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PAH concentrations simulated with the AURAMS-PAH chemical transport model over Canada and the USA

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Abstract. The offline Eulerian AURAMS (A Unified Regional Air quality Modelling System) chemical transport model was adapted to simulate airborne concentrations of seven PAHs (polycyclic aromatic hydrocarbons): phenanthrene, anthracene, fluoranthene, pyrene, benz[a]anthracene, chrysene + triphenylene, and benzo[a]pyrene. The model was then run for the year 2002 with hourly output on a grid covering southern Canada and the continental USA with 42 km horizontal grid spacing. Model predictions were compared to \sim 5000 24 h-average PAH measurements from 45 sites, most of which were located in urban or industrial areas. Eight of the measurement sites also provided data on particle/gas partitioning which had been modelled using two alternative schemes. This is the first known regional modelling study for PAHs over a North American domain and the first modelling study at any scale to compare alternative particle/gas partitioning schemes against paired field measurements. The goal of the study was to provide output concentration maps of use to assessing human inhalation exposure to PAHs in ambient air. Annual average modelled total (gas + particle) concentrations were statistically indistinguishable from measured values for fluoranthene, pyrene and benz[a]anthracene whereas the model underestimated concentrations of phenanthrene, anthracene and chrysene + triphenylene. Significance for benzo[a]pyrene performance was close to the statistical threshold and depended on the particle/gas partitioning scheme employed. On a dayto-day basis, the model simulated total PAH concentrations

to the correct order of magnitude the majority of the time. The model showed seasonal differences in prediction quality for volatile species which suggests that a missing emission source such as air–surface exchange should be included in future versions. Model performance differed substantially between measurement locations and the limited available evidence suggests that the model's spatial resolution was too coarse to capture the distribution of concentrations in densely populated areas. A more detailed analysis of the factors influencing modelled particle/gas partitioning is warranted based on the findings in this study.

1 Introduction

Polycyclic aromatic hydrocarbons (PAHs) are ubiquitous air pollutants that tend to be most concentrated in areas of dense human population (Hafner et al., 2005) but are also detected at locations remote from local sources (Hung et al., 2005). Many PAH species have been classified as carcinogens (IARC, 2010) and they are implicated routinely as toxicants in airborne particulate matter (Kelly and Fussell, 2012). They are regulated under international agreements such as the Aarhus Protocol on Persistent Organic Pollutants. Benzo[*a*]pyrene, a commonly reported PAH species, is subject to ambient air guidelines in many jurisdictions.

In Canada, PAHs meet the criteria for inclusion on the toxic substances list of the Canadian Environmental 4066

Protection Act (Environment Canada and Health Canada, 1994), and the resulting government obligation has been to reduce or minimise their release into the environment. Nationwide anthropogenic emissions of benzo[a]pyrene, a commonly reported species, fell by 70 % between 1990 and 2010 according to estimates made by the National Pollutant Release Inventory (Environment Canada, 2012). Though there are no federal guidelines for PAHs in Canadian air, a recent analysis of ambient monitoring data found that measured PAH concentrations regularly exceed the health-based guidelines set by the Canadian province of Ontario (Galarneau and Dann, 2011).

In the USA, PAHs are listed in the Clean Air Act as hazardous air pollutants as part of the polycyclic organic matter (POM) class of compounds (US EPA, 2012a) and have been identified as a regional cancer concern in the National-Scale Air Toxics Assessment (US EPA, 2012b). Industrial releases to air reported to the Toxics Release Inventory (TRI) fell by 35 % between 1995 and 2010 (US EPA, 2012c). There is no federal US guideline for PAHs in ambient air.

PAH measurements are labour-intensive compared to those of criteria air contaminants such as ozone and particulate matter, and the processes governing their atmospheric fate are not yet well-understood. In an attempt to elucidate the spatiotemporal distributions of PAH sources and ambient concentrations, several numerical modelling studies have been published. Lagrangian frameworks have been used for Europe (Van Jaarsveld et al., 1997; Halsall et al., 2001) and China (Liu et al., 2007; Lang et al., 2007, 2008). Others studies have used box modelling (Prevedouros et al., 2004) and multimedia fate approaches (Yaffe et al., 2001; Prevedouros et al., 2008). Eulerian chemical transport models (CTMs) have been developed for Europe (Shatalov, 2005; Aulinger et al., 2007; Matthias et al., 2009; Gusev et al., 2011; Bieser et al., 2012) and east Asia (Zhang et al., 2009, 2011a, b; Inomata et al., 2012), and three such studies on a global scale have also been published in recent years (Sehili and Lammel, 2007; Lammel et al., 2009; Friedman and Selin, 2012).

The aforementioned studies differ in many respects relating to the PAH species examined, the temporal variability of their emissions, and the spatial resolutions and process representations in the models. None has focussed exclusively on North America at the regional scale. As well, although several particle/gas partitioning mechanisms have been explored in other models, including Junge–Pankow adsorption (Junge, 1977; Pankow, 1987), organic matter sorption (Finizio et al., 1997), and combined adsorption/absorption (Dachs and Eisenreich, 2000), no previous studies have evaluated model output against paired phase-distributed measurements for alternative partitioning expressions on the same domain.

This study presents the results of a chemical transport model, AURAMS (A Unified Regional Air quality Modelling System)-PAH, run over North America at 42 km horizontal grid spacing with hourly output for the year 2002. Seven PAH species were simulated with the model. Three isomer pairs of decreasing volatility and increasing particulate fraction comprise six of the species: phenanthrene (PHEN) and anthracene (ANTH) (178 g mol⁻¹), fluoranthene (FLRT) and pyrene (PYR) (202 g mol⁻¹), and benz[*a*]anthracene (BaA) and chrysene/triphenylene (C + T) (228 g mol⁻¹). The seventh PAH, benzo[*a*]pyrene (BaP) (252 g mol⁻¹), is not generally considered to be semivolatile but has been included due to its common use as a representative PAH species. Two particle/gas partitioning schemes, Junge-Pankow (JP: Junge, 1977; Pankow, 1987) and Dachs-Eisenreich (DE: Dachs and Eisenreich, 2000), were tested.

Model performance was evaluated against ~ 5000 measurements from 45 stations in established networks in Canada and the USA. This is the first published model to be run and evaluated for PAH concentrations and their distributions between the particle and gas phases using two partitioning methods. It is also the first such model to be evaluated over a regional North American domain.

2 Methods

2.1 Model description

AURAMS is an Eulerian CTM originally developed to simulate criteria air contaminants. The standard version of the model uses a sectional approach to represent the size distribution of airborne particles: 12 size bins from 0.01 to 40.96 µm in diameter and 9 particulate species (sulphate, nitrate, ammonium, elemental carbon, primary organic aerosol, secondary organic aerosol, crustal material, sea salt, and aerosol water) are usually considered. The model includes process representation for tropospheric gas-phase oxidative chemistry, the absorptive formation of secondary organic aerosols, inorganic heterogeneous chemistry, particle microphysics (nucleation, condensation, coagulation, etc.), cloud processing of aerosols, advective transport, vertical diffusion, and gas and particle emissions and deposition. A detailed overall description of AURAMS appears in Gong et al. (2006) while a description of the aerosol sectional approach and the microphysics modules of the model can be found in Gong et al. (2003a, b). Performance evaluation and model intercomparison results for AURAMS appear in Mc-Keen et al. (2007), Smyth et al. (2009), Makar et al. (2010), Kelly et al. (2012) and Solazzo et al. (2012) among other publications.

A modified version of the AURAMS CTM known as AURAMS-PAH was developed to incorporate primary semivolatile organic compounds that are subject to sorptive partitioning. Starting from the standard AURAMS CTM had the advantage that a number of required fields for modelling PAHs were already available. These included hydroxyl concentration, total particle surface area, and fractions of particle elemental carbon and organic carbon. The modifications made to AURAMS version 1.3.2 in order to simulate PAHs are described below. Physico-chemical property values used for each PAH in the modified code are found in Table S1.1 of the Supplement.

2.1.1 Dry deposition of gases

Within AURAMS, gaseous dry deposition velocities are modelled using the inverse resistance analogy for several land-use categories (Zhang et al., 2002). Three resistances are assessed in AURAMS and only the first of these (aerodynamic resistance) is independent of the chemical species under consideration. The species-dependent resistances are the quasi-laminar sublayer resistance and the surface or canopy resistance. The latter two depend on the gas-phase diffusivity of the compound in question, and this quantity was calculated in the model according to the Fuller et al. method described in Reid et al. (1987).

Surface or canopy resistance is the most complex of the three gaseous dry deposition component resistances and tends to dominate total dry deposition (Zhang et al., 2002). One of its subcomponents, mesophyll resistance, was set to $100 \,\mathrm{s}\,\mathrm{m}^{-1}$ for species that are relatively insoluble in water and have small oxidising capacities, as is the case for PAHs. The remaining subcomponents (cuticle and ground resistances) are determined by scaling to O₃ and SO₂ settings based on physico-chemical qualifications. For the PAHs, scaling factors to O_3 and SO_2 for both acetaldehyde and C_3 carbonyls, the least soluble organic compounds considered in AURAMS aside from the PAHs, were used. Unsubstituted compounds such as PAHs are generally considered to have high resistances to deposition whereas carbonyl resistances are thought to be lower (Zhang et al., 2002). However, published observations of PAH deposition led us to assume that deposition velocities would be greater than zero (low resistances) and we therefore used the best-available homologues in AURAMS to represent PAHs. This is an uncertainty in the model that merits future attention.

Volatilisation of gaseous PAHs can occur from exposed water (Hoff et al., 1996), soil (Jones, 1994), and impervious urban surfaces (Diamond et al., 2000). Net gaseous deposition to the Great Lakes in 2002 was downward (Blanchard et al., 2008), suggesting that PAH fugacities in air exceeded those in surface compartments at the regional scale. Volatilisation was not included in this first-generation version of AURAMS-PAH and the effect of this omission is presented in Sect. 3.1.1.

2.1.2 Gas-phase reactions

Reactions of gas-phase PAHs with hydroxyl radicals are considered in this model. Since these reactions consume relatively little hydroxyl due to the trace concentrations of PAH, their reactions were simulated outside the AURAMS gasphase chemistry solver. PAH oxidative loss was estimated as a first-order process using the model-predicted OH concentration immediately preceding particle-gas partitioning. Only seven new gas-phase concentration fields were added to the CTM; PAH reaction products were not tracked in the model, either as individual gas-phase species or as contributors to SOA.

Hydroxyl reaction rate constants were taken from the program AOPWIN (atmospheric oxidation program for Microsoft Windows) which is part of the US EPA's (Environmental Protection Agency) EPI (Estimation Programs Interface) Suite (US EPA, 2006). Measured constants are available for three low-molecular-weight PAHs considered here (phenanthrene, anthracene, fluoranthene) and these values were represented in AOPWIN. However, measurements for the remaining four PAHs are not available and the software predicted the same hydroxyl reaction rate constant of $50 \times 10^{-12} \text{ cm}^3 \text{ molec.}^{-1} \text{ s}^{-1}$ for these species.

2.1.3 Particle representation of PAHs

Seven additional particle species, each with 12 size bins as in the original AURAMS configuration, were added to the model to represent the particle-bound PAH mass.

2.1.4 Particle/gas partitioning of PAHs

A new algorithm was developed for AURAMS-PAH to account for the sorptive particle/gas partitioning of PAHs. It is fully adaptable to other semivolatile species with similar atmospheric partitioning behaviour to PAHs such as dioxins and furans, PCBs (polychlorinated biphenyls), and organochlorine pesticides. The partitioning of PAHs to airborne particles was assumed to be fully reversible.

Two instantaneous equilibrium sorptive partitioning expressions were incorporated in the new partitioning subroutine. The first treated particle/gas partitioning as a Langmuirian adsorption process on a uniform particle surface (JP: Junge, 1977; Pankow, 1987). The model calculations began by adding the particulate PAH concentrations in all size bins (ΣC_p) and the gas-phase PAH concentration (C_g) to give a total PAH concentration (C_{TOT}) for each species. An updated bulk particulate fraction (φ) was then assigned according to the first part of Eq. (1):

$$\varphi = \frac{c\,\Sigma\theta}{c\,\Sigma\theta + p_L^0} = \frac{\Sigma C_{\rm p}}{C_{\rm TOT}},\tag{1}$$

where c is a constant set at 0.173 J m⁻² (estimated from Fig. 3 in Junge, 1977), $\Sigma \theta$ is the total particle surface area concentration (m² m⁻³) and p_L^0 is the saturated vapour pressure of the subcooled liquid (Pa) taken from the temperaturedependent values measured by Offenberg and Baker (1999; see Table S1.1). We have selected Junge's (1977) value of c over that estimated by Pankow (1987) since the latter was based on assumptions that have not been revisited in light of the numerous observations of PAH partitioning published since. The total particulate PAH concentrations dictated by φ were then redistributed among the particle size bins by prorating to the proportion of total aerosol surface area concentration within each size bin. The redistributed gas-phase PAH concentration was determined by difference between C_{TOT} and ΣC_{p} .

The second equilibrium partitioning expression available in the partitioning subroutine developed a partition coefficient (K_p , m³ g⁻¹) based on the contributions of two additive processes: absorption into particulate organic matter and adsorption onto particulate soot (DE: Dachs and Eisenreich, 2000):

$$K_{\rm p} = 10^{-12} \left(1.5 f_{\rm OC} / \rho_{\rm oct} K_{\rm OA} + f_{\rm EC} K_{\rm SA} \right)$$
$$= \frac{\left(\Sigma C_{\rm p} / C_{\rm TSP} \right)}{C_{\rm g}}, \qquad (2)$$

where ρ_{oct} is the bulk density of octanol (0.82 kg L⁻¹), f_{OC} is the organic carbon fraction of the particulate matter (the 1.5 multiplier converts organic carbon to organic matter which is assumed to be well-represented by octanol), K_{OA} is the octanol–air partition coefficient (dimensionless), f_{EC} is the elemental carbon fraction of the particulate matter, K_{SA} is the soot–air partition coefficient (L kg⁻¹), ΣC_p is the particulate PAH concentration across all the size bins (ng m⁻³), C_{TSP} is the total particulate matter concentration (µg m⁻³), and C_g is the gas-phase concentration (ng m⁻³).

Soot-air partition coefficients $(K_{SA} L kg^{-1})$ were estimated as the ratios of soot-water (K_{SW}) to air-water partition (K_{AW}) coefficients since direct K_{SA} measurements are not available for PAHs. K_{SW} values from Jonker and Koelmans (2002) were used in this model. These values vary substantially (up to a factor of 47) between relevant soots for each PAH considered here. Since a single K_{SW} was needed for each PAH in the model, representative values were determined by weighting the reported K_{SW} values by the contribution of their related combustion processes to the total emitted fine particulate matter (PM_{2.5}) used in the inventory of Galarneau et al. (2007). Temperature-dependent K_{AW} values were taken from Bamford et al. (1999). K_{OA} values were taken from the temperature-dependent expressions determined by Odabasi et al. (2006).

PAH partition coefficients were calculated according to the first part of Eq. (2). By determining the contribution of each size bin's organic matter and soot carbon to the totals across all size bins, the total particulate PAH was apportioned to each size bin. For example, if a total partition coefficient had contributions from the organic matter and soot carbon of 20 and 80%, respectively, and size bin 1 held 10% of the total particulate organic matter and 15% of the total soot carbon, the fraction of total particulate PAH assigned to size bin 1 would be 14% (viz., $0.2 \times 0.1 + 0.8 \times 0.15$). Gas-phase concentrations were then determined by the difference between C_{TOT} and ΣC_{p} .



Fig. 1. Map of modelled (JP) annual average total (gas + particle) fluoranthene concentrations (ng m⁻³).

2.1.5 Below-cloud (precipitation) scavenging

Scavenging of gas and particle PAHs by liquid precipitation was calculated as per Gong et al. (2006). Particle scavenging assumed that particle-bound PAHs do not dissolve in falling rain; particle-bound PAHs were thus treated as passive aerosol tracers. Snow scavenging of gaseous PAHs was not considered in this version of AURAMS though particlebound PAHs are scavenged by snow in the model as passive components of airborne particles.

2.1.6 Cloud processing

Cloud processing in the model was treated in a similar manner to precipitation scavenging whereby gas-phase mass transfer to cloud water is species-dependent, whereas particulate interactions with cloud droplets are only affected by the presence of PAHs in terms of the size (mass and volume) that they represent as part of the overall aerosol. Solid-phase densities used to relate aerosol PAH mass to volume were taken from Mackay et al. (2006; see Table S1.1).

2.2 Model domain, emissions, and boundary conditions

The model domain included southern Canada and the continental USA (see Fig. 1). It was run on a 42 km polar stereographic grid using offline meteorology generated with the Global Environmental Multiscale numerical weather prediction model (GEM v 3.2.0: Côté et al., 1998a, b).

Emissions of PAHs were taken from the inventory of Galarneau et al. (2007) that had been updated from 2000 to 2002 and to which benzo[*a*]pyrene had been added using identical methods and data sources. As discussed in Galarneau et al. (2007), hourly PAH emissions fields were estimated with an emissions processing system using source-specific temporal profiles. The temporal profile library included 3020 month-of-year, 64 day-of-week, and 2672 hour-of-day temporal profiles for Canada and 1500, 49, and 680



Fig. 2. Map of measurement stations used in the AURAMS-PAH evaluation.

analogous temporal profiles for the USA. The overall temporal profile thus varies from grid cell to grid cell due to the different mixtures of source types found in each one.

All PAHs were emitted exclusively in the gas phase. Particle/gas partitioning took place at each 15 min CTM time step according to the partitioning module described in Sect. 2.1.5. As mentioned in Sect. 2.1.2, no emissions of previously deposited PAHs were considered in this first-generation version of the model and the implications of this are discussed in Sect. 3.1.1. Emissions of SO₂, NO_x, NH₃, CO, volatile organic compounds (VOCs), and particulate matter were derived using Environment Canada and US EPA databases and methods for the year 2002.

Initial PAH concentrations at all lateral boundaries were set to zero in anticipation of pronounced spatial gradients away from localised source regions. As a result, modelled concentrations in Mexico and near its border with the USA are not expected to be reliable, particularly since PAH emissions from Mexico have not been included in the model. Model output along the northern edge of the domain over western Canada is similarly expected to be unreliable since emission sources are located close to the model boundary in that region. The development of representative nonzero boundary concentrations is anticipated as part of future model development.

2.3 Evaluation data

Observational PAH data used for comparison with model output were collected from four measurement networks: NAPS (Canada), IADN (Canada-USA), CARB (California), and Rio Tinto Alcan (Kitimat, British Columbia, Canada). The measurement stations are depicted in Fig. 2 and described in Sect. 2 of the Supplement.

Measurement data were available from a total of 45 stations, 23 in Canada and 22 in the USA, all of which collected samples integrated over periods of 24 h. Particle/gas partitioning was assessed at eight stations, three in Canada and five in the USA, all of which were operated by IADN.

The IADN phase-distributed data were also combined to yield total concentrations. These combined IADN data, along with NAPS and Rio Tinto data, yielded a total of 28 sites at which the total PAH concentration for all the modelled PAHs could be assessed. Particulate PAH measurements from the latter networks were determined from samples of total suspended particles (TSP). CARB provided data for benzo[*a*]pyrene in particles smaller than 2.5 μ m in diameter (PM_{2.5}) at a further 17 locations.

Four model grid squares (Kitimat, Toronto, Hamilton, and Montreal) contained two or more measurement stations thus allowing for an assessment of the adequacy of modelling all seven PAHs at 42 km grid spacing.

3 Results

3.1 Total PAH concentration

3.1.1 Overall spatiotemporal domain

Total PAH concentration refers to the sum of the gas and particulate concentrations whether these have been analysed together (e.g. NAPS) or separately (e.g. IADN). For stations at which the gas and particle phases were analysed separately, a valid total concentration was assumed to exist if at least one of the gas and particle phase concentrations was greater than the detection limit. Non-detectable values were assumed equal to zero for the calculation of total concentrations.

A representative plot of the spatial distribution of modelled annual average concentrations is presented in Fig. 1 for fluoranthene. The remaining PAHs show similar spatial distributions and maps of their modelled concentrations are found in Sect. 3 of the Supplement. All the PAHs show spatial distributions of their modelled concentrations that are consistent with regional dispersion of their emissions as depicted in Galarneau et al. (2007).

A summary of annual mean modelled and measured values over the entire spatiotemporal model domain is shown in Table 1. Only modelled values for which there was a corresponding measurement were included.

In comparing modelled results to measurements, the annual means were statistically indistinguishable at the 95% confidence level for FLRT, PYR, BaA, and BaP (JP) whereas they were statistically different for PHEN, ANTH, C + T, and BaP (DE). For PHEN, ANTH, and C + T, modelled values were underestimated relative to measurements whereas they were overestimated for DE BaP.

The model's temporal variability tended to be smaller than that of the corresponding measurements; the relative standard deviations of the measurements were 1.3–2.7 times greater than those of the modelled values. A similar observation has

PAH	Modelled-JP	Modelled-DE	Measured	n^*
PHEN	12.75 (36.44)	12.76 (36.44)	36.06 (131.8)	790
ANTH	0.9123 (1.757)	0.9104 (1.759)	2.804 (11.56)	701
FLRT	6.781 (14.40)	6.888 (14.66)	9.179 (32.44)	789
PYR	5.727 (12.23)	6.009 (13.40)	5.733 (21.57)	785
BaA	1.227 (2.438)	1.328 (2.704)	1.326 (6.081)	610
C + T	1.511 (3.964)	1.473 (3.569)	3.303 (21.95)	721
BaP	1.173 (2.002)	1.424 (2.455)	0.9047 (3.238)	595

Table 1. Summary of the 2002 annual modelled and measured total PAH concentration mean (standard deviation) values (ng m^{-3}).

* n = number of modelled-measured data pairs

been made in the modelling of particulate matter with AU-RAMS and other regional air-quality models (Solazzo et al., 2012) For PAHs, this effect was also seen by Matthias et al. (2009), who concluded that temporal variability in PAH emissions was not adequately represented by their inventory. This is a plausible contributing factor in the current study as well. Furthermore, meteorological parameters vary over a scale much finer than that used for regional air quality models. As a result, observed concentrations from point locations can be expected to exhibit greater variability than modelled concentrations determined for entire grid squares.

Differences in mean modelled total concentrations between the two partitioning versions (JP and DE) were statistically indistinguishable at a 95 % significance despite the finding that the two BaP model results differed in their comparison to measured values. The latter anomaly indicates that the BaP distributions were close to the 95 % confidence threshold. As a result, no conclusion can be drawn about which partitioning mechanism was superior in simulating overall total PAH concentrations. Phase partitioning of semivolatile organic compounds (SVOCs) is a major determinant of their potential for long-range transport (Bidleman, 1988), yet it does not appear to have a large effect on the simulation of their total concentrations at the regional scale. Model performance in simulating phase partitioning is discussed in Sect. 3.2.

The model's performance was also more closely evaluated by examining the pertinent data distributions. Figure 3 depicts frequency distributions of the ratios of modelled-tomeasured concentrations for all of the valid data pairs available for the model evaluation. Four PAH species (ANTH, FLRT, PYR, and C+T) yielded median values of the modelled-to-measured concentration ratio that were close to the ideal value of unity (1.1, 1.1, 1.5, and 1.4, respectively). PHEN showed an overall tendency toward underestimation by the model (0.2), whereas BaA and BaP tended toward overestimation (3.2/3.5 and 3.0/3.5 JP/DE, respectively).

BaA and BaP are reactive PAHs (e.g. Behymer and Hites, 1985; Pöschl et al., 2001; Kwamena et al., 2004; Esteve et al., 2006, Shiraiwa et al., 2009) and the exclusion of particlebound reactions in this first-generation model may explain a portion of their overestimation in AURAMS-PAH as sug-



Fig. 3. All-site ensemble of modelled-to-measured concentration ratios for total (gas + particle) PAHs using JP and DE partitioning expressions. Box boundaries are 25th, 50th and 75th percentile values; whiskers are 10th and 90th percentile values. JP – Junge– Pankow partitioning; DE – Dachs–Eisenreich partitioning.

gested in a comparable model for Europe (Matthias et al., 2009). However, BaP is subject to losses during sampling (Menichini, 2009) and some portion of the apparent model overestimation may in fact be due to measured concentrations that are biased low since the samplers used were not equipped with oxidant denuders. This presents a priority for future research since many jurisdictions use BaP as an indicator PAH when setting air quality standards.

As presented in Sect. 2.1.2, volatilisation of gaseous PAHs from surface compartments such as water and soil was not included in this first-generation version of AURAMS-PAH. If such volatilisation were significant to the balance of PAHs in ambient air relative to the other processes simulated, one would expect an overall bias in model results whereby the most volatile PAHs, which are found predominantly in the gas phase, would be underestimated and the least volatile particulate species would be unaffected. Summary results provide indefinite evidence. Volatile PHEN is systematically underestimated yet its similarly volatile isomer, ANTH, shows an ambiguous central tendency whereby its mean concentrations are underestimated by the model (Table 1) but its median concentrations are not (Fig. 3). Less volatile but nonetheless predominantly gaseous FLRT and PYR show no tendency toward underestimation.

Though results are equivocal on an annual basis, monthly patterns observed in the model output are consistent with the absence of a seasonal source (e.g. air–surface exchange). Volatilisation from a variety of environmental compartments is typically stronger in warmer periods than in cooler ones (e.g. Nelson et al., 1998; Smith et al., 2001; Motelay-Massei et al., 2005; Bozlaker et al., 2008; Wang et al., 2011). Figure 4 shows the monthly distribution of modelled-to-measured concentration ratios for PHEN and PYR. Both exhibit higher values in winter than in summer as do ANTH and FLRT whereas this seasonality is not observed for the higher molecular BaA, C+T or BaP (not shown). These findings are consistent with a missing volatilisation source that emits during warmer weather. However, other factors could also be involved including overestimated loss terms



Fig. 4. All-site ensemble of modelled-to-measured concentration ratios for total (gas + particle) PHEN (a) and PYR (b) using the JP partitioning expression plotted by month.

(e.g. oxidation, deposition) or underestimated emissions (e.g. forest fires) during warmer periods. The investigation of the relevant causes is a priority for future model development. Regardless of the causes, the seasonal effect on model output appears to be compounded by further, as yet unidentified factors whereby PHEN is underpredicted throughout the year and ANTH, FLRT and PYR are overpredicted through some seasons, potentially due to air–surface exchange that leads to net deposition during cooler months.

The range of modelled-to-measured concentration ratios shown in Fig. 3 varied substantially by species. The ratios of 90th to 10th percentile values for PHEN, FLRT, PYR, BaA, and C + T spanned fewer than, or close to, two orders of magnitude (55, 59, 67/68, 67/63, and 100/93, respectively). The ratio for BaP was larger (180/270) and that for ANTH was very large (5900/7400), with extreme values tending toward underestimation for the latter species. As seen with the comparison of means, the two partitioning parametrisations used by AURAMS-PAH led to similar model performance overall when considering the distribution of total PAH concentrations.

Additional quantitative performance metrics are presented for the two particle/gas partitioning parametrisations in Tables S4.1 and S4.2 of the Supplement. The normalised mean bias and error have been included for completeness, but their utility in this evaluation is questionable given the large range of concentrations. Measured maximum to min-



Fig. 5. Site-specific modelled-to-measured concentration ratios for total (gas + particle) fluoranthene for JP partitioning. N.B. Box boundaries are 25th, 50th and 75th percentile values; whiskers are 10th and 90th percentile values.

imum concentration ratios range from 4.7×10^6 (PYR) to $1.3 \times 10^9 / 1.4 \times 10^9$ (C + T). Therefore, the mean measured concentrations used to normalise the bias and error do not represent the data set well.

The correspondence between individual modelledmeasured data pairs is weak as demonstrated by the low coefficients of determination, non-unity slopes, and high intercepts listed in Tables S4.1 and S4.2 of the Supplement. However, the ability of the model to simulate observed concentrations within a certain tolerance is reasonable, especially when considering that PAHs are trace organic compounds subject to numerous sampling artefacts (Mc-Dow, 1999) and poor measurement precision (Galarneau, 2008). Depending on PAH species, 22-34% of modelledmeasured data pairs fell within a factor of 2 of each other. This increased to 61–86% when considering a factor of 10. As a result, it can be stated with confidence that, on average, AURAMS-PAH was able to simulate atmospheric PAH concentrations in North America for rural to urban locations to the correct order of magnitude.

3.1.2 Site-specific performance

Model performance was not spatially uniform. Figure 5 depicts the variation in distributions of individual modelledto-measured concentration ratios across measurement sites for fluoranthene, the PAH species for which overall performance was best as determined by the median and spread in modelled-to-measured concentration ratios. Note that only JP partitioning values have been plotted since these are visually indistinguishable from those for DE partitioning.

Of the 30 sites depicted in Fig. 5 (CARB sites could not be considered since only benzo[a]pyrene was reported there), the median modelled-to-measured concentration ratio ranged from 0.061 (St. John's) to 4.0 (Hamilton – Confederation Park), whereas the median value for all sites was 1.1. The variability at individual sites is itself highly variable, with

Station	PHEN	ANTH	FLRT	PYR	BaA	C + T	BaP	O ₃ ²	TSP
Kitimat	106	101	96.5	93.7	87.4	91.3	93.2	N/A	N/A
Hamilton	52.1	59.0	59.2	58.8	73.1	117	62.8	24.9	30.5
Toronto3 ¹	36.2	48.7	42.5	39.6	42.2	32.9	43.0	N/A	22.1
Toronto2 ¹	36.5	45.4	39.3	35.7	32.9	25.0	38.0	12.2	24.6
Montréal	49.2	52.9	45.8	44.6	55.2	55.3	51.6	35.3	29.0

Table 2. Average coefficient of variation (%) between contemporaneous measurements at sites falling within the same 42 km AURAMS-PAH grid square.

¹ Toronto3 includes data from all three Toronto measurement sites. Toronto2 includes only data from the Gage

Institute and Judson & Etona because O3 data were not available from Junction Triangle.

² Ozone data have been aggregated to 24 h concentrations contemporaneous with PAH measurements.

ratios of 90th to 10th percentile values of the modelled-tomeasured concentration ratio ranging from 5.8 (Toronto – Junction Triangle) to 105 000 (Haul Road, near the Rio Tinto Alcan smelter in Kitimat, British Columbia). A low value of 1.1 was observed for Saint John, but this was based on only two modelled–measured data pairs. Sixteen of the 30 sites (53%) had median modelled-to-measured ratios that fell within a factor of two of the median value for all sites.

The other compounds varied spatially in a manner similar to fluoranthene with the following exceptions. ANTH exhibited atypically large underestimation at the three sites near the Rio Tinto Alcan smelter in Kitimat, suggesting that inaccurately low ANTH emissions are associated with the dominant source there. The reporting threshold for point-source ANTH emissions through the Canadian National Pollutant Release Inventory (NPRI) system is higher than the thresholds for other commonly measured PAHs and no ANTH emissions were reported to the NPRI by Rio Tinto Alcan for 2002. The C + T performance at Jonquière, home to aluminum smelting facilities, suggests that reported emissions there are also too low. Emissions for other PAHs were reported from this location for 2002 but not so for chrysene, which is called benzo[*a*]phenanthrene in the NPRI.

3.1.3 Model grid squares containing multiple measurement sites

The smoother the spatial distribution of a pollutant, the coarser the model resolution that can be used to simulate it. Four AURAMS-PAH model grid squares contain more than one measurement site, thus allowing for an assessment of the 42 km spatial resolution used for the evaluation runs. The multisite grid squares are all located in Canada, and from west to east, they encompass sites in Kitimat (2 sites), Hamilton (2), Toronto (3), and Montreal (2) (see Tables S2.1 and S2.4 in the Supplement).

Kitimat is a town 650 km northwest of Vancouver with approximately 9000 residents whose largest employer is the aluminum smelter complex operated by Rio Tinto Alcan (District of Kitimat, 2009). Two measurement sites (Haul Road and Kitamaat Village) are located in the same model

grid square and a third site (Whitesail) lies in an adjacent square even though it is only a few kilometres away. Hamilton is a city at the western end of Lake Ontario that is known colloquially as the "Steel Capital of Canada" and had a population of approximately 700 000 in 2010. It is part of the so-called "Golden Horseshoe" conurbation at the western end of Lake Ontario whose 2010 population, estimated as the sum of the populations of Oshawa, Toronto, Hamilton, and St. Catharines – Niagara, was over 7 million (Statistics Canada, 2011). Toronto and Montreal are the largest cities in Canada having 2010 populations of 5.7 and 3.9 million, respectively.

Table 2 lists the variability in contemporaneously measured concentrations at the four grid squares as represented by their coefficients of variation (COV). At any given site, the average COVs for the different PAH species tend to be similar to each other. Substantial differences exist between sites, however, particularly when grouping the urban sites (Hamilton, Toronto, and Montréal) against the industrial site at Kitimat. This is not unexpected. Urban areas include complex mixtures of point, area, and mobile sources that are distributed over distances similar to the scale of the model. Kitimat houses industrial operations within a relatively small area of otherwise rural land and wilderness. Steep spatial gradients in pollutant concentrations are expected there as a result.

The COVs for ozone and total suspended particles (TSP) have also been included in Table 2 as comparative gaseous and particulate pollutants, respectively. Both vary less between sites in the same grid square than do PAHs. Ozone and a portion of TSP are secondary pollutants created by the mixing and reaction of precursor compounds. The atmospheric residence times required for their creation is consistent with a smoothing of the spatial variability in their concentrations though ozone variability is further complicated by reactions with NO_x near emissions from mobile sources. Conversely, unsubstituted PAHs are primary pollutants whose concentrations would be expected to vary in space over a finer resolution when multiple sources are found close by.

The results presented above suggest that a 42 km spatial resolution is not sufficiently fine to represent PAH concentrations in areas close to sources such as cities and industrial



Fig. 6. All-site ensemble of modelled-to-measured PAH particulate fraction ratios for JP and DE partitioning expressions. N.B. Box boundaries are 25th, 50th and 75th percentile values; whiskers are 10th and 90th percentile values.

areas if an average model accuracy better than an order of magnitude is desired. AURAMS modelling of fine particulate matter has shown substantial improvement when grid spacing has been reduced to 2.5 km (Stroud et al., 2011), and similar results can be expected for the modelling of PAHs. No 42 km model grid squares in rural or background areas away from sources contain multiple measurement stations and, as a result, a comparison cannot be made for these areas. However, it is expected that spatial variation in PAH concentrations will be less in such areas and, as such, a 42 km resolution model may be sufficient there.

3.2 Particle/gas partitioning

As noted in Sect. 3.1, the choice of partitioning expression (JP or DE) had little effect on the simulation of total PAH concentrations. This implies that the partitioning from each approach is sufficiently similar that regional-scale differences in removal rates between gaseous and particulate PAHs have little effect. However, differences between the two expressions with respect to simulating phase-resolved concentrations were noted.

3.2.1 Overall spatiotemporal domain

Figure 6 shows frequency distributions of the ratios of individual modelled-to-measured particulate fractions for all data pairs available to the model evaluation. Note that only the eight IADN stations are included since the gas and particle phases are analysed separately only at those sites.

Figure 6 shows that PAH particulate fraction is underestimated for all species except BaP. The degree of underestimation decreases with increasing molecular weight. The particulate fractions of volatile PHEN and ANTH (178 g mol⁻¹) are underestimated by approximately two orders of magnitude whereas equipartitioning BaA and C + T (228 g mol⁻¹) have particulate fractions that are underestimated by only a factor of two. A similar pattern appears when examining the partition coefficient, K_p (not shown).



Fig. 7. Site-specific modelled-to-measured partition coefficients for fluoranthene for DE partitioning for eight IADN sites. N.B. Box boundaries are 25th, 50th and 75th percentile values; whiskers are 10th and 90th percentile values.

For all species other than BaP, Dachs-Eisenreich partitioning performs slightly better than Junge-Pankow partitioning in simulating measured particulate fractions. The allsite median particulate fraction simulated using DE is between 1.1 (PYR) and 2.9 (ANTH) times higher than that using JP. However, the performance of the partitioning expressions is highly dependent on the physico-chemical property values used. For example, estimated soot-air partition coefficients vary by more than an order of magnitude (Galarneau et al., 2006) and translate directly to variations in predicted partitioning by the Dachs-Eisenreich expression. For Junge-Pankow partitioning, the value of the constant, c, in Eq. (1) and the estimation of aerosol surface area also introduce uncertainties. A full analysis of the sensitivity of modelled partitioning is beyond the scope of this paper and is explored in a separate publication (Galarneau, 2014).

3.2.2 Site-specific performance

As was the case for total concentration, there is substantial variability in the simulation of partitioning between sites. Figure 7 shows the variation in frequency distribution of individual modelled-to-measured particulate fractions for fluoranthene using Dachs-Eisenreich partitioning. Model performance for particulate fraction simulation is better at urban (Chicago) or urban-influenced (Sturgeon Point, Egbert) sites than at those that are remote (Eagle Harbor). An analysis of measured partitioning at IADN stations (Galarneau et al., 2006) found that the proportionality between partitioning and volatility varied between sites, and in some cases, over the annual cycle. Volatility is included in both the JP (through $p_{\rm L}^0$) and DE (through $K_{\rm OA}$ and $K_{\rm SA}$) partitioning expressions and the proportionality between it and partitioning magnitude is much smaller in model outputs than in measurements. As noted earlier, factors involved in the performance of model partitioning such as modelled particulate matter concentration and composition are explored in a separate publication (Galarneau, 2014).

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4 Conclusions

This study described the first-known modelling results for atmospheric PAHs at the regional scale over North America. Predictions from the AURAMS-PAH model were compared to roughly 5000 24 h-average PAH measurements from 45 sites, eight of which also provided data on particle/gas partitioning which had been modelled using two different partitioning schemes.

The evaluation of the model is key to determining its potential utility as an input for estimating the impacts of PAH inhalation exposure on human health. Annual average modelled total (gas + particle) concentrations were statistically indistinguishable from measured values for fluoranthene, pyrene and benz[a]anthracene, indicating the model's potential utility for providing inputs to health impact estimation for these species. The model's annual average concentrations for phenanthrene, anthracene and chrysene + triphenylene were biased low. For these species, the negative bias would have to be considered if used as inputs to human health impact estimates as the model in its present form underestimates long-term exposure.

The utility of the model for prediction purposes may also be considered on a day-to-day basis though this is less relevant to the chronic health effects associated with carcinogenic PAHs. The model simulated total PAH concentrations to the correct order of magnitude 64–86 % of the time. That level of accuracy must be considered when assessing human health impacts; annual exposure estimates are likely of more utility with the model in its current state.

The partitioning approach chosen did not have a significant impact on the model results for total concentrations though differences resulting form the choice of parametrisation approached the 95% significance level for benzo[a]pyrene. At this time, neither of the two approaches used here provided a clear advantage for simulation accuracy of total concentrations.

As a first work of this nature, the analysis has suggested several avenues for further model development and improvement. Improved temporal emission estimates for PAHs are key to improving model simulations of these species; simulated PAHs showed less temporal variability than the measurements. The reactions of particulate PAH species with atmospheric oxidants should be given further consideration since the more reactive species were overestimated in the current model. The addition of an air-surface exchange parametrisation should be evaluated as a potential response to the seasonally varying prediction capability of the model for the most volatile compounds. Model resolution has been shown to be a key factor in improving air-pollution estimates in areas with high human exposures. While the 42 km horizontal grid spacing used in this study is finer than that used in global models, it was insufficient to capture the distribution of concentrations in densely populated areas. A more detailed analysis of the factors influencing modelled particle/gas partitioning is needed to improve the distribution of PAHs between the gas and particle phases in the atmosphere given that both partitioning schemes used here showed increasing negative biases for particle-bound PAH concentrations of increasing volatility.

Supplementary material related to this article is available online at http://www.atmos-chem-phys.net/14/ 4065/2014/acp-14-4065-2014-supplement.pdf.

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