehyde. Sampling is performed by using modified solid phase extraction tubes as diffusive, tube-type samplers (the Passive Aldehydes and Ketones Sampler (PAKS)).⁶¹

3.3 PFPH-coated sorbents

Ho and Yu⁶⁴ also developed a technique based on a derivatization agent, but using coupled gas chromatography/mass spectrometry after thermal desorption (TD-GC/MS) for analysis. Pentafluorophenyl hydrazine (PFPH) was introduced as a coating for Tenax® TA (60/80 mesh) as a solid sorbent. Even though the authors describe the method as highly sensitive with a LOD of ~0.8 µg m^{−3} for a sampling volume of 24 L, which led to the detection of higher background values, the approach has not been adopted widely.⁶⁴

3.4 PFBHA sampling

In order to trap ambient acrolein at trace levels (ppt) with short sampling times, a method based on a mist chamber was developed which is filled with a sodium bisulfite solution to absorb volatile acrolein. After dissociation of the sulfonates, the formed free carbonyls are derivatized with *o*-(2,3,4,5,6-pentafluorobenzyl)hydroxylamine (PFBHA) resulting in thermally stable oxime adducts which can be separated by GC followed by electron capture negative ionization mass spectrometry (GC/MS). Short term fluctuations of acrolein shall be detected by this technique at very short sampling times and a high sensitivity (LOD: 0.012 µg m^{−3}). Drawbacks are the facts that this approach requires intensive work and custom-made mist chambers as well as new calibration at each sampling day.^{30,41,68,77} Destailats *et al.*⁴⁶ sampled indoor air in impingers filled with a PFBHA solution (0.25 mg mL^{−1}). The formed PFBHA derivatives were extracted in dichloromethane after acidification with H₂SO₄. Analysis was performed by high-



resolution gas chromatography coupled with ion trap mass spectrometry (HRGC/ITMS). The method offers low limits of detection (LOD) at low sampling volumes, see Table 3.⁴⁶ Ho *et al.*⁵⁴ compared PFPH and PFBHA sampling in their study of acrolein concentrations in exhaust air of Chinese restaurants, but without highlighting similarities and differences of these methods.

3.5 Headspace and SPME-techniques

Headspace-based techniques have been published already in 1987 and later in 2013. In the 1987 study, the morpholine derivative of acrolein, 3-morpholinopropanal, was used to determine acrolein emissions. Analysis of the headspace was performed by GC coupled with a thermionic detector (TID),³² and in the 2013 experiments a GC/MS system was used.³⁹ In addition, solid phase microextraction (SPME fibers) was also used with DNPH or PFPH as the derivatization agent and subsequent GC/MS-analysis.^{67,78} Fullana and co-workers⁴² collected fumes of heated oil in Tedlar bags and analyzed them by GC/MS without a prior derivatization step. Identification of acrolein was confirmed by laboratory standards, and quantification was performed using MS response factors. A combination of headspace and needle trap microextraction (NTME) with subsequent GC/MS-analysis was applied by Lomonaco *et al.*²⁷

3.6 Solid sorbent sampling

Solid sorbents were just applied in very few cases without prior derivatization. Singer *et al.*⁷⁹ used Tenax® TA filled sorbent tubes with an extra section of Carbosieve® S-III, a carbon molecular sieve (CMS). Due to its very strong sorbent strength, Carbosieve® S-III is recommended to sample very volatile organic compounds (VOCs) in the range C2 to C5.^{35,79} However, the results indicated a possible sample breakthrough or incomplete recovery of acrolein. In addition, acrolein concentrations happened to be below the LOQ and, thus, complicated the quantification of background values.³⁵

3.7 On-line monitoring

As a fast on-line technique with high resolution, proton-transfer-reaction time-of-flight (PTR-TOF) mass spectrometry was introduced to detect and quantify acrolein in air samples with a LOD of 21 pptv, based on a 1 min integration period, and a precision of 1.5%.⁶⁵ For acrolein, the nominal target $M + 1$ ion is m/z 57, the proton affinity of acrolein is 797.0 kJ mol⁻¹⁸⁰ and the calculated proton transfer reaction rate constant between H₃O⁺ and acrolein is 3.35×10^{-9} cm³ s⁻¹.^{65,66,81} Lunderberg *et al.*⁵⁰ used a proton-transfer-reaction time-of-flight mass spectrometer (PTR-TOF-MS) to investigate occupant exposure patterns indoors in two California residences. During occupancy the mean acrolein concentrations were below 1 ppb.

4. Experimental methodology

4.1 Solid sorbent sampling without derivatization

A VVOC method developed earlier and published in 2018¹⁹ served as the basis for the experiments described here. Single-

bed stainless steel desorption tubes (Markes International Ltd., 89 mm length, 6.4 mm O.D.) were filled with the graphitized carbon black (GCB) Carbograph™ 5TD (20/40 mesh, Markes International Ltd.) as a solid sorbent. ~300 mg of the sorbent was placed between glass wool end plugs. Environmental test chamber air was sampled actively by means of a pump with a flow rate of 125 mL min⁻¹ and a total sampling volume of 4 L. After sampling, the tubes were sealed with Swagelok brass end caps fitted with PTFE ferrules. For analysis, the tubes were thermally desorbed (TD-100, Markes International Ltd.) into a coupled GC/MS system (Agilent 7890A/5975C). The conditions for thermal desorption were prepurge 3 min with a flow rate of 50 mL min⁻¹, primary desorption at 300 °C for 6 min at a flow rate of 20 mL min⁻¹, no inlet split, cold trap low 25 °C, pretrap fire purge 3 min at 50 mL min⁻¹, trap heating rate 40 °C s⁻¹, cold trap high at 300 °C for 6 min, outlet split 10 mL min⁻¹, and the flow path temperature at 200 °C. The cold trap contained quartz wool/Carbograph™ 1TD (40/60 mesh) and Carboxen® 1000 (80/100 mesh) with a ratio of 1 : 4.

The compounds were separated on a fused silica capillary column (6%/94% cyanopropylphenyl/dimethylpolysiloxane) of medium polarity (DB 624, 60 m, 0.32 mm, 1.8 µm, Agilent J&W). The initial column oven temperature was 30 °C (6 min), increased in a first step to 45 °C (1 °C min⁻¹) and in a second step to 240 °C with an increasing rate of 40 °C min⁻¹ (25.8 min run). The GC was operated in scan mode with a mass range of 20–450 amu, the MS source temperature was 230 °C, and the quadrupole temperature was 150 °C.

4.1.1 Calibration. Calibration was carried out by using an analytical grade acrolein standard (purity ≥90%, Sigma-Aldrich), which contained hydroquinone (~0.2%) and water (~3%) as stabilizers. Absolute grade methanol used for the preparation of the liquid standard solution was supplied by Sigma-Aldrich. The standard solution was prepared by weighing 10 mg acrolein into a glass flask, which was filled up with 10 mL methanol to obtain a standard concentration of 1 mg mL⁻¹. The limit of detection (LOD) and the limit of quantitation (LOQ) were calculated from the linear calibration curve $y = a \times x + b$ based on the approach given by Einax *et al.*⁸² and in accordance with DIN 32645.⁸³ For an analytical uncertainty of 33.3%, a significance level of 95% ($\alpha = 0.05$) and a sampling volume of 4 L, a LOD of 0.1 µg m⁻³ and a LOQ of 0.4 µg m⁻³ were calculated for a low concentration range (0.01 mg mL⁻¹ to 0.001 mg mL⁻¹). For a high concentration range (0.1 mg mL⁻¹ to 0.002 mg mL⁻¹) a LOD of 0.2 µg m⁻³ and a LOQ of 0.8 µg m⁻³ were obtained. At a total sampling volume of 4 L at an air flow rate of 125 mL min⁻¹, no breakthrough occurs.

Further information regarding calibration and breakthrough of acrolein is provided (ESI, Section S1†).

4.2 Comparison with other analytical methods

4.2.1 Derivatization methods. Sampling of acrolein during chamber tests (see below) was additionally carried out by using cartridges filled with silica gel impregnated with DNPH and with DNSH. For DNPH-coated cartridges (S10, Supelco, Inc.),



the flow rate was 1 L min⁻¹ at a total sampling volume of 75 L. DNSH-cartridges were sampled with a flow rate of 0.5 L min⁻¹ at a total sampling volume of 60 L. DNPH-coated cartridges were analyzed after extraction with acetonitrile by HPLC (Kinetex® 2.6 µm C18 100 Å, LC column 100 × 4.6 mm) with DAD-detection (Agilent 1260 Infinity HPLC DAD) in accordance with ISO 16000-3.⁵⁵ For calibration the acrolein-2,4-DNP-hydrazone was used. In addition to UV detection, analysis of the formed derivatives was also performed by HPLC Triple Quad MS (Agilent 6470 Triple Quad LCMS) in order to evaluate if a more precise identification of the peaks and the reaction products might be possible.

For preparing DNSH-cartridges, 100 mg DNSH (98%, 500 mg; Sigma Aldrich) were diluted with 100 mL acetonitrile (Rotisolv Ultra LC-MS; Carl Roth GmbH + Co. KG). After shaking, the solution was placed into an ultrasonic bath for 5 min until the hydrazine was fully dissolved. 100 µL acetic acid (100%; Merck) were added to the solution. The cartridges (S10; Supelco, Inc.) were cleaned with acetonitrile and subsequently eluted with the DNSH-solution for coating the silica gel bed. The prepared cartridges were dried in a nitrogen stream (100 mL min⁻¹) for one hour. Afterwards, the cartridges were closed with end caps and stored under laboratory conditions. After extraction with acetonitrile, samples were measured by HPLC (Zorbax RRHD Eclipse Plus, C18, 95 Å, 1.8 µm, 2.1 × 50 mm, Agilent Technologies) followed by MS-detection (Agilent 6470 Triple Quad LCMS). Quantification was performed against a liquid standard solution of DNSH in acetonitrile (50 ng mL⁻¹).

4.2.2 PTR-QMS. Throughout the chamber tests, the acrolein concentration in air of the 24 m³ stainless steel chamber was additionally monitored by quadrupole proton-transfer-reaction mass spectrometry (PTR-QMS, 07-06/48, IONICON Analytik GmbH).⁸⁴ The PTR-QMS was operated in the multiple ion detection mode (MID) and was placed outside the chamber to avoid interferences with the chamber atmosphere.

4.2.3 Sampling on Tenax® TA. A frequently applied method for identifying and quantifying VOCs in indoor air is described in ISO 16000-6.²⁰ During chamber experiments, air samples were drawn actively with a flow rate of 125 mL min⁻¹ through stainless steel desorption tubes (Markes International Ltd.) filled with the polymeric sorbent Tenax® TA (60/80 mesh, Chrompack). The total sampling volume was 4 L. The tubes were subsequently analyzed by GC/MS (Agilent 7890A/5975C) after thermal desorption (290 °C, 8 min; TD-100, Markes International Ltd.). The cold trap contained Carbograph™ 2/ Carbograph™ 1 (Markes International Ltd.), cold trap low -25 °C. The transfer line temperature was 180 °C. Separation was performed on a nonpolar DB-5 MS column (60 m × 0.25 mm, 0.25 µm) using a starting temperature of 32 °C, followed by a 5 °C min⁻¹ ramp to 150 °C, and a final 10 °C min⁻¹ ramp to 300 °C. The MS was operated in scan mode with a mass range of 25–550 amu, MS source temperature of 250 °C, and quadrupole temperature of 150 °C.

The mass spectra and retention data of acrolein were identified with its reference compound. For quantification, its own response factor was used. The limit of quantitation (LOQ) was ≤1 µg m⁻³.

Adsorbent media, analytical methods and sampling parameters chosen are also summarized in ESI, Section S4†.

4.3 Experimental set-up

4.3.1 Recovery of acrolein on Carbograph™ 5TD. The recovery of acrolein on the solid sorbent Carbograph™ 5TD was evaluated by using an acrolein permeation tube in order to obtain a stable concentration of acrolein in air. In the permeation tube (G-Cal, VICI Metronics), the permeant gas diffuses through a membrane and is mixed with a carrier gas at a controlled flow rate in order to obtain a known target concentration. According to the manufacturer, the permeation rate was given as 578 ± 29 ng min⁻¹ at a temperature of 45 °C. The permeation tube was inserted into a dosing apparatus in order to dilute the target substance released by the permeation tube. The temperature of the dosing apparatus was set to 45 °C. An air flow was directed through the heated oven and, thus, over the permeation tube with a flow rate of 100 mL min⁻¹. The outlet air stream was diluted with an air flow rate of 1000 mL min⁻¹ and a split ratio of 3 : 1 (0.825 L min⁻¹ : 0.275 L min⁻¹). Before the air flow passed the sampling sorbent tube, the smaller fraction was diluted with nitrogen (N₂; 5000 mL min⁻¹). For verification of the method, the test was also carried out with a toluene permeation tube (G-Cal, VICI Metronics) as a reference. The permeation rate of toluene was experimentally determined by the manufacturer and given with 2590 ± 648 ng min⁻¹ at a temperature of 60 °C (333.15 K). The permeation rate q_{d1} = 992 ng min⁻¹ of toluene at 45 °C was calculated according to empirical eqn (1):⁸⁵

$$\ln q_{d1} = \ln 2590 - 6794 \left(\frac{1}{318.15} - \frac{1}{333.15} \right) \quad (1)$$

Additionally, the permeation rate was checked by measuring the weight loss of the permeation tubes over the experimental period. The target concentration [µg m⁻³] of each substance in the gas flow was calculated by considering the permeation rate given by the manufacturer and calculated on the weight loss [ng min⁻¹], the carrier gas flow [mL min⁻¹] and the split ratio [1 : 3]. The test lasted 28 days. For safety reasons, the experiments were performed under an exhaust hood.

4.3.2 Recovery of acrolein concentrations dosed into chamber air. The analytical methods described above were compared on the basis of experiments under controlled climatic conditions, which were carried out in a 1 m³ glass emission test chamber and in a 24 m³ stainless steel emission test chamber, respectively. In the first step, acrolein test gas (Linde GmbH) was continuously dosed into the empty chamber in four increasing concentration stages: (i) 5 µg m⁻³, (ii) 10 µg m⁻³, (iii) 20 µg m⁻³ and (iv) 40 µg m⁻³. In the 1 m³ environmental test chamber, another level of 80 µg m⁻³ as the highest concentration was added. The calculated theoretical levels differed slightly (see the ESI, Table S4†). According to the manufacturer, the test gas had a target concentration of 20 mg m⁻³, a certified concentration of 22.6 mg m⁻³ and a tolerance of ±20%. The first sample was taken before dosing acrolein into chamber air



in order to determine background levels. Constant acrolein concentration levels were achieved approximately 3–4 h in the 1 m³ chamber and approximately 8–10 h in the 24 m³ chamber after the dosing started. As soon as the steady-state was achieved, sampling was performed with the analytical methods described above at specific time intervals (2 h, 4 h, 5.5 h and 7 h) at each acrolein concentration level. Sampling on DNSH-coated cartridges was only carried out after 2 h and 5.5 h due to the long sampling time. All presented results were obtained from duplicate analysis. The test method, steady-state acrolein concentrations and sampling intervals are provided in the ESI, Section S4†. In addition, online-monitoring of the acrolein chamber concentration was carried out by PTR-QMS in the 24 m³ test chamber and started simultaneously with the test gas dosing.

The test method was repeated in a second test series in order to verify if contaminants in the chamber air might influence the results due to chemical interactions. Therefore, the 24 m³ chamber was loaded with upholstered furniture and furnishings. The loading factor was not determined. The 1 m³ test chamber was loaded with different materials (*i.e.* leather, foam, wood-based materials, and bamboo) with a loading factor of 4.8 m² m⁻³.

Environmental conditions in the test chambers during both test series were set as follows: 23 °C ± 1 °C temperature, 50% ± 3% relative humidity and an air exchange rate of 1 h⁻¹ (1 m³ test chamber) and 0.5 h⁻¹ (24 m³ test chamber), respectively (see also the ESI, Table S4†).

The background value was determined before each experiment. If this value was above the LOQ, the measurement data

Table 2 Acrolein outdoor and indoor concentration (arithmetic mean ± standard deviation). *N* = number of measurements

Country, location		<i>T</i> [°C]/RH [%]	<i>N</i>	<i>C</i> [μg m ⁻³]	References
California, Oakland-San Francisco Bay area	Outdoor	—	12	0.032–0.100	Destailats <i>et al.</i> ⁴⁶
California, non-smoking residences	Outdoor	12–25/49–100	—	0.09–1.7	Seaman <i>et al.</i> ⁴¹
	Indoor	17–22/52–79	—	2.1–12.2	Seaman <i>et al.</i> ⁴¹
California, coastal	Outdoor	—	20	0.041–0.13	Cahill ⁴⁷
California, remote	Outdoor	—	20	0.041–0.11	Cahill ⁴⁷
California, urban	Outdoor	—	14	0.101	Cahill ⁴⁷
California, urban, LA Basin	Outdoor	—	8	0.32	Cahill ⁴⁷
California, single-family residences, summer	Indoor	—	1	1.3 (0.57 ppb)	Lunderberg <i>et al.</i> ⁵⁰
California, single-family residences, winter	Indoor	—	2	0.8–1.0 (0.36–0.44 ppb)	Lunderberg <i>et al.</i> ⁵⁰
United states, urban areas and private homes	Outdoor	—	353	0.46	Liu <i>et al.</i> ⁴⁸
	Indoor	—	353	0.59	Liu <i>et al.</i> ⁴⁸
Canada	Indoor	—	59	1 ± 1	Gilbert <i>et al.</i> ⁵¹
Portugal	Outdoor	25 (298 K)/—	51	0.2–2.3 (0.1–1.0 ppb)	Cerqueira <i>et al.</i> ⁴⁹
UK, Northern London, Mall, non-smoking	Indoor	20/—	—	0.5 ± 1.1 (0.2 ± 0.5 ppb)	Williams <i>et al.</i> ⁵³
UK, Northern London, student lounge	Indoor	20/—	—	6.2 ± 2.5 (2.7 ± 1.1 ppb)	Williams <i>et al.</i> ⁵³
Germany, prefabricated wooden house, naturally ventilated, after shell completion	Indoor	15–16/67–71	8	8.6 ± 6.3	Schieweck ⁴⁰
Germany, prefabricated wooden house, naturally ventilated, at delivery and in use	Indoor	23–26/58–63	15	18.3 ± 13.7	Schieweck ⁴⁰
Germany, prefabricated wooden house, mechanically ventilated, at delivery	Indoor	20–25/31–56	8	5.1 ± 2.0	Schieweck ⁴⁰
China, temple during incense burning	Outdoor	—	—	75 (32.5 ppbv)	Ho and Yu ²⁸
	Indoor	—	—	63–214 (27.3–92.9 ppbv)	Ho and Yu ²⁸
China, home during incense burning	Indoor	—	—	39 (16.8 ppbv)	Ho and Yu ²⁸
China, general Chinese restaurants (large, medium and small sized), exhaust	Indoor	21.0–36.6/38.5–62.1	9	~15.7–800.5 (6.8–347 ppb) ^a	Ho <i>et al.</i> ⁵⁴
China, Sichuan spicy food, exhaust	Indoor	43.0/36.3	2	~387.6–567.5 (168–246 ppb) ^a	Ho <i>et al.</i> ⁵⁴
China, HK style fast food, exhaust	Indoor	29.3/46.0	2	~58.8–100.1 (25.5–43.4 ppb) ^a	Ho <i>et al.</i> ⁵⁴
China, Chinese BBQ manufacturer, exhaust	Indoor	24.4–33.0/43.7–82.0	4	~7.4–218.9 (3.2–94.9 ppb) ^a	Ho <i>et al.</i> ⁵⁴
China, Korean BBQ, exhaust	Indoor	34.4/64.4	2	~148.3–168.2 (64.3–72.9 ppb) ^a	Ho <i>et al.</i> ⁵⁴
China, Western fast-food shops, exhaust	Indoor	24.0–32.2/28.5–67.9	12	~21.9–205.1 (9.5–88.9 ppb) ^a	Ho <i>et al.</i> ⁵⁴
China, Western Restaurant, exhaust	Indoor	22.2/51.0	2	~107.3–111.9 (46.5–48.5 ppb) ^a	Ho <i>et al.</i> ⁵⁴
Finland, restaurant kitchens and bakeries	Indoor	180–300 °C cooking temperature	10	10–590	Vainiotalo and Matveinen ³¹
Norway, restaurant kitchens	Indoor	—	44	10 ± 10	Svendsen <i>et al.</i> ²⁹

^a Conversion of ppb to μg m⁻³ at *T* = 23 °C.





Table 3 Analytical procedures and applied sorbent media for detecting acrolein in the gas phase, as reported in the literature

Sorbent type/media	Derivatization agent	Analytical technique	Column	Minimum detection limit (MDL)	Sampling volume (flow rate)	Reference
Mist chamber/bisulfite solution (0.1 M)	Hydrogen peroxide, pentafluorobenzylhydroxyl amine (PFBHA), sulfuric acid	GC/MS	—	—	190 L (19 L min ⁻¹)	Seaman <i>et al.</i> ^{30,41,68}
Tedlar bags	—	GC/MS and GC/FID	DB-5 (30 m, 0.25 mm, 0.25 µm)	41 ng m ⁻³	—	Cahill and Okamoto ²⁵
Impinger	—	GC/MS	Supel-Q Plot column (30 m, 0.53 mm)	—	200 L (10 L min ⁻¹) 0.5 L	Cahill ⁴⁷ Fullana <i>et al.</i> ⁴²
Silica gel	Hydrogen peroxide, pentafluorobenzylhydroxyl amine (PFBHA), 20 mL (0.25 mg mL ⁻¹) 2,4-Dinitrophenylhydrazine (DNPH)	GC/MS	DB-XLB HRGC (30 m, 0.25 mm, 0.25 µm)	0.91 ± 0.26 ng m ⁻³	1.8–4 L (0.45–1 L min ⁻¹)	Destailats <i>et al.</i> ⁴⁶
		HPLC/UV	C18 reverse phase	—	—	Svendsen <i>et al.</i> ²⁹
			Supelcosil LC-18 (250 mm, 4.6 mm, 5 µm)	—	— (1 L min ⁻¹)	Cerqueira <i>et al.</i> ⁴⁹
			ChromSep OmniSphere 5, C18 (150 mm × 5.6 µm)	—	—	Gilbert <i>et al.</i> ⁵¹
			—	—	30 L (0.5 L min ⁻¹)	Wang and Morrison ¹⁰³ according to TO-11A ¹⁰⁴
			Novapak C18 (150 mm, 3.9 mm, 4 µm)	—	5–10 L (50 mL min ⁻¹)	Machado Corrêa and Arbilla ²³
			C18 reverse phase	—	60–120 L	Jiang and Zhang ¹⁰⁵ according to TO-11A ¹⁰⁴
			—	—	—	Ruprecht <i>et al.</i> ³⁷ according to TO-11A ¹⁰⁴
			—	—	5 mL	da Silva and Pereira ⁴³
Impinger	2,4-Dinitrophenylhydrazine (DNPH), solution (10 mL)	LC-DAD-MS	Xterra MS C18 (250 mm, 2.1 mm, 5 µm)	—	Max. 80 L (0.5 L min ⁻¹)	Williams <i>et al.</i> ⁵³ according to TO-5 ⁶⁹
		HPLC/UV	L1, C18 (250 mm, 4.6 mm, 5 µm)	—	—	Guariero <i>et al.</i> ²⁴
			x-Terra MS column (250 mm, 2.1 mm, 5 µm)	—	—	Zhang <i>et al.</i> ⁶⁰
Solid/personal aldehydes and ketones sampler (PAKS)	Silica-based bonded C18 packing material, coated with DNSH	HPLC-fluorescence	Waters Nova-Pak C18 column (3.9 × 150 mm)	26 pg 0.25 µg m ⁻³	—	Herrington <i>et al.</i> ⁶¹
Tenax® TA/ Carbosieve® S-III Tenax® TA	—	TD-GC/MS	—	—	Max. 3 L (5 mL min ⁻¹)	Singer <i>et al.</i> ⁷⁹ according to TO-1 ¹⁰⁶
	Pentafluorophenyl hydrazine (PFPH)	TD-GC/MS	HP-5 MS (30 m, 0.25 mm, 0.25 µm)	LOD: ~0.8 µg m ⁻³ (<0.3 ppb) LOD: ~13.2–3.2 µg m ⁻³ (5.7–1.4 ppb)	0.72–3.89 L (20–30 mL min ⁻¹) 0.3–1.2 L (20 mL min ⁻¹)	Ho and Yu ⁶⁴ Ho <i>et al.</i> ⁵⁴
	Pentafluorobenzylhydroxyl amine (PFBHA)			PFPH, LOD: ~3 µg m ⁻³ (1.3 ppb) and ~1.6 µg m ⁻³ (0.7 ppb)	1.5 L and 3 L (20 mL min ⁻¹)	Ho <i>et al.</i> ⁵⁴
Solid phase microextraction (SPME, 85 µm polyacrylate)	2,4-Dinitrophenylhydrazine (DNPH)	GC/MS	HP-5 MS (30 m, 0.25 mm, 0.25 µm)	—	—	Osório and de Lourdes Cardeal ⁷⁸

Table 3 (Contd.)

Sorbent type/media	Derivatization agent	Analytical technique	Column	Minimum detection limit (MDL)	Sampling volume (flow rate)	Reference
Solid phase microextraction (SPME, 65 μm polydimethylsiloxane-divinylbenzene) Headspace	Pentafluorophenyl hydrazine (PFPH)	GC/MS	5% diphenyl, 95% dimethylpolysiloxane (30 m, 0.25 mm, 0.25 μm)	—	—	Dias <i>et al.</i> ⁶⁷
	—	HS-GC/TID	DB-1 (30 m, 0.25 mm)	—	—	Umano and Shibamoto ³² Endo <i>et al.</i> ³⁹
	—	HS-GC/FID	DB-5 or WAX fused silica capillary columns (18 m, 0.18 mm) DB-624 ultra inert (60 m, 0.25 mm, 1.4 μm)	—	—	Lomonaco <i>et al.</i> ²⁷
Headspace/needle trap microextraction (NTME) Carbograph TM 5TD	—	GC/MS TD-GC/MS PTR-QMS (scrubber) PTR-TOF-MS	DB 624 (60 m, 0.32 mm, 1.8 μm)	LOQ: 0.4 $\mu\text{g m}^{-3}$ LOQ: 0.3 $\mu\text{g m}^{-3}$ LOD: ~2 ppb LOD: 21 ppt	4 L (0.125 mL min ⁻¹) 6 L (0.125 mL min ⁻¹)	This work Knighton <i>et al.</i> ⁶⁶ Graus <i>et al.</i> ⁶⁵

were corrected accordingly. In order to avoid memory effects, the respective chamber was heated to 80 °C for 12–16 h after each experiment.

The experiments were performed under conditions which are representative for the indoor environment (23 °C, ~50% RH). As can be seen in Tables 1 and 2, cooking is a major emission source of acrolein indoors. Moreover, many cooking processes cause a relative humidity higher than 50%. The potential problem of water retention by solid sorbents has been addressed by several studies with a main focus on carbon molecular sieves (CMS) which are known to retain water due to their high sorbent strength.^{86–89} Thus, water management options are often needed when using CMS for indoor air sampling.⁹⁰ CarbographTM 5TD as graphitized carbon black (GCB) is classified as a hydrophobic adsorbent medium by different manufacturers. It is also used together with Tenax® TA as a hydrophobic combination for solid sorbent sampling.⁹⁰ Richter *et al.*⁹¹ determined recovery rates of 85 ± 14% and 116 ± 8% under dry (0% RH) and humid (50% RH) sampling conditions, respectively. The water adsorption capacities of different solid sorbents were comprehensively examined by Helmig and Vierling⁸⁹ including different types of Tenax, Carbotrap and Carbosieve® S-III as well as Carboxen® 569. Regarding the two CMS Carbosieve® S-III and Carboxen® 569, they proposed several steps to reduce the water uptake during sampling. However, the Carbotrap types (20/40 mesh), which are comparable to CarbographTM 5TD, showed no water uptake rates so they were recommended for air sampling even at high humidities. Fastyn *et al.*⁸⁸ also come to the conclusion that the water uptake of CarbographTM 5TD is low in comparison to other sorbents.

5. Results and discussion

5.1 Recovery of acrolein on the solid sorbent

As shown in Fig. 2 and supported by the measurement data, a constant target concentration in the carrier gas flow was achieved after approx. 14 days. Mean values obtained by active air sampling on CarbographTM 5TD and Tenax® TA, respectively, were 20 ± 1 $\mu\text{g m}^{-3}$ and 7 ± 1 $\mu\text{g m}^{-3}$. The permeation rates of acrolein and toluene, the calculated target concentrations and the measurement data (mean ± standard deviation) from active sampling on CarbographTM 5TD and Tenax® TA are summarized in the ESI, Table S6†. The target concentrations for acrolein calculated on the basis of the manufacturer information and on the basis of the weight differed to a small extent of ~9.4% (24 $\mu\text{g m}^{-3}$ vs. 22 $\mu\text{g m}^{-3}$). Measurement data obtained by active sampling on CarbographTM 5TD were in the same range (20 ± 2 $\mu\text{g m}^{-3}$) and quite good considering the tolerance given for the permeation rate (578 ± 29 ng min⁻¹). Therefore, the findings show that the recovery of acrolein on CarbographTM 5TD is good and reproducible with a low standard deviation. The recovery of acrolein on Tenax® TA was poor and a factor of 3 lower (7 ± 1 $\mu\text{g m}^{-3}$) than on CarbographTM 5TD.

However, the obtained data on Tenax® TA for toluene (mean value: 40 ± 1 $\mu\text{g m}^{-3}$) were in the range of the target concentration based on the calculated permeation rate according to



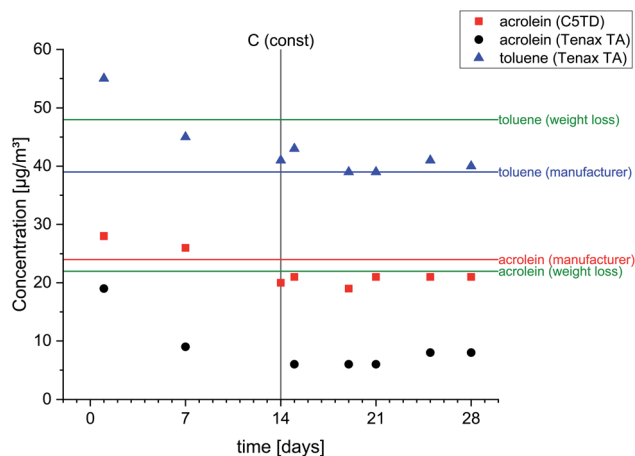


Fig. 2 Concentrations of acrolein and toluene obtained by permeation tubes and active air sampling on Carbograph™ 5TD (C5TD) and Tenax® TA.

the manufacturer information ($39 \mu\text{g m}^{-3}$), but differed from the permeation rate calculated using the weight loss ($48 \mu\text{g m}^{-3}$). The difference between both permeation rates might be explained by a less exact weight loss measurement. In general, it also cannot be excluded that the permeation rate might be influenced by external parameters during the experiment due to different materials and designs of the permeation tubes.

5.2 Recovery of acrolein concentrations dosed into chamber air

5.2.1 Recovery in empty emission test chambers. Fig. 3A and B show the acrolein concentrations dosed into the air of an empty 1 m^3 glass chamber and an empty 24 m^3 stainless steel test chamber. In the 24 m^3 chamber the acrolein concentration

was also monitored continuously by PTR-QMS (see Section 4.3.2). Data obtained by sampling chamber air on Carbograph™ 5TD showed the best agreement with the dosed concentrations. In both experiments, the deviations at all concentration levels were less than 10%.

The poor recovery on Tenax® TA was again underlined by the chamber test results. Both the data obtained by sampling on Tenax® TA and on DNPH-cartridges showed a broad variation and no correlation with the dosed acrolein concentrations. Analysing the cartridges by MS-detection did not lead to better results. In contrast, a reliable quantification was not possible due to the fact that the formed peaks could neither be exactly identified nor their formation was reproducible. By applying DNSH-cartridges, the di-derivatized acrolein (dimer) with a molecular mass of $568.71 \text{ g mol}^{-1}$, as recommended for quantification by Herrington and co-workers,⁶¹ could not be detected in the present study. The areas of the two stereoisomers were summed for quantification, but did not lead to reliable results. Also, calibration was not possible, as neither a reproducible peak distribution nor a reproducible peak ratio were obtained. The differences between the findings achieved in this study and those published might be due to the different analytical set-up. Herrington *et al.*⁶¹ used tube-type diffusive samplers, which were analyzed by HPLC and subsequent fluorescence detection. Moreover, their HPLC system was equipped with a different analytical C18 column ($3.9 \times 300 \text{ mm}$, $60 \text{ \AA}$, $4 \mu\text{m}$). The set-up of the system was not modified to that used by Herrington *et al.*^{61,62} as the DNSH-method was tried to be included into daily routine work according to ISO 16000-3,⁵⁵ which is even more difficult as one analytical run lasts 80 min. Due to these analytical difficulties, analysis of both DNPH-cartridges and DNSH-cartridges by HPLC/MS was not possible so that no results are included in the figures.

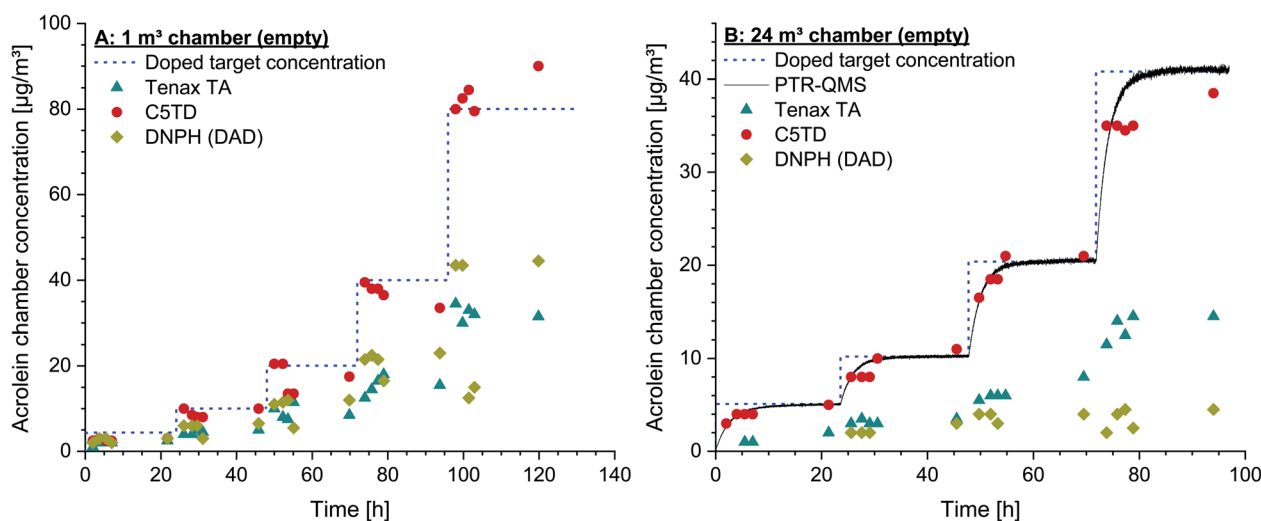


Fig. 3 (A) Acrolein target concentrations in chamber air of an empty 1 m^3 glass test chamber. Concentration range $4.4\text{--}80.1 \mu\text{g m}^{-3}$. Measurement data obtained by active air sampling on Carbograph™ 5TD (C5TD), Tenax® TA and DNPH (analysis by HPLC/DAD). (B) Dosed acrolein target concentrations in chamber air of an empty 24 m^3 stainless steel test chamber. Concentration range $5.1\text{--}40.8 \mu\text{g m}^{-3}$. Measurement data obtained by active air sampling on Carbograph™ 5TD (C5TD), Tenax® TA and DNPH (analysis by HPLC/DAD). Continuous online-monitoring by PTR-QMS.



In order to check the reproducibility of the results, the experiment in the 1 m³ glass chamber was repeated by using a new test gas with the same specifications and from the same manufacturer. The same concentrations could also be set with this new test gas under identical conditions, which indicates a high degree of accuracy of the given specifications. The results of the C5TD analyses were also completely analogous to the data shown in Fig. 3A.

A further test series was carried out in an empty 1 m³ glass chamber to examine the recovery of low acrolein concentrations (0.5–5 µg m⁻³). In Fig. 4 the acrolein concentrations measured after active sampling on Carbograph™ 5TD are plotted *versus* the dosed target concentration. The data were corrected for the background concentration in the empty chamber (duplicate measurements, 0.83 ± 0.02 µg m⁻³). The figure visualizes a linear dependency between the measurement data and the target concentrations. The respective data and standard deviations are provided in the figure captions.

5.2.2 Recovery in loaded emission test chambers. The examinations carried out in the empty test chambers were repeated after loading both the 1 m³ glass chamber and the 24 m³ stainless steel chamber with furnishing and various indoor materials as described above (see Fig. 5A and B). In both chambers, the summed VOC-concentration at the first acrolein steady-state concentration (1 m³: 4.4 µg m⁻³, 24 m³: 5.1 µg m⁻³) was approx. 845 µg m⁻³. A variety of alcohols, aromatic hydrocarbons, aldehydes, terpenes, carboxylic esters and glycol esters were identified in chamber air. Among these, C₄- and C₅-alcohols (1-butanol, 2-methylpropan-1-ol and 1-pentanol), aldehydes (pentanal, nonanal), carboxylic esters (*n*-butyl acetate) and carboxylic acids (propanoic acid) were detected in low to moderate concentrations (5–50 µg m⁻³) in the 24 m³ chamber

and in minor concentrations in the 1 m³ chamber (3–10 µg m⁻³). The Carbograph™ 5TD data were corrected for the background value of acrolein due to the loading of the chambers. The recoveries were in accordance with the findings obtained during the test series in the empty chamber. In the loaded 24 m³ chamber measured data matched the dosed acrolein levels. However, systematic deviations were observed in the loaded 1 m³ chamber at the two highest concentration levels.

5.3 Monitoring of acrolein by PTR-QMS

The experimental design was also used to examine the suitability of the PTR-QMS for indoor air measurements. The mass of the mainly occurring product ion C₃H₅O⁺ is *m/z* 57.033,⁹² while the low-resolution quadrupole only allows the setting of the nominal mass *m/z* 57. Zhao and Zhang⁸¹ published a reaction rate constant of $k = 3.35 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1}$ for acrolein and estimated an uncertainty of ±15%. They used the theoretical approach of Su and Bowers,⁹³ which is based on the polarizability α and the dipole moment μ_D of the respective compound. However, Zhao and Zhang⁸¹ used $\alpha_{cis} = 6.15 \times 10^{-9} \text{ cm}^3$ and $\mu_{D,cis} = 2.636 \text{ D}$ for the *cis* conformation of acrolein, which is significantly more unstable at room temperature than that of the *trans* conformer.^{92,94} Both parameters α and μ_D are significantly higher for the *trans* conformer.^{95,96} Cappellin *et al.*⁹⁷ used the values $\alpha_{cis} = 6.48 \times 10^{-9} \text{ cm}^3$ and $\mu_{D,trans} = 3.41 \text{ D}$ to calculate reaction constants between $3.20 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1}$ and $3.85 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1}$ for different settings of their PTR-TOF-MS.

The PTR-QMS experiments took place in the 24 m³ chamber. If only acrolein was dosed (empty chamber), the theoretical curve could be accurately fitted to the PTR-QMS signal (see Fig. 3B). The calibration was carried out using the steady-state concentrations; the reaction constant determined according to eqn (2) was $k = 2.95 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1}$.

$$[\text{RH}^+] = \frac{A}{k t_R} \frac{[\text{RH}_{\text{cps}}^+]}{[\text{H}_3\text{O}_{\text{cps}}^+]} \quad (2)$$

$[\text{RH}^+]$ is the concentration in µg m⁻³, k is the reaction rate constant, t_R is the reaction time, $[\text{RH}_{\text{cps}}^+]$ and $[\text{H}_3\text{O}_{\text{cps}}^+]$ are the ion signals in counts per second and A is a system specific parameter, which involves the pressure in the drift tube, the temperature, the ion transmission coefficients and factors for the conversion from ppb to µg m⁻³ (see the ESI, Section S3,† for details).

This experimentally determined k for acrolein is at the lower end but still in good agreement with previously reported data. Taking into account the experimental uncertainties, a reaction constant of $3\text{--}4 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1}$ appears to be sufficiently accurate for the standard applications of PTR-QMS.

In the loaded chamber, however, there were clear positive deviations from the respective concentration (not shown). Although there are few substances like butene and its isomers with an $M + 1$ mass *m/z* 57, it can be assumed that fragments of propionic acid, 1-butanol or alkyl residues interfere with the signal for acrolein.^{98,99} For this reason, the PTR-QMS technology

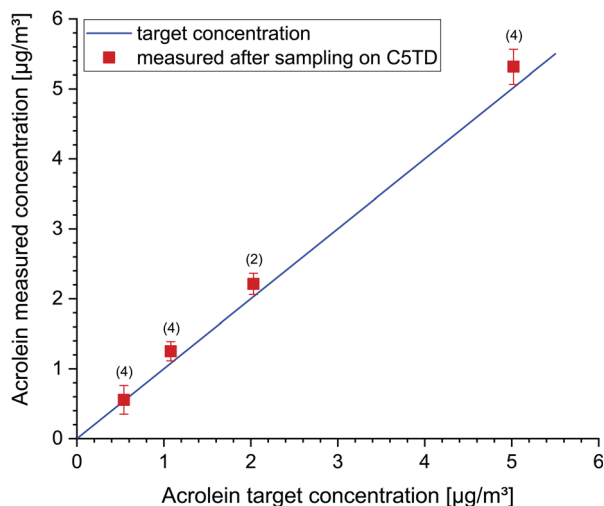


Fig. 4 Acrolein target concentrations in the air of an empty 1 m³ glass test chamber *versus* acrolein concentration measured after active sampling on Carbograph™ 5TD (C5TD). The whiskers show the respective standard deviations, the numbers in brackets indicate the number of repeated measurements. Individual data (target/measured in µg m⁻³): 0.54/0.56 ± 0.21; 1.08/1.25 ± 0.14; 2.03/2.21 ± 0.15; 5.02/5.31 ± 0.25.



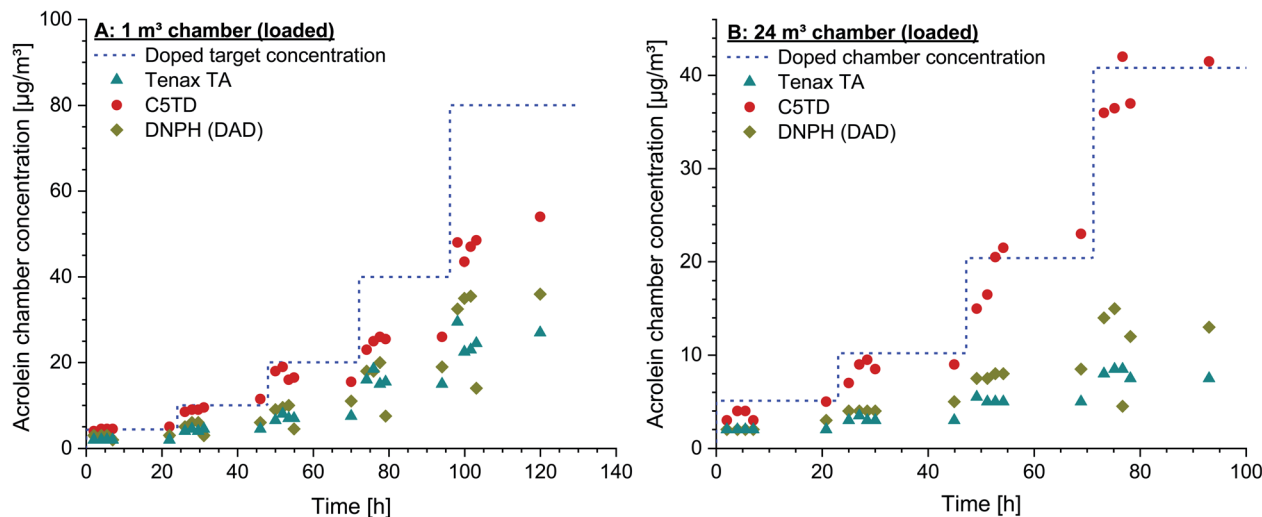


Fig. 5 (A) Acrolein target concentrations in the air of a 1 m³ glass test chamber, loaded with indoor materials. The concentration range is 4.4–80.1 µg m⁻³. Measurement data obtained by active air sampling on Carbograph™ 5TD (C5TD), Tenax® TA and DNPH (analysis by HPLC/DAD). (B) Dosed acrolein target concentrations in chamber air of a 24 m³ stainless steel test chamber, loaded with furniture. Concentration range 5.1–40.8 µg m⁻³. Measurement data are obtained by active air sampling on Carbograph™ 5TD (C5TD), Tenax® TA and DNPH (analysis by HPLC/DAD).

appears to be hardly suitable for monitoring acrolein in indoor air. Therefore, Knighton *et al.*⁶⁶ developed a scrubber system specifically for the detection of acrolein by PTR-QMS. They reported a reaction constant of $k = 4.2 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1}$, which is applicable for their particular instrument. However, a PTR-TOF-MS, as used by Lunderberg *et al.*⁵⁰ is much better suited to correct for interferences and, thus, for detecting acrolein in real indoor environments.

6. Conclusions and outlook

The analytical method described here enables the selective determination of low acrolein concentrations in the air of test chambers. Other VVOCs and VOCs do not interfere, so that the method is also suitable for the analysis of indoor air. No breakthrough occurs with a total sampling volume of 4 L and an air flow rate of 125 mL min⁻¹. Storing of sampled C5TD tubes under laboratory conditions over a time period of 7 days is possible without analyte loss, even at a higher sampling volume of 10 L. A limit of quantitation (LOQ) of 0.4 µg m⁻³ could be achieved for the low concentration range when calibrating with a liquid standard solution for a sampling volume of 4 L and can be lowered to a LOQ of 0.3 µg m⁻³ when sampling 6 L. Even though the liquid analytical grade standard of acrolein was discontinued in Europe, acrolein dissolved in methanol can be used for analytical purposes in future. The DNPH-method according to ISO 16000-3⁵⁵ with sampling on cartridges showed the previously reported problems, making its application for acrolein analysis impossible, even with detection by mass spectrometry. Trapping the target substance on DNSH-coated silica gel cartridges was tested as an alternative approach, but without success under routine conditions. In order to obtain a method which can be integrated into daily routine, an existing HPLC system was used, whose settings

complied with ISO 16000-3.⁵⁵ The preparation of DNSH-cartridges was sophisticated and the restrictions during sampling might be unacceptable when using a heavier weight sorbent bed. Moreover, not all of the three chromatographic peaks (acrolein-DNSH-stereoisomers and acrolein-DNSH-dimer) which are described in the literature^{58,61} could be detected. Due to interfering substances, the PTR-QMS technology at m/z 57 appears to be hardly suitable for monitoring acrolein in indoor environments. PTR-TOF-MS has been shown to be better suited to overcome interference problems.⁵⁰

In summary, collecting airborne acrolein on GCB Carbograph™ 5TD followed by TD-GC/MS has proven to be a promising approach regarding discontinuous sampling and analysis. It offers a fast, easy-to-handle and robust solution for routine analyses and indoor surveys in the relevant concentration range.¹⁹ The recovery of acrolein on Carbograph™ 5TD is good and reproducible with a low standard deviation. As even on freshly packed and conditioned tubes acrolein can be found in trace amounts, it is strongly recommended to document carefully the contamination substances and their amounts detected by the first analysis of a fresh tube. The method is suitable for assessing indoor air quality with regard to the acute and chronic Reference Exposure Limits (RELs) published by OEHHA.⁸ However, assessment using the U.S. EPA Reference Concentration for Inhalation Exposure of $2 \times 10^{-5} \text{ mg m}^{-3}$ ¹⁰⁰ is beyond the scope of this method. In general, such a low detection limit in the range of 0.02 µg m⁻³ can only be achieved with specially designed methods,⁴⁶ which, however, are hardly suitable for routine measurements. This underlines the impracticability of non-accessible guide values as discussed before.^{45,101}



Author contributions

The authors designed and planned the test series presented in this manuscript. All authors read and approved the final manuscript.

Conflicts of interest

The authors declare that they have no conflict of interest.

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