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Concentrations, Trends, and Air-Water Exchange of PAHs and PBDEs Derived from Passive Samplers in Lake Superior in 2011

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Concentrations, Trends, and Air-Water Exchange of PAHs and PBDEs Derived from Passive Samplers in Lake Superior in 2011

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Abstract

 Polycyclic aromatic hydrocarbons (PAHs) and polybrominated diphenylethers (PBDEs) are both currently released into the environment from anthropogenic activity. Both are hence primarily associated with populated or industrial areas, although wildfires can be an important source of PAHs, as well. Polyethylene passive samplers (PEs) were simultaneously deployed in surface water and near surface atmosphere to determine spatial trends and air-water gaseous exchange of 21 PAHs and 11 PBDEs at 19 sites across Lake Superior in 2011. Surface water and atmospheric PAH 28 concentrations were greatest at urban sites (up to 65 ng L^{-1} and 140 ng m⁻³, respectively, averaged from June to October). Near populated regions, PAHs displayed net air-to- water deposition, but were near equilibrium off-shore. Retene, probably depositing following major wildfires in the region, dominated dissolved PAH concentrations at most Lake Superior sites. Atmospheric and dissolved PBDEs were greatest near urban 33 and populated sites (up to 6.8 pg L^{-1} and 15 pg m⁻³, respectively, averaged from June to October), dominated by BDE-47. At most coastal sites, there was net gaseous deposition of BDE-47, with less brominated congeners contributing to Sault Ste. Marie and eastern open lake fluxes. Conversely, the central open lake and Eagle Harbor sites generally displayed volatilization of PBDEs into the atmosphere, mainly BDE-47.

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Introduction

active-sampler derived concentrations agree within a factor of two to three. **8,9** When

89 industrial applications and landfills,²⁴ suggesting continued release to the environment and deposition into Lake Superior.

 The objective of this study was to assess the spatial distribution and environmental cycling of PAHs and PBDEs across Lake Superior. We deployed polyethylene passive samplers (PEs) in the air and water at 19 sites across Lake Superior from April-October 2011 with the aim to (1) enhance the spatial coverage of air and water sampling stations across Lake Superior; (2) determine the concentrations and spatial trends of PAHs and PBDEs; (3) discern their air-water exchange; and (4) evaluate the efficacy of PEs as tools to monitor regional POP distributions.

Materials and Methods

Sampling Methodology

101 Low density PE (2 mil) was cut into approximately 10x40 cm strips from commercial sheeting (Berry Plastics Corporation, Evansville, IN) yielding passive samplers ~2 g each. PEs were cleaned successively in dichloromethane and hexane for 24 h each. PEs were spiked with performance reference compounds (PRCs) to account for mass-transfer limitations during deployment. Deuterated PAHs 106 (naphthalene-d₈, pyrene-d₁₀), brominated biphenyls (dibromobiphenyl, tetrabromobiphenyl, pentabromobiphenyl) and octachloronaphthalene (Ultra Scientific, North Kingstown, RI, USA; Cambridge Isotopes, Andover, MA, USA) were equilibrated with the PEs in 80:20 methanol:water on a shaker table for 4 weeks 110 based on the procedure reported by Booij et al. (2002) ²⁵ PEs were woven onto stainless steel wire and distributed to collaborators and volunteers. Air PEs were

 deployed in inverted bowl stations at 11 coastal and three open-lake sites (see TOC). Surface water PEs (unprotected) were deployed in tandem with the coastal air samplers, as well as at three open-lake sites (slightly offset from air deployments) and two additional coastal sites (in-cage deployments, see SI for details). Samples were collected in three deployment periods to evaluate temporal trends: April-June, June- August, and August-October 2011 (Table SI 3). Duplicate air samples and triplicate water samplers were deployed at two sites during each deployment. Field blanks were taken from 2-3 sites per deployment period. After retrieval, samplers were wrapped in 120 foil, shipped to the lab and stored at 4° C until analysis.

Analytical Methodology

 PEs were wiped clean with Kimwipes and extracted for 24 hours in ethyl acetate and condensed down. Extracts were spiked with 45-50 ng of labeled PAH 125 (acenaphthalene-d₁₀, phenanthrene-d₁₀, chrysene-d₁₂, perylene-d₁₂ from Ultra 126 Scientific, North Kingstown, RI USA) and 50 ng PBDE $(^{13}C_{12}$ -BDE28, $^{13}C_{12}$ -BDE47, $127 \text{ }^{13}C_{12}$ -BDE99, $^{13}C_{12}$ -BDE153, $^{13}C_{12}$ -BDE183 from Cambridge Isotope Laboratories) surrogates to determine analyte recovery during sample processing (see SI page 5). PAHs were analyzed on an Agilent 6890 Series GC - Agilent 5973 MS (mass spectrometer), PBDEs on a Waters Quattro micro GC Micromass MS-MS. *Quality Assurance/Quality Control* PE matrix spikes were prepared in each batch of approximately 20 samples.

Analyte concentrations were recovery corrected. Average surrogate recoveries for

Calculations/Data Analysis

 At each site, 21 PAHs and 11 PBDEs were determined (for details, see SI page 148 6). Truly dissolved concentrations of POPs in water, C_W (pg L^{-1}), were calculated as 26 . 149 $C_W = C_{PEw} / K_{PEW}$ (1)

150 where C_{PEw} is the concentration of the POP in the PE when in equilibrium with water

151 (pg kg_{PE}^{-1}), and K_{PEW} is the POP partitioning coefficient from water into polyethylene

152 (L kg⁻¹). K_{PEW} values were obtained from Lohmann $(2012)^{27}$ and temperature-

corrected according to a modified form of the Van't Hoff equation

$$
K_{PEW(T)} = K_{PEW(298K)} * e^{(-(\Delta H vap/R) * [(1/298K) - (1/T)])}
$$
(2)

155 where ΔH_{vap} is the enthalpy of vaporization (kJ mol⁻¹) (Table SI 5, Table SI 14), R is

156 the gas constant (kJ/K*mol) and T is the average water temperature during

deployment (Kelvin) (Table SI 4). Gas-phase atmospheric POP concentration, CA,

was determined with the same calculations, substituting the air-PE partitioning

159 coefficient, KPEA, for KPEW.

178 where k_{ol} is mass transfer coefficient (m day⁻¹) (see SI page 7) and K_{PEW} is the POP's 179 temperature-corrected PE-water partitioning coefficient $(L kg^{-1})$. Equation (4) derives

- air-water exchange gradients without needing to rely on Henry's Law Constants,
- 181 which are arguably the least constrained physico-chemical constants. ^{10,29}
-

Results and Discussion

Polycyclic Aromatic Hydrocarbons (PAHs)

185 **Spatial Distributions in Air:** Averaged Σ₂₁PAH gas-phase concentrations 186 (June – October) ranged from <1 to >100 ng m⁻³ (Figure 1A, Table SI 10).

Concentrations were always greatest at Sault Ste. Marie, (June-October mean 140 ng

188 m⁻³), and lowest at Eagle Harbor, MI (June-August 0.054 ng m⁻³) and the open lake

189 sites (mean 0.13-0.95 ng m⁻³) (Table 1). An urban-rural gradient, as observed in this

study, can occur due to particle scavenging near urban sources, precipitation

scrubbing, and OH radical degradation.**30,31** PAHs tend to have a strong urban

signature from vehicle emissions, power generation plants, and industrial

193 activity.^{31,32,33} At some locations, atmospheric concentrations increased with

increasing temperature; yet mostly the changes were within the overall uncertainty of

the passive sampling (factor of 2; see Table SI 4 and Table SI 10). Atmospheric

concentrations of semi-volatile compounds generally have a seasonal component due

to the effect of ambient temperature on the direction of pollutant transfer across the

198 air-water interface.³⁴ It has been suggested, however, that atmospheric PAH

199 concentrations are a function of emissions source rather than temperature³⁵, in-line

with the results presented here.

 A previous sediment study measured the greatest PAH concentrations near 202 Lake Superior's second largest city, Duluth, MN,¹² however, the study did not include

- 223 resulting from biomass burning and the decomposition of conifer trees, was also
- 224 significantly present (ca. 4% of Σ_{21} PAHs, 1.1 ng m⁻³). Retene is associated with wood

225 varnish and lumber treatment,³² therefore, the lumber industry in Ashland could be contributing to the retene signal in that region.

 Atmospheric PAH concentrations at other urban locations, while a tenth and a third, or less, of those at Sault Ste. Marie and Ashland, respectfully, were still ca. two- to ten-times greater than rural coastal sites, and several times the open-water air concentrations. Duluth had a lower atmospheric PAH concentrations (June-October 231 – average 12 ng m⁻³) than Sault Ste. Marie and Ashland despite being an urban center and possible emissions linked to shipping traffic in the Duluth/Superior harbor, the 233 busiest port in the Great Lakes system.³⁸ This may be because the PE was deployed at the regional EPA office northeast (upwind) of the city center. Following Duluth at 235 slightly lower PAH concentrations were Marquette (June-August average 10 ng m⁻³) 236 and Thunder Bay/Welcome Isle (June-October average 5.4 ng m^{-3}). The sampler at Thunder Bay was also several km removed from the city center. Concentrations were characterized by many of the same petroleum- and biomass-derived PAHs measured at shipping sites: phenanthrene, methyl phenanthrenes, fluoranthene, and pyrene. Thunder Bay and Duluth have relatively modest populations (110,000 and 100,000 people, respectively), and these samples demonstrate the effect even small cities have on local atmospheric PAH concentrations. Eagle Harbor, MI atmospheric PAH concentrations were below the detection 244 limit for April-June and <0.1 ng m⁻³ for June-August; August-October data were not available, however, temporal trends for other sites suggest atmospheric concentrations

may have been greater during the late summer. The IADN master station for Lake

Superior is located at Eagle Harbor and has been collecting high volume air samples

261 Northern, rural coastal sites (Sturgeon Bay: 1.4 ng m⁻³; Foster Island: 0.89 ng 262 m^3), and eastern open lake (0.95 ng m⁻³) were all at background levels similar to 263 Eagle Harbor of around 1 ng m⁻³. The central and western open lake samples displayed lower PAHs concentrations, as could be assumed for overwater sites. This comparison also serves to support the choice of Eagle Harbor as a rural background site for IADN; yet it might not be a good representation of over-water concentrations. **Spatial Distributions in Water:** PAH concentrations in the water followed similar spatial patterns as the atmospheric PAH distributions (Figure 1B, Table SI 11). 269 Seasonally averaged Σ_{21} PAH concentrations in Lake Superior were greatest at Sault

270 Ste. Marie (June-October 65 ng L^{-1}), followed by Ashland (26 ng L^{-1}). Concentrations

289 concentrations of PAHs across most sites were dominated by retene. It is the

290 combustion byproduct of abietic acid, a natural product mainly found in coniferous

291 trees.^{32,40} Retene can be produced in urban areas from municipal waste incinerators,

292 tire burning, and incineration of building waste; however, it is primarily emitted by

293 burning wood.⁴¹ Rural households consume about three times more wood than urban

294 homes.³² It is therefore not surprising that retene accounts for ca. 20-60% of the total PAHs at the northern coastal sites and ca. 40-70% of total PAHs present in open lake waters. Atmospheric retene concentrations typically peak in the winter months when wood is burned for residential heating, but lower-magnitude spikes can occur between 298 April and September due to naturally occurring wildfires.³²

 Several sites exhibited elevated retene concentrations in both air and water during deployment 3 (August-October) relative to deployment 2 (June-August) (Figure 2). Dissolved retene increased strongly at Sturgeon Bay (from 0.92 to 26 ng 302 L⁻¹), station 113 (0.13 to 8.3 ng L⁻¹), and Station 139 (0.92 to 6.9 ng L⁻¹). Because PE sampling was limited to one season, we cannot confirm whether these trends occur in annual cycles or are due to unique events. However, sites with the greatest retene concentrations (Sturgeon Bay and Station 113) were northern and central lake locations, which probably received enhanced retene inputs following elevated forest f fire intensity from July – September 2011 northwest of the Lake (see Figure SI 3).⁴² Retene could also enter the Lake's water through tributaries, but Sturgeon Bay and Station 113 were not close to river plumes.

Comparison to active sampling: Venier et al. (2014)⁴³ reported apparently dissolved water concentrations of various POPs, including PAHs across the Great Lakes using active GFF - XAD sampling in 2011/12. For the open Lake Superior, apparently-dissolved PAH concentrations were, on average, dominated by 314 fluoranthene (310 pg L⁻¹), followed by pyrene (140 pg L⁻¹), benz(a)anthracene (100 pg $\rm L^{-1}$) and retene (50 pg $\rm L^{-1}$). These concentrations were much lower than what we determined here. Venier et al. (2014) collected samples during the spring season,

 before Lake Superior stratified during the summer (see temperature profiles for May and July in Figure SI 2). Most likely, PAH concentrations across Lake Superior increased strongly during the summer of 2011, probably due to wildfires, which 320 peaked in July (see Figure SI 3): retene was present in the ng L^{-1} range at all sites 321 during this study, while Venier et al. (2014) only reported 50 pg L^{-1} . The comparison between active and passively sampled PBDEs (see below) similarly displayed significant differences between both sampling approaches.

 PAH Air-Water Exchange: Atmospheric deposition is a major source of 325 pollutants to the Great Lakes.^{44,45,12} Chemical exchange across the air-water interface is a key process controlling concentrations and residence times of semi-volatile 327 pollutants in these water bodies.^{46,47,34} PE sampling improves upon previous estimates of air-water PAH exchange by (i) a greater spatial coverage; (ii) simultaneously measured time-averaged concentrations for both the air and water dampening diurnal effects and minimizing anomalous signals; (iii) by only measuring gas- or dissolved- phase compounds directly available for exchange across the air-water interface, and (iv) by providing concentrations from open Lake locations. To our knowledge these are the first open lake passive measurements for PAHs in the Great Lakes. The direction of air-water exchange was determined by the ratio of the concentration of an analyte in the PE at equilibrium with the air to the concentrations of the analyte in the PE at equilibrium with the water: 337 Flux direction = C_{PEa}/C_{PEw} (5) Uncertainty propagation indicated that ratios needed to be < 0.14 and >3.9 to determine net deposition or volatilization (see SI page 9). Hence, a ratio >3.9 indicated a higher activity in the air, and therefore, net deposition, while a ratio <0.14 resulted in net volatilization.

Polybrominated Diphenylethers (PBDEs)

Spatial Distributions: Σ_{11} BDE concentrations in air were more than three orders of magnitude lower than PAHs across Lake Superior (Figure 4A, Table SI 19). BDE-47 was generally the most abundant congener present (approximately 20 to 60% in air and 40-75% in water samples) throughout the sampling period, followed by BDE-28, 49, 99, and 100. BDE-153 concentrations were low or below detection at all most sites and BDE-154 was not detected in any samples. Changes in atmospheric and

 aqueous PBDE concentrations between deployments at a given site were smaller than variations among sites. In other words, the proximity to emission sources was more important than geochemical variations (currents, winds, temperature).

 Atmospheric PBDE concentrations were greatest at Marquette (June-October 368 average 15 pg m⁻³) (Table 2). This station is located at a dock for a coal-fired power plant and in proximity to a number of sites registered with the EPA Toxic Release 1870 Inventory, including facilities that manufacture polyurethane and explosives,⁴⁸ where PBDEs might have been used in the past. Sault Ste. Marie and Duluth also had 372 elevated atmospheric PBDE concentrations from June-October, 7.4 and 5.7 pg m⁻³, 373 respectively. Atmospheric concentrations at nearly all other sites were ≤ 1 pg m⁻³. The eastern open lake site (ODAS45004) exhibited concentrations >10 times higher than the other open lake sites, perhaps receiving input downwind from Marquette. Water concentrations had similar spatial distributions as air concentrations (Figure 4B, Table SI 20). Concentrations were greatest at Marquette (June-August 378 average 6.8 pg L^{-1}), followed by Thunder Bay/Welcome Island, Pointe Aux Pins, and 379 Sault Ste. Marie (June-October average 2.1, 1.8, and 1.7 pg L^{-1} , respectively). Dissolved PBDE concentrations were lowest at rural Canadian sites: Sturgeon Bay 381 (0.20 pg L⁻¹), Foster Island (0.21 pg L⁻¹), and Michipicoten Bay (0.46 pg/L). Eastern and western open lake sites exhibited PBDE concentrations similar to coastal rural 383 sites (0.46 and 0.24 pg L^{-1} , respectively), with the exception of Eagle Harbor (0.97 pg L⁻¹). Average central open lake PBDE concentrations were slightly higher than the 385 other open lake sites (0.86 pg L^{-1}), similar to Station 139 (0.75 pg L^{-1}) near Thunder Bay/Welcome Isle.

410 elevated active sampling results), and the efficient retention of PBDEs associated with 411 DOC and possibly small particles by active sampling.

 PBDE Sources: PBDE concentrations in both the air and water were correlated with industrial and urban sites, reflecting their origin as flame retardants used widely in consumer, commercial, and industrial products. Atmospheric deposition is an important source of PBDEs to Lake Superior and is probably the main 436 pathway to open-lake sites.⁴⁵ Physico-chemical properties indicate that lighter (less 437 brominated) congeners will travel farther from PBDE sources.¹⁸ Tri-, tetra-, and pentaBDEs were most commonly detected at Lake Superior sites, whereas lighter and heavier congeners were typically below detection. These congeners would typically stem from the penta-BDE commercial mixture. We cannot exclude that this is a reflection of the environmental fractionation of BDEs, their use, and our sampling approach. Waning PBDE sources may explain the reduced atmospheric PBDE 444 concentrations found in this study compared to IADN-derived measurements. Σ_7BDE

445 gas-phase concentrations from 2005-2006 were 2.8 ± 0.28 pg m⁻³.² Our atmospheric concentrations were an order of magnitude lower at Eagle Harbor, possibly indicating the positive effect of reduced emissions in the region. Likewise, our Eagle Harbor 448 fluxes indicate reduced gas deposition (1.5 kg y^{-1}) since 2005-2006 $(18\pm9.5 \text{ kg y}^{-1})$,² however, urban locations may contribute to greater deposition. It should be noted that the passive samplers deployed in our study only detect truly gas-phase compounds, while IADN operates high-volume samplers. The observed decrease in PBDE concentrations over time could thus at least partially be due to methodological differences.

Implications

 Although long-range atmospheric transport is a major pathway for PAH and PBDE distribution across Lake Superior, point sources along the lake perimeter significantly impact local and regional concentrations. Fluxes for both currently- emitted POPs were generally into the lake near industrial and urban sources, whereas the open lake sites appeared to volatilize POPs back into the atmosphere at much lower rates. Thus, Lake Superior combines primary source-driven deposition of POPs at its periphery with serving as a secondary source of POPs back to the atmosphere in its interior. Enhancing spatial and temporal coverage of the Great Lakes region will provide meaningful trends in PAH and PBDE concentrations and fluxes as use and regulations change. We anticipate PAH concentrations to remain constant or even increase with time, but PBDE concentrations may decrease as use is phased out. Changes in atmospheric concentrations could lead to an equilibrium reversal and the volatilization of PBDEs out of Lake Superior into the atmosphere. Such trends can be monitored effectively and affordably with passive samplers. There is evidence of a strong discrepancy between active and passive sampling results for POPs in water, which will impact our ability to derive air-water fluxes accurately.

Supporting Information Available

Additional information on sampling locations, physicochemical constants, sampling

rates, QA/QC, analysis, calculations and concentrations of PAHs and PBDEs and their

air-water exchange available free of charge via the Internet at http://pubs.acs.org/.

Acknowledgements

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643 **Tables and Figures**

Table 1: Σ_{21} PAH concentrations in air (ng m⁻³) and water (ng L⁻¹) by deployment period

645 Σ_2 PAH = sum of biphenyl, acenaphthylene, acenaphthene, fluorene, phenanthrene, anthracene, methyl phenanthrenes, fluoranthene, pyrene, retene, benz(a)anthracene, chrysene, benzo(b)fluoranthene, benzo(b)fluoranthen benzo(k)fluoranthene, benzo(a)pyrene, benzo(e)pyrene, perylene, benzo(j)fluoranthene, indeno(1,2,3-cd)pyrene, dibenzo(a,h)anthracene, benzo(g,h,i)perylene

 $bd =$ below detection limit of $\overline{GC}/\overline{MS}$

 N/A = not available due to lost polyethylene samplers

Figure 1: Average Σ_{20} PAH (as per Table 1 minus retene) concentrations at each station. (A) Air concentrations; average from June to October 2011 for all stations except Marquette and Eagle Harbor, which only represent June-August. (B) Water concentrations; average from June to October 2011 for all stations except Marquette, which only represents June-August, and Pointe Aux Pins, which represents August-October 2011.

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Figure 2: Changes in retene concentration (Δ) between June-August and August-October 2011.

Figure 3: (a) PAH and (b) PBDE fluxes (Σ_{21} PAHs as per Table 1, Σ_{11} PBDE as per Table 2) across the air-water interface of Lake Superior. Positive values indicate volatilization from the surface water, while negative values indicate atmospheric deposition. Only values from significant C_{PEa}/C_{PEw} ratios were averaged. PAH averages (Figure 3A) from June to October 2011 for all stations except Marquette and Eagle Harbor which only account for the second deployment (June-August 2011), and Pointe Aux Pins which only accounts for the third deployment (August-October 2011). Eastern refers to buoys Station 23 and ODAS45004, Central is Station 113 and ODAS45001, and Western is Station 169 and ODAS45006. PBDE averages (Figure 3B) from June to October 2011 for all stations except Marquette which was from June-August 2011.

Figure 4: Average Σ11PBDE (as defined by Table 2) concentrations at each station. (A) Air concentrations (pg m⁻³) are averaged from June to October 2011. (B) Water concentrations (pg L^{-1}) are averaged from June to October 2011 for all stations except Marquette, which represents June-August.

	Air (pg m^{-3})				Water (pg L^{-1})			
Site	April-	June-	August-	June-October	April-	June-	August-	June-October
	June	August	October	Average	June	August	October	Average
Sault Ste. Marie	1.4	12	2.6	7.4	3.7	1.4	1.9	1.7
Pointe Aux Pins		0.54	0.49	0.51		1.8	1.8	1.8
Ashland	0.56	1.5	0.77	1.1	2.7	0.93	1.1	1.0
Thunder Bay/Welcome Isle		0.77	0.84	0.80		3.0	1.3	2.1
Station 139						0.73	0.76	0.75
Duluth	1.4	6.0	5.3	5.7		1.2	0.86	1.1
Marquette	0.92	24	6.3	15	3.5	6.8	N/A	N/A
Ontonagon		1.3	0.26	0.75		0.37	0.64	0.50
Michipicoten Bay		0.27	0.38	0.32		0.69	0.23	0.46
Sturgeon Bay		0.14	0.38	0.26		0.31	0.086	0.20
Foster Island		0.25	0.44	0.35		0.19	0.23	0.21
Eagle Harbor	0.046	0.21	bd	0.11		0.99	0.94	0.97
Eastern Open Lake		0.48	0.87	0.68		0.34	0.58	0.46
Central Open Lake		0.030	bd	0.015		0.53	1.2	0.86
Western Open Lake		0.031	bd	0.015		0.42	0.065	0.24

Table 2: Σ_{11} PBDE concentrations in air (pg m⁻³) and water (pg L⁻¹) by deployment period

 Σ_{11} PBDE = sum of BDE-2, 8, 15, 28, 30, 47, 49, 99, 100, 153, 154

 $bd = below detection limit of GC/MS$

N/A = not available due to lost polyethylene samplers