1	Sources and fate of polycyclic aromatic compounds (PAHs,							
2	oxygenated PAHs and azaarenes) in forest soil profiles opposite of an							
3	aluminium plant							
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31 Abstract

Little is known about oxygenated polycyclic aromatic hydrocarbons (OPAHs) and azaarenes 32 33 (AZAs) in forest soils. We sampled all horizons of forest soils from five locations at increasing distances from an Al plant in Slovakia, and determined their polycyclic aromatic compound 34 35 (PACs) concentrations. The Σ 29PAHs concentrations were highest in the Oa and lowest in the Oi horizon, while the Σ 14OPAHs and Σ 4AZAs concentrations did not show a consistent 36 37 vertical distribution among the organic horizons. The concentration ratios of PAHs and OPAHs between deeper O horizons and their overlying horizon (enrichment factors) were positively 38 correlated with the octanol-water partition coefficients (K_{OW}) at several locations. This is 39 attributed to the slower degradation of the more hydrophobic PACs during organic matter 40 decomposition. PACs concentrations decreased from the organic layer to the mineral horizons. 41 The concentrations of $\sum 29$ PAHs (2400 – 17000 ng g⁻¹), $\sum 140$ PAHs (430 – 2100 ng g⁻¹) and 42 Σ 4AZAs (27 – 280 ngg⁻¹) in the mineral A-horizon generally decreased with increasing 43 distance from the Al plant. In the A horizons, the concentration of Σ 29PAH were correlated 44 45 with those of $\sum 140PAH$ (r = 0.95, p = 0.02) and $\sum 4AZA$ (r = 0.93, p = 0.02) suggesting that 46 bioturbation was the main transport process of PACs from the organic layer into the mineral soil. At each location, the concentrations of PACs generally decreased with increasing depth 47 48 of the mineral soil. Enrichment factors of PAHs in the mineral horizons were not correlated with Kows, pointing at colloid-assisted transport and bioturbation. At some sites, the 49 50 enrichment factors of OPAHs were negatively correlated with K_{OW} indicating that these compounds were leached in dissolved form. Compared to a study, 13 years before, the 51 52 concentrations of PAHs had decreased in the O horizons but increased in the A and B horizons because of soil-internal redistribution after emissions had been reduced. 53

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59 **1. Introduction**

Environmental contamination with polycyclic aromatic compounds (PACs) such as polycyclic 60 aromatic hydrocarbons (PAHs), oxygenated PAHs (OPAHs) and nitrogen heterocyclic PAHs 61 (azaarenes, AZAs) is a source of concern because of their (eco)toxicological properties, 62 persistence and bioaccumulation. PAHs have been the main focus of previous studies on soils 63 (Wilcke, 2007; 2000). However, recent studies show that fractions of soil extracts that contain 64 the much less studied polar derivatives of PAHs (e.g., OPAHs and AZAs) and pure OPAHs 65 and AZAs can induce (eco)toxicological effects of similar or higher magnitude as the PAH-66 67 containing fractions or pure PAHs (Anyanwu and Semple, 2016; Bandowe and Wilcke, 2011; Brinkmann et al., 2014; Lundstedt et al., 2007; Misaki et al., 2016; Wincent et al., 2016; 2015). 68 69 Some OPAHs and AZAs are also more persistent and bioaccumulative than PAHs (Anyanwu and Semple, 2016, 2015; Arp et al., 2014; Lundstedt et al., 2007). 70

All PACs are produced and emitted to the environment from incomplete combustion of fossil 71 fuels and biomass (Bleeker et al., 2002; Booth and Gribben, 2005; Lima et al., 2005; Lundstedt 72 73 et al., 2007). OPAHs are also formed by post-emission transformation of PAHs via photolysis, photooxidation, chemical degradation, thermal degradation and biodegradation (Bandowe and 74 Wilcke, 2011; Lundstedt et al., 2007; Walgraeve et al., 2010). AZAs and OPAHs have higher 75 water solubility and lower vapour pressure than their related PAHs. They have functional 76 groups that allows for more diverse mechanisms of partition, sorption and transport in air and 77 78 soil.

Soil is the major final sink of PACs in the environment, with some estimates indicating that up to 90% of PAHs in the United Kingdom environment are stored in soils (Wild and Jones, 1995).
Such estimates are however, to date not available for the polar PACs such as OPAHs and AZAs. PACs emitted by anthropogenic activities can be deposited to soils by dry/wet deposition and direct air-soil exchange from contaminated air. PACs can also be scavenged by 84 vegetation and reach the soil via litterfall (Cousins et al., 1999; Moeckel et al., 2008). Forest soils are a major sink for atmospheric organic micropollutants because of the scavenging effect 85 of the tree canopy with its large surface area (Horstmann and McLachlan, 1999; Matzner, 86 87 1984). PACs in forest soils undergo sorption-desorption, transformation, degradation, transport (leaching, dissolved organic (DOM) and colloid-assisted 88 matter transport), volatilization/condensation and bioaccumulation that eventually determine their spatial 89 90 distribution (Krauss et al., 2000; Moeckel et al., 2008; Semple et al., 2003). Concentrations of PAHs in soils generally decrease exponentially with distance from a point source (Van 91 92 Brummelen et al., 1996a; Wilcke et al., 1996). The contributions of higher molecular weight PAHs (HMW-PAHs, >4 benzene rings) to the total PAHs concentrations usually progressively 93 94 decrease with distance from point sources because of their stronger association with particles 95 and rapid deposition to the soil (Van Brummelen et al., 1996a; Yang et al., 1991). The spatial 96 distribution of OPAHs and AZAs in forest soil with distance from point emissions sources has not yet been investigated. It was, however, shown in grassland soils near a power and industrial 97 98 plants of the Angren region, Uzbekistan, that the spatial distribution of OPAHs was similar to those of PAHs (Bandowe et al., 2010). 99

100 The processes that influence the concentrations, composition patterns and vertical distribution of PAHs in the three organic (O) horizons of the organic layer on top of mineral soils are 101 different from those that influence the vertical distribution in the mineral soil (Krauss et al., 102 2000; Pichler et al., 1996; Wilcke, 2000). The organic layer of forest soils consists of up to 103 three distinct horizons (Oi, Oe and Oa) of increasing depth and degree of decomposition. Oi, 104 Oe and Oa horizons are defined as fresh undecomposed litter, partially decomposed and 105 completely decomposed organic matter, respectively (Krauss et al., 2000). Depending on soil 106 properties and climate, these horizons show different states of development and might even be 107 absent. The concentrations of total PAHs in the organic horizons of forest soils generally 108

increase in the order Oi < Oe < Oa (Krauss et al., 2000; Pichler et al., 1996; Wilcke, 2000).
The proportion of HMW-PAHs in organic horizons of forest soils also increase in the order Oi
< Oe < Oa. This is because of the higher persistence of HMW-PAHs (than the LMW-PAHs)
and hence higher proportions of the LMW-PAHs than of HMW-PAHs being biodegraded
during the decomposition of organic matter (Krauss et al., 2000; Pichler et al., 1996; Wilcke et al., 1996).

115 In the mineral horizons, total PAHs concentrations and the contribution of HMW-PAHs to total PAHs concentrations generally decrease with increasing depth (Guggenberger et al., 1996; 116 Wilcke, 2000). The vertical distribution in the mineral horizon is driven by the higher 117 118 concentrations of organic matter in topsoils (A horizons) of forest soils that serve as a strong sorbent for the PAHs. The depth distribution of PAHs can be explained by the vertical transport 119 via leaching as truly dissolved solute (mainly LMW-PAHs) and DOM/colloid-assisted 120 121 transport and retention (mainly HMW-PAHs, Guggenberger et al., 1996; Totsche et al., 1997). The stronger leaching of truly dissolved LMW-PAHs explains the decreasing contributions of 122 the more hydrophobic HMW-PAHs to total PAHs concentrations with increasing mineral soil 123 depth. The individual OPAH/related-PAH concentration ratios in the mineral horizons were 124 125 found to increase with increasing depth because of the higher water solubility of the OPAHs 126 than of their related PAHs (Bandowe et al., 2011, 2010). Like the OPAH/related-PAHs ratios, it is expected that the AZA/related PAHs ratios will increase with increasing depth of the 127 mineral soil. 128

To study the distribution and transport of OPAHs and AZAs, we selected a beech forest (*Fagus sylvatica* L.) near an aluminium plant in Žiar nad Hronom (Central Slovakia). Aluminium smelting on industrial scale releases significant amounts of PAHs (and possibly OPAHs and AZAs), trace metals and other pollutants from both combustion of fossil fuels used to produce energy that powers the smelting process and from the carbonaceous electrode material used in

aluminium smelting itself (Booth and Gribben, 2005; Maňkovská and Steinnes, 1995; Wilcke 134 et al., 1996). This site offers an ideal location to study the distribution of PACs (OPAHs, AZAs 135 and PAHs) originating from a point source as well as the depth distribution and the underlying 136 processes that drive these distributions. The distribution and composition of PAHs in soils 137 sampled from these sites in 1994 was reported earlier (Wilcke et al., 1996). Shortly before the 138 study of Wilcke et al. (1996), the Al smelter company had undergone restructuring and installed 139 140 new technologies for both production and emission control (Jamnická et al., 2007; Maňkovská and Steinnes, 1995; Wilcke et al., 1996). This study will therefore also provide useful data 141 142 about the effect of the introduction of modern technology in aluminium production and emission control as well as about the long-term behaviour of PAH in soils. The overall goal of 143 this study is to gather knowledge about the sources (direct emission vs. post-depositional 144 145 formation), spatial distribution, cycling and transport of PACs (PAHs, OPAHs and AZAs) in forest soils. Our work is guided by the following research questions: 146

- What does the distribution of PAHs, OPAHs and AZAs with increasing distance to the
 smelter tell us about its sources and mechanism of transport?
- 2. What does the (depth) distribution of PACs tell us about the mobility, transport
 mechanism of the different PAC groups, and the post emission formation of OPAHs?
 We will test the hypothesis that the concentrations of OPAHs and the OPAH/relatedPAH concentration ratios will increase in the order Oi < Oe < Oa because of the
 increasing formation (and accumulation) of OPAHs from the transformation of PAHs
 which accompany the decomposition of organic matter.
- 3. What does the change in concentration and spatial distribution of PAHs from 1994 to
 2007 reveal about the long-term behaviour of PAHs in soils?
- 157 2. Materials and methods

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158 2.1 Sampling and soil characterization

In September 2007, we sampled all horizons of the soils at five sites with increasing distances (ca. 500, 1000, 1500, 2000, 2500 m) from an aluminium plant in Žiar nad Hronom (Central Slovakia, Figure 1). The Al plant has been in operation since 1953 and was reconstructed in 1986, when the emissions were substantially lowered (Jamnická et al., 2007; Maňkovská and Steinnes, 1995). The operation of the smelter caused a severe contamination of its surroundings with PAHs, heavy metals and fluoride (Maňkovská and Steinness, 1995; Wilcke et al., 1998, 1996).

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Figure 1

At each location, we collected samples from all horizons of Typic Dystrudepts or Dystric 167 Cambisols (sites 1, 2, 4 and 5) and Aquic Dystrudepts or Stagnic Dystric Cambisols (site 3, 168 IUSS Working Group WRB, 2014; Soil Survey Staff, 2014), that were accessible by hand 169 digging (Oi, Oe, Oa, A, B and in some soils C horizons) from a soil pit. All samples were 170 171 collected on the same day. Samples were kept in closed aluminium containers and stored fresh at -18 °C in a freezer until analysis. Several physical and chemical properties (texture, soil pH, 172 effective cation-exchange capacity (ECEC), total organic carbon (TOC) and inorganic carbon 173 174 (IC) were determined on sub-samples that were dried in an oven at 40 °C, then gently crushed and homogenized (organic layers) or sieved (< 2 mm, mineral soil). The procedures for the 175 determination of the physico-chemical properties are described in detail in previous papers 176 (Bandowe et al., 2010; Bandowe and Wilcke, 2010; Wilcke et al., 2014a, b). Properties of the 177 soil are reported in Table S1 (supplementary information). 178

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180 2.2 Analysis of PAHs, OPAHs and AZAs

181 About 2-5 g (ground organic horizons) and 13-16 g (mineral soil horizons) from each sampling site were weighed, mixed with inert bulk sorbent (Isolute HM-N, Uppsala, Sweden) and 182 transferred into 33-mL extraction cells (accelerated solvent extractor, ASE 200, Dionex, 183 Sunnyvale, CA). Samples were spiked with 11 perdeuterated PAHs (naphthalene-D₈, 184 acenaphthene- D_{10} , phenanthrene-D₁₀, benzo[a]anthracene-D₁₂, 185 pyrene-D₁₀, benzo[k]fluoranthene-D₁₂, benzo[a]pyrene- D_{12} , chrysene- D_{12} , 186 pervlene- D_{12} , dibenzo[a,h]anthracene-D14, and benzo[ghi]perylene-D₁₂: 150 μ L of a 10 μ g mL⁻¹ solution of 187 each of the compounds in toluene) as internal standard for PAHs and benzophenone-D₅ (150 188 μ L of a 10 μ g mL⁻¹ solution of the compound in toluene) as internal standard for OPAHs and 189 AZAs. Each sample was extracted twice by pressurized liquid extraction with the ASE 200. 190 191 The first extraction was with dichloromethane (DCM), while the second extraction was with 192 acetone/DCM/1% trifluoroacetic acid (250:125:1 v/v/v). The ASE was operated with the same 193 settings as reported previously (Bandowe and Wilcke, 2010). The two extracts from each sample were combined and passed through a Na₂SO₄ column to remove water. Hexane (1 mL) 194 was added to each extract, rotary evaporated and again 10 mL hexane was added and rotary 195 evaporated to < 1 mL. The extracts of organic horizons were then cleaned up by gel permeation 196 chromatography (GPC). The GPC system consisted of a PrepLinc GPC, equipped with a UV 197 detector (254 nm), an autosampler and glass column (750 mm x 25 mm ID glass column) filled 198 with swelled Bio-beads S-X3 (J2-Scientific, PA, USA). The GPC was run with 199 200 hexane:dichloromethane (1:1 v/v) as the mobile phase at a flow rate of 5 mL min⁻¹.

Extracts of mineral and organic horizons (cleaned on GPC) were transferred onto a column made of 3 g silica gel (10% deactivated). PAHs were eluted with 15 mL hexane:dichloromethane followed by the elution of OPAHs and AZAs with 8 mL dichloromethane and 5 mL acetone. The PAHs fractions and OPAHs+AZAs fractions were collected in separate flasks, spiked with toluene, rotary evaporated, spiked with fluoranthene-

 D_{10} (50 µL of 22 a µg mL⁻¹ solution) before being transferred into vials for measurements of 206 their PACs concentrations with a gas chromatograph coupled to a mass spectrometer (GC-MS, 207 GC 7890A with a 5975 MSD, Agilent technologies, Palo Alto, CA). Target compounds were 208 209 quantified with a set of calibration standards that were measured together with samples in the same GC-MS sequence. The internal standard calibration functions were used to quantify 210 masses of all PAC target compounds in our extracts. GC-MS signals, their processing, 211 calibration and quantification were realized with the Agilent ChemStation Software. Further 212 details of the GC-MS parameters and the limits of determinations of our analytical method can 213 214 be found in previous papers (Bandowe et al., 2014; 2011; 2010; Bandowe and Wilcke, 2010; Lundstedt et al., 2014; Wei et al., 2015). 215

216 We determined the concentrations of the following compounds:

29 PAHs: 1,2,3,4-tetrahydronaphthalene (THNAPH), naphthalene (NAPH), 217 2-218 methylnaphthalene (2-MNAPH), 1-methylnaphthalene (1-MNAPH), biphenyl (BP), 1,3-dimethylnaphthalene (1,3-DMNAP), acenaphthylene (ACENY), acenaphthene 219 (ACEN), fluorene (FLUO), phenanthrene (PHEN), anthracene (ANTH), 1-220 221 methylphenanthrene (1-MPHEN), 3,6-dimethylphenanthrene (3,6-DMPHEN), fluoranthene (FLUA), pyrene (PYR), 1-methyl-7-isopropylphenanthrene (RET), 222 benz[a]anthracene $(\mathbf{B}(\mathbf{A})\mathbf{A}),$ chrysene+triphenylene 223 (CHRY), 224 benzo[b+i+k]fluoranthenes (B(BJK)), benzo[e]pyrene (B(E)P), benzo[a]pyrene (B(A)P), perylene (PERY), indeno[1,2,3-cd]pyrene (IND), dibenz[a,h]anthracene 225 (DIBE), benzo[ghi]perylene (B(GHI)) and coronene (COR); 226

14 OPAHs: 1-indanone (1-INDA), 1,4-naphthoquinone (1,4-NQ), 1-naphthaldehyde
(1-NLD), 2-biphenylcarboxaldehyde (2-BPCD), 9-fluorenone (9-FLO), 1,2acenaphthylenedione (1,2-ACQ), 9,10-anthraquinone (9,10-ANQ), 4Hcyclopenta[d,e,f]phenanthrene-4-one (CPHENone), 2-methyl-9,10-anthraquinone (2-

MANQ), benzo[a]fluorenone (B(A)FLUone), 7H-benz[d,e]anthracene-7-one
(BANTone), benzo[a]anthracene-7,12-dione (7,12-B(A)A), 5,12-naphthacenequinone
(5,12-NACQ) and 6H-benzo[c,d]pyrene-6-one (BPYRone)

4 AZAs: quinoline (QUI), benzo[h]quinoline (BQI), acridine (ACR) and carbazole
(CBZ)

236 2.3 Quality assurance and control

All used organic solvents were of high purity for pesticide residue analysis or HPLC grade, 237 purchased from Carlo Erba (Milan, Italy). All labware were machine-washed with soap, rinsed, 238 239 and dried in a clean oven. All the non-volumetric glassware and metallic labware were also baked at 250 °C for 24 h to remove contaminants. Three procedural blanks made of bulk inert 240 sorbent (Isolute, HM-N) were extracted and their PACs concentrations measured. Most of the 241 target compounds were not detected in the blanks or their amounts in the blanks were 242 insignificant compared to their amounts in the real samples. The mean amounts of PACs the 243 244 three blanks (where detected) was then subtracted from those measured in the samples to correct for laboratory contamination. We also extracted and analysed replicates (n = 2) of the 245 certified reference material ERM-CC013a-PAHs in soils (Federal Institute for Materials 246 Research Testing [BAM], Berlin, Germany). The relative standard deviation (RSD) of 247 Σ 29PAH, Σ 14OPAHs and Σ 4AZAs concentrations in the replicates (n = 2) of ERM-CC013a 248 determined with our procedures were 8%, 9% and 12%, respectively. The recoveries of the 249 250 sum of PAH concentrations for which indicator and certified values were provided was $135\pm12\%$ (average \pm standard deviation). For the OPAHs and AZAs, there are no certified 251 reference materials. We therefore compared our OPAHs and AZAs concentrations to the grand 252 means determined for this same reference material in a recent intercomparison exercise 253 254 (Lundstedt et al., 2014). The recoveries compared to the grand means in the intercomparison

exercise were $101 \pm 9\%$, $83 \pm 10\%$ for the OPAHs and AZAs respectively. At the location 256 2500 m from the Al plant, the A-horizon soil was analyzed in replicates (n = 2). The RSD of 257 the $\sum 29$ PAHs, $\sum 14$ OPAHs and $\sum 4$ AZAs were 24%, 7.4% and 7.1%, respectively. At location 258 500 m from the Al plant the Oe and A-horizons were analyzed in replicates (n = 2). The RSD 259 of the replicates analysis of Oe horizon were 53%, 5% and 20%, respectively for $\sum 29$ PAHs, 260 $\sum 14$ OPAHs and $\sum 4$ AZAs. For the A horizon at this location the RSD for the replicate analysis 261 were 31%, 16% and 24% respectively for $\sum 29$ PAHs, $\sum 14$ OPAHs and $\sum 4$ AZAs, respectively.

262 2.4 Calculations and statistical analyses

The sum of the concentrations of all analyzed PAHs, OPAHs and AZAs are called Σ 29PAHs, 263 \sum 14OPAHs and \sum 4AZAs, respectively. The sum of the concentrations of all non-alkylated 264 PAHs, the sum of the 16 US-EPA PAHs, the sum of all non-alkylated PAHs with 2-3 rings, 265 266 and 4-7 rings are called Σ 21PAHs, Σ EPA-PAHs, Σ LMW-PAHs and Σ HMW-PAHs, respectively. We calculated the enrichment factors e for each compound as the concentration 267 of a compound in a soil horizon, divided by the concentration of that same compound in the 268 horizon above it within the same soil profile. Concentration data were $\log (x+1)$ (where x is 269 concentration) transformed to improve normal distribution of the data sets. Statistical analysis 270 271 was performed with STATISTICA 7.0 software (Statsoft, Hamburg, Germany). Statistical tests were set to be significant at a p < 0.05. Octanol-water partition coefficients (K_{OW}) for PAHs 272 were taken from the literature (Mackay et al., 2006; Neff et al., 2005), while those of OPAHs 273 and AZAs were estimates with K_{OW}WIN version 1.67 EPI SuiteTM (US EPA: 274 http://www.epa.gov/opptintr/exposure/pubs/episuitedl.htm). 275

276 **3. Results**

The concentration of $\sum 29$ PAHs in the Oi horizon was lowest at the location farthest from the Al plant, but the $\sum 29$ PAHs concentrations at the other locations were similar (Table 1, Figure

279	2). The PAHs mixtures in the Oi horizons were dominated by PHEN (25%), PYR (10%),
280	NAPH (8%) and ANTH (8%) (Figure 3, Table S2). The 2+3-ring PAHs dominated the non-
281	alkylated PAHs mixtures in the Oi horizon, with a mean contribution of 57% to the $\sum 21$ PAHs
282	concentrations (Figure S1).

- 283
- 284 Figure 2
- 285

Figure 3

Table 1

At most locations, the Oe horizons showed higher concentrations of Σ 29PAHs than the Oi 286 horizons. The highest concentrations of Σ 29PAHs in the Oe horizons occurred at the location 287 farthest from the Al plant, with clearly elevated concentrations at the other locations (Table 1, 288 Figure 2). Moreover, PAHs mixtures were dominated by PHEN (13%), B(BJK) (12%), PYR 289 (10%) and FLUA (10%), showing a higher contribution of HMW-PAHs to the PAHs mixtures 290 291 than in the Oi horizons (Figure 3, Table S2). The 4-ring PAHs (41%) contributed a higher proportion to the $\sum 21$ PAHs compared to that of $\sum 2+3$ -ring PAHs (25%) (Figure S1). The 292 ∑HMW-PAHs/∑LMW-PAHs concentration ratios in the Oe horizons averaged 3.4 (range: 1.5-293 6.8) with the highest and lowest values at locations closest and farthest from the Al plant, 294 respectively. 295

The concentrations of $\sum 29$ PAHs in the Oa horizons were much higher than in the two other O horizons. The highest concentration of $\sum 29$ PAHs in the Oa horizons was at the location closest to and the lowest farthest away from the Al plant (Table 1, Figure 2). The PAHs mixtures were dominated by 2-MPHEN (14%), B(BJK) (12 %), PHEN (12%) and RET (11 %). The contribution of the $\sum 2+3$ ring PAHs, $\sum 4$ -ring PAHs, $\sum 5$ -ring PAHs and $\sum 6+7$ -ring PAHs to the $\sum 21$ PAH were 38%, 33%, 21% and 8%, respectively (Figure 3 & S1, Table S2). The \sum HMW- 302 PAHs/∑LMW-PAHs concentration ratios were 3.8 and 2.4, at the locations closest and farthest
303 from the Al plant, respectively.

- 304 In the mineral A horizon, the concentrations of $\sum 29$ PAHs were highest closest to and lowest
- farthest away from the Al plant (Table 1, Figure 2). The relationship between the concentration
- of $\sum 29$ PAHs and distance from the emission source was exponential (R² = 0.95) (Figure 4).
- 307 The PAHs mixtures in the A horizons were dominated by B(BJK) (18%), PHEN (15%), IND
- 308 (10%) and CHRY (9%) (Figure 3, Table S2). The Σ HMW-PAHs/ Σ LMW-PAHs concentration
- ratio in the A horizon averaged 2.9 (range: 2.6 3.3) and was independent of the distance from
- the Al plant and higher than in the O horizons.

The concentrations of $\sum 29$ PAHs in the deeper mineral horizons (B, Bg and C) did not show consistent decreases with increasing distance from the Al plant. The concentrations of $\sum 29$ PAHs were consistently lower with increasing depth of the mineral horizon (i.e. A > B > C) (Table 1).

315

Figure 4

The concentrations of the \sum 140PAHs in the Oi, Oe and Oa horizons were generally higher at 316 locations closest to the plant but did not show consistent changes with distance from the Al 317 plant (Table 1, Figure 2). At the location closest (500 m from Al plant) to the Al plant, the 318 \sum 14OPAHs concentrations in the Oi horizon were higher than in the Oe horizon, while at 319 320 location 2000 m from the Al plant, the reverse was true (Table 1). The concentrations of Σ 14OPAHs in the three O horizons at location 2500 m from the Al plant decreased in the order, 321 322 Oi > Oe > Oa. The composition of the OPAHs mixtures in the Oi horizon was dominated by 9-FLO (28%), B(A)FLUone (21%) and BPYRone (13%). The OPAHs mixtures in the Oe 323 horizon were dominated by 9,10-ANQ (26 %), 9-FLO (14%) and BANTone (11%) and those 324

in the Oa horizon by BPYRone (39%), 1,2-ACQ (19%) and 9,10-ANQ (17%) (Figure 3B,
Table S3).

The \sum 14OPAHs concentrations in the mineral A horizon (Table 1, Figure 2) were highest 327 closest to and lowest farthest away from the Al plant. Similar to those of the Σ 29PAHs, the 328 concentrations of $\sum 150$ PAHs in the A horizon decreased exponentially with distance from the 329 Al plant ($R^2 = 0.90$; Figure 4). The OPAHs mixtures in the A horizons were on average 330 dominated by 9,10-ANQ (29%), 7, 12-B(A)A (25%), B(A)FLUone (17%) and 9-FLO (8%) 331 (Figure 3B, Table S3). Interestingly, the Σ 14OPAH/ Σ 29PAH ratio was highest in the one 332 horizon, showing clear visible indications of redox changes (location 3, Bg horizon; Table 2). 333 Here the ratio was higher than in all other horizons, except the Oi horizons closest to the 334 smelter. 335

Similar to those of the $\sum 14$ OPAHs, the concentrations of $\sum 4$ AZAs in the Oi horizons were higher than in the Oe horizons at location 1 (500 m from the Al plant) and 5 (2500 m from the Al plant), while the reverse was true at location 4 (2000 m from the the Al plant) (Table 1, Figure 2). The AZAs mixtures in the Oi horizons were dominated by BQI (43%) and ACR (41%), in the Oe horizons by BQI (47%) and CBZ (21%), and in the Oa horizons by BQI (47%) and ACR (44%; Figure 3B, Table S3).

The concentrations of $\sum 4AZAs$ in the A horizon were the highest at locations close to and the lowest farthest away from the Al plant (Table 1, Figure 2). The concentrations of $\sum 4AZAs$ decreased exponentially with distance (R² = 0.93; Figure 4). The composition of the AZAs mixtures in the A horizons were dominated by CBZ (40%) and BQI (37%; Figure 3B, Table S3).

The $\sum 29$ PAHs, $\sum 14$ OPAHs and $\sum 4$ AZAs concentrations in the B horizons were highest at the sampling location farthest from the smelter and decreased with decreasing distance. At the location farthest from the Al plant the $\sum 29$ PAHs and $\sum 14$ OPAHs concentrations were only slightly lower in the B compared to the A horizons, while $\sum 4$ AZAs concentrations were identical (Table 1).

To identify the drivers of the vertical distribution, we correlated the enrichment factors e at 352 each site with the octanol-water partition coefficients [log K_{OW}]. For the PAHs, significant 353 correlations occurred between enrichment factors and their K_{OW} values at site 2 $e_{Oe/Oi}$ (r = 0.57, 354 p < 0.001), site 3 $e_{Oe/Oi}$ (r = 0.64, p < 0.001), site 3 $e_{A/Oe}$ (r = 0.42, p = 0.01), site 4 $e_{A/Oe}$ (r = 355 0.44, p = 0.03) and site 5 e_{Oe/Oi} (r = 0.60, p = 0.01) (Figs. S2 & S3). For the OPAHs, significant 356 correlations between enrichment factors (Fig S4) and K_{OW} occurred at site 1 e_{Oe/Oi} (r = 0.76, p 357 = 0.007), site 2 $e_{B/A}$ (r = -0.71, p = 0.020) and site $5e_{Oa/Oe}$ (r = 0.59, p = 0.01). For the AZAs 358 the low number of compounds (n = 4) was not suitable for a correlation analysis between 359 enrichment factors and Kow values. 360

361

362 **4. Discussion**

4.1. Concentrations and distribution of PACs with distance from the Al plant

364 The concentrations of Σ 29PAHs in the O horizons of the soils were in the range of those of Σ US-EPA PAHs (105 – 14889 ng g⁻¹) reported in an extensive (n = 447) nationwide survey of 365 German forest soils covering both background and contaminated sites (Aichner et al., 2013; 366 2015). The concentrations of PAHs in the A horizons of the forest soils near Žiar nad Hronom 367 were elevated compared to mineral horizons of forest soils from background and rural areas of 368 the USA (Obrist et al., 2015). The extensive survey of Aichner et al. (2015) in Germany found 369 a concentration range of 20-9038 ng g⁻¹ in the 0-5 cm layer of mineral soils under forest 370 (Aichner et al., 2015), while in mineral horizons of forest soils from northern Czech Mountains 371 the concentration of Σ 20PAHs was 238 – 7476 ng g⁻¹ (Wilcke and Zech, 1997; Wilcke et al., 372

1996). The mean and median concentrations of PAHs in mineral forest soils in the review of 373 Wilcke (2000) were 904 and 410 ng g⁻¹, respectively. Thus, the concentrations of the Σ 29PAHs 374 in the O horizons at the sites nearest (500 - 1000 m) to the Al plant were at the upper end or 375 even above the range reported for forest soils and thus can be considered as strongly 376 contaminated. Even the three sites further away from the Al plant still show concentrations 377 above background. Interestingly, the A horizon was dominated by higher proportions of the 378 379 HMW-PAHs compared to the O horizons suggesting that the PAHs did not reach the A-horizon by leaching (which would have favoured the LMW-PAHs). Instead, we suggest that the main 380 381 mechanism of transfer of PAHs from the Oa to the A-horizon was biological mixing.

The concentrations of $\sum 14$ OPAHs in our study were much higher than those of $\sum 9$ OPAHs measured in organic layers of remote forest soils of USA, the only other study of OPAHs concentration in organic layers of forest soils, we are aware of (i.e. 6 ± 6 to 39 ± 25 ng g⁻¹) (Obrist et al., 2015) indicating the substantial enrichment of OPAHs in the forest soils near Žiar nad Hronom, because of the emissions from the Al plant. The OPAHs could have been directly emitted together with PAHs from the Al plant, with additional OPAHs formed by postemission transformation of PAHs (Lundstedt et al., 2007; Wilcke et al., 2014b).

The dominance of 9-FLO and 9,10-ANQ is similar to findings from organic horizons and mineral forest soils in the USA (Obrist et al., 2015). 9-FLO and 9,10-ANQ were also found to be the most abundant compounds in mineral soils at other locations and in combustion emissions (Bandowe et al., 2014; 2011;2010; Layshock et al., 2010; Vicente et al., 2016).

We are not aware of any measurements of AZAs concentrations in forest soils near an Al plant. However, AZAs were detected in ambient air samples near an Al smelter in Canada, so that the presence of AZAs in our study soils can be expected (Roussel et al., 1992). The concentrations of the Σ 4AZAs in surface soils from the cities of Bangkok (Thailand) and Xi'an

(China) were 0.1-31 ng g⁻¹ and 784 ng g⁻¹, respectively (Bandowe et al., 2014; Wei et al., 397 2015a). The concentrations of Σ 9AZAs in the mineral topsoil (0-10 cm) of an industrial site in 398 the Czech Republic were 640-694 ng g⁻¹, while the concentration in a reference rural site was 399 6.9 ng g^{-1} (Švábenský et al., 2009). The concentrations in the A horizons at our study site were 400 higher than in soils from background and remote locations (e.g., Argentina Σ 4AZA: n.d – 0.97 401 ng g⁻¹, Wilcke et al., 2014a) but comparable to those from industrial sites as summarized in 402 Bandowe et al. (2014). The dominance of CBZ is similar to that observed in a study of urban 403 soils in Bangkok (Bandowe et al., 2014). Carbazole was also measured at concentration of 8-404 268 ng g^{-1} , not detected (n.d.) – 37 ng g^{-1} and n.d. – 10 ng g^{-1} in the organic horizon, 0-5 cm 405 and 5-10 cm of mineral horizons, respectively of German forest soils (Mumbo et al., 2016). 406 The study of Mumbo et al. (2016) also showed the trend organic horizon > 0.5 cm > 5.10 cm 407 408 in the concentrations of carbazole, which is contrary to the trend we observe in the vertical distribution of the Σ 4AZA. In our study the Σ 4AZAs concentration in the A-horizon was 409 higher than in the O-horizons at most locations. 410

The distribution of PAHs, OPAHs and AZAs in the organic horizons demonstrate that the Al 411 plant was the main point source for the PACs in O horizons of soils. Although the location 412 2500 m away from the emission source was clearly least affected, there was no systematic 413 increases in PACs concentrations in each of the organic horizons with decreasing distance to 414 the Al plant. Several factors could explain the lack of distance-concentration relationships 415 including differences in adsorbing surface area among the sampled organic layers, age 416 differences between similar layers at different locations, soil properties (Table S1), 417 composition of tree species, topography, turnover times and vertical transport into deeper 418 horizons. 419

420 In the A horizons, the concentrations of $\sum 29$ PAHs correlated significantly with those of 421 $\sum 14$ OPAHs (r = 0.95, p = 0.016) and $\sum 4$ AZAs (r = 0.93, p = 0.02), which is in contrast to the

O horizons. The concentrations of several individual PAHs, OPAHs and AZAs were also 422 significantly correlated with each other in A horizons. Such correlations have been observed 423 424 in other studies in soils and are interpreted as indication of the common origin (emissions from combustion processes taking place in the Al plant) and similar post-emission fates of these 425 three groups of PACs (Bandowe et al., 2014; Obrist et al., 2015; Wilcke et al., 2014). In the 426 study soils, the close correlation of many PACs concentrations with each other can likely be 427 428 attributed to biological mixing from the organic layer into the mineral A horizon. The higher pH at the two sites nearest to the Al plant could be indicative of a higher biological activity and 429 430 thus faster biological mixing of the organic layer into the mineral soil. However, these sites were at the same time more strongly contaminated with a number of organic and inorganic 431 pollutants and the higher pH is attributable to the deposition of alkaline dusts associated with 432 433 the pollutant emissions (Maňkovská et al., 1995; Wilcke et al., 1998; 1996). Because the stronger contamination of these sites likely reduces biological activity in spite of the higher pH, we do not expect 434 435 marked differences in biological mixing or O with A horizons.

The concentrations of each of the PAC groups in the A-horizons showed a close correlation 436 with distance from the Al plant, indicating that emissions from the Al plant were the major 437 source of most of the studied PACs. Similar exponential relationships between distance from 438 emission source and concentrations of PAHs were also observed in previous studies in forest 439 440 soils at the same and other locations (Van Brummelen et al., 1996a; Wilcke et al., 1996). In the B horizon, the concentration of PACs did not consistently decrease with increasing distance 441 from Al plant (or indeed even tended to show an opposite trend to what was observed for the 442 A horizons, Table 1). This points to the importance of soil properties for the transport of PACs 443 in the mineral soil, which is overprinting the higher deposition close to the smelter. With 444 increasing distance from the Al plant, the organic matter content as well as the ECEC decreases 445 446 (Table S1), offering less sorption sites, particularly for the more polar PACs and thus favour

greater translocation of PACs into deeper horizons of the mineral soils. Another reason for the stronger depth transport at the farther distances might be the higher sand concentrations allowing for a faster percolation of water and solutes, particularly at site 5.

The ∑HMW-PAHs/∑LMW-PAHs concentration ratios in the Oi horizons averaged 0.85, with the highest value at the location closest to the smelter and lowest at the location farthest away probably indicating a distillation effect, i.e. faster deposition of HMW-PAHs into soils, and longer-range atmospheric transport of LMW-PAHs because of the higher proportion of HMW-PAHs that is partitioned to the particle phase of air (Wei et al., 2015b: Wilcke et al., 1996; Yang et al., 1991).

The ratios of individual OPAHs or AZAs to their related PAHs (and presumed precursors of 456 the OPAHs) are used to indicate whether transformation and transport processes taking place 457 with increasing soil depth lead to the accumulation of OPAHs or AZAs. Several of the 458 OPAHs/related PAH ratios were > 1, indicating the higher abundance of the OPAHs relative 459 to their related and often monitored PAHs (Table 2). Similar observations have been 460 461 consistently reported in soils, air and combustion emissions (Bandowe et al., 2014, 2011, 2010; Vicente et al., 2016; Walgraeve et al., 2010; Wei et al., 2015a,b; Wilcke et al., 2014; Layshock 462 et al., 2010) and can result from primary emission sources or from post-emission 463 transformation of the PAHs to OPAHs. 464

465

Table 2

466 4.2 The depth distribution as indicator for sources and transport of PACs.

In general, the concentration of $\sum 29$ PAHs in the O horizons increased in the order Oi < Oe < Oa (Figure 2, Table 1). The concentrations of $\sum 29$ PAHs in the Oa horizons were higher or similar to the concentration of the underlying A horizons (Figure 2). The vertical trends are in line with previous studies (Krauss et al., 2000; Pichler et al., 1996). The mineral horizon 471 showed consistent decreases of $\sum 29$ PAH concentration with increasing depth largely 472 consistent with previous studies (Guggenberger et al., 1996; Krauss et al., 2000). The organic 473 matter concentrations in the mineral horizons, which followed similar trends, can partly explain 474 the decreasing PAH concentrations, because organic matter is the main sorbent of PAHs in 475 soils (Table S1, Wilcke, 2000).

In the O horizons, the enrichment factors of PAHs were at several study sites significantly 476 positively correlated (p < 0.05) with the K_{OW} values (Figure S2) reflecting the accumulation of 477 the more hydrophobic PAHs with increasing organic matter transformation (Pichler et al., 478 1996; Wilcke and Zech, 1997). The HMW-PAHs are less bioavailable, less degraded (more 479 480 persistent) compared to the more hydrophilic, more bioavailable LMW- PAHs, which are increasingly degraded with increasing organic matter transformation (Guggenberger et al., 481 1996; Krauss et al., 2000; Pichler et al., 1996). We did not find significant correlations of the 482 483 enrichment factors with K_{OW} values in the mineral horizons suggesting that the translocation of PAHs from the organic layer into the mineral soil horizons and between the mineral horizons 484 were partly controlled by DOM- and colloid-assisted transport (Guggenberger et al., 1996; 485 Krauss et al., 2000; Wilcke et al., 1996). We interpret the fact that none of the PAH enrichment 486 factors at the location closest to the Al plant correlated with K_{OW} as strong indication that the 487 higher impact of fresh emissions from the Al plant might be influencing the vertical 488 distribution. 489

In contrast to the $\sum 29$ PAHs concentrations, we could not identify a consistent pattern of increasing concentrations of $\sum 14$ OPAHs and $\sum 4$ AZAs from the Oi to the Oa horizons (Table 2, Figure 2). We attribute this finding to the fact that OPAHs can both be degraded and formed during advancing decomposition of organic matter (Wilcke et al., 2014b), both processes compensating each other. 495 In few cases the enrichment factors of the OPAHs in the O horizons were significantly positively correlated with the K_{OW} values (Figure S4). In these soils at sites 1 and 5, the 496 preferential degradation of the OPAHs with advancing organic matter decomposition seems to 497 498 overwhelm the formation of new OPAHs from parent PAHs. Similarly, there were weak but significant correlations of the enrichment factors of OPAHs between the A and Oa horizons 499 and the K_{OW} values (Fig. S4) at site 2 suggesting that at this site leaching of OPAHs from the 500 501 organic layer into the mineral soils played a role, in spite of the dominating transfer of PACs from the O into the A horizons by biological mixing. This underlines the higher water solubility 502 503 and resulting leaching of OPAHs than of PAHs in soil. In the mineral horizons, the enrichment factors of OPAHs were significantly negatively correlated with the Kow values at the location 504 1000 m from the Al plant (Figure S4), suggesting that the vertical translocation of OPAHs at 505 506 this site was mainly driven by leaching as truly dissolved compound.

507 At locations 1 (500 m from the Al plant) and 5 (2500 m from the Al plant), the only two sites in which we detected OPAHs in two to three O horizons, the $\sum 140PAHs/\sum 29PAHs$ 508 509 concentration ratios and the individual OPAHs/related-PAHs decreased from the Oi via the Oe 510 to the Oa horizons (Table 2). The OPAH/PAH ratio in the Oi horizon reflects the ratios in the primary emission (or transformation in the atmosphere/on plant surfaces) (Weiss, 2000). This 511 512 suggests that the degradation of organic matter (Oi to Oa) mainly reduces the concentrations of OPAHs relative to PAHs, which may indicate faster degradation of OPAHs (as a result of 513 higher bioavailability) compared to PAHs or faster leaching of OPAHs out of the organic 514 horizon (as a result of higher water solubility) relative to the PAHs. In the mineral horizons, 515 we found increases in the $\sum 140PAHs/\sum 29PAHs$ concentration ratios with increasing depth. 516 However, this trend was only clear in the profiles in which a Bg or C horizon was present (500 517 m and 1500 m from the Al plant). In the other profiles and generally from the A to the B horizon 518 the Σ 14OPAH/ Σ 29PAH concentration ratios only showed minor changes. The 1,2-519

520 ACQ/(ACENY+ACEN) concentration ratios increased consistently in all study soils. This was occasionally also true for some other individual OPAH/related PAH concentration ratios (Table 521 2). Such vertical concentration trends suggest faster translocation of the OPAHs, particularly 522 of 1,2-ACQ into deeper mineral horizons by leaching as truly dissolved solute. We attribute 523 the lack of consistent results for other OPAHs or AZAs than 1,2-ACQ to the fact that a 524 multitude of site-specific soil properties interact with OPAHs and AZAs in a way that either 525 inhibits or enhances the transport independent of their water solubility (Bi et al., 2006; Weigand 526 et al., 2002). The increased Σ 14OPAHs/ Σ 29PAHs ratios and some individual OPAHs or 527 528 AZAs/related PAHs ratios in the B and C horizons, might point at a higher risk of ground water resources to be contaminated with PAC mixtures that contain high fractions of OPAHs and 529 AZAs (Schlangers et al., 2008; Weigand et al., 2002), which could have possible 530 531 (eco)toxicological consequences, considering the higher toxicity of some of these OPAHs and AZAs than of their related PAHs (Anyanwu and Semple, 2016; Bleeker et al., 2002; Lundstedt 532 et al., 2007). 533

Another explanation for the high $\sum 140$ PAHs/ $\sum 29$ PAHs, particularly in the Bg horizon at site 534 3 might be the fact, that this soil horizon showed visible influence of redox changes (mottled 535 pattern of bleached and red zones). These redox changes might also influence the formation of 536 OPAHs, relative to PAHs. Redox conditions are known to influence decomposition of PAHs 537 (Leduc et al., 1988) and the alternating redox conditions, affect and change the microbial 538 community (Hanke et al. 2013). The alternating redox conditions in soils might affect 539 formation of OPAHs, when the soil is reoxidized. However, as we have only one sample with 540 an indication for redox changes, these hypotheses needs further testing. 541

4.3 What does the change of PAH distribution from 1994 to 2007 tell us about thetransport of PAH in soils.

22

544 In the previous study from 1994, when three soils were sampled along the same deposition gradient (distance 1320 to 2180 m), the concentrations of $\sum 20$ PAHs were 38596 – 72523, 545 57273 - 93785 and 35409 - 73003 ng g⁻¹ in the Oi, Oe and Oa horizons (Figure 5; Wilcke et 546 al., 1996), respectively indicating marked decreases in the concentrations of PAHs in samples 547 taken in 2007 compared to those taken in 1994 (Wilcke et al., 1996). In the previous study, the 548 concentrations of Σ 20PAHs in the A horizons ranged from 1172 to 2485 ng g⁻¹ and in the B 549 horizon from 151 to 376 ng g⁻¹ (Figure 5) and thus were higher in 2007 than in 1994. These 550 results illustrate that the organic layers recovered from their high PACs load in 1994, because 551 552 of their limited residence time and the emission control measures established from 1986 on (Jamnická et al., 2007; Maňkovská and Steinness, 1995). The PAHs in the highly contaminated 553 organic layers, however, were not fully degraded but incorporated into the mineral horizons by 554 555 biological mixing leading to higher concentrations in the A and B horizons in 2007 than in 1994. Our results illustrate that the disappearance of a contaminant load from soil organic 556 layers in forests does not necessarily mean that the compounds were degraded. The reductions 557 we see are also consistent with the emission of PAHs from the Žiar nad Hronom Al smelter 558 which has decreased from 68 t a^{-1} (in 1990), to 12 t yr⁻¹ (in 1995) and 0.8 t yr⁻¹ (in 2001) 559 (Jamnická et al., 2007). 560

561

Figure 5

562 **5. Conclusions**

This study showed significant inputs of OPAHs, AZAs and PAHs originating from
an Al plant into forest soils. The spatial distribution of PAHs, OPAHs and AZAs in
the A and some organic horizons identified the Al plant as common source. The lack
of a systematic relationship between the distance from the Al plant and the
concentrations of PACs in the B underlined the importance of soil properties for

vertical PACs redistribution in soils, because the study soils varied particularly withrespect to their pH values and sand concentrations.

The vertical distribution of PAHs and OPAHs in the organic horizons could be 570 • explained by the relative accumulation of the more hydrophobic PACs with 571 increasing organic matter transformation. The concentrations of PAHs in the A 572 horizons were significantly correlated with those of OPAHs and AZAs, which is 573 attributable to their common primary source and mixing from the organic layer into 574 the mineral soil by biological activity. The vertical distribution of PAHs in the 575 mineral soil was not significantly correlated with their octanol-water partition 576 coefficients, because of DOM- and colloid-assisted leaching. At several locations, 577 the vertical distribution of OPAHs in the soils was related with the octanol-water 578 partition coefficient demonstrating that leaching in truly dissolved form played a 579 more important role for OPAHs than for PAHs. This was further backed by increases 580 in the ratios of individual OPAHs to PAH ratios with increasing depth of the mineral 581 soil at several locations. 582

• The PAHs concentrations in the O horizons have significantly decreased between 1994 and 2007, because of improved emission control, turnover of the O horizons at a decadal scale and mixing into the mineral soil. Despite decreasing emissions in recent times, the concentrations of PAHs in the A and B horizons have increased from 1994 to 2007 as a result vertical transport.

588

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- 596 chemical characterization of the soils.

Sampling point	Distance from the Al plant	Horizon designation	∑29PAHs	∑14OPAHs	∑4AZAs	
#	[m]			ng g-1		
1	500	Oi	1100	1800	110	
1	500	Oe	6200	43		
1	500	Oa	21000	na	na	
1	500	А	17000	2873	280	
1	500	В	670	670 130		
1	500	С	61	33	0.10	
2	1000	Oi	1500	na	na	
2	1000	Oe	6500	950	45	
2	1000	Oa	17000	na	na	
2	1000	А	8400	2000	200	
2	1000	В	520	90	5.2	
3	1500	Oi	1900	na	na	
3	1500	Oe	2200	840	28	
3	1500	А	4800	740	58	
3	1500	В	1000	210	17	
3	1500	Bg	47	61	1.9	
4	2000	Oi	na	300	4.7	
4	2000	Oe+Oa	1900	620	32	
4	2000	А	2600	740	60	
4	2000	В	980	320	14	
5	2500	Oi	800	350	54	
5	2500	Oe	680	150	14	
5	2500	Oa	1800	120	17	
5	2500	А	2400	430	27	
5	2500	В	1900	320	27	

Table 1. Sum of the concentrations of the three studied PACs groups.

^ana is not available.

Location/		1,4-	1-NLD/1-	9-	1,2-	9,10-	7,12-				
Horizon	∑140PAH/∑29PAH	NQ/NAPH	MNAPH	FLO/FLUO	ACQ/(ACENY+ACEN)	ANQ/ANTH	B(A)A/B(A)A	QUI/NAPH	BQI/PHEN	ACR/ANTH	CBZ/FLUO
1-Oi	1.6	0.04	1.3	36	6.2	1.2		0.3	0.40	0.41	0.20
1-Oe	0.12		0.01	1.5	0.36	1.1		0.03	0.02	0.05	0.09
1-Oa											
1-A	0.17	0.01	0.35	6.3	0.04	1.6	1.1	0.01	0.04	0.11	3.1
1-B	0.19			3.7		1.5	1.7		0.03	0.09	0.95
1-C	0.53					0.90	0.81			0.01	
2-Oi											
2-Oe	0.15		0.06	4.1	0.65	1.1	0.38	0.05	0.05	0.05	0.39
2-Oa											
2-A	0.24		0.47	9.7	0.42	2.1	2.1	0.07	0.04	0.17	4.3
2-B	0.17			5.7	0.80	0.78	0.16		0.02		2.0
3-Oi											
3-Oe	0.39	0.01	0.12	4.6	4.5	1.1	0.65	0.14	0.01	0.03	0.26
3-A	0.15	0.01	0.12	5.0	0.14	0.92	1.3	0.01	0.05	0.02	2.1
3-B	0.21			10	0.83	0.76	2.3	0.02	0.03	0.06	2.5
3-Bg	1.3		0.14	10	7.6	1.0	7.4	0.02	0.01	0.03	5.1
4-Oi											
4-Oe	0.33	0.01		3.1	1.6	1.6	3.0	0.02	0.14	0.01	0.29
4-A	0.29		0.20	9.5	0.06	1.5	3.7		0.07	0.12	2.4
4-B	0.33			8.9	0.20	1.5	4.3		0.04		2.4
5-Oi	0.43			2.2	13	0.96		0.04	0.15	0.52	
5-Oe	0.22			1.0	5.2	0.95		0.07	0.08	0.02	
5-Oa	0.07		0.02	1.1	2.7	0.31		0.02	0.05	0.11	
5-A	0.18		0.42	3.6	0.16	0.82	1.9	0.01	0.02	0.01	1.6
5-B	0.17		0.16	7.8	0.18	1.8	2.1	0.01	0.04	0.15	2.7

Table 2: Ratios of the concentration of OPAHs or AZAs to related PAHs.

Figure Legends

Figure 1: Study sites with sampling locations.

Figure 2: Concentrations of PACs in five soil profiles at sites [a] 500 m [b] 1000 m [c] 1500 m [d] 2000 m and [e] 2500 m from the Al plant.

Figure 3: Average composition pattern of the mixtures of [a] PAHs and [b] OPAHs and AZAs.

Figure 4: Relationships between the concentration of PACs in the A horizon and distance from the Al plant. The continuous line is the best fit for PAHs, the interrupted one for OPAHs and the dotted one for AZAs.

Figure 5: Comparisons of the mean concentrations of PAHs in [a] Oi [b] Oe [c] Oa, [d] A and [e] B horizons between the sampling campaigns in the years 2007 and 1994. The boxes show the range, whisker show 5th and 95th percentile, the central line is the median and the squares dots is the mean of the data set





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Figure 3









Figure 5



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