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Use biocidal products (insect sprays and electro-vaporizer) indoor scenarios modeling areas Exposure and exposure ^a, Susanne а Edith Berger-Preiß , Wolfgang Koch Gerling b Heiko Kock a. Klaus E. Appel

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Abstract

Five commercially available insect sprays were applied in a model room. Spraying was performed in accordance with manufacturers' in an overdosed to simulate the instructions and manner in order worst-case conditions or an TM aerosol The Respicon monitoring unforeseeable misuse. In addition, we examined electro-vaporizers. system was (10 seconds) the following applied to determine inhalation exposure. During normal spraying and during 2-3 minutes, 70 to 590 mg/m ³ for the d-phenothrin, exposure concentrations ranged from pyrethroids tetramethrin, cy...uthrin, the pyrethrins. Calculated bioallethrin. inhalable doses were 2-16 mg. A concentration of approximately and ³ (inhalable 850 mg chlorpyrifos/m dose: approximately 20 mg) was determined when "Contra ...y spray" the insect mg/m^{3} Highest exposure concentrations (1100-2100 were measured for piperonyl butoxide (PBO). corresponding to an inhalation intake of 30-60 mg. When simulating worst-case conditions, concentrations exposure mg/m³ and inhalable of 200-3400 doses of 10-210 mg were determined for the various active substances. Highest mg/m³) were measured (4800-8000 for PBO concentrations (inhalable: 290-480 mg).

"Nexa the electro-vaporizer Lotte" plug-in mosquito killer concentrations for d-allethrin were in the of 5–12 mg/m 3 and 0.5–2 mg/m 3 for PBO while "Paral" with the plug-in mosquito killer concentrations of 0.4-5 mg/m ³ for pyrethrins and 1-7 mg/m 3 for PBO were measured

Potential using dermal exposures determined exposure Between 1000 mg active were pads. 80 and substance bioallethrin. (tetramethrin, chlorpyrifos) on the clothing phenothrin, cy...uthrin, pyrethrins, were deposited of the total user. Highest levels (up to 3000 mg) were for PBO. Worst-case surface area of the spray determined uses of the sprays led to 5-9 times higher concentrations.

nearby an operating electro-vaporizer led to a contamination of the clothing (total a 2-hour stav amounts body were 450 mg d-allethrin and 50 mg PBO for "Nexa Lotte" mosquito killer and 80 mg pyrethrins the whole plug-in and 190 mg PBO for "Paral" plug-in mosquito killer).

biomonitoring (E)-trans-chrysanthemum Human data revealed urine concentrations of the metabolite dicarboxylic acid ((E)-trans-CDCA) between 1.7 mg/l and 7.1 mg/l after 5 minutes of exposure to the different sprays. Also the use of electro-vaporizers led to (E)-trans-CDCA concentrations in the urine in the range of 1.0 mg/l to 6.2 mg/l (1-3 hours exposure period).

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The exposure data presented can be used for performing human risk assessment when these biocidal products were applied indoors.

airborne The concentrations of the non-volatile active chemical compounds could be predicted from ...rst principles а deterministic exposure model (SprayExpo). using

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Keywords: Insect sprays; Electro-vaporizers; Pyrethroids; Pyrethrum; Piperonyl butoxide; Chlorpyrifos; Indoor use; Dermal exposure; Inhalation exposure; Exposure modeling

Introduction

of possible health Human risk assessment hazards is characterization based on two basic elements: hazard exposure leading to an integration of and quanti...cation. the exposure ...ndings and toxicological effects of the substance in question order reach a characterizain to tion of possible health risks. Assessment of chemical risks by inhalation as the only exposure route takes consideration the concentrations of substances in the air which can be inhaled. Biocidal products like sprays Or control electro-vaporizers are used to pests in indoor They contain insecticidal substances like pvreareas. pyrethroids, organophosphates, and carbamates. thrum. When they are applied the consumer is exposed to these biocides and various other ingredients because inhalation of the aerosol will take place. In particular a non-foreseeable misuse can lead to high indoor concentrations which may affect human health.

involving conditions, scenarios such exposure risk need address those effects assessment mav to which are single potential acute concern (e.g. toxicity. irritation) and repeated exposures. With respect to single exposures the scenarios presented here are certainly realistic and probably also representative But the data may also be valuable for an exposure estimation. when be applied. biocidal products will repeatedly

aim to quantify The of this study was therefore and exposure characterize the extent and level of to those applied biocidal products which are indoors. It was not comprehensive human risk intended perform а assessment. which would have to take into account the toxicological effects these biocides. Therefore, this of in research project we present data for various exposure scenarios in conjunction with the self-use of biocidal consumer products (sprays, electro-vaporizers) priindoor These data describe indoor vate areas. the air contamination and the exposure of users and occupants with to inhalation potential regard and dermal expometabolite Furthermore. concentrations sures. are determined urine of reported which were in the sprav users or bystanders after exposure.

There that careful consideration is good agreement subnecessarv in the exposure assessment for consumer populations with a particular exposure pattern, and this

with regard to children a foreseeable misuse has to be conditions considered. Therefore. of exposure representing worst-case conditions were also simulated. Furthermore. it in mind biocidal must be borne that numerous substances are persistent in indoor which means that frequent areas. indoors. in an accumulation use may result

For the assessment will consumers risk procedure modeled estimates often relv on exposure which are on product/article speci...cations the content based (e.g. substance in the product/article) and assumptions on intended and other reasonably foreseeable uses experimental obtained Therefore. the data here connection with spraying were compared with results using a deterministic from calculations model on spray the use biocidal dispersion of consumer spravs for private indoor areas

Materials and methods

Model rooms

The were aerosol sprays and electro-vaporizers 16 m², applied in equally sized model rooms (area 40 m³) which volume about were furnished like normal living rooms.

Walls and ceilings were covered with woodchip wall paper, and with textile carpets. In each room there was a cupboard shelves, a sofa, a coffee table. a chair. dining table well as a window and radiator.

During the experiments. ventilator constantly а was in the operated rooms in order to simulate air circulation. temperature, air humidity, and The relative air pressure monitored continuously the rooms. were in

Each sprav product was applied in the middle of the the door and window. The electroroom between vaporizers were plugged into a power outlet in the wall.

Applications

Insect sprays

Five different insect sprays, which are intended for use as room sprays, were used:

"Amisia insect spray", "Blattanex ...y spray", "Blat-

on

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...y spray". Detailed "Contra insect information ingredient active concentrations in Table is given 1. simulating For the correct use. each product was using applied in a separate amount room the recommended the manufacturers ("Amisia insect spray" was sprayed in the for 20 seconds, the other sprays room for 10 seconds.) Then the rooms were kept closed for minutes and after that they were ventilated

In a second experiment an overdose of each spray was applied (simulating worst-case conditions). In this case products were sprayed for 2 minutes and the rooms were not ventilated

Electro-vaporizers

minutes.

Two different electro-vaporizers were used in separate model for 6 hours day rooms per according to the manufacturers instructions.

pyrethrum "Paral", containing 20 ma extract and 40 mg PBO per pad and "Nexa containing 4.6% Lotte". d-allethrin 4.6% **PBO** per pad (manufacturer declaration) or about 38 mg of each substance per pad.

Spray characterization

The sprays were characterized with respect to the particle diameter of the released droplets. Measurements diffraction carried out with a laser spectrometer were (HELOS (1269),Sympatec GmbH). The size distribucalculated using tion was а mathematical inversion method. The cumulative volume distribution Q_3 as a in Fig. of the droplet 1. Table function size is shown percentiles for the distribution of the presents the particles in the sprays. aerosol

Inhalation exposure

(0.06 g/100 g)

During the spraying procedure (see above), the inhalation exposure of the spray user was recorded with exposure the personal monitor Respicon 3-F aerosol (Hund, Wetzlar, Germany), which was developed at the Fraunhofer Institute of Toxicology and Experimental the Medicine (Koch et al., 1999). This device allows simultaneous sampling and on-line concentration monitoring of the three health-relevant particle size fractions according to the CEN convention 481: the respirable (C $_{Th}$), and the inhalable fractions (C_{l}) (C_R) , the thoracic (CEN, 1992, 1993). Using glass ...lters three ...ber (37 mm), in a special case followed by polyurethane foam (PUF) plugs (sampling chlorpyrifos), different aerosol fractions were separated. Online detection was performed by three built-in light-scattering photometers.

according the manufacturers' During application instructions personal measurements were done during the spraying operation and 2-3 minutes for up to thereafter.

When spraying under worst-case conditions, measure-TM with ments the personal Respicon were carried of a total of 5 minutes. addition, а period In over TM was positioned in the stationary Respicon the room, and sampling was done over a period minutes.

analyzed **Filters** below). were extracted and Results concentrations in are aiven as exposure ma ³ inhaled active substance/m air and as doses (respiratory

100

75

[%] 50

25

0 10 100 1000 aerosol particle diameter [µm]

> Paral **Amisia Blattanex** Blattanex new Contra insect

Fig. 1. Spray characterization using laser diffraction metry. Cumulative volume distribution Q₃ as a function of the aerosol particle diameter.

Overview of active substance concentrations in the sprays used.

(0.2 g/100 g)

| "Amisia insect spray" | "Blattanexy spray" | "Blattanexy spray" (new) | "Paral insect spray" | "Contra insecty spray" |
|-----------------------|-----------------------|-----------------------------|----------------------|------------------------|
| d-Tetramethrin | Cyuthrin | d-Tetramethrin | Pyrethrum extract | Chlorpyriphos |
| (0.04 g/100 g) | (0.04 g/100 g) | (0.15 g/100 g) | (0.25 g/100 g) | (0.5 g/100 g) |
| d-Phenothrin | Tetramethrin | d-Phenothrin | PBO (1 g/100 g) | Bioallethrin |

(0.15 g/100 g)

(0.15 g/100 g)

Table 2. Percentiles (as mm) and geometric standard deviations (distribution range) of the distributions given in Fig. 1 (sg is the geometric standard deviation of the droplet size distribution).

| Product | ^x 16 | ^x 50 | ^x 84 | sg |
|------------------------|-----------------|-----------------|-----------------|------------|
| Amisia insect spray | 13.4570.19 | 27.9470,73 | 44.0071.57 | 1.8170.035 |
| Paral insect spray | 14.6170.31 | 28.4170.44 | 48.7978.47 | 1.8370.160 |
| Blattanexy spray | 8.5170.15 | 18.1470.35 | 28.6770.74 | 1.8470.029 |
| Blattanexy spray (new) | 15.7570.28 | 33.0270.76 | 55.0872.01 | 1.8770.038 |
| Contra insecty spray | 2.6770.13 | 12.0770.33 | 19.9870.52 | 2.7270.074 |

minute volume: 12 l/min) related to the duration of exposure (see Table 3).

Potential dermal exposure

Exposure pads made from ...lter paper (Macherev-Nagel, type 1640 m, size 10 10 cm, back covered with aluminum foil) were positioned on the sprayer's overall at the following positions (1/4 8% of total body surface): head, chest, upper left arm, upper right arm, left left thigh, forearm, right forearm, right thigh, left shin, to ECB. 2006 and right shin according 1998, DIN CEN, OECD. 1997.

After respective the spraying procedure and a short the pads (2-3)minutes stay in room the were removed after 5 minutes the after correct use. and in case of active overdose). extracted. and substance concentrations were measured (see below).

substance amount per pad was related to the area of the respective part of the body using standard factors and the resulting amounts were added up to determine The the potential total dermal body dose results are duration οf given as dermal doses related to the exposure.

Extraction and analysis of samples

The ...Iters, PUF plugs, or the cut exposure pads were using (10 minutes) extracted three times n-hexane or ethyl acetate in an ultrasonic bath. The extracts were combined, reduced in volume using N 2 (TurboVap II), glass wool. and ...ltered through silanized Final volumes of 1 or 5 ml were adjusted

Pyrethrins were determined using a gas chromatograph with an electron capture detector (GC/ECD). GC analysis was carried out employing an instrument from (5890 II), a DB5.625 Hewlett Packard series using 10–12 m, i. d. 0.25 mm, $d_f \frac{1}{4}$ 0.25 mm), a column (length: gap, and helium as carrier gas. The deactivated retention iniected on-column. The following temperasample was ture program was used: 60 1C (1 minutes), 10 1C/min up to 270 _{1C} (3 minutes). Analytical to 180 1C. 6 1C/min details were already described previously (Berger-Preiß

The individual spray ingredients (phenothrin, tetra-PBO) methrin, cy...uthrin, chlorpyrifos, bioallethrin, were measured using a gas chromatograph which was with a mass-selective detector (MSD/EI). GC equipped analysis was carried out employing an instrument from Agilent **Technologies** (6890N). using an HP-5MS d_f ¼ 0.25 mm) and column (30 m length, i.d.: 0.25 mm, helium as carrier gas (constant ...ow, 1.4 ml/min). The 50 1C following temperature program was used: up to $280 \ 1C$ (5 The (0 minutes), 10 1C/min minutes). sample split/splitless injected in mode (injector 250 _{1C)} and detected temperature: using an Agilent 5975 detector. mass-selective

MSD The ...tted with a quadrupole ...lter mass was used in electron impact (EI) mode. FI mass spectra were obtained 70 eV. MSD temperatures were as follows: 280 1C: 1C; quadrupole: transfer line: ion source: 230 150 1C. The MSD was run in selected ion monitoring The (SIM) mode. following target and quali...er ions (m/z)were monitored: phenothrin (123/183),tetramethrin (164/123),cy...uthrin (163/215)chlorpyrifos (197/314),bioallethrin (123/79),and PBO (176/119).

Quanti...cation was done by means of a characteristic target mass using external calibration curves for the individual compounds.

Biomonitoring

Metabolite concentrations in the urine of the spray and second person (bystander) were determined after 10 seconds of spraying and staying in the for room 5 minutes. Twenty-four-hour urine was (E)-cis/trans-chrysanthemum collected. The metabolites acid ((E)-cis/trans-CDCA), dicarboxylic cis/trans-3-(2,2--2,2-dimethylcyclopropane dichlorovinyl) carboxylic (cis/trans-DCCA), acid and 4-...uoro-3-phenoxybenzoic acid (4-F-3-PBA) were analyzed usina gas chromatography-mass spectrometry (GC/MS (NCI)) (El...ein et al., 2003; Barr et al., 2007).

Exposure modeling

The model SprayExpo was used for exposure model—
ing. The model calculates the airborne concentrations of

Table3. Description findividuals prayscenario and results of inhalation exposur exposur concentration and calculate dose for the inhalable thoracic, and respirable fractions of active substance an earn values of 2 spray application in each case and potential total dermal exposures.

| Insectspray | Spraytime [s] | Active | Time ^a | Averageexp | osurœoncentra | tion[mga.s./m] | Inhalabled | ose[mga.s./a | pplication) | Dermaldos € | |
|--------------------|---------------|----------------|-------------------|------------|---------------|----------------|------------|--------------|-------------|----------------------|-----------------------|
| scenari d D | Amount[g] | substanca.s. | [min] | Inhalable | Thoracic | Respirable | Inhalable | Thoracic | Respirable | [mga.s./application] | |
| | Amount[g] | | [] | IIIIaabic | THOTAGIC | Козрігавіс | milalabic | THOTAGIC | Кезрпавіс | [mga.s./application] | |
| Amisia | | | | | | | | | | | |
| S1(A) | 20 | d-Tetramethrin | 3 | 67.7 | 49.4 | 20.1 | 2.4 | 1.8 | 0.7 | 80 | |
| | 23.8 | d-Phenothrin | | 85.6 | 66.7 | 26.3 | 3.1 | 2.4 | 0.9 | 100 | |
| S2(A) | 120 | d-Tetramethrin | 5 | 231.2 | 163.7 | 93.6 | 13.9 | 9.8 | 5.6 | 267 | E. |
| | 114.4 | d-Phenothrin | | 306.2 | 229.7 | 131.2 | 18.4 | 13.8 | 7.9 | 360 | Berger-Pr |
| S3(A) | 120 | d-Tetramethrin | 60 | 114.1 | 87.8 | 49.4 | 82.2 | 63.2 | 35.6 | NA ^d | |
| ` , | 111.4 | d-Phenothrin | | 176.8 | 136.8 | 77.3 | 127.3 | 98.5 | 55.6 | NA | |
| Blattanex | | | | | | | | | | | et al. ARTICI / |
| S1(B) | 10 | Cyuthrin | 2.2 | 120.9 | 82.6 | 34.2 | 3.2 | 2.2 | 0.9 | 114 | Int. |
| 01(2) | 11.4 | Tetramethrin | | 587.1 | 397.7 | 166.6 | 15.5 | 10.5 | 4.4 | 334 | J. |
| | | РВО | | 2122 | 1425 | 598.9 | 56.0 | 37.6 | 15.8 | 1138 | Hyg. |
| | | | | | | | | | | | En vin ion. |
| S2(B) | 120 | Cyuthrin | 5 | 424.4 | 329.0 | 172.6 | 25.5 | 19.7 | 10.4 | 712 | PRESS |
| | 110.3 | Tetramethrin | | 3223 | 2342 | 1212 | 193.4 | 140.5 | 72.7 | 1973 | Health |
| | | PBO | | 8014 | 5629 | 2952 | 480.8 | 337.7 | 177.1 | 6328 | Tleatti |
| S3(B) | 120 | Cyuthrin | 60 | 267.4 | 245.8 | 177.3 | 192.5 | 177.0 | 127.7 | NA | 212 |
| | 110.3 | Tetramethrin | | 1992 | 1681 | 1108 | 1434 | 1210 | 798.0 | NA | (2009) |
| | | РВО | | 4518 | 3614 | 2305 | 3253 | 2602 | 1659 | NA | |
| | | | | | | | | | | | 505–518 |
| Blattanex(ne | ew) | | | | | | | | | | |
| S1(BN) | 10 | d-Phenothrin | 3 | 144.5 | 85.6 | 25.7 | 5.2 | 3.1 | 0.9 | 722 | |
| | 16.3 | d-Tetramethrin | | 147.4 | 87.8 | 28.8 | 5.3 | 3.2 | 1.0 | 752 | |
| S2(BN) | 120 | d-Phenothrin | 5 | 1416 | 815.0 | 265.0 | 84.9 | 48.9 | 15.9 | 4017 | |
| | 189.2 | d-Tetramethrin | | 1935 | 1104 | 357.9 | 116.1 | 66.2 | 21.5 | 3931 | |
| S3(BN) | 120 | d-Phenothrin | 60 | 538.3 | 403.0 | 181.4 | 387.6 | 290.1 | 130.6 | NA | |
| - | 189.2 | d-Tetramethrin | | 695.6 | 520.0 | 236.5 | 500.8 | 374.4 | 170.3 | NA | |
| | | | | | | | | | | | |

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Table 3. (continue)d

| Insectspray scenaridD | Spraytime [s] | Active substance.s. | Time ^a | Averageexpo | sur c oncentral | tion[mga.s./m] | Inhalabled | dose[mga.s./ap | pplication) | Dermaldose | |
|-----------------------|---------------|---------------------|-------------------|-------------|------------------------|----------------|------------|----------------|-------------|----------------------|----------------------|
| | Amount[g] | | [min] | Inhalable | Thoracic | Respirable | Inhalable | Thoracic | Respirable | [mga.s./application] | |
| Paral | | | | | | | | | | | |
| S1(P) | 10 | Pyrethrins | 3 | 344.7 | 208.3 | 99.4 | 12.4 | 7.5 | 3.6 | 1086 | |
| | 19.8 | РВО | | 1467 | 917.8 | 381.0 | 52.8 | 33.0 | 13.7 | 2969 | |
| S2(P) | 120 | Pyrethrins | 5 | 2616 | 1573 | 736.5 | 156.9 | 94.4 | 44.2 | 4736 | |
| - (·) | 162.4 | PBO | • | 5920 | 3343 | 1594 | 355.2 | 200.6 | 95.6 | 20895 | E. |
| | | | | | | | - | , - | | | Berger-Pr |
| S3(P) | 120 | Pyrethrins | 60 | 753.3 | 541.3 | 305.8 | 542.4 | 389.7 | 220.2 | NA | |
| | 162.4 | РВО | | 2466 | 1658 | 802.6 | 1775 | 1193 | 577.9 | NA | |
| | | | | | | | | | | | et |
| Contrainsec | ot | | | | | | | | | | ^{al.} ARTIC |
| S1(C) | 10 | Chlorpyrifos | 2.2 | 845.5 | 661.4 | 397.3 | 22.3 | 17.5 | 10.5 | 249 | / Int. |
| | 9.5 | Bioallethrin | | 426.9 | 303.8 | 178.6 | 11.3 | 8.0 | 4.7 | 124 | J. |
| | | РВО | | 1167 | 944.0 | 569.9 | 30.8 | 24.9 | 15.0 | 264 | Hyg. |
| | | | | | | | | | | | Env in on. |
| S2(C) | 120 | Chlorpyrifos | 5 | 3444 | 2907 | 1723 | 206.6 | 174.4 | 103.4 | 1817 | |
| | 89.2 | Bioallethrin | | 1565 | 1329 | 793.3 | 93.9 | 79.7 | 47.6 | 571 | PRESS |
| | | PBO | | 4831 | 4021 | 2356 | 289.9 | 241.2 | 141.4 | 1792 | Health |
| /0\ | | -·· <i>w</i> | 33 | | | | | | | | |
| S3(C) | 120 | Chlorpyrifos | 60 | 2036 | 1750 | 1236 | 1466 | 1260 | 890.1 | NA | 212 |
| | 91.6 | Bioallethrin | | 888.6 | 779.2 | 555.8 | 639.8 | 561.0 | 400.1 | NA | (2009) |
| | | PBO | | 3578 | 3159 | 2315 | 2576 | 2274 | 1667 | NA | |

 $^{{}^{}a}\!\text{Time} of applicationairs amplint \textit{im} eor duration of exposu \textit{(e} prayin \textit{g} roceduzen dminute \textit{th} ereafter).$

d. . . .

 $^{^{\}text{C}}$ mgactiv \mathbf{s} ubstan $\mathbf{p}\mathbf{e}$ rapplicatio \mathbf{n} yhol \mathbf{e} odywithou \mathbf{h} ands.

mg.ouv@abotanperappiloationignol@odywithouthand

other meaningful size fractions of aerosols generated The model is a short-term during the spraying processes exposure model covering time scales typical for the release process. Long-term emissions of vapors from walls and other surfaces are not included

It is assumed that the sprayed product is composed of a non-volatile substance active dissolved in а solvent with known volatility. The model based is on taking the released droplets simulation of motion Ωf settling, mixing into gravitational turbulent account with the surrounding and droplet evaporation. In air. the model continuous spatial release patterns be can distribution need simulated. No arti...cial volumes to be the calculation of inhaled and dermal de...ned. the spatial distribution of the concentration is doses the explicitly taken into account.

The main parameters are: released droplet input the spectrum. release rate, concentration of active substance. spatial and temporal pattern of the release (surface spraying ceiling, wall; process against ...oor. of the spraying, pressure liauid. size of room y.), vapor ventilation of the the room. and rate. The path spraver be explicitly included into the model. can

Results

Inhalation exposure

In order to describe the inhalation exposure three fractions the inhalable determined: the aerosol were fraction mass fraction inhaled particles respirable (the of C_Rp5 penetrating to non-cilicated airwavs (alveoli), thoracic fraction ٥f inhaled mm), the fraction (the mass p10 C_{Th} larynx, particles penetrating beyond the mm) inhalable fraction (the fraction and the mass Ωf total airborne particles that are inhaled through the mouth C₁) according CEN 481, 1993. The data and nose. to shown Table 3 are given as exposure concentrations ³ (average active substance (a.s.)/m concentration in mg (2-5)during spraying and the following time minutes 60 minutes. as described 4 of Table 3)) and as in column inhalable active substance/application doses in mq of exposure). (duration

"Amisia During correct the use Ωf insect spray" 68 mg/m 3 concentrations of and 86 mg/m ³ exposure d-phenothrin, determined d-tetramethrin were for and and the inhalable doses were 2-3 mg of active sub-When simulating worst-case conditions stances. the active substance concentrations were 231 mg/m ³ 3

5 minutes: 18 mg of 14 and each substance) However. in the hour) inhalable after a longer stay room (one doses higher. the active substances are 6-7 times

On average, 36% of the active substance from the inhalable aerosol particles were found in the respirable fraction.

When "Blattanex spraying the ...y spray" (10 seconds to the exposure according manufacturer's instructions). PBO cy...uthrin, and concentrations tetramethrin. for mg/m 3. $587 \text{ mg/m} ^{3}$ 2122 mg/m were 121 and respectively. In the Ωf use Ωf the sprav and case correct short-term in the approximately 3 and 16 mg stav room. (cy...uthrin, 56 mg PBO inhalable. tetramethrin) were and

The scenario (2 minutes worst-case sprav application of spraying and 3 minutes thereafter) led to 4-5 times higher exposures. In the ...rst 5 minutes, 26 mg cy...uthrin, 193 mg tetramethrin, and 481 mg PBO were inhalable. In the case of 1-hour stay, active substance intake is correspondingly higher.

For "Blattanex ...y spray", on average 33% of the active substance-carrying particles were in the respirable fraction.

a diffe-Furthermore. a "Blattanex spray" with ...y (d-phenothrin rent active substance composition and d-tetramethrin) was studied. Exposure concentrations 145 mg/m ³ 3 were between and 147 mg/m determined for the active substances during correct use of the spray. conjunction with this procedure and during a stay of 3 minutes in the room, approximately 5 mg of each active substance inhalable were

After worst-case application of the the sprav. 1416 mg/m ³ and were 1935 mg/m (shortconcentrations time measurement). Overdosing the spray intake of active substance doses between 85-116 mg Active conjunction with a 5-minute stay in the room higher when substance doses were approximately 4 times one hour. the spray user stayed in the room for

On average, 19% of the active substance-carrying particles were in the respirable fraction.

In the "Paral" the case Ωf consumer expospray 3 concentrations of 345 mg/m (pyrethrins) and sure 3 1467 mg/m (PBO) were determined for correct use during the spraying process the 3 minutes. and next inhalable doses during this period were calculated to be 12 mg pyrethrins and 53 mg synergist.

conditions Worst-case led to far higher exposure concentrations Inhalable doses from 157 mg to increased mg (PBO) 542 mg (pyrethrins) 355 mg to 1775 and from during an exposure time from 5 minutes to one hour.

"Paral In the of the insect spray", case on average 28% of the active substance-carrying particles were respirable fraction. the

When "Contra ...y spray" (10 seconds applying insect during 2.2 minutes), active spraving and the next of 846 mg/m ³ substance concentrations of (chlorpyrifos),

inhalable measured. doses during this period substances calculated to be were mg. During the worst-case use of the times exposure concentrations could measured be which decreased only slightly during one hour. Inhalable doses of the active substance were 94-290 mg during the 5 minutes and seven times higher when staying in the room for one hour.

the "Contra of spray", In the case insect ...у on 48% of the active substance-carrying particles average in the respirable fraction were

Fig. 2 To summarize. all results are presented in Fig. inhalconcentrations) 3 (calculated (exposure and 10 able doses). During correct spraying (normally and the 2-3 minutes, average seconds of spraving) next exposure concentrations of the pyrethroids tetramethrin, d-phenothrin, cy...uthrin, bioallethrin, and the mg/m 3. pyrethrins were between 70-590 Calculated inhalable active substance doses ranged from 2-16 mg The exposure organophosphate concentration 850 mg/m ³ (inhalable chlorpyriphos approximately was dose: 20 mg). Highest concentrations were measured for mg/m^{3}), the synergist PBO (1100-2100 leading to an inhalation of 30-60 ma. intake

During sprav applications under worst-case condisubstance concentrations measured for tions active mg/m 3 minutes ranged from 200-3400 (inhalable doses: 10-210 mg). The highest values (concentrations: mg/m 3, 4800-8000 inhalable doses: 290-480 mg) were obtained PBO. In the case of longer stavs approximately overdosed rooms hour) up

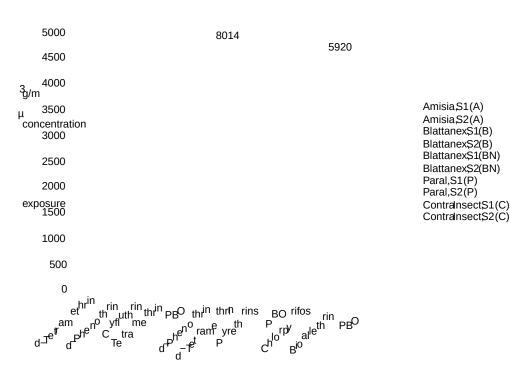
1500 mg active substance and 3250 mg synergist may be inhaled.

As an example, Fig. 4 represents the behavior of spray particles the air during release of "Blattanex ...у spray" (worst-case conditions). As shown by the timeresolved concentration curves recorded with the Respicon TM. concentrations of nonor semivolatile comof the (d-phenothrin, d-tetramethrin) pounds aerosol up to the increased from the beginning of spraying (120 seconds) Thereafter. due to air exchange of release particle deposition on surfaces. the concentrations and at different of the individual particle fractions decreased rates.

Summarizing all results. it concluded that can be between 19-48% of the active substance-carrying particles were in the respirable fraction and 59-80% in thoracic fraction. the

Measurements were also performed during use of biocide-containing electro-vaporizers Measurements which determine inhalation exposure carried out to TM when the aerosol monitorina system Respicon using because could not be reliably evaluated. active substance concentrations too low. For this reason particulate were collected small ...lter instrument matter the was by GS050 equipped with а glass ...ber ...lter. **Applying** this method indoor concentrations could be measured. air

During operation of the "Nexa Lotte" pluq-in mosquito killer in the above mentioned model room several days, indoor air concentrations ranged from 3 (PBO). 3 (d-allethrin) mg/m mg/m 0.5 - 2When 5-12 using the "Paral" plug-in mosquito killer, pyrethrin



active substance

Fig. 2. Exposure concentrations of active substances during application of different consumer sprays after correct use and under

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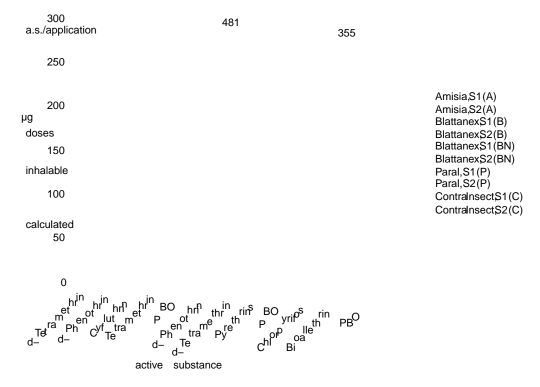
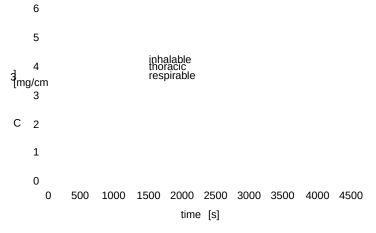


Fig. 3. Calculated inhalable doses of active substances per application during use of different consumer sprays after correct use and under worst-case conditions in model rooms (for a scenario description, see Table 3).



Time-resolved concentration curves (respirable, thorand inhalable fractions) for nonor semivolatile compounds (d-phenothrin, d-tetramethrin) measured with Respicon TM during worst-case application of the "Blattanex ...y spray" (new).

concentrations were between 0.4–5 mg/m 3 and PBO concentrations between 1–7 mg/m 3 .

Dermal exposure

The potential was determined during dermal exposure during under correct use of the sprays and spraving After user remained worst-case conditions spraying the period 3). in the room for a short of time (see Table **Table** 3 and Fig. 5 report the results for potential total

of the sprays, between 80 and 1086 mg active use substance/application (d-tetramethrin, d-phenothrin, cy...uthrin, bioallethrin, pyrethrins, chlorpyrifos) were deposited on the clothing (whole body) the sprayer. Highest 3000 mg/application) deter-(up to were mined the synergist PBO. substance concentrations, which considerably in the different vary spray formulations, had an important in...uence on the level contamination.

The potential total dermal exposure was far higher condiwhen spraying was performed under worst-case dermal tions. In most cases the calculated potential of single doses active substances were about 5-7 times higher application (exception: than during correct "Amisia 3 times). insect spray",

Regarding individual body the contamination of parts, the results showed that upper body were parts In contaminated more strongly than lower ones. most cases. contamination decreased in the order: arms 4back or chest, head4thighsXforearms4shins. Fig. 6, as an example, gives the distribution of the active substance tetramethrin during spraying of "Blattanex spray".

The deterpotential dermal exposure was also after operation of mined electro-vaporizers in the the model room. Α 2-hour stay in close the room to potential total vaporizer led to dermal doses of approximately 450 mg d-allethrin 50 mg PBO ("Nexa and killer), approximately Lotte" plug-in mosauito and of 80 mg pyrethrins and 190 mg **PBO** ("Paral" plug-in

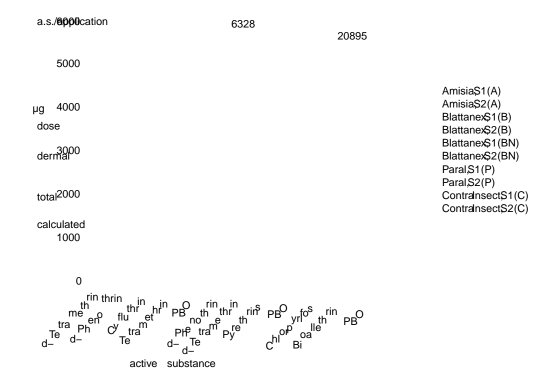


Fig. 5. Potential dermal exposure (whole body) to active substances during application of different consumer sprays after correct use and under worst-case conditions in model rooms (for a scenario description, see Table 3).

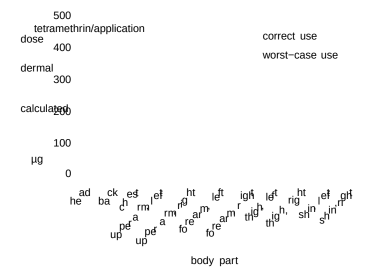


Fig. 6. Contamination of individual body parts after application of "Blattanex ...y spray" during correct use and under worst–case conditions.

Biomonitoring

Metabolite determined the concentrations were in urine after of the spray and/or bystanders users correct "Amisia spray", "Blattanex use of insect ...у spray", "Paral "Blattanex spray" (new). and insect sprav". The contained in active substances these consumer Table After sprays shown in 1. exposure are pyrethrum tetramethrin, phenothrin, allethrin, and

((E)-cis/trans-CDCA), dicarboxylic acid is suitable mum as a biomarker for internal exposure. Furthermore, cis/trans-3-(2,2-dichlorovinyl)-2,2-dimethylcyclopropane carboxylic acid (cis/trans-DCCA) and 4-...uoro-3phenoxybenzoic acid (4-F-3-PBA) should be detectable in urine cy...uthrin exposure.

The results regarding metabolite concentrations in 24-hour urine are given in Table 4. After exposure to the different concentrations of (E)-trans-CDCA in (E)-cis-CDCA and 7.1 mg/l. urine 1.7 was samples. not in anv the urine Concentrations in the urine of the were slightly lower than in the urine of bystanders. The metabolites cis/trans-DCCA and 4-F-3-PRA of detection. were below the limit This may be cy...uthrin to the fact that the in due concentration product "Blattanex" the is ...ve times lower than the tetramethrin concentration.

Furthermore, metabolite concentrations deterwere mined urine of consumers after operating electrovaporizers which contain d-allethrin and pyrethrum. (E)-trans-CDCA concentrations were between 1.0 mg/l and 6.2 mg/l, depending on the period spent in the room, which was between 1-3 hours

Exposure modeling

Results modeling for Blattanex release of the exposure pattern shown Fia 7. The concentration is are characterized an increase in concentration during the

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8

| Table | 4. | С | Concentrations of | | of | (E)-trans-chrysanthemum | | | dicar- | |
|---------|----|-----|-------------------|-------|----|-------------------------|-------|-----|-------------|--|
| boxylic | a | cid | in | urine | of | spray | users | and | bystanders. | |

| Insect spray | | 24-h urine volume [l] | (E)-trans-CDCA concentration [mg/l] |
|-----------------|----------------------|--------------------------|-------------------------------------|
| Amisia | Sprayer Bystander | 1.3 1.2 | 1.7 2.3 |
| Blattanex | Sprayer | 1.3 | 2.4 |
| Blattanex | Sprayer | 1.5 | 4.4 |
| (new) | Bystander | 1.3 | 7.1 |
| Paral | Sprayer | 1.2 | 4.7 |
| | | | |

mainly due to air exchange and particle settling. 4) shows comparison with the experimental curves (Fig. reasonable agreement. Further comparisons were made basis of the average concentrations during correct in Fig. use of the sprays. These are shown 8. Except the largest droplets, agreement the with between the ab initio model calculations and the experimental results is quite reasonable.

Discussion

Applying household insecticide products raises several important considerations concerning safety. These related the dif...culty of controlling these products. Especially the extent and duration user's potential exposure to the active ingredients can hardly be controlled. Insecticidal substances vaporizers tained in are released slowly and have potential low-level particular for long-term exposure with (minimum 6 hours per day). On the other hand. formulations shortregard to spray cans and aerosol high-level exposure be of more But term may concern. it should with products also these be taken into account that spraying is frequently performed over a longer time period (e.g. summer time). The duration extent and the exposure are therefore also highly product-speci...c. Exposure concentrations have be safe, when such are applied household products by the consumer. As for assessment, concentration-dependent as well risk both time-related effects as concentration have to be considered. Knowledge about the extent time period and is of paramount importance. It should exposure he that it was not the intention of this study pointed out to perform comprehensive risk evaluations. Therefore anv toxicological data referring the active substances to Risk should tested were not considered. assessment be a separate task for each of the insecticide for which

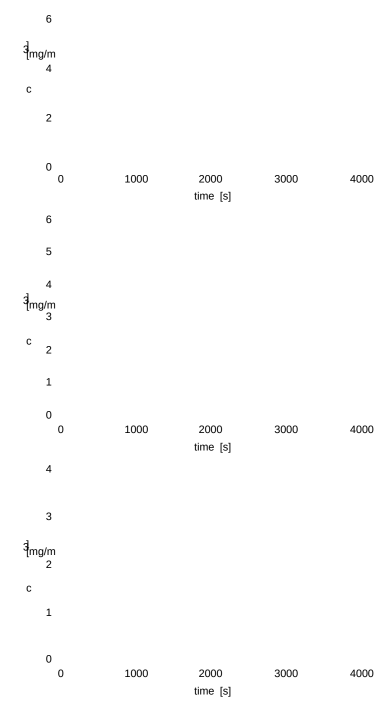


Fig. 7. Concentration curves of the respirable, thoracic, and inhalable fractions of the active substances released during worst-case use of the "Blattanex ...y spray" (new), as calculated in the model based on the measured spray droplet distribution and the release rate.

With inhalation there regard to exposure are only reports literature in the about indoor air concentrafew tions after using aerosol sprays. For instance. spraving of two different for 10-15 seconds in a aerosol sprays 50-m 3 room led indoor air concentrations to average (during spraying and within the following 15 minutes)

5 inhalable 4 thoracic experiment respirable 3 / model 1 0 5 10 15 20 25 30 35 x50 [µm]

8. Ab initio model prediction and measurement results for the concentration of respirable, and inhalable thoracic. fractions during the correct use of insect sprays. X50 is the median particle diameter of the spray droplet distribution.

125 mg/m ³ for pyrethrins, and 55 mg/m ³ for tetramethrin 1991a). (Class and Kintrup, The authors calculated an inhalation instance, tetramethrin for intake of. 70 mg for within the ...rst hour of spraying. Some studies about from aerosols in indoor are also available use of areas Matoba. use of biocidal room sprays (10 seconds) (23 m^3) in room led to average concentrations 148 mg/m 3 (20 minutes including spraying) in the air of 20.5 mg/m 3 for d-tetramethrin d-resmethrin, and and using respectively (Matoba et al., 1998a). When treatment (2.5)minutes sprayaerosol spray for surface 752 mg/m ³ for ing) average indoor concentrations of $1040 \text{ mg/m}^3 \text{ for}$ d-phenothrin and d-tetramethrin were determined (Matoba et al., 1998b). **Further** data were reported, e.g., when using the insectides chlorpyrifos and permethrin surface in various pump sprays for ...ea control in a private (Koehler Moye, 1995). area and indoor concentrations measured (during Average application to 2 hours in the spray and up thereafter) 21-52 mg/m ³ for chlorpyrifos apartments and were 32-54 mg/m ³ for permethrin.

Durina spraying various in an aircraft cabin aerosols the purpose of disinsection (Berger-Preiß et al., 2006), concentrations of d-phenothrin (average concentrations during spraying and in the next 20 minutes) to be 234-1313 mg/m³ (no air exchange were reported mg/m ³ (air and 116-348 conditionthe passenger cabin) operating, air exchange (1/h): approximately 22). ing Inhalable d-phenothrin relevant the doses Ωf for e.g. 29-235 were calculated between mg spray user to he 100 g The with (when using spray). of а spray use pyrethrum and PBO for aircraft disinsection when the conditioning operating (Berger-Preiß al.. air was et 2004) led to inhalable doses of 3-12 mg pyrethrins and mg PBO the spray user who spent 20 minutes for in the passenger cabin (calculations based on the use of

conclusion, biocide concentrations our data on after and inhalation exposure applying household comparable insecticide with those sprays are well after which have been reported bv other authors using insecticidal indoors. The exposure concentrations sprays for spraying determined according to the manufacturers' instructions during the period of the actual spraying and shortly thereafter (2 - 3)minutes) are also on the same scale like e.g. the concentrations (average values 20 obtained over minutes) for sprays used for aircraft disinsection (air conditioning switched As on). expected, simulating worst-case conditions the concentrations of biocidal substances in the indoor air are far higher.

Furthermore concentrations of active substances in the of active air are strongly dependent by level substances in the different formulations. sprav

the private sector electro-vaporizers In are frequently applied for controlling ...ying insects. After operating authors reported electro-vaporizers. other have 2-5 mg/m 3 for air of allethrin indoor concentrations 4 mg/m ³ (Class. 1991b) and approximately coniuncwith the staging of simulation experiments (Matoba 1994). The indoor air concentrations reported by the authors during operation of the "Nexa Lotte' and "Paral" the plug-in mosquito killer were in the same "Paral" range. In case of the vaporizer the concentration ratio of pyrethrins and PBO in the air was similar as active substance composition the expected from the Ωf the "Nexa Lotte" pad. In for vaporizer the contrast PRO the d-allethrin level. level lower than lt was seems that **PBO** completely vaporized during was not the Summarizing the results obtained operating process. for electro-vaporizers. one can conclude that the active substance concentrations in the air are far lower than during biocide spraying. For this reason, measurements

the Respicon aerosol monitoring system, could not be reliably evaluated.

Only a very few publications have presented results on potential dermal exposure during application of the aerosol sprays. Some studies were performed by our group When usina different biocidal aerosol sprays methods aircraft disinsection. and for values of 1700-4100 for d-phenothrin, 200-830 ma ma for pyre-2140-8840 PBO mg for could determined thrins. and be for the potential total dermal dose of the spraying person (using 100 g of the product) (Berger-Preiß et al.. 2004. 2006).

Other available information dermal on exposure during spray use by consumers are based on tracer experiments and model estimates (Popendorf and Selim. Roff. 1996: Roff Baldwin. 1995: **Thompson** and and 1997). with Any comparison the results reported is however dif...cult. study

From our data it can be concluded that the concentrations of the active substances, which vary the considerably in various spray formulations. have а impact level of dermal As decisive the contamination. on expected worst-case applications sprays lead to far of higher dermal exposure levels. Dermal exposure levels after normal use of the sprays are comparable with those after application of electro-vaporizers, а prolonged the room is assumed. stay in

With regard to human biomonitoring have our investigations on chrysanthemum dicarfocused (CDCA), boxylic acid which is а metabolite of pyrethrins, tetramethrin, phenothrin, and allethrin. Also literature metabolite in the some data on this have been reported. Exposure of а pest controller to (S)-bioallethrin led а **CDCA** concentration in the to urine 204 mg/l 1999) After (Leng et al.. of spraying а d-phenothrin-containing aerosol sprav for aircraft disconcentrations of 0.6 insection. **CDCA** and 1.2 mg/l were found in the urine of persons who had entered cabins immediately after spraying (Berger-Preiß aircraft et al.. 2006).

(E)-trans-CDCA another study concentrations in 00.05 of to 54 mg/l (mean: 1.1 mg/l7 the range up 4.35 mg/l, 95th percentile: 9.95 mg/l) were reported in 30 individuals after they had used sprays containing pyrethrum (Lena et al., 2006). E-cis-CDCA could not in anv At the same time, the metabolite be detected case. concentrations in the urine of 45 test persons without biocide known exposure were determined. anv demonstrating background concentrations with of 00.05 (E)-trans-CDCA values a maximum mg/l up to 0.82 mg/l. Furthermore. metabolite concentrations test persons determined of the urine after operating electro-vaporizers containing d-allethrin and pyrethrum 2003). (E)-trans-CDCA concentrations et al., 1.0 mg/l to 6.2 mg/l, depending ranged from on the time hours) Values

metabolites of pyrethroid insecticides in urine are also reported by Heudorf (Heudorf et al., 2006).

results obtained Comparison of the in our study with data in the literature shows that the (E)-trans-CDCA concentrations found in urine after correct use of four different consumer spravs were on the same scale as the in the values reported literature after spraying or using electro-vaporizers.

Conclusions

A characterization of the exposure scenarios during self-use of biocidal consumer products like aerosol sprays and electro-vaporizers in indoor areas was **Products** according the undertaken. were applied to manufacturers' instructions and in a worst-case scenario simulating Differforeseeable unforeseeable ormisuse ent active substances were determined: various pyrethroids and pyrethrins as well as chlorpyrifos and the PBO. Breathable air and dermal svneraist concentrawere measured, and inhalable and doses tions dermal the sprayers were quanti...ed. In addition. biomonitoring was performed by determining urine metabolites of the bystanders. sprayers and The data presented allow а better understanding and assessment of the potential exposure levels for occupants of room in which biocides are sprayed or vaporized. Based on the calculated levels realistic risk exposure а assessment should be able to be performed with regard to normal exposure conditions. and worst-case

In addition, the time-dependent air concentrations of the active substances after spraying were simulated by a deterministic model (SprayExpo). The model calculations mainly led to an acceptable correlation with the experimental data.

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