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## Semivolatile Organic Compounds in Homes: Strategies for Efficient and Systematic Exposure Measurement Based on Empirical and Theoretical Factors

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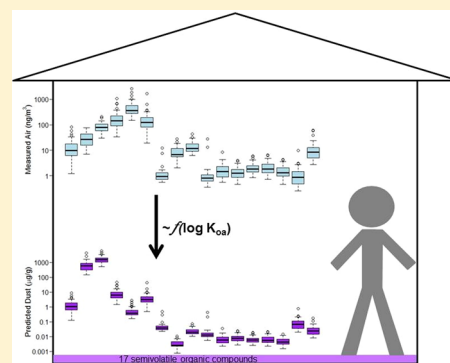
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### Supporting Information

**ABSTRACT:** Residential exposure can dominate total exposure for commercial chemicals of health concern; however, despite the importance of consumer exposures, methods for estimating household exposures remain limited. We collected house dust and indoor air samples in 49 California homes and analyzed for 76 semivolatile organic compounds (SVOCs)—phthalates, polybrominated diphenyl ethers (PBDEs), polychlorinated biphenyls (PCBs), polycyclic aromatic hydrocarbons (PAHs), and pesticides. Sixty chemicals were detected in either dust or air and here we report 58 SVOCs detected in dust for the first time. In dust, phthalates (bis(2-ethylhexyl) phthalate, benzyl butyl phthalate, di-*n*-butyl phthalate) and flame retardants (PBDE 99, PBDE 47) were detected at the highest concentrations relative to other chemicals at the 95th percentile, while phthalates were highest at the median. Because SVOCs are found in both gas and condensed phases and redistribute from their original source over time, partitioning models can clarify their fate indoors. We use empirical data to validate air-dust partitioning models and use these results, combined with experience in SVOC exposure assessment, to recommend residential exposure measurement strategies. We can predict dust concentrations reasonably well from measured air concentrations ( $R^2 = 0.80$ ). Partitioning models and knowledge of chemical  $K_{oa}$  elucidate exposure pathways and suggest priorities for chemical regulation. These findings also inform study design by allowing researchers to select sampling approaches optimized for their chemicals of interest and study goals. While surface wipes are commonly used in epidemiology studies because of ease of implementation, passive air sampling may be more standardized between homes and also relatively simple to deploy. Validation of passive air sampling methods for SVOCs is a priority.



## INTRODUCTION

Exposures at home can dominate total exposure for a variety of consumer product chemicals, including some flame retardants, phthalates, and pesticides.<sup>1–4</sup> These observations are consistent with Wambaugh et al.'s study that found consumer products to be a strong predictor of biological levels, based on a high throughput exposure model of over 1900 commercial chemicals.<sup>5</sup> These studies show the importance of being able to predict exposures to consumer product chemicals but also highlight the limited measurement data and gaps in our understanding of these exposure pathways.<sup>6</sup> Given the vast number of chemicals in commercial use, improving models to predict exposure levels and pathways is a priority in order to evaluate and manage health risks.

Fate and transport models have been widely developed and validated to estimate exposure from ambient emissions, but models relevant to indoor exposures are less developed. In a recent comparison of available exposure models, researchers identified the lack of data concerning near-field exposures to

consumer product chemicals as a major gap in knowledge needed for better exposure-based chemical prioritization.<sup>7</sup> Specifically, exposure measurements from indoor environments are needed to improve and validate models for near-field exposure.

Many consumer product chemicals of current and emerging health concern are classified as semivolatile organic compounds (SVOCs), including flame retardants, phthalates, pesticides, and perfluorinated compounds. SVOCs are found in both the gas and condensed phases and redistribute from their original source over time to indoor air, house dust, and other indoor surfaces.<sup>8</sup> Their distribution in the indoor environment determines how people are exposed, so characterizing this

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distribution informs model development, sampling approaches, and strategies for intervention to reduce exposure.

A few exposure studies have measured a broad range of SVOCs indoors. We previously analyzed house dust and indoor air samples from 120 homes on Cape Cod for 89 semivolatile endocrine disrupting compounds (EDCs), including phthalates, alkylphenols, parabens, flame retardants, polychlorinated biphenyls (PCBs), and pesticides.<sup>9</sup> We extended this research to 50 homes in northern California<sup>10</sup> and found that indoor air concentrations were substantially higher than outdoor concentrations for most of the 104 SVOCs measured, indicating that indoor sources dominate total exposure.<sup>11</sup> Building on this work, Blanchard et al. recently reported concentrations of 57 SVOCs in air and dust in 30 French homes.<sup>12</sup> Comprehensive residential exposure measurement studies such as these are resource intensive, so environmental health researchers and chemical regulators have called for validated methods for estimating exposures.<sup>4,6,13,14</sup> In response, Weschler and Nazaroff developed a series of equilibrium partitioning models based on physical-chemical properties to describe the fate of SVOCs in indoor environments.<sup>8,15–18</sup>

In this paper, we analyze relationships between simultaneously measured air and dust levels in the 49 northern California (CA) homes. The dust measures are reported here for the first time. We use indoor air and dust measures to evaluate the equilibrium partitioning models developed by Weschler and Nazaroff. This validation supports U.S. Environmental Protection Agency (EPA) efforts to develop high throughput exposure models.<sup>5–7,19</sup> In addition, we use these partitioning models and our measurement data to demonstrate how chemical behavior can be anticipated based on chemical properties, and to provide other researchers with guidelines for efficient sampling design. For example, we demonstrate how these partitioning models can be used to select sampling methods based on chemical properties and to predict concentrations in different media.

## METHODS

**Sampling and Analytical Methods.** The California Household Exposure Study (CAHES) collected indoor and outdoor air and house dust in 2006 from 50 nonsmoking homes in Richmond and Bolinas, California. Richmond is a predominately low income urban community on the northeast coast of San Francisco Bay, whereas Bolinas is more rural and located on the Pacific coast north–northwest of San Francisco. Additional information about the study communities and participant selection can be found elsewhere.<sup>10</sup> We were able to collect 49 dust samples from the 50 study homes. We previously reported concentrations of polybrominated diphenyl ethers (PBDEs) and 36 other flame retardants in dust samples collected in a subset of homes from this study ( $n = 16$ ) in 2006 and then again in 2011.<sup>20</sup> Indoor and outdoor air samples were also collected and have previously been reported.<sup>11</sup> Briefly, air samples comprising gas and particle-bound phases were collected using parallel 160 mm URG personal pesticide samplers (Universal Research Glassware (URG); Chapel Hill, NC) at a target flow rate of 8–9 L/min over 24 h. Each sampler contained a 10  $\mu\text{m}$  at 4 L/min impactor-equipped inlet followed by a 25 mm quartz fiber filter and 3 g XAD-2 sandwiched between two 1 13/16 in. polyurethane foam plugs. We collected respirable particulate ( $\text{PM}_{2.5}$ , particulate matter less than 2.5  $\mu\text{m}$  in diameter) on Teflon filters at a flow rate of approximately 5 L/min in 42 homes.

Dust samples were collected using a 9 A Eureka Mighty-Mite vacuum cleaner modified to collect dust into a 19  $\times$  90 mm<sup>2</sup> cellulose extraction thimble (Whatman, Inc.; Clifton, NJ) within a polytetrafluoroethylene Teflon crevice tool. Dust sampling began immediately following termination of the air sampling. Sample collection was accomplished by slowly dragging the crevice tool just above the surface of rugs, upholstery, wood floors, windowsills, ceiling fans, and furniture in the primary living areas of the home for  $\sim 30$  min. At the completion of sampling, thimbles containing the collected dust were removed and placed in precleaned, certified glass jars with Teflon-lined lids and stored at  $-4$  °C prior to shipping to the laboratory.

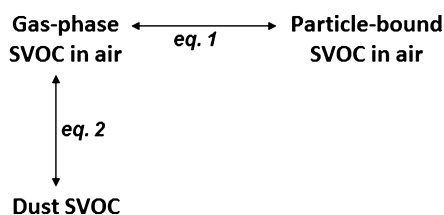
Chemical analysis of the dust and air samples was conducted at the Southwest Research Institute in San Antonio, TX. Samples were stored for less than 6 months at  $< -4$  °C prior to extraction and analysis. A total of 79 compounds were targeted in the dust samples, including pesticides, phthalates, polycyclic aromatic hydrocarbons (PAHs), PBDEs, and polychlorinated biphenyls (PCBs). Dust samples passed through a  $< 150$   $\mu\text{m}$  sieve prior to analysis. Approximately 0.5 g of dust (median) was analyzed by GC/MS in selected ion monitoring mode after Soxhlet extraction using 6% diethyl ether in hexane, concentration to 2.5 mL, florisil column cleaning, and final concentration to 2 mL in 1% ether. Air samples were extracted across the entire URG setup and represent both gas and particle-bound phases. Additional details on extraction and analytical techniques are provided elsewhere.<sup>9,21</sup>

Quality assurance/quality control (QA/QC) measures were conducted to ensure accuracy and reliability of measurements. To estimate precision, we split three dust samples. To evaluate contamination from the laboratory, we analyzed solvent blanks ( $n = 3$ ). Matrix spikes ( $n = 3$ ) and surrogate recoveries were also used to characterize accuracy, compound recovery from the matrix, and extraction efficiency. Additional QA/QC information and results are presented in Supporting Information (SI).

**Statistical Methods.** For each analyte, the method reporting limit (MRL) was defined as either the analytical detection limit or the 90th percentile of the solvent blank concentrations, whichever was larger. Values reported by the laboratory but below the MRL were not included in the detection frequencies but were treated as estimated values to visualize distributions. Summary statistics use blank-corrected values, whereas modeling uses uncorrected values.

Data tended to be left censored due to detection limits from the laboratory analysis, which means that Pearson or Spearman correlation estimates using arbitrary substitutions for non-detects (e.g., detection limit/2) will result in poor correlation estimates. Therefore, we calculated Kendall's  $\tau$  rank correlation coefficients, adjusted for censored data,<sup>22</sup> to explore linear relationships between dust and air concentrations, with  $p$ -values obtained from 10 000 bootstrap replications. Kendall  $\tau$  correlation estimates were calculated for air and dust analytes with at least three detected paired concentration values.

**Partitioning Theory.** We used equilibrium partitioning concepts to explore relationships between concentrations in air and mass fractions in household dust (see Figure 1). Weschler and Nazaroff tested this model using aggregate data (e.g., published means and medians) available in the literature.<sup>15</sup> Partitioning theory tells us that gas-phase chemicals will readily and predictably partition between airborne particles, surfaces, and house dust in the home. Air measurements in this study



**Figure 1.** Schematic illustration of SVOCs in air and dust indoors. Equations 1 (partitioning between gas-phase and total SVOCs in air) and 2 (partitioning between gas-phase air and dust) are described in the text.

combined gaseous and particle phases; however, for equilibrium conditions, partitioning applies to gas-phase air concentrations and mass fractions in dust. We estimate gas-phase concentrations ( $C_g$ ,  $\mu\text{g}/\text{m}^3$ ) from measured total air concentrations in a manner similar to Weschler and Nazaroff<sup>15</sup> using the following equation:

$$C_g = \frac{C_t}{1 + \frac{\text{TSP}}{\rho_{\text{part}}} \times f_{\text{om\_part}} \times K_{\text{oa}}} \quad (1)$$

where  $C_t$  is the measured total air concentration ( $\mu\text{g}/\text{m}^3$ ), TSP is the total suspended particles or average indoor concentration of particles ( $\mu\text{g}/\text{m}^3$ ),  $\rho_{\text{part}}$  is the density of airborne particles, assumed to be  $1 \times 10^6 \text{ g}/\text{m}^3$ ,  $f_{\text{om\_part}}$  is the fraction of organic matter associated with particles, assumed to be 0.4 (unitless), and  $K_{\text{oa}}$  is the octanol-air equilibrium partitioning coefficient. We assumed the same values for particle density and fraction organic matter as Weschler and Nazaroff.<sup>15</sup> Values for  $\log K_{\text{oa}}$ , the ratio of the concentration of a chemical in octanol (a surrogate for organic matter) to the concentration in air at equilibrium were obtained from the KOAWIN program in Estimation Programs Interface (EPI) Suite Version 4 developed by U.S. EPA and Syracuse Research Corporation (see SI Table S2).<sup>23</sup> Estimated rather than experimental  $K_{\text{oa}}$  values were used because authoritative experimental values were not available for all SVOCs we measured.

We estimated home-specific TSP—concentration of particles with the same cut-point as particles represented in total SVOC air concentrations—from  $\text{PM}_{2.5}$  concentrations measured for each home. In this study, the approximate particle size cut-point, based on URG sampler design and average flow rate, was  $6.9 \mu\text{m}$ . Since we measured indoor  $\text{PM}_{2.5}$  for the duration of SVOC air sample collection, we used this measurement to estimate average particle concentration for  $<6.9 \mu\text{m}$  particles using published data. Specifically, we estimated concentrations of particulate matter across multiple size fractions from a study conducted in the greater Boston area.<sup>24</sup> Using these data, we developed a linear regression model between measured  $\text{PM}_{2.5}$  and  $\text{PM}_{6}$ , the closest size fraction to  $6.9 \mu\text{m}$  available. We applied measured  $\text{PM}_{2.5}$  data from this study (median =  $5.6 \mu\text{g}/\text{m}^3$ ) to the linear model [ $\log(\text{PM}_6 \text{ concentration, } \mu\text{g}/\text{m}^3) = 0.43 + 0.89 \log(\text{PM}_{2.5} \text{ concentration, } \mu\text{g}/\text{m}^3)$ ] derived from the Long et al. data to estimate average indoor mass of  $\text{PM}_6$  in the CA residences (SI Table S3). When residence-specific  $\text{PM}_{2.5}$  concentrations were not available ( $n = 8$ ), we used the estimated average  $\text{PM}_6$  concentration for the study ( $12 \mu\text{g}/\text{m}^3$ ).

In a manner similar to work by Weschler and Nazaroff, we used SVOC air concentrations to predict dust concentrations ( $X_{\text{dust}}$ ,  $\mu\text{g}/\text{g}$ ) assuming equilibrium partitioning between gas-phase air concentrations and settled dust:

$$X_{\text{dust}} = \frac{C_g \times f_{\text{om\_dust}} \times K_{\text{oa}}}{\rho_{\text{dust}}} \quad (2)$$

where  $C_g$  is mass concentration of gas-phase SVOC ( $\text{ng}/\text{m}^3$ ),  $f_{\text{om\_dust}}$  is the fraction of dust that is organic matter (unitless), and  $\rho_{\text{dust}}$  is the density of dust (typically in the range of  $1\text{--}2.5 \times 10^6 \text{ g}/\text{m}^3$ ). As Weschler and Nazaroff assert, this theoretical relationship assumes equilibrium partitioning that is dominated by the physical-chemical process of absorption in the organic fraction of dust and that octanol is an appropriate chemical model for the organic matter in dust, as it relates to sorption.<sup>15</sup> Like Weschler and Nazaroff, we assumed house dust to have an organic matter fraction of 0.2 and a density of  $2 \times 10^6 \text{ g}/\text{m}^3$ .<sup>15</sup> We compared predicted and measured dust concentrations using regression models with median concentrations for each chemical.

## RESULTS AND DISCUSSION

**Measured Dust Concentrations.** Overall, we detected 58 target analytes in house dust, which represents a long-term reservoir for chemicals in the residential environment. Table 1 presents CAHES house dust concentrations and includes all chemicals analyzed, even if they were not observed above the MRL. Phthalates and flame retardants are particularly abundant; DEHP, BBP, PBDE 99, PBDE 47, and DBP were the five chemicals with the highest concentrations at the 95<sup>th</sup> percentile. We found flame retardants at some of the highest concentrations in the world; whereas, several phthalates—DEHP and DBP—and PAHs were generally found at lower levels than reported elsewhere (see Discussion below). PCB and pesticide levels were not substantially different compared to other available studies. Chemicals that have been banned for years (e.g., DDT) persist in homes where exposure continues because of limited degradation. SI Figure S2 shows measured air and dust concentrations for chemicals concurrently detected in both media. Below, we summarize the dust concentrations by chemical group, and compare concentrations measured in this study to the Cape Cod Household Exposure Study (1999–2001), which used the same sampling and analytical methods. We also compare concentrations to levels reported in other peer-reviewed studies, but note that differences may be due to different sampling methods as well as geographic and temporal variation in chemical use.

**Phthalates.** All 11 target phthalates were detected in house dust samples. Most phthalates were detected in  $>90\%$  of the dust samples, with the exception of DPeP, DPP, and DCP. The median concentrations of phthalates were lower than those measured in Cape Cod in 1999–2001 for all but one phthalate, DIBP (range 1.1 to 320, median  $4.4 \mu\text{g}/\text{g}$ ).<sup>9</sup> Concentrations of DBP, BBP, DEHP, and DEP were substantially lower at the median than those observed in 30 Berlin apartments tested in 2000 and 2001<sup>25</sup> and Canadian homes collected between 2007 and 2010,<sup>26</sup> but similar to U.S. concentrations reported in 2013 by Shin et al.<sup>27</sup> Kolarik et al. and Blanchard et al. recently summarized house dust concentrations of DEHP, BBP, and DBP across different countries.<sup>12,28</sup> DEHP and DBP dust concentrations in our study were generally lower than those reported in other countries; whereas, median BBP concentrations in this study fall within the wide range of concentrations reported in the literature.<sup>12,28,29</sup>

**Flame Retardants.** PBDEs (PentaBDE congeners) were detected in nearly 100% of dust samples at levels higher than

Table 1. Summary Statistics for Semivolatile Organic Compounds in Household Dust in California Homes ( $\mu\text{g}/\text{g}$ ) ( $n = 49$ )

compound	abbrev.	%>MRL (MRL) <sup>b</sup>	min.	median	95 <sup>th</sup> percentile	max. <sup>e</sup>
phthalates						
benzyl butyl phthalate <sup>c</sup>	BBP	98 (0.2)	–	19	220	330
bis(2-ethylhexyl) adipate	DEHA	100 (0.04)	1.1	5.1	14	24
bis(2-ethylhexyl) phthalate <sup>c</sup>	DEHP	100 (0.4)	50	140	460	800
di- <i>n</i> -butyl phthalate <sup>c</sup>	DBP	98 (0.9)	–	11	35	56
di- <i>n</i> -hexyl phthalate	DHP	96 (0.04)	–	0.66	5.7	110
di- <i>n</i> -octyl phthalate	DOP	100 (0.02)	0.42	1.6	3.9	9.4
di- <i>n</i> -pentyl phthalate <sup>d</sup>	DPeP	12 (0.04)	–	–	0.59	2.2
di- <i>n</i> -propyl phthalate	DPP	2 (0.06)	–	–	–	0.095
dicyclohexyl phthalate <sup>d</sup>	DCP	16 (0.04)	–	–	7.4	13
diethyl phthalate <sup>c</sup>	DEP	96 (0.1)	–	2.1	6.3	85
diisobutyl phthalate <sup>c</sup>	DIBP	100 (0.2)	1.1	4.4	12	320
flame retardants						
polybrominated diphenyl ether 47	PBDE47	100 (0.03)	0.11	2.7	39	110
polybrominated diphenyl ether 99	PBDE99	100 (0.03)	0.098	3.8	47	170
polybrominated diphenyl ether 100	PBDE100	94 (0.04)	–	0.68	9.1	31
tris(2,3-dibromopropyl) phosphate <sup>d</sup>	TrisBP	8 (0.01)	–	–	0.032	0.072
polychlorinated biphenyls						
polychlorinated biphenyl 52 <sup>d</sup>	PCB52	35 (0.02)	–	–	0.13	0.32
polychlorinated biphenyl 105 <sup>d</sup>	PCB105	33 (0.02)	–	–	0.18	0.27
polychlorinated biphenyl 153 <sup>d</sup>	PCB153	55 (0.02)	–	0.022	0.32	0.54
polycyclic aromatic hydrocarbons						
acenaphthene <sup>d</sup>	AcNThe	24 (0.02)	–	–	0.033	0.034
acenaphthylene	AcNThy	0 (0.02)	–	–	–	–
anthracene <sup>d</sup>	Anth	29 (0.02)	–	–	0.043	0.064
benzo(a)anthracene <sup>d</sup>	BaA	86 (0.02)	–	0.047	0.13	0.2
benzo(a)pyrene <sup>d</sup>	BaP	90 (0.007)	–	0.085	0.19	0.26
benzo(b&j)fluoranthene <sup>d</sup>	BbjFluAn	98 (0.01)	–	0.14	0.33	0.35
benzo(k)fluoranthene <sup>d</sup>	BkFluAn	92 (0.007)	–	0.06	0.17	0.37
benzothiophene	BThPhe	0 (0.03)	–	–	–	–
chrysene/ <i>iso</i> -chrysene <sup>d</sup>	Chrys	96 (0.1)	–	0.15	0.34	0.45
dibenz(a,e)pyrene	DBaePyr	2 (0.03)	–	–	–	0.057
dibenz(a,h)anthracene <sup>d</sup>	DBahA	39 (0.02)	–	–	0.081	0.079
3,6-dimethyl phenanthrene	DMPhenan	2 (0.02)	–	–	–	0.023
fluoranthene	FluAn	100 (0.007)	0.078	0.18	0.39	0.62
fluorene <sup>d</sup>	Flu	76 (0.007)	–	0.023	0.057	0.086
indeno(1,2,3- <i>cd</i> )pyrene <sup>d</sup>	IcdPyr	57 (0.02)	–	0.072	0.18	0.18
1-nitropyrene	1NPyr	0 (0.04)	–	–	–	–
phenanthrene <sup>c</sup>	Phenan	98 (0.009)	–	0.19	0.4	0.62
pyrene	Pyr	100 (0.007)	0.064	0.18	0.42	0.48
dibenzothiophene	DBTPhe	2 (0.03)	–	–	–	0.85
4,6-dimethyl dibenzothiophene <sup>d</sup>	DMDBTPhe	20 (0.03)	–	–	0.26	1
2-methyl dibenzothiophene <sup>d</sup>	2MDBTPhe	22 (0.03)	–	–	0.27	2.1
1-methyl phenanthrene <sup>d</sup>	1MPhenan	96 (0.01)	–	0.05	0.2	0.38
2-methyl phenanthrene <sup>d</sup>	2MPhenan	98 (0.007)	–	0.083	0.27	0.38
3-methyl phenanthrene <sup>d</sup>	3MPhenan	96 (0.01)	–	0.081	0.3	0.58
9-methyl phenanthrene <sup>d</sup>	9MPhenan	96 (0.01)	–	0.058	0.3	0.69
pesticides						
alachlor	Alach	0 (0.04)	–	–	–	–
aldrin	Aldr	0 (0.04)	–	–	–	–
atrazine	Atraz	0 (0.02)	–	–	–	–
bendiocarb <sup>d</sup>	Bendio	4 (0.1)	–	–	–	0.35
carbaryl	Carb	14 (0.06)	–	–	0.85	1.8
carbofuran	Crbfur	0 (0.04)	–	–	–	–
$\alpha$ -chlordane <sup>d</sup>	aChlor	61 (0.02)	–	0.02	0.16	0.2
$\gamma$ -chlordane <sup>d</sup>	gchlor	61 (0.02)	–	0.021	0.15	0.2
chlorothalonil	Chorth	43 (0.02)	–	–	0.53	1.2
chlorpyrifos <sup>d</sup>	ChlPy	51 (0.02)	–	0.022	0.24	0.61
cyanazine	Cyan	0 (0.06)	–	–	–	–
cypermethrin	Cyper	16 (0.1)	–	–	7.5	140
4,4'-dichlorodiphenyldichloroethane <sup>d</sup>	DDD	59 (0.02)	–	0.027	0.21	0.32

Table 1. continued

compound	abbrev.	%>MRL (MRL) <sup>b</sup>	min.	median	95 <sup>th</sup> percentile	max. <sup>e</sup>
		pesticides				
4,4'-dichlorodiphenyldichloroethylene <sup>d</sup>	DDE	76 (0.01)	–	0.046	0.17	0.29
4,4'-dichlorodiphenyltrichloroethane	DDT	86 (0.02)	–	0.33	1.9	2.4
diazinon <sup>d</sup>	Diaz	16 (0.02)	–	–	0.26	5.3
dicofol	Dico	0 (0.04)	–	–	–	–
dieldrin	Dield	0 (0.04)	–	–	–	–
endrin	Endr	0 (0.04)	–	–	–	–
ethyl parathion	Parath	0 (0.1)	–	–	–	–
heptachlor	Hept	0 (0.02)	–	–	–	–
lindane	Lind	2 (0.04)	–	–	–	0.41
malathion	Malth	0 (0.02)	–	–	–	–
methoxychlor <sup>d</sup>	MX	57 (0.04)	–	0.073	0.92	1.9
methyl parathion	MePthion	0 (0.04)	–	–	–	–
metolachlor	Metol	0 (0.02)	–	–	–	–
nitrofen	Nitrof	0 (0.04)	–	–	–	–
<i>cis</i> -permethrin	cPerm	98 (0.02)	–	0.87	17	160
<i>trans</i> -permethrin	tPerm	98 (0.03)	–	1	28	280
piperonyl butoxide	PipBO	88 (0.02)	–	0.14	8.3	110
<i>o</i> -phenyl phenol <sup>d</sup>	oPPh	96 (0.01)	–	0.082	0.52	0.65
prometon	Prom	0 (0.04)	–	–	–	–
propoxur <sup>d</sup>	PrPx	57 (0.08)	–	0.12	1.5	2
simazine	Simz	0 (0.04)	–	–	–	–
trifluralin	Trifl	0 (0.02)	–	–	–	–
4-nitrotoluene	4NT	0 (0.04)	–	–	–	–

<sup>a</sup>– indicates insufficient number of detects to calculate summary statistic. <sup>b</sup>MRL = method reporting limit (defined as either the analytical detection limit or the 90<sup>th</sup> percentile of the solvent method blanks, whichever is larger). <sup>c</sup>Values subject to blank correction by subtracting the median blank concentration. <sup>d</sup>Indicates that elevated nondetect values (due to analytical interferences) are included in the summary statistics presented. <sup>e</sup>Nondetect with elevated detection limit (due to analytical interferences) excluded.

levels reported outside of California, likely the result of California's unique furniture flammability standard, and at levels that may pose a health risk, especially for children.<sup>20,30,31</sup> The concentrations we detected were similar—within a factor of 2—to other California studies,<sup>29,32</sup> and we have previously published detailed analysis and discussion of these findings, including an analysis of time trends in relation to phase out of PentaBDE and its substitution with other flame retardant chemicals.<sup>20</sup>

**Polychlorinated Biphenyls.** Three PCBs were targeted in this analysis: PCB 52, PCB 105, and PCB 153. PCBs were detected at higher concentrations in the Cape Cod study versus CAHES, but were more frequently detected in CAHES because the MRL was approximately 10 times lower (MRL = 0.02  $\mu\text{g/g}$ ) than in the Cape Cod study.<sup>33</sup> Median levels of PCB 153 in this study were higher than the geometric mean (GM) and arithmetic mean (AM) from dust in 212 California residences collected in 2001–2006.<sup>34</sup> Ninety-fifth percentile concentrations of PCB 105 and 153 in the CAHES were also higher than concentrations reported in the review by Roberts et al.<sup>35</sup>

**Polycyclic Aromatic Hydrocarbons.** PAHs were detected frequently in dust samples, with 13 PAHs having detection frequencies >75%. Dust concentrations of benz(a)anthracene, benzo(a)pyrene, and pyrene in this study were approximately 10 times lower at the median than the reported concentrations in the Cape Cod study;<sup>9</sup> however, benz(a)anthracene and benzo(a)pyrene concentrations are comparable to samples collected in Los Angeles using similar methods.<sup>36</sup> While the concentrations of PAHs in settled house dust in a review of 18 published studies revealed substantial variation, summarized GMs were generally 2–3 times higher than those observed in this study.<sup>37</sup>

**Pesticides.** Eighteen of the 36 pesticides analyzed were detected. Several historic use pesticides, including DDT, DDE, DDD, chlordane, and methoxychlor, were detected in the majority of homes. Common pyrethroids in household pesticides, *cis*- and *trans*-permethrin, were detected in almost all dust samples (98%). The maximum dust concentrations for historic use pesticides were generally substantially lower than those in the Cape Cod study. Methoxychlor, an organochlorine insecticide banned between the sampling periods for the Cape Cod study and CAHES in 2003,<sup>38</sup> was detected at concentrations approximately 3 times lower at the median (0.24 versus 0.07  $\mu\text{g/g}$ ) in this study and approximately 7 times lower at the maximum (12.9 versus 1.9  $\mu\text{g/g}$ ). Bendiocarb, voluntarily canceled in 1999,<sup>39</sup> had a substantially lower maximum concentration compared to the Cape Cod study (40.7 versus 2.3  $\mu\text{g/g}$ ). DDE concentrations measured in the CAHES study were higher than those measured in vacuum bags collected from Davis, CA apartments; however, the chlordane concentrations were comparable.<sup>29</sup>

**Relationships between Indoor Air and Household Dust.** We used measured dust and air data to describe the relationship between air and dust concentrations in the home and to determine if one measure can be used to predict the other, which would simplify and reduce costs related to residential exposure assessments. Seventy-six compounds were analyzed in both dust and air: 11 phthalates, 3 PBDEs, 3 PCBs, 24 PAHs, and 36 pesticides. Of these, 60 were detected in either dust or air; 40 were simultaneously detected in dust and air in at least one home; and 17 were detected in greater than 50% of air and 50% of dust samples (SI Figure S2).

**Correlation between Air and Dust.** We first estimated correlations between measured air and dust concentrations for

each compound with sufficient pairs of detected concentrations. We show these correlations on scatterplots sorted by  $\log K_{oa}$ , which is a predictor of air-dust partitioning (SI Figure S3). Of the 34 analytes with sufficient data for comparison, 25 had significant positive Kendall's  $\tau$  correlation estimates ( $p < 0.05$ ). Sorting correlation estimates by  $\log K_{oa}$  did not reveal any clear patterns across chemical classes. Phthalates with significant correlation estimates were DEP, DIBP, DBP, and BBP. Three phthalates with the highest  $\log K_{oa}$  values (DHP, DEHP, and DEHA) were not correlated. The only flame retardant with sufficient data for comparison (PBDE 47) was significantly positively correlated. Correlation estimates for PCB52 and PCB105 were similar; however, the estimate for PCB52, which has a lower  $\log K_{oa}$ , was significant ( $p = 0.03$ ), while the estimate for PCB105 was only marginally significant ( $p = 0.08$ ). PAHs with significant correlations were as follows: acenaphthene; fluoranthene; phenanthrene; dibenzothiophene; 3-, 9-, and 1-methylphenanthrene; 2-methyl dibenzothiophene; and fluoranthene. PAHs without significant correlations were as follows: anthracene; 2-methylphenanthrene; pyrene; and 4,6-dimethyl dibenzothiophene. Pesticides with significant positive correlation estimates were as follows: chlorothalonil, *o*-phenylphenol, chlorpyrifos,  $\alpha$ -chlordane, diazinon, DDE,  $\gamma$ -chlordane, DDT, *cis*-permethrin, and piperonyl butoxide. Propoxur is the only pesticide without a significant correlation.

**Dust/Air Ratios.** To see the relative concentrations in house dust and air across chemicals, we calculated ratios of measured dust and total (measured) air concentrations for 40 compounds with simultaneous detects (SI Figure S4). Ratios span 6 orders of magnitude, with compounds with lower  $\log K_{oa}$  values having smaller ratios and compounds with higher  $\log K_{oa}$  values having higher ratios. There was a moderately strong correlation ( $\rho \approx 0.8$ ) between  $\log K_{oa}$  and ratio of dust to air concentrations. Also, compounds with higher  $\log K_{oa}$  values ( $>10$ ) had lower detection frequencies in air, except for the two phthalates DEHP and DEHA, which are so abundant that they were detected in most air and dust samples.

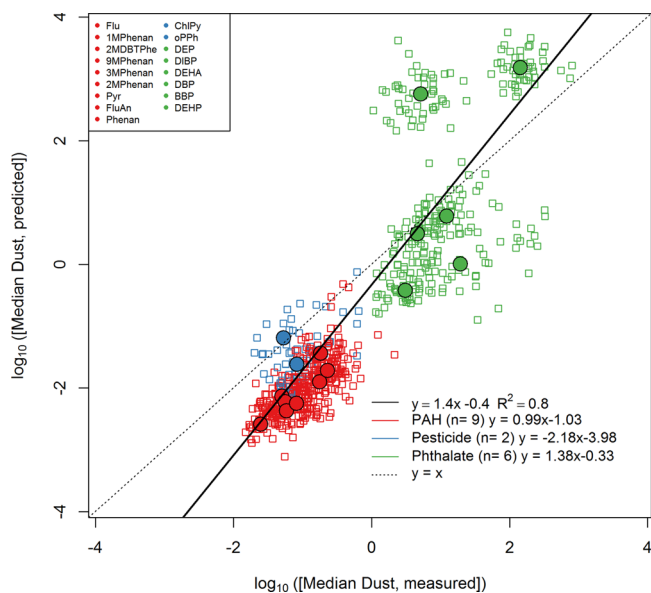
**Exposure Rate by Pathway.** We also estimated the relative importance of dust exposure versus other household exposure pathways by calculating exposure rates (ng/day) for dust ingestion, dermal exposure through dust adherence, and inhalation of indoor air (equations in SI). We assumed an inhalation rate of 14.9 m<sup>3</sup>/day, an absorption fraction for air of 0.5, a dust ingestion rate of 0.064 g/day, a gastrointestinal absorption fraction of 0.9, a dermal loading of 3.55 g/m<sup>2</sup>, a dermal transfer coefficient of 0.06 m<sup>2</sup>/h, and a dermal absorption fraction of 0.05 averaged over a 70 year lifespan.<sup>40,41</sup> We calculated the fraction exposure to dust relative to total indoor exposure for 40 compounds with simultaneous detects (SI Figure S5). Dermal exposure via gas-phase absorption<sup>16,42</sup> is not accounted for in this model, potentially underestimating the air exposure contribution, especially for compounds with lower  $K_{oa}$  values. For lower molecular weight compounds, such as fluorene and DEP, inhalation is the main route of exposure; whereas, for compounds with larger  $\log K_{oa}$  values such as DEHP and DEHA, exposure to dust (dermal and ingestion) dominates. However, for the majority of the SVOCs with a wide range of  $\log K_{oa}$  values, exposure to dust and air are both important.

**Partitioning between Gas and Particle Phase in Air.** Our analysis of dust:air concentrations in relation to  $\log K_{oa}$  (SI Figure S4) shows that chemicals with high  $\log K_{oa}$  tend to be more abundant in dust, while those with low  $\log K_{oa}$  are more

abundant in air. We next used the Weschler and Nazaroff partitioning model to see whether indoor air and dust relationships are predictable so that one can be estimated from the other. To do this, we used the partitioning model to separate the measured air concentrations into gas-phase and particle-phase concentrations. SI Figure S6 shows the relationship of estimated gas to measured total air concentration by  $\log K_{oa}$ . Using our data, we see that measured total air concentrations comprise mostly gas-phase concentrations (gas-phase to total air concentration ratios near 1) for  $\log K_{oa}$  values up to approximately 10, consistent with Weschler and Nazaroff and Schossler et al.<sup>15,18</sup> At higher  $\log K_{oa}$  values, there is increased variability in the contribution of gas-phase to total air concentrations, and, at the highest  $\log K_{oa}$  values, total air concentrations comprise mostly particulate phase concentrations. This is expected since higher  $\log K_{oa}$  values indicate greater partitioning to the octanol phase, a proxy for organic matter in particulates.

**Relationship of Gas Phase Air to Dust Concentrations.** Looking at the relationship between estimated gas-phase air concentrations and dust concentrations, we found a significant positive correlation across all compounds ( $\tau = 0.37$ ,  $p < 0.05$ ). A mixed-effects model of  $\log$  dust and  $\log$  gas-phase air concentrations with chemical designated as a random effect also revealed a significant relationship between air and dust ( $\beta = 0.34$ ,  $p < 0.001$ ). The estimated  $R^2$  value for the mixed-effects model ( $R^2 = 0.05$ ) indicates that a substantial portion of variation in the gas-phase air concentrations cannot be explained by measured dust concentrations alone. We expanded the mixed-effects model to include  $\log K_{oa}$ . As expected based on the apparent relationship between dust/air concentrations and  $\log K_{oa}$  (SI Figure S4), there is a significant negative association between  $\log K_{oa}$  estimates and  $\log$  gas-phase air concentrations. Approximately 48% of the variation in air concentrations is explained by dust and  $\log K_{oa}$  estimates. SI Figure S7 shows dust and gas-phase air concentrations categorized by rounded  $\log K_{oa}$  values. Compounds with lower relative  $\log K_{oa}$  values (5–7) tend to lie toward the upper left portion of the graph, with higher relative air concentrations and lower relative dust concentrations; whereas, compounds with higher relative  $\log K_{oa}$  values (10–12) tend to lie toward the bottom right portion of the graph, with lower relative air concentrations and higher relative dust concentrations.

**Partitioning Model Validation.** This is the first use of a SVOC partitioning model applied to a range of SVOCs measured simultaneously in indoor air and house dust in the same study. We used our air and dust measurements for 17 chemicals to evaluate Weschler and Nazaroff's air-dust partitioning model (eq 2). We used our estimated gas-phase concentrations to predict dust concentrations and then compared predictions to measured dust concentrations. We predicted dust concentrations for 17 compounds, mostly PAHs and phthalates, detected in at least 50% of air and 50% of dust samples (Figure 2). The predicted and measured concentrations are well correlated ( $R^2 = 0.8$ ), with a slope slightly greater than one ( $\beta = 1.4$ ). On the basis of the intercept ( $-0.4$ ), the predicted dust concentrations are, on average, 2.5 times lower than measured concentrations. In fact, almost all of the compounds are under-predicted at the median with the exception of DEHA, DEHP, and chlorpyrifos. When DEHP and DEHA are excluded from the model, the model fit improves ( $R^2 = 0.85$ ), but the predicted concentrations are, on average, now 6 times lower than the measured concentrations



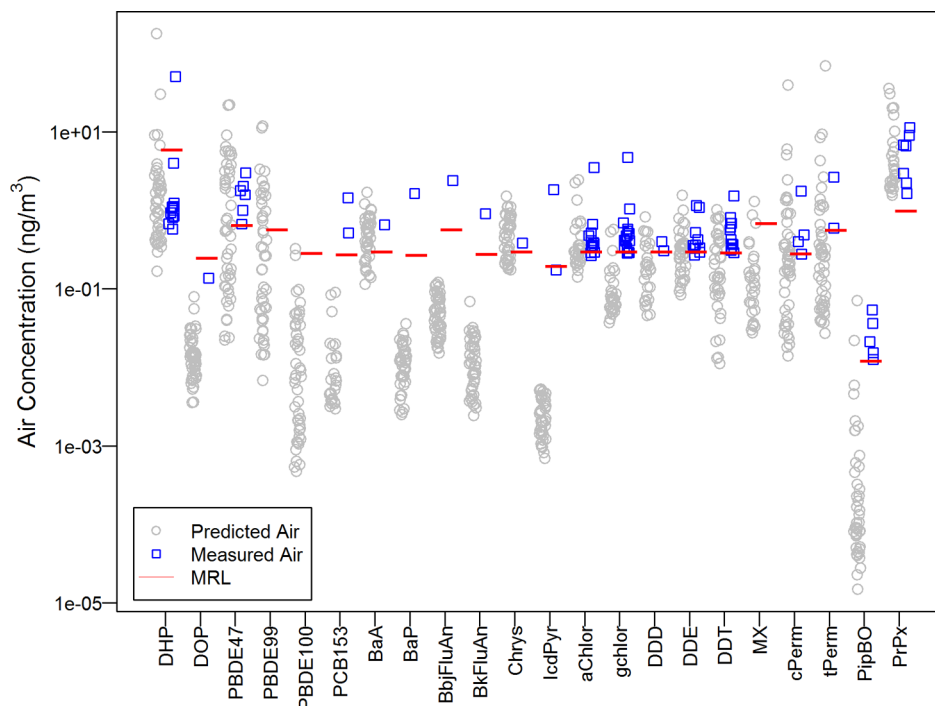
**Figure 2.** Predicted dust concentrations (micrograms/gram) versus measured dust concentrations (micrograms/gram) for 17 SVOCs with at least 50% detection frequency in both indoor air and dust. Predictions made using measured air concentrations in the same homes. Dashed line represents 1:1 line or perfect prediction. Individual points shown (unshaded symbols); however, regression model fit to median concentrations (large shaded circles).

(results not shown). These data were also visualized by plotting the ratios of predicted to measured dust concentrations sorted by  $\log K_{oa}$  (SI Figure S8). There is variability in the ratios of predicted and measured for each compound, with over a 10-

fold range for some compounds. Also apparent is the under-prediction for PAHs and variability for phthalates.

The model performed reasonably well by predicting 80% of the variability in dust concentrations. On average, dust levels were under-predicted for most chemicals and this is particularly true for PAHs. There are several possible explanations for this: (1) sampling artifacts that result in underestimate of air concentrations; (2) oxidation of PAHs on the air samples over the 24 h sampling period; (3) the origin of the measured dust particles since particles of ambient origin may have higher levels of PAHs relative to other chemicals with primarily indoor sources; and (4) sorption behaviors of PAHs to dust particles, which is dependent upon the organic carbon vs black carbon nature of the particle. Phthalates were both over- and under-predicted in dust depending on the compound. DEHP and DEHA, which have  $\log K_{oa}$  values  $>12$ , were overpredicted, whereas the other phthalates—DEP, DIBP, DBP, and BBP—which have  $\log K_{oa}$  values  $<10$ , were under-predicted.

A potential explanation for imprecise prediction is inaccurate  $K_{oa}$  values. We used estimated  $\log K_{oa}$  values obtained from the KOAWIN program in EPISuite. These estimated values likely have the largest potential uncertainty in the prediction models.<sup>15</sup> While the overall accuracy of  $K_{oa}$  estimation is good ( $R^2 = 0.957$ ),<sup>43</sup> estimates may be quite discrepant for specific chemicals. For example, the estimated  $\log K_{oa}$  value for BBP from KOAWIN is 9.018; however, Weschler and Nazaroff, using the SPARC online calculator report an estimated  $\log K_{oa}$  value for BBP of 11.6.<sup>15</sup> These different  $K_{oa}$  values lead to substantially different dust concentration predictions (over 300-fold). Similarly, perhaps the under-predictions for PAHs result from systematic under-predictions of  $\log K_{oa}$  values for PAHs due to algorithm limitations for these types of chemicals.



**Figure 3.** Predicted air concentration (nanograms/cubic meter) from measured dust and measured air concentration for chemicals with at least 50% detection frequency in dust. Predictions are presented next to measured concentrations to compare range of concentrations expected and observed. Predictions are only made using detected dust concentrations. Red line represents study-specific maximum method reporting limit (MRL) for each chemical (MRL calculated on mass basis; variations in MRL attributed to volume differences). Note log-scale.



**Partitioning Model As a Tool for Study Design.** We then used the partitioning model to predict total air concentrations for a range of plausible dust concentrations and log  $K_{oa}$  values (SI Figure S9). This prediction model can support study design by indicating detection limits that are needed to generate detectable values. For example, lower molecular weight phthalates (e.g., DEP) are typically detected in dust at central tendency concentrations ranging from 10 to 100  $\mu\text{g/g}$  and have log  $K_{oa}$  values of 7–9. Predicted air concentrations would range from 100 to >10 000  $\text{ng/m}^3$ .

Finally, we used partitioning theory to predict air concentrations for 22 compounds detected in the majority of dust samples but detected less frequently in air (<50%) (Figure 3). We can use the predictions to evaluate reporting limit requirements for future studies. For all but one compound (propoxur), the minimum predicted air concentration was lower by 2- to 100-fold than the MRL in the CAHES. In general, we captured the upper end of the predicted air concentration distributions. Instances when we did not detect chemicals in air despite predictions greater than the MRL may result from model failure or analytical issues. Model failure includes violations of the equilibrium assumption or sorption assumptions, whereas analytical issues may arise from poor absorption onto sampling matrix or desorption during sampling or storage.

**Limitations.** While a partitioning model provides some important insights into the dynamics of SVOCs indoors and provides opportunities to make more accurate exposure estimates using fewer measurements, there are some limitations to this work. First, in this analysis, we modeled only compounds, mostly phthalates and PAHs, detected in  $\geq 50\%$  of air and  $\geq 50\%$  of dust samples. This is because we wanted to use central tendency estimates based on observed rather than estimated data and because if we include all individual points in the model, we bias toward chemicals with the highest detection frequencies. While we were able to use house-specific estimates for total suspended particles (TSP) in estimating the gas-phase concentrations, we had to assume values for dust and particle density and fraction organic matter.

There are also some limitations with the partitioning model concept. Since the partitioning model assumes equilibrium between air and dust, short-term variation in air concentrations, usually measured over hours or days, may not be reflected in dust measurements, which are thought to be stable over months to years.<sup>44</sup> Similarly, chemicals with high  $K_{oa}$  may not partition readily enough to achieve equilibrium. Also, by using the octanol-air partition coefficient, we assume that octanol is representative of organic matter in dust and particles, and that chemicals will readily partition to the organic matter. This assumption may not hold for chemicals found in nonorganic material in the dust, such as observed by Webster et al. for BDE 209.<sup>45</sup>

**Implications and Recommendations.** Our analysis has implications for modeling exposure pathways as well as for sampling study design. For example, chemicals with large  $K_{oa}$  values (>10), especially if they are not found at relatively high concentrations like phthalates, may be best measured using dust sampling or air sampling that captures both the gas and particulate phases. In contrast, chemicals with lower  $K_{oa}$  values (<10) would be readily measured in air samples. For many SVOCs with  $K_{oa}$  values between 5 and 10, partitioning theory suggests, and our analysis confirms, that measurements in one media (e.g., air) can be used to predict concentrations in other

media (e.g., dust) reasonably well. Our estimate of the relative importance of dust versus air exposure pathways also revealed that one particular medium does not dominate total exposures. If a researcher is interested in a large number of chemicals with a wide range of physical-chemical properties, then either air or dust sampling may be able to provide information on exposure source concentrations and route-specific exposures.

Researchers need reliable measurement methods that represent household exposures to SVOCs; however, a standard approach has not been established. Many comprehensive exposure studies have used active air sampling, which collects gas and particle-phase air, and field-technician vacuum dust collection; however, both methods rely on trained personnel and require electricity, making them infeasible in some situations and more costly, particularly in large cohort studies. Many large health studies that have a residential exposure measure for SVOCs are using dust wipes in conjunction with biological samples, despite relatively little validation.<sup>4,13,46</sup> Dust wipe samples involve wiping a specified area to collect dust and surface SVOCs and analytical results are presented as chemical mass per area. There are several disadvantages to dust wipes. First, dust wipes cannot be collected as duplicates because once a particular surface has been wiped another sample cannot be retrieved, requiring that a second surface must be wiped. Second, discretion in selecting wipe locations leads to a lack of standardization, since dust loading as well as chemical concentrations may vary across the surface. Studies of the variations in dust wipe samples have mostly focused on lead, with one studying finding coefficients of variation for dust wipes in different areas of the home ranging from 0.55 to 1.53.<sup>47</sup> Cost, standardization, precision, and relevance to exposure and/or health effect all need to be considered when selecting a residential exposure method.

Passive sampling, or diffusive sampling without the aid of a pump, using semipermeable membrane devices, polyurethane foam (PUF), or other matrices, has been explored by a few researchers, generally demonstrating that they have potential as low-cost versatile measurement tools for SVOCs.<sup>48–53</sup> Despite the promise of this technology, there has been limited development or application of passive indoor sampling as an exposure tool for environmental health studies. Our findings that gas-phase air concentrations can be used to predict total air and house dust concentrations support the potential utility of this approach. A next step is to compare measured air concentrations from passive air samples with measured dust concentrations to investigate and validate this recommendation.

SVOCs from consumer products and building materials are common in residential air and house dust. We found 40 simultaneously detected phthalates, flame retardants, PCBs, PAHs, and pesticides in 49 homes in northern CA, and some chemical classes such as phthalates and PAHs were abundant in both air and dust. Because of their physical-chemical properties, SVOCs redistribute throughout the indoor environment. Resource-intensive techniques such as active air sampling and field-technician collected vacuum dust have been used successfully in the past; however, these methods may be too costly for large-scale cohort studies. Simpler methods are needed. We applied a theoretical partitioning model to empirical data measured in this study and found that we can predict dust concentrations reasonably well from measured air concentrations. On the basis of these findings and ease of implementation, we conclude that passive air sampling, in particular, may be a more standardized measure compared to

surface wipes, and is less resource intensive to collect than vacuum dust and active air samples. Validation of passive air methods for SVOCs is a priority.

## ■ ASSOCIATED CONTENT

### 5 Supporting Information

Specific details for analytical methods; Quality Assurance/Quality Control methods and results; exposure modeling; measured air and dust concentrations; measured total air concentrations versus dust concentrations with correlation estimates; ratio of dust to air concentrations; relative exposures; ratio of gas-phase air to total air concentrations; gas-phase air versus dust concentrations with log  $K_{oa}$ ; ratios of predicted to measured dust concentrations; and contour plot of predicted air concentrations. This material is available free of charge via the Internet at <http://pubs.acs.org/>

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### Notes

The authors declare no competing financial interest.

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