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Here was tested how well a NF/FF dispersion model predicts particulate matter concentrations when source emission potency was estimated using material dustiness index.



Environmental impact statement

The REACH requires that manufacturers or importers within European Union must estimate human exposure by all routes for each potential exposure scenario which will then be used in risk assessment. To fulfill this regulation several exposure assessment models/tools were developed but their performances to predict particulate matter exposures have not been tested in work environments. In this study we tested how well a NF/FF dispersion model predicts particulate matter concentrations when source emission potency was estimated using material dustiness index. It was found that dustiness index did not describe source emission potency well. To overcome this problem, particle emissions from different sources should be studied with well-controlled work simulations where emission rates are assessed with indoor aerosol modelings.

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Cite this: DOI: 10.1039/x0xx00000x

Received ooth January 2012, Accepted ooth January 2012

DOI: 10.1039/x0xx00000x

www.rsc.org/

Testing a Near Field/Far Field model performance for prediction of particulate matter emissions in a paint factory

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A Near Field/Far Field (NF/FF) model is a well-accepted tool for precautionary exposure assessment but its capability to estimate particulate matter (PM) concentrations is not well studied. The main concern is related to emission source characterization which is not as well defined for PM emitters than *e.g.* for solvents. One way to characterize PM emission source strength is by using the material dustiness index which is scaled to correspond to industrial use by using modifying factors, such as handling energy factors. In this study we investigate how well the NF/FF model predicts PM concentration levels in a paint factory. PM concentration levels were measured during big bag and small bag powder pouring. Rotating drum dustiness indices were determined for the specific powders used and applied in the NF/FF model to predict mass concentrations. Modeled process specific concentration levels were adjusted to be similar than the measured concentration levels by adjusting the handling energy factor. The handling energy factors were found to vary considerably depending on the material and process even-though they have the same values as modifying factors in the exposure models. This suggests that the PM source characteristic and process-specific handling energies should be studied in more detail to improve model-based exposure assessment.

Introduction

The Registration, Evaluation, Authorization, and Restriction of Chemicals regulation (REACH)¹ requires that manufacturers or importers must report or estimate human exposure for all relevant routes to determine the appropriate risk management measures and prevent excessive exposure.²⁻³ By the end of May 2018, this applies to all chemicals that are manufactured or imported in quantities over 1 metric ton per year within the European Union. In principle, this means that exposure should be assessed for hundreds of thousands chemicals for all potential routes in different exposure scenarios including occupational and consumer exposures.

The one way to overcome this requirement is through mathematical exposure modeling. Thus, in the recent years several exposure assessment models were developed, such as the ECETOC TRA⁴, the Stoffenmanager⁵, the EMKG-Expo-Tool⁶, and the Advanced REACH Tool⁷⁻¹¹ (ART,

https://www.advancedreachtool.com). The ART is an advanced model designed to assess absolute exposure levels for different exposure scenarios. It is based on a Near Field/Far Field (NF/FF) model which is a well-accepted exposure assessment model within the scientific community and occupational hygienists.¹² The ART exposure assessment model also uses calibration factors for inhalable dust, vapors, and mists that are defined from existing occupational exposure data.^{9,13}

One key parameter in the NF/FF model is characterization of the source emission rate and how well it describes emissions in occupational environment. The NF/FF model has been found to predict occupational exposure levels well from solvent mixtures.^{3,12,14-17} For a particulate matter (PM) generated by powder handling, the source term is usually characterized by a dustiness index. The concept of the dustiness index was developed to classify bulk materials according to their relative dust producing capacities. Dustiness is not a well-defined

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tools, such as those listed above, use the dustiness index to estimate particle source strength and the potential PM exposure level. However, because the powder properties that control the specific powder dustiness levels are still not fully understood, therefore it is still difficult to predict workplace exposure levels accurately. Previous correlation studies between dustiness index and measured concentrations correlate relatively well (R² 20 m ~70 %) at small scale powder handling^{22,25} but at large scale material bagging and dumping, less consistent results have been found.^{26,27} See also commentary by Lidén.²¹ Here, we studied how well the NF/FF model predicts PM exposure concentrations at a paint factory where powders were poured into a mixing tank. The emission source strengths were characterized by dustiness indices, which were determined by using a down-scaled EN15051 dustiness drum^{28,29}, and by using a handling energy factor and localized control factor. Particle concentration measurements and gravimetric PM sampling were made in both the NF and FF. Particles origin and nature were analyzed by scanning and transmission electron microscopy. The NF/FF model concentrations were adjusted to match measured respirable mass concentrations by adjusting the handling energy factor. The handling energy factors were

physical or chemical property but an empirically defined value

which depends on the test methods¹⁸⁻²⁴. Exposure assessment

Experimental

example in the ART.

Work environment and processes

Particle measurements were carried out in a paint factory mixing hall (20 m wide, 30 m long and 2.5 m height; Figure 1). The air-exchange in the mixing room was in this special case maintained by natural ventilation where most of the air was assumed to exchange through two pairs of doors which were open all the time to a loading ramp. The mixing room ventilation rate was estimated to be 5 h⁻¹. In a mixing station, raw materials in liquid and powder form were poured through a quadratic opening (0.8 x 0.8 m²) into a mixing tank located below the mixing room. The powders, which are listed in Table 1, were in 25 kg small bags (SB) or 500 kg big bags (BB) with a discharge cone at the bottom. Bags were transported with an electrical forklift. SBs were opened with a knife and poured at a height of 1 m into the mixing chamber whereas BBs were lifted with the electrically powered crane above the mixing chamber where the sackøs bottom discharge cone was opened and the powder fell down into the mixing chamber by its own weight.

then compared with previously assigned values used for

Workplace measurements

Particle concentrations were measured with stationary instruments at a height of 170 cm. A near field (NF) measurement location was always approximately 0.5 m from the mixing station and 1 m from the worker. A far field (FF) measurement location was 7.7 m from the mixing station (Figure 1).



Fig. 1 Layout of the mixing room.

The NF and FF mobility particle size distributions were measured from 5.6 nm to 560 nm in 1s intervals with two Fast Mobility Particle Sizers (FMPS, TSI model 3091, TSI Inc., Shoreview, MN, USA). The NF optical particle size distributions were measured from 250 nm to 30 μ m in 6 s intervals with a Grimm Dust Monitors (DM, Grimm model 1.109, Grimm Aerosoltechnik, Ainring, Germany). The NF aerodynamic particle size distributions were measured from 6 nm to 10 μ m in 1s intervals with an Electrical Low Pressure Impactor (ELPI, Dekati model ELPI+, Dekati Ltd., Tampere, Finland).

Gravimetric samples of respirable particulate matter were collected from the NF, FF, breathing zone (personal sampler), and outdoor air. The respirable fraction has a D_{50} cut size at 4 μ m and is defined in detail by the European Committee for Standardization³⁰ and the ACGIH³¹. Samples were collected on 37 mm Teflon filters with a 0.8 μ m pore size (Millipore, Billerica, MA, USA) using BGI Model SCC1.062 ($Q_s = 1.05$ L min⁻¹) and BGI Model GK2,69 ($Q_s = 4.2$ L min⁻¹)³² Triplex cyclones (BGI Inc., Waltham, MA, USA). Three blind filters were used as controls to correct for handling and environmental factors. Filter weighing was completed in a climate controlled weighing room (50 % relative humidity and 22°C) after at least 24-hour acclimatization.

For the purpose of single particle analysis with electron microscopes, particles were collected on Ni-TEM grids and carbon substrates using a micro-inertial impactor equipped with two stages, covering a size range from 50 nm to $3.5 \,\mu$ m projected area diameter. For further description of sampling see Kandlera *et al.*³³ and their references. Particle samples that correspond to NF were sampled for a short time (minutes) during pouring processes, and reference samples were taken outdoors, where background conditions are considered.

easured P, Resp.,] [%] .0 99.7

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Table 1. Commonly used powders in the paint factory with product information according to material suppliers, respirable occupational exposure limits (OEL) according to Danish regulations⁷³, and measured dustiness indices in respirable mass and with ART multipliers in brackets, and number of emitted particles per kg with ultrafine particle (UFP; $D_p < 100$ nm) and respirable (Resp.) fractions.

	Formula	CAS-No.	Bulk density, [kg m ⁻³]	D ₅₀ , [μm]	OEL, [mg m ⁻³]	GK2.69	ELPI			FMPS		
Product						DI_m^{a} , [mg kg ⁻¹] (ART multiplier)	$DI_N,$ [×10 ⁸ kg ⁻¹]	UFP, [%]	Resp., [%]	$DI_N,$ [×10 ⁸ kg ⁻¹]	UFP, [%]	Resp., [%]
RD3 (Sachtleben Pigment GmbH, Pori, Finland)	TiO ₂ (93%), Al ₂ O _{3,} ZrO ₂ , organic	13463-67-7	800	0.22	6	5.3±2.3 (0.03)	1.6	66.1	99.0	0.8	64.0	99.7
Micro Mica W1 (Norwegian Talc AS, Knarrevik, Norway)	KAI ₂ (AlSi ₃ O ₁₀)(OH) ₂	12 001-26-2	-	9	5 ^b	22.4±5.5 (0.1)	7.7	20.3	97.7	10.7	39.0	99.6
TR92 (Huntsman, Hartlepool, UK)	TiO ₂ (94%), Al ₂ O ₃ , ZrO ₂	13463-67-7	1200 (tamped)	0.24	6	1.6±1.0 (0.01)	0.6	76.9	99.4	0.1	52.5	99.6
Satin Tone W (Whitetex; BASF Florham Park, NJ, U.S.A.)	Al ₂ Si ₂ O ₅ (OH) ₄	92704-41-1	256	1.4	2	2.4±5.5 (0.01)	0.7	75.4	99.1	0.7	83.3	99.8
Microdol Norwegian Talc AS, Knarrevik, Norway)	CaMg(CO ₃) ₂	16389-88-1	700	7.5	5 ^b	12.1±1.2 (0.03)	0.4	16.0	97.6	0.8	73.4	99.8
R-KB-6 (Sachtleben Pigment GmbH, Krefeld, Germany)	TiO ₂ (94%), Al ₂ O ₃ , ZrO ₂	13463-67-7, 21645-51-2	500	0.37	6	9.9±1.1 (0.03)	2.5	35.9	98.5	1.5	32.2	99.6
Plastorit, Talc (Imerys Talc, Andritz, Austria)	SiO ₂ (33%), chlorite (33%), mica (33%)	1318-59-8, 12001-26-2, 14808-60-7	860	-	5 ^b	20.3±7.0 (0.1)	3.6	25.6	97.1	6.5	21.5	99.6
Diafil 530 (Imerys, San Jose, CA, U.S.A.)	SiO ₂ (93%)	61790-53-2	470	11	5 ^b	108.9±11.1 (0.3)	9.8	8.8	95.6	7.3	19.0	99.5

^aMean ± standard deviation; Classification of the Art multipliers are described by Fransman et al.⁸; ^bInorganic respirable dust

and Jensen.28

a)

b)

11+ Lpm

11+ lpm

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The mixing room relative humidity and temperature were monitored by using Gemini TinytagPlus Data Loggers (Gemini Data Loggers Ltd, West Sussex, UK).

Unit conversion from particle number to mass.

Mass concentrations were calculated from particle number size distributions measured by the DM and ELPI by assuming spherical particles. Respirable mass concentration was calculated by multiplying the mass distributions with the simplified respirable fraction penetration efficiency according to Hinds.³⁵ Particles average density was selected so that the DM respirable mass concentration correspond the average mass concentration measured by the gravimetric samplers (Table 2) over the respective time period. This results to density of 1.7 g cm⁻³ while bulk densities varied from 0.256 to 1.2 g cm⁻³ (Table 1).

Electron Microscopy

Particles were analyzed by means of scanning (SEM) and transmission electron microscopy (TEM), using a FEI ESEM Quanta 200 FEG with an acceleration voltage of 20 kV and a FEI TEM Tecnai T20 G2 (FEI, the Netherlands) operating at 200 kV. Chemical characterization was derived by spot analysis and mapping with energy dispersive X-ray detection (EDS, Oxford Instruments 80 mm² X-Max SDD detector, Oxford Instruments, UK).

Characterization of dustiness indices

Dustiness measurements were performed for Table 1 materials, which were collected during the measurement campaign from the bags. Dustiness indices were determined using the downscaled EN15051 dustiness drum and procedures described therein^{28,29} with a sampling train for the FMPS, ELPI, and an Aerodynamic Particle Sizer (APS, model 3321, TSI Inc., Shoreview, MN, USA) particle size distribution measurements. Sample for gravimetric measurements were collected on Teflon filters with the GK2.69 (BGI Inc., Waltham, MA, USA) respirable cyclone (Figure 2). Here, we do not show results from the APS measurements.

The drum was ventilated with a HEPA-filtered air with relative humidity of 50 % at a rate of 11 L min⁻¹. Prior to each quantitative test, the drum surfaces and sampling lines were õsaturatedö with dust using 2 grams of the powder and rotating it for 60 seconds. After this saturation run, the drum was emptied and 6 grams of test material were loaded into the drum for experiments using setup 1 as shown in Figure 2a. Before the experiment, the background particle concentration was measured for 60 seconds to ensure a particle free test atmosphere. The experiment was then started through 60 second of drum rotation followed by 120 seconds sampling to

an additional two times to ensure repeatability. Thereafter an additional testing was performed using setup 2 as shown in Figure 2b. Each dustiness index was determined as the average of three repeats. The drum and sampling system was thoroughly cleaned between each test material as described in Schneider 9 lpm Excess air 10 Lom FMPS M Rotating Drum 1 Inn Respirable Filter sampling 4.2 Lpm Excess air 4.8 lpm 4 lpm Excess ai 5 lpm 20X APS M HEPA Rotatina Drum 10 lpm Excess air FIP 4 lpm

Fig. 2 Setups to measure dustiness index with a small rotating drum using (a) the FMPS and (b) the ELPI.

5 Lpm

Dilutor

20X

APS

HEPA

collect the entire dust cloud. The experiment was then repeated

Description of the NF/FF model

The NF/FF model that was used in this study is described in general by Cherrie et al.³⁶ and in detail by Zhang et al.¹⁷ The model assumes that (1) all mass entering the model volume is created at a source inside the NF volume, (2) particles are fully mixed at all times in the NF and FF, (3) there is limited air exchange between NF and FF volumes, and (4) there are no other particle losses than the FF ventilation. In the NF/FF model, the NF volume can be fixed either onto a source (see e.g. Jayjock et al.¹² and its references) or the workerøs head which is moving around in the FF volume with an enveloping NF volume.⁷ Here, the NF was defined as a cube with 2 m sides $(V_{NF} = 8 \text{ m}^3)$ centered on the quadratic opening where material was poured into the mixing tank, at a height of 1 m from the ground, which was considered to be the main source for particulate matter. Thus, the NF volume covered the workerøs breathing zone and the NF instruments inlets during the pouring process. In the ART model, this corresponds to a situation where a worker is within 1 m of the source. One critical parameter in the NF/FF model is the airflow between the NF and FF which is discussed in detail by Zhang et al.¹⁷ Our modelings were made using three different air flow rates

Excess nin

between NF and FF, Q_{NF} (m³ min⁻¹), which represents minimal (3 m³ min⁻¹), intermediate (10 m³ min⁻¹), and maximal (30 m³ min⁻¹) likely convective airflows as described by Cherrie.³²

The emission from process *i* by handling of powder *j* is described with the potential emission rate $E_{i,j}$ (units min⁻¹):

$$E_{i,j} = DI_j \cdot H_i \cdot \frac{dM_j}{dt} \cdot LC , \qquad (1)$$

where DI_j is the respirable dustiness index of material *j* expressed here in units of particle number (kg⁻¹) or mass (mg kg⁻¹), H_i (-) is the handling energy factor for the process *i*, dM_j/dt (kg min⁻¹) is the mass-flow of powder *j* in the process, and *LC* (-) is the total protection factor of localized controls.

The handling energy factor of the process, H_i , relates the mechanical energy used in the process *i* to the mechanical energy applied in dustiness index measurement.²⁸ Thus, the handling energy factor of the process equals 1 if the applied mechanical energy equals to the energy that was used to measure dustiness index or has the same effective dispersion effect. If de-agglomeration was complete in the dustiness index measurement, then the handling energy factor is $0 \le H_i \le 1$, otherwise $H_i \ge 0$.²¹ Determinants underlying material dustiness are listed with references by Fransman *et al.*⁸ and the potential emission source modifying factors are described by Van Tongeren *et al.*³⁸

In this study, pouring-process-specific handling energy factors were defined by adjusting the NF/FF pouring-process-specific concentration levels at similar concentration levels measured by the DM.

The ART model parameterization

In the ART model, the handling energy factor multiplier depends on the mass-flow (varies from 0.03 to 30; here 3 corresponding to a transfer rate of 10 ó 100 kg min⁻¹), handling carefulness (0.3 or 1; here 1 corresponding routine transfer), and drop height (1 or 3; here 3 corresponding > 0.5 m drop height).8 Other modifying factors are related to powder properties, such as the dustiness index for which the corresponding ART multipliers are listed in Table 1. Powder moisture content was assumed to be 0 % in this study, which has a multiplier of 1.8 The dispersion multiplier would be close to 0.7 which in the ART model is used for a 3000 m³ room with 3 air exchanges per hour. In the SB pouring there was no localized control why we used a multiplier of 1 whereas low specification containment by the discharge cone was present in the BB pouring, leading us to use a localized control multiplier of 0.1. The ART concentrations, that were calculated according to Fransman et al.⁸, were corrected by using calibration factor of $\ln() = 3.01$ as defined by Schinkel *et al.*⁹ and using dustiness multipliers given in Table 1.

Results

Dustiness measurements

Table 1 shows measured dustiness indices in units of mass and particle number. By following the EN 15051 standard classification²⁹, Diafil amorphous silica was ranked in dustiness as *Moderate*; Microdol, Pastorit, and Micro Mica were ranked as *Low*; and Satin Tone and titanium dioxides RD3, TR92, and R-KB-6 were ranked as *Very Low*.

Currently there are no ranking levels for dustiness in particle number. According to the ELPI, in particle number Diafil emitted $9.8 \times 10^8 \text{ kg}^{-1}$, Microdol, Pastorit, and Micro Mica from 0.4×10^8 to $7.7 \times 10^8 \text{ kg}^{-1}$, and Satin Tone and titanium dioxides RD3, TR92, and R-KB-6 from 0.6×10^8 to $2.5 \times 10^8 \text{ kg}^{-1}$. According to the ELPI, nearly all particles (> 95.6 %) were respirable particles, and 8.8 to 76.9 % were ultrafine particles (Table 1). The correlation between the ELPI and FMPS dustiness index measurements was 0.88.

Concentrations

In the workplace, relative humidities and temperatures measured at the NF and FF varied between 40 to 55 % and 19.5 to 23.5 °C. Figure 3 shows that powder pouring increased the NF concentrations of particles in between 0.1 to 20 μ m. Figure 4 shows that the respirable mass distributions were similar for both days when averaged over NF gravimetric sampling periods shown in Figure 5. The averaged mass distribution mode was 2.8±0.3 μ m with a geometric standard deviation of 2.16±0.12 (95 % of the mass was between 0.6 to 13.1 μ m).



Fig. 3 NF particle size distribution time series for (a) day 1 and (b) day 2. Material pouring time intervals are shown with horizontal white lines.



Fig. 4 Respirable mass distribution averages calculated over the gravimetric NF sampling period and P_{RF} shows the simplified respirable fraction penetration efficiency for particles over 1 μ m in diameter.

The NF mass concentrations were above the detection limit only for GK2.69 during day 1 and SCC1.062 during day 2 (Table 2) where the average mass concentration of 149 μ g m⁻³. The average respirable mass concentration for the respective measurement periods estimated from converted the DM and

ELPI data were 142.7 μ g m⁻³ and 219.7 μ g m⁻³, respectively. Personal respirable dust exposure levels sampled from the worker breathing zone was 5.1 and 3.94 times higher than the respective NF concentrations (Table 2). Outdoor respirable mass concentrations were clearly below detection limits.



Fig. 5 NF respirable mass concentration time series for (a) day 1 and (b) day 2. Gravimetrical personal and NF samplers show the mean mass concentration level defined for the sampling time interval.

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Figure 5 shows how powder pouring clearly elevates respirable mass concentrations measured with the ELPI and DM but do not elevate significantly the NF number concentrations as measured by the ELPI or FMPS (Figure 6). Particle number concentrations increase was difficult to detect because the background particle number concentration was high as compared to the process particle emissions. Thus, the NF/FF concentration ratios measured by the FMPSøs were

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nearly the same (1.06 and 1.10 for days 1 and 2, respectively; Table 2).

Table 2. Work shift mean mass concentrations measured with gravimetric samplers (SCC1.062 and GK2.69), calculated from particle measurements assuming spherical particles with mean density of 1.7 g cm⁻³ (DM, ELPI, and FMPS), estimated with the NF/FF model where $Q_{NF} = 10 \text{ m}^3 \text{ min}^{-1}$, estimated with the ART, and mean number concentrations measured with the FMPS, DM, ELPI, and estimated with the NF/FF model. Note that the FF mass concentrations sampled with SCC1.062 were clearly below detection limits, and in number concentration, the NF/FF model concentrations account only process particle concentrations.

	Day 1		Day 2					
Instrument/ method	<i>m</i> , (µg m ⁻³)	N, (cm ⁻³)	<i>m</i> , (µg m ⁻³)	N, (cm ⁻³)				
Near Field								
SCC1.1062	101.0 ^a	-	125.0	-				
GK2.69	173.0	-	19.0 ^b	-				
DM	99.1	210	186.3	230				
ELPI	253.0	11800	170.9	8300				
FMPS	6.6	6500	5.6	4980				
The NF/FF model	86.4	420	145.9	230				
The ART ^e	770 (410 ó 1500)	-	550 (290 ó 1100)	-				
Far Field								
SCC1.1062	0.02 ^c	-	0.05 ^d	-				
FMPS	6.1	6140	5.3	4520				
The NF/FF model	5.3	24	9.3	15				
Personal sampling								
SCC1.1062	515.0	-	493.0	-				

^{a-d}Values below detection limit which were for ^a177, ^b31, ^c109, and ^d100 µg m⁻³; ^ebrackets shows the 75th percentile inter-quartile confidence interval.

Microscopy analysis

Particles on the samples collected for electron microscopy were mainly mineral pigment/filler particles. Even though the sampling time spanned more than one process in most cases, the single particles can be allocated to the powders as described by the manufacturer data sheets. However, most particles appear to be agglomerated and coatings are observed on many of them. Generally coatings were assumed to be the inorganic/ organic as specified in the technical data sheets (Table 1) usually few tens of nanometers thick on single particles. Thicker layers of coatings appeared on heavily agglomerated particles collected during a pouring process (Figure 7e). While the primary particles sizes as measured from microscopy images (data not shown) were usually close to the D_{50} indicated by the manufacturers data sheets, the agglomerate size might vary. Figure 7 shows examples of pigment/filler particles (a, b, and c) sampled from the NF and soot (d) sampled outdoors. Internally mixed agglomerates and thick-coated particles were

found only on those samples corresponding to the NF measurements.

The TEM samples also suggest that soot was present in the background in high abundances. Soot was found in high number of individual agglomerates (e.g. 6 soot agglomerates per 16 pigment/filler particles in respirable fraction) on the samples corresponding to the NF. TEM high resolution (Figure 7d) on individual soot particles revealed the typical onion shelllike structure in primary particles of 20 to 50 nm in diameter.



Fig. 7 Micrographs of collected particles: a) SE image showing Dolomite (Ca+Mg+C), Mica (Al+Si) and TiO₂ (Ti) particles and HRTEM images of b) agglomerated TiO₂, c) Mica sheets, d) soot, and e) overview of mixed particles.

NF/FF modelings

The handling energy factors (H_i) were defined separately for each pouring process by adjusting H_i so that modeled and measured mass concentrations covered from the DM measurements were similar during pouring. The DM was selected as the reference instrument because 95 % of the respirable mass emitted during pouring was between 0.6 to 13.1 µm which is the range covered by the DM. Average mass Table 3. Defined handling energy factors for BB (LC = 0.1) and SB (LC = 1) pouring processes: *n* shows number of pouring repeats, H_i is the handling energy

concentration ratios with standard deviations calculated for modeled and measured concentrations during pouring events using different NF ventilation rates were $m_{NF}/m_{DM} = 1.00\pm0.02$ and $m_{NF}/m_{ELPI} = 1.6 \pm 1.1$. This means that the NF/FF modelled concentrations were very similar to the values measured by the DM but differ quite much from the ELPI mass concentrations when using the adjusted handling energy factors shown in Table 3.

factor, averages of mass and number concentrations measured during pouring events by the DM, and mass concentration predicted by Fransman et al. ⁸							
Pouring process	n ^a	H_i , ($Q_{NF} = 3 \text{ m}^3 \text{ min}^{-1}$)	H_i , $(Q_{NF} = 10 \text{ m}^3 \text{ min}^{-1})$	H_i , $(Q_{NF} = 30 \text{ m}^3 \text{ min}^{-1})$	<i>m_{DM}</i> , (µg m ⁻³)	N_{DM} , (cm ⁻³)	Fransman <i>et al.</i> ⁸ , (mg m ⁻ ³)
BB RD3	4	0.86	2.35	6.30	82.7	210	0.38
BB TR92	5	8.30	18.60	49.00	364.0	420	0.128
BB Microdol	2	2.60	7.40	19.70	767.8	400	0.38
SB RD3	1 (10)	0.26	0.78	2.12	167.1	250	3.8
SB Micro Mica	1 (17)	0.06	0.19	0.51	305.4	280	12.8
SB SatinTone	1 (16)	0.36	1.09	2.93	97.6	190	1.3
SB Microdol	1 (11)	0.09	0.23	0.60	245.2	220	3.8
^a Brackets shows	number	of poured SBs during pou	ring process.				

Table 2 shows mass and number concentrations predicted by the NF/FF model using the adjusted H_i multipliers defined with $Q_{NF} = 10 \text{ m}^3 \text{ min}^{-1}$. The NF/FF model mass concentration was 0.5 times lower and 1.2 times higher than gravimetric measurements for days 1 and 2, respectively (Table 2). If gravitational settling was taken into account at the NF volume as in Cherrie et al.36, the NF and FF concentrations would decrease approximately 0.3 % when using settling velocities for 2.8 μ m particles as defined by Schneider *et al.*³⁹

Average mass concentrations predicted by the ART for days 1 and 2 were approximately 4.4 times higher for both days than measurements (Table 2; Supplementary gravimetric information Reports 1 and 2 show the ART reports for days 1 and 2, respectively). For individual pouring activities, the ART predicted average mass concentrations with 75th percentile inter-quartile confidence interval of 900 (470 \acute{o} 1700) $\mu g~m^{-3}$ for BB pouring and 2700 (1400 ó 5100) µg m⁻³ for SB pouring (Supplementary information Reports 3 to 10 show the ART reports for each pouring). Table 3 shows individual pouring time concentrations calculated according to Fransmann et al.⁸

Discussion

Dustiness indices of powders used for the paint manufacturing were mainly Low or Very Low (Table 1). Diafil was found to have moderate dustiness even-though the primary particle median size was 11 µm. Here we did not find any correlation between primary particle size given by the manufacturer and dustiness index either in units of particle number or mass (Table 1). Similar findings were made by Evans *et al.*⁴⁰ for fine and nanoscale powders.

The NF respirable mass concentrations at the mixing station were 2.9 and 3.9 times lower than personal exposure measured

at the breathing zone for days 1 and 2, respectively (Table 2). This means that the inlets of the NF instruments underestimated personal exposure concentrations, or alternatively, the worker was exposed to particles mainly outside of NF volume. Such exposure may occur for example when empty bags were folded outside of the NF volume and approximately 3 m from FF instruments so that the emission was not detected by instruments. It has also previously been reported that personal exposure levels exceed the values determined by stationary measurements.⁴¹ Personal exposure levels were clearly below occupational exposure limits, which varied from 2 to 6 mg m⁻³ for respirable fraction (Table 1), even if the workersø use of respirators was not taken into account. However, especially PM exposure is known to relate to a wide range of occupational diseases as shown e.g. by Omland et al.⁴² Also, exposure to urban fine particulate matter $< 2.5 \ \mu m \ (PM_{2.5})$ is recognized to be globally the 9th most powerful risk factor for burden of disease.⁴³ Thus, it is recommended minimize the exposure by using local exhaust ventilations, enclose the emission source, or wear respirators in case such engineered controls are not available.

Source emission characteristics are affected by several modifying factors which are discussed in general by Cherrie and Schneider⁴⁴ and protection factors of localized controls by Fransman et al.⁴⁵ Modifying factor multipliers that are used in the ART model are shown by Fransman et al.^{7,8} and McDonnell et al.¹³ Here the SB pouring was an open process (LC was 1) and in the BB pouring the discharge cone in the big bag was assumed to be a low specification containment (LC was 0.1). In this case, pourings were performed in the same company and by the same person in similar environmental conditions. Thus, it is reasonable to assume that pouring process repeatability was

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good and modifying factors should remain similar. The handling energy factor multiplier for pouring from over 0.5 m is 3 and the modifying factor multiplier for a transfer rate between 10 \circ 100 kg min⁻¹ is 3.⁸ Thus, the modifying factor multipliers for both pouring processes are 9.

However, assuming an average aerosol density of 1.7 g cm⁻³, it was found that the handling energy factor varied for H_{BB} from 2.35 to 18.60 and for H_{SB} from 0.19 to 1.09 when Q_{NF} , was 10 m³ min⁻¹ (Table 3). The NF air flow rate Q_{NF} reduces or increases the handling energy factor approximately 2.7 times if Q_{NF} is either 3 or 30 m³ min⁻¹ (Table 3). In this study, it was possible to assess only the handling energy factor range because Q_{NF} was not measured and ventilation Q_{FF} was not well known. However, in this study, Q_{FF} does not have high effect on results. For example, if air exchange ratios would be 2 or 8 h⁻¹ (i.e. 5±3 h⁻¹), the NF/FF model NF concentration during day 1 would be 91.3 or 84.6 µg m⁻³, respectively. Thus, the Q_{FF} has only minor effect when the handling energy factor is adjusted from the NF concentrations.

A critical finding in this study was the variation of the handling energy factor between similar processes and systematic difference between H_{BB} and H_{SB} . Depending on the Q_{NF} , the H_{BB} was from 6 to 49 times higher than the H_{SB} even though they should be at a similar level according to previously assigned modifying factors. Also H_{SB} was clearly below 9 even at maximal likely convective airflow (Table 3). Critical parameters for scaling by the modifying factors are the values defined for the handling energy factors, drop-heights and definition of localized controls. These immediately cause a significant offset in scaling. In addition, gravimetric mass samples consist of background particles (e.g. soot) and pigment/filler particles with vapours condensed on them (Figure 7a). The amount of condensed vapor may vary depending on the particle type and cause relative error in pigment/filler mass concentration assessment. However, none of these parameters can explain the high variability observed in the derived handling energy factors. It is obvious that the observed variability is partly related to measurement uncertainties and low amount of pouring repeats but those factors alone cannot explain such high and systematic differences between the two types of pouring.

An important issue in the estimation of the PM emission source strength from the dustiness index, is how well forces applied to the powder in the specific dustiness measurement corresponds to forces in occupational powder handling.^{21,24,46} Bach *et al.*²⁴ discuss in detail the forces acting on the material during the dustiness measurement process and show their relative influence for the two reference methods according to the EN15051 (rotating drum and continuous drop)²⁹ and two other methods (UNC Dustiness Tester and single drop). They conclude that material dustiness depends on the interacting forces used in the test procedure and a testing method should be selected so that it reproduces energy used in the corresponding material handling. This is also emphasized in the EN15051 and one of the key topics in the development of the new dustiness standard under development for nanomaterials.^{47,48} Dustiness index characterized with the continuous drop method might provide better correlation with the both pouring processes and reduce deviation between handling energy factors that were assigned here.

Another issue is how well the dustiness index, defined by EN15051, is scalable for large mass flows. Higher mass flow is assumed to increase emissions but this may not always be true because the product to air interface may decrease.⁴⁹ However, here it looks as if material pouring through the big bag discharge cone increased the handling energy factor. Therefore, one might consider that careful powder pouring from bags or pouring from the edge into a semi-closed mixer would not necessarily result in a higher exposure risk when assessed from the stationary measurements. The situation, however, may be very different, if the exposure is measured in the workerøs inhalation zone as the personal exposure levels were notable higher than the stationary PM concentrations. This is likely due to the fact that the worker is in closer to the source than the stationary measurement station during the powder handling events; especially in handling small bags.

The NF/FF model comparison with the ART is not straightforward because the ART exposure model is using inhalable, and not respirable, fractions as in this study. Calibration multipliers which are defined from occupational hygienic measurements, and the interpretation of parameters is not always straightforward. However, individual pouring concentrations were overestimated by methods described by Fransman et al.⁸ 1.5 times in BB pouring and 56 times in SB pouring (Table 3). Subsequently, the work shift exposure including non-exposure time during day 1 was 2.2 mg m⁻³ which is nearly 3 times more than the ART estimated (Table 2). Individual pouring concentrations were also overestimated by the ART 4.7 and 28 times, respectively, for BB pouring and SB pouring. In this study, the ART predicts the work shift inhalable concentrations well when compared to the gravimetrically measured NF and personal respirable concentrations. Discrepancy may be caused by differences in the source emission potency calculation methods between the ART and Fransman et al.⁸ However, this shows that in this case the model in the method by Fransman et al.⁸ estimates well the NF concentrations for BB pouring but overestimates the NF concentrations during SB pouring.

In the number concentration modeling, we used the same handling energy factors that were characterized from the mass concentration modelings with $Q_{NF} = 10 \text{ m}^3 \text{ min}^{-1}$ (Table 3). Comparison of the modeled NF values with the measurements is challenging due to high background number concentrations compared to the number concentration increase by pouring processes. However, Figure 6 shows that number concentration measured by the DM was below 200 cm⁻³ when there were no pouring processes and Table 3 shows that during pouring processes, the average number concentrations in the NF varied from 190 to 420 cm⁻³. Thus the pouring process increased the NF number concentrations roughly up to 200 cm⁻³ on average, which is less than the average modeled process particle

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concentration values of 420 cm⁻³ and 230 cm⁻³, respectively, for days 1 and 2 (Table 2).

One way to study the modifying factors in the PM source characteristics is to measure and assess particle emission rates from various processes in laboratory conditions and at work-places where environmental conditions and processes are well known. As pointed out by Zhang *et al.*¹⁷, the air exchange Q_{NF} between the NF and FF volumes is one poorly known parameter. The PM dispersion at work places could be studied with flux measurements.⁵⁰

Emission source characterization should be extended from powder handling to also cover PM emissions from other processes such as ultrafine particle emissions from high energy processes (*e.g.* Koponen *et al.*⁵¹) and processes where new particle formation may occur (*e.g.* Nørgaard *et al.*⁵²). This requires measurement of size and time-resolved process particle concentrations where source emission rates are resolved with indoor aerosol modeling.^{53,54} There are various source characterization studies made both for consumer products and occupational processes.⁵⁵⁻⁷¹ Similarly particle sinks can be assessed according to the model Mølgaard *et al.*⁷² made for indoor air cleaners. Some of these scenarios could potentially be covered with exposure assessment tools such as the ConsExpo REACH tool, but this tool may not be applicable to larger scale industrial exposure scenarios.

Conclusions

Development of aerosol modeling for exposure assessment is compulsory in order to fulfil the REACH regulation. Several quantitative exposure assessment tools are available. However, their performance for particulate matter exposure assessment has not been previously evaluated in detail by comparing modeling with measured exposure in a real work environment.

In this study, we evaluated the performance of the NF/FF model using a source emission strength that was based on the rotating drum dustiness test. We showed that particle emission rates for a powder pouring process from big bags and small bag were not well predicted when using the dustiness index. We suggest that emission source characterization based on dustiness index should be further studied by defining processspecific emission rates with indoor aerosol models in wellcontrolled environments and processes which are then linked to dustiness indices determined using different types of dustiness tests.

We propose that a generalized particle emission rate term (units *e.g.* in particles min⁻¹, μ m min⁻¹, or mg min⁻¹) should be implemented in REACH inhalation exposure assessment tools. Such emission terms could be defined also for sources emitting ultrafine particles and exposure analysis could thereby be assessed also in other units than mass, which could also be useful for assessment of the biologically effective dose.

Acknowledgements

The research leading to these results has received funding from the European Community's Seventh Framework Programme (FP7/2007-2013) under NanoValid (grant agreement N°263147) and this work was supported by a grant from the Danish Centre for Nanosafety (grant agreement N°20110092173/3) by the Danish Working Environment Research Fund.

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Electronic Supplementary Information (ESI) available: Reports of the predicted exposure levels by the Advanced REACH Tool (ART). See DOI: 10.1039/b000000x/

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