1 Nitro-polycyclic aromatic hydrocarbons - gas-particle partitioning, mass size distribution, and formation along transport in marine and continental background air 2 3 Gerhard Lammel ^{1,2*}, Marie D. Mulder ¹, Pourya Shahpoury ², Petr Kukučka ¹, Hana Lišková 4 ¹, Petra Přibylová ¹, Roman Prokeš ¹, Gerhard Wotawa ³ 5 6 ¹ Masaryk University, Research Centre for Toxic Compounds in the Environment, Brno, 7 Czech Republic 8 ² Max Planck Institute for Chemistry, Multiphase Chemistry Department, Mainz, Germany 9 ³ Zentralanstalt fuer Meteorologie und Geodynamik, Wien, Austria 10 11 *lammel@recetox.muni.cz 12 13 **Abstract** 14 15 Nitro-polycyclic aromatic hydrocarbons (NPAH) are ubiquitous in polluted air but little is 16 known about their abundance in background air. NPAHs were studied at one marine and one 17 continental background site i.e., a coastal site in the southern Aegean Sea (summer 2012) and a site in the central Great Hungarian Plain (summer 2013), together with the parent 18 compounds, PAHs. A Lagrangian particle dispersion model was used to track air mass history. 19 Based on Lagrangian particle statistics, the urban influence on samples was quantified for the 20 first time as a fractional dose to which the collected volume of air had been exposed to. 21 At the remote marine site, the 3-4 ring NPAH (sum of 11 targeted species) concentration was 22 23.7 pg m⁻³ while the concentration of 4-ring PAHs (6 species) was 426 pg m⁻³. 2-23 nitrofluoranthene (2NFLT) and 3-nitrophenanthrene were the most abundant NPAHs. Urban 24 fractional doses in the range <0.002–5.4% were calculated. At the continental site, the Σ_{11} 3-25 4rNPAH and Σ_6 4rPAH were 58 and 663 pg m⁻³, respectively, with 9-nitroanthracene and 26 2NFLT being highest concentrated amongst the targeted NPAHs. The NPAH levels observed 27 in the marine background are the lowest ever reported and remarkably lower, by more than 28 one order of magnitude, than one decade before. Day-night variation of NPAHs at the 29

continental site reflected shorter lifetime during the day, possibly because of photolysis of 30 some NPAHs. The yields of formation of 2NFLT and 2-nitropyrene (2NPYR) in marine air 31 seem to be close to the yields for OH-initiated photochemistry observed in laboratory 32 experiments under high NO_x conditions. Good agreement is found for prediction of NPAH 33 34 gas-particle partitioning using a multi-phase poly-parameter linear free energy relationship. Sorption to soot is found less significant for gas-particle partitioning of NPAHs than for 35 PAHs. The NPAH levels determined in the southeastern outflow of Europe confirm 36 intercontinental transport potential. 37

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Keywords: long-range transport potential, semi-volatile organic compounds, PAH photochemistry,

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1. Introduction

PAHs may undergo chemical transformations in the gaseous and in the particulate phase 43 (Finlayson-Pitts and Pitts, 2000; Keyte et al., 2013). Nitro-PAHs (NPAHs) observed in urban 44 and rural areas (Nielsen et al., 1984; Feilberg et al., 2001; Finlayson-Pitts and Pitts, 2000; 45 Keyte et al., 2013) and predicted based on smog-chamber experiments (Atkinson and Arey, 46 1994), seem to be most significant derivatives: Mutagenicity of atmospheric aerosols in 47 general is mostly related to NPAHs (Grosjean et al., 1983; Garner et al., 1986; Finlayson-Pitts 48 and Pitts, 2000; Claxton et al., 2004; Hayakawa, 2016). A large part, more than one third, of 49 the mutagen potential of ambient aerosols may be attributable to NPAHs (Schuetzle, 1983). 50 Secondary formation of NPAH from PAHs is thought to occur on short time scales (hours). 51 52 This has been observed for PAHs collected on filters (Ringuet et al., 2012a; Zimmermann et al., 2013; Jaryasopit et al., 2014a, 2014b), and also in in urban plumes (Bamford and Baker, 53 2003; Arey et al., 1989; Reisen and Arey, 2005). Although many NPAHs are emitted from 54 road traffic, only a few are abundant in this source type (Arey, 1998; Keyte et al., 2013 and 55 2016; Inomata et al., 2015; Alves et al., 2016). The occurrence of various isomers of 56 nitrofluoranthene (NFLT) and nitropyrene (NPYR) can be used to study PAH sources, PAH 57 chemical transformations and the role of the photo-oxidants hydroxyl radical (OH) and nitrate 58

radical (NO₃) (Ciccioli et al., 1996; Finlayson-Pitts and Pitts 2000). E.g., 3- and 2-

nitrofluoranthene (3-, 2NFLT) are indicative of primary and secondary sources, respectively.

These substances have been suggested as tracers for air pollution on the time scales of hours

- 62 to days (Ciccioli et al., 1996; Finlayson-Pitts and Pitts 2000; Keyte et al. 2013), but their
- atmospheric lifetimes are still unknown.
- 64 Like their precursors, NPAHs are semivolatile organic compounds (SVOCs), partitioning
- between the phases of the atmospheric aerosol. Similar to other SVOCs, the NPAHs' phase
- distribution was found to depend on temperature (summer and winter campaigns in the Alps;
- 67 Albinet et al., 2008b) and results from both absorptive as well as adsorptive contributions
- 68 (Tomaz et al., 2016). NPAHs have primarily been observed in polluted areas (e.g. Pitts et al.,
- 69 1985; Ramdahl et al., 1986; Garner et al., 1986; Albinet et al., 2007 and 2008a; Ringuet et al.,
- 70 2012a and 2012b; Zimmermann et al., 2012; Barrado et al., 2013; Li et al., 2016). Though
- there are a few studies in rural environments i.e., in Germany (Ciccioli et al., 1996), in the
- French Alps (100-1000 pg m⁻³ range for the sum of 10 NPAHs; Albinet et al., 2008a) and in
- northern China (Li et al., 2016). Very few measurements have been performed in the remote
- atmospheric environment i.e., in the Mediterranean (Tsapakis and Stephanou, 2007), high
- altitude sites in the Himalayas (single data; Ciccioli et al. 1996) and French Alps (Albinet et
- al., 2008a), and in the Arctic (with so-called Arctic haze; Masclet et al. 1988; Halsall et al.
- 77 2001). With regard to the long-range transport potential, the state of the knowledge is that at
- least some NPAHs are expected to go into intercontinental transport (Lafontaine et al., 2015)
- and might be ubiquitous in the global atmosphere (Ciccioli et al., 1996).
- 80 However, there is limited NPAH data from remote atmospheric environments is obvious and
- 81 little is known about their long-range transport potential. The aim of this study was to
- characterise the long-range transport potential of NPAHs by measurements at remote sites of
- 83 Europe, addressing the continental background and the outflow of the continent.

85 **2. Methodology**

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86 2.1 Sampling

- 87 High-volume air sampling was conducted at a marine background site, Finokalia
- 88 (35.3°N/25.7°E, 250 m a.s.l.), in the context of a coordinated field experiment July 2-13 2012
- 89 (Lammel et al., 2015) and at a continental background site in central Europe, K-puszta
- 90 (46°58'N/19°33'E, 125 m a.s.l.; Degrendele et al., 2016), August 5-16 2013. The Finokalia site
- 91 is located on a cliff at the northern coast of Crete, some 70 km east of major significant
- anthropogenic emissions (Iraklion, a city of 100000 inhabitants with airport and industries;
- 93 Mihalopoulos et al., 1997; Kouvarakis et al., 2000). The K-puszta site is located on a clearing,
- 94 characterised by uncultivated grassland, in a mostly coniferous forest in the Hungarian

- 95 (Pannonian) Great Plain, ca. 70 km and 270 km southeast of Budapest and Vienna,
- 96 respectively (≈2 mn inhabitants each). The background site character of both observatories
- 97 was demonstrated (Borbély-Kiss et al., 1988; Kouvarakis et al., 2000; Vrekoussis et al., 2005).
- 98 Meteorological parameters and trace gases are measured at both observatories, which are
- 99 stations of the EMEP network (EMEP, 2015).
- 100 High volume air samples were collected using a HV-100P (Baghirra, Prague, Czech
- 101 Republic), equipped with a multi-stage cascade impactor (Andersen Instruments Inc.,
- Fultonville, New York, USA, series 230, model 235) with five impactor stages, corresponding
- to 10–7.2, 7.2–3, 3–1.5, 1.5–0.95 and 0.95–0.49 μm of aerodynamic particle size, D, (spaced
- roughly equal $\Delta log D$), a backup filter collecting particles < 0.49 μm and, downstream, two
- polyurethane foam plugs (PUFs, Molitan, Břeclav, Czech Republic, density 0.030 g cm⁻³,
- placed in a glass cartridge), together 10 cm high. Particles were sampled on slotted QFF
- substrates (TE-230-QZ, Tisch Environmental Inc., Cleves, USA, 14.3 × 13.7 cm) and glass
- fibre filters (Whatman, 20.3×25.4 cm). The filters were cleaned prior to use by heating
- 109 108 (330°C). PUFs were cleaned (8 hour-extraction in acetone and 8 hours in
- dichloromethane (DCM)), wrapped in two layers of aluminum foil, placed into zip-lock
- polyethylene bags and kept in the freezer prior to deployment. The sampler was operated at
- 112 constant flow rate of 68 m³ h⁻¹. Day/night sampling (changing at sunset and sunrise) of
- gaseous samples (PUF) was performed at both sites ($V = 600-1000 \text{ m}^3$), while at the marine
- site the impactor filter (QFF) samples were collected over 24 h (5) or 48 h (3).
- PUFs were cleaned (8 hour-extraction in acetone and 8 hours in dichloromethane (DCM)),
- wrapped in two layers of aluminum foil, placed into zip-lock polyethylene bags and kept in
- the freezer prior to deployment. The sampler was operated at constant flow rate of $68 \text{ m}^3 \text{ h}^{-1}$.
- Day/night sampling of gaseous samples (PUF) was performed at both sites (12 h, $V \approx 700 \text{ m}^3$),
- while at the marine site the impactor filter (QFF) samples were collected over 12 h (n = 1), 24
- 120 h (4) or 48 h (3).
- Particle number concentration, N, was determined by an optical particle counter (Grimm
- model 107, Ainring, 31 channels between 0.25 and 32 mm of aerodynamic particle diameter,
- 123 D). Aerosol surface concentration, S (cm⁻¹), was derived as $S = \pi \Sigma_i N_i D_i^2$ assuming
- sphericity. Hereby, true S will be underestimated, in particular if particles of irregular form
- were abundant (e.g. Jaenicke, 1988). Comparisons with absolute methods (e.g. Pandis, et al.
- 126 1991) suggest that the discrepancy may reach up to a factor of 2-3. The mass median diameter

127 (D_m , μm), was derived as $\log D_m = \Sigma_i m_i \log D_i / \Sigma_i m_i$ with m_i denoting the mass in size class

i, D_i being the geometric mean diameter collected on stage i of the cascade impactor.

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2.2 Chemical analysis

- All air samples were extracted with DCM using an automatic warm Soxhlet extractor (Büchi
- B-811, Switzerland). Deuterated PAHs (D8-naphthalene, D10-phenanthrene, D12-perylene;
- Wellington Laboratories, Canada) were used as surrogate standards for both PAHs and
- NPAHs. Deuterated PAHs proved to be suitable surrogate standards for NPAHs. These were
- spiked on each PUF prior to extraction. The extract was split in two parts, 1/9 for PAHs and
- Nitro-PAHs analysis, 9/10 for PBDEs, PCBs and OCPs. The PAHs and Nitro-PAHs aliquot
- was a subject to open column chromatography clean-up. A glass column (1 cm i.d.) was filled
- with 5 g activated silica (150°C for 12 h), sample was loaded and eluted with 10 mL n-
- hexane, followed by 40 mL DCM. The cleaned sample was evaporated under a stream of
- nitrogen in a TurboVap II apparatus (Biotage, Sweden), transferred into a conical GC vial and
- spiked with recovery standard, terphenyl, the volume was reduced to 100 µL.
- 142 GC-MS analysis of 4-ring PAHs (fluoranthene (FLT), pyrene (PYR), benzo(b)fluorene
- 143 (BBN), benzo(a)anthracene (BAA), triphenylene (TPH) and chrysene (CHR)) and 2-4 ring
- NPAHs (1- and 2-nitronaphthalin (1-, 2NNAP), 3- and 5-nitroacenaphthene (3-, 5NACE), 2-
- nitrofluorene (2NFLN), 9-nitroanthracen (9NANT), 3- and 9-nitrophenanthren (3-, 9NPHE),
- 146 2- and 3-nitrofluoranthene (2-, 3NFLT), 1- and 2-nitropyrene (1-, 2NPYR), 7-
- nitrobenz(a)anthracene (7NBAA), 6-nitrochrysene (6NCHR) was performed using a gas
- 148 chromatograph atmospheric pressure chemical ionization tandem mass spectrometer (GC-
- 149 APCI-MS/MS) instrument, Agilent 7890A GC (Agilent, USA), equipped with a 60m ×
- 150 0.25mm × 0.25mm DB-5MSUI column (Agilent, J&W, USA), coupled to Waters Xevo TQ-S
- (Waters, UK). Injection was 1 µL splitless at 280°C, with He as carrier gas at constant flow
- 1.5 mL min⁻¹. The GC oven temperature program was as follows: 90°C (1 min), 40°C/min
- to150°C, 5°C/min to 250°C (5 min) and 10°C/min to 320°C (5 min). APCI was used in
- 154 charge transfer conditions. The isomers 2- and 3NFLT were not separated by the GC method,
- but co-eluted and are reported as sum.
- Recovery of native analytes varied 72-102% for PAHs and deuterated PAHs, 70-110% for
- NPAHs (details see supplementary material (SM), Table S1a). The results were not recovery
- 158 corrected. The mean of field blank values was subtracted from the sample values. Values
- below the mean + 3 standard deviations of the field blank values were considered to be

160 < LOQ. Field blank values of some analytes in air samples were below the instrument limit of

quantification (ILOQ), which corresponded to 0.004-0.069 pg m⁻³ for NPAHs (except for

162 1NNAP for which it ranged 0.60-0.87 pg m⁻³) and 0.010-0.126 pg m⁻³ for 4-ring PAHs

163 (except for FLT and PYR for which it ranged 0.17-0.59 pg m⁻³) (Table S1).

Higher LOQs were determined for some of the NPAHs and for all 4-ring PAHs in gaseous air

samples (PUFs), namely 0.006-0.009 ng (corresponding to 3.5-8.0 pg m⁻³) for 3NACE and

2NPYR, 0.028-0.097 (corresponding to 16-86 pg m⁻³) for 2NNAP, 2NFLT and 1NPYR, and

167 0.10-0.27 ng (corresponding to \approx 60-240 pg m⁻³) for 4-ring PAHs (except for FLT and PYR for

which it was 1.71 and 1.05 ng, respectively, corresponding to $\approx 600-1500$ pg m⁻³). In

particulate phase samples, where separate field blanks for the 2 different QFFs were

determined (on the impactor stages on one hand side and the backup filter on the other hand

side), higher LOQs were determined for some of the NPAHs and for all 4-ring PAHs, namely

172 0.008-0.089 ng (corresponding to 4.6-79 pg m⁻³) for 2NNAP, 2NFLT, 1NPYR and 2NPYR,

173 0.26-0.31 ng (corresponding to 150-274 pg m⁻³) for 9NANT, and 0.05-0.22 ng (corresponding

to \approx 30-200 pg m⁻³) for 4-ring PAHs (except for FLT and PYR for which it was 0.79 and 0.36

175 ng, respectively, corresponding to \approx 200-700 pg m⁻³).

176 The breakthrough in PUF samples was estimated (Pankow, 1989; ACD, 2015; Melymuk et

al., 2016), and as a consequence, 2-3 ring PAHs and 2-ring NPAHs results were excluded

from this study as their sampling may have been incomplete. We, therefore, report $\sum_{\epsilon} 4rPAH$

179 and Σ_{11} 3-4rNPAH.

180 Particulate matter mass (PM_{10}) was determined by gravimetry (microbalance, filters

accommodated to stable temperature and humidity, 3 replicate weighings), and organic matter

182 (OM) and elemental carbon (EC) contents of PM by a thermal-optical method (Sunset Lab.,

183 USA; EUSAAR protocol).

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2.3 Gas-particle partitioning

186 Gas-particle partitioning was studied by applying a multiphase ppLFER model, which was

recently introduced (Shahpoury et al., 2016). In brief, partitioning of semivolatile compounds

in air can be described (Yamasaki et al., 1982), by

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190 (1)
$$K_p = c_{ip} / (c_{ig} \times c_{PM})$$

where K_p (m³air (g PM)⁻¹) is the temperature dependent partitioning coefficient, c_{PM} (g m⁻³) is the concentration of particulate matter in air, c_{ip} and c_{ig} are the analyte (i) concentrations (ng m⁻³) in the particulate and gas phase, respectively. K_p can be predicted using models based on single- and poly-parameter linear free energy relationships (spLFER, ppLFER). spLFER's relate the partitioning coefficient to one physic-chemical property i.e., assume one process to determine the sorption process, while ppLFER's in principle account for all types of molecular interactions between solute and matrix (Goss and Schwarzenbach, 2001). The observed particulate mass fraction data, $\theta = c_p / (c_g + c_p)$ (Table 2), were tested with both a spLFER and a ppLFER model. The spLFER chosen is the widely used K_{oa} model of Finizio et al., 1997 (results presented in the Supplementary material (SM), S2.3). The ppLFER is a multi-phase model recently presented (Shahpoury et al., 2016) and applied for NPAHs (Tomaz et al., 2016). It is based on linear solvation energy relationships (Abraham, 1993; Goss, 2005):

206 (2)
$$\log K_p = eE + sS + aA + bB + lL + c$$

207 (3)
$$log K_p = sS + aA + bB + vV + lL + c$$

where capital letters E, S, A, B, L, and V are solute-specific Abraham solvation parameters for excess molar refraction (describes interactions between π - and lone (n-) electron pairs), polarizability/ dipolarity, solute H-bond acidity, solute H-bond basicity, logarithm of solute hexadecane-air partitioning coefficient (unitless), and McGowan molar volume (cm³ mol⁻¹)/100, respectively (Endo and Goss, 2014). The corresponding parameters e, s, a, b, l, and v reflect matrix-specific solute-independent contribution to K_p. In lack of experimental data, the solute descriptors for NPAHs were taken from M.H. Abraham (personal communication). The multi-phase ppLFER considers adsorption onto soot, (NH₄)₂SO₄, and NH₄Cl, and absorption into particulate organic matter (OM). OM is assumed to be constituted of two separate phases, low to mid molecular mass, both organic soluble and water soluble OM. For these, ppLFER equations for dimethyl sulfoxide-air (representing the low molecular mass range) and for polyurethane ether-air (representing the high molecular mass OM) are used, respectively (Shahpoury et al., 2016).

222 A conventional single-parameter LFER (K_{oa}) model is applied, too.

2.4 Air mass history analysis

225 The HYSPLIT (Draxler and Rolph, 2003) and FLEXPART (Stohl et al., 1998, 2005) models were used to identify air mass histories over 10 and 2 days, respectively. The possible 226 influence of polluted air on samples was quantified using a novel method of applying 227 228 Lagrangian particle statistics (FLEXPART, see SM, S2.2). To this end, for the entire sampling period, one particle per second was released. The model output is generated at 0.062° (≈7 229 km), every 30 minutes and expressed as 'residence time' i.e., a measure of the time particles 230 resided in grid cells. ECMWF meteorological data (0.125°×0.125° resolution, hourly) were 231 used as input. 232

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2.5 Quantification of urban influence on samples

- 235 The potential urban influence for individual samples collected at the marine site was based on
- 236 the fraction of released Lagrangian particles which travelled through an urban boundary layer.
- 237 A backward run from the sampling site was performed with Lagrangian particles (i.e. air
- parcels) being released during the entire sampling period. Three urban areas were considered,
- 239 i.e. Izmir (≈300 km direct distance, 38.2-38.8°N/26.2-27.3°E), Athens (≈300 km, 37.8-
- 240 38.1°N/23.5-23.8°E) and Istanbul (≈500 km away, 40.8-41.1°N/28.6-29.5°E).
- The urban fractional dose, D_u i, an air mass collected in sample i had received for a given
- simulation period Δt can be derived as:

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(4)
$$D_{ui} = \sum_{t} N_{blua}(t) \times \Delta t_{Rblua} / (N_{tot}(t) \times \Delta t_{i})$$

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- with $N_{blua}(t)$ = number of virtual particles within the urban boundary layer during the specific
- 247 time step, model output time resolution $\Delta t_{Rblua} = 0.5$ h, and $N_{tot}(t)$ = number of virtual
- particles present during the specific time step. Under the given flow conditions in the region, a
- 249 2-day time horizon is considered her. Hence, the simulation period is given as:

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251 (5)
$$\Delta t_i = \Delta t_{\text{sample}} + 48h$$

with Δt_{sample} being the sampling time. $D_{u\ i}$ takes values between 0 and 1, corresponding to

254 none or all, respectively, of the entire sample air having crossed the urban boundary layer. The

255 D_u time series with allocation to 3 urban areas is shown in the SM, Fig. S3.

256 The comparison of urban influence in samples of various sample volume, V, requires

normalisation to V, a relative dose (equ. (5), with n = total number of samples collected).

Values of $D_{ru i}$ may exceed 1.

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(6)
$$D_{ru i} = \left[\sum_{n} V_{n} / (nV_{i})\right] \times D_{u i}$$

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262 The urban fractional dose, D_u i, accuracy is limited by the meteorological input data (here

263 0.125°×0.125° resolution, hourly) and boundary layer depth calculation. In the FLEXPART

model, the latter is done according to Vogelezang and Holtslag, 1996.

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3. Results and discussion

The NPAH levels are distinctly lower at the marine than at the continental site, $\Sigma_{11.3-4r}$ NPAH =

268 22.5 and 58.5 pg m⁻³, respectively (Table 1). The NPAHs showing the highest concentrations

were 2NFLT and 3NPHE at the marine (Fig. 1b) and 9NANT and 2NFLT at the continental

site (Fig. 1d, Table 2). The substance patterns (composition of NPAH mixture) at both sites

are similar, though ($R^2 = 0.76$, P > 0.99, t-test). At the marine site, advection was northerly,

with air masses originating (time horizon 10 days) in eastern and central Europe and, towards

the end of the campaign, in the western Mediterranean. The site was placed into the

southeastern outflow of Europe. NO_x (0.2-0.6 ppbv), EC (0.2-0.8 μg m⁻³) and PM_{10} (18.3-39.3

 $\mu g \ m^{-3}$) reflect background conditions. Air mass history analysis suggests that the somewhat

elevated concentration in the first sample collected at the marine site (Fig. 1a) is related to

long-range transport influenced by passage over the urban areas of Izmir and Istanbul (urban

fractional dose $D_u = 5.0\%$, in contrast to the mean which was 1.6%; Fig. S3). Overall, urban

279 fractional dose in the range <0.002-5.4% was received at the marine site. Across all samples

at the marine site, Du is found to be significantly correlated with the pollutant sum

concentrations $\Sigma_{6.4r}^{}$ PAH and $\Sigma_{11.3-4r}^{}$ NPAH (R² = 0.61 and 0.69, respectively, both P > 0.99).

From the marine site data set, subsets of each two samples are formed, representing minimum

283 (i.e., almost no influence from industrialised area 48 hours prior to arrival (hereforth called

'marine background', urban fractional dose $D_u = 0.4\%$) and maximum observed influence 284 (hereforth called 'background with urban influence', $D_u = 3.1\%$; SM Table S2, Figure S3). 285 The results for these subsets are listed in Tables 1-3. Such classification was not deemed 286 meaningful for the samples collected at the continental site, as the relevant source distribution 287 288 in central Europe was too homogeneous during this episode. Advection was mostly from northwest and partly from easterly directions, with air mass origin (time horizon of 10 days) 289 mostly in central Europe and, to a lesser extent in eastern Europe and the western Balkans. 290 The NO₂ (1.2-2.6 ppbv), total carbon (3-6 μ g m⁻³) and PM₁₀ (10.7-46.3 μ g m⁻³) levels during 291 the campaign reflect continental background conditions. 292 The 4-ring PAH concentrations in samples from the continental site on the one hand, and in 293 background air with urban influence collected at the marine site (urban areas 300-500 km 294 295 away) on the other hand, are similar (Table 2). Also, the substance patterns are more similar than when relating all samples at the marine site i.e., $R^2 = 0.88$ (P > 0.999, t-test) instead of R^2 296 297 = 0.76. The investigation of the diffusive air-surface exchange processes during the 298 measurements presented here showed that 4rPAHs were in fact influenced by secondary emissions, namely throughout day and night from the soil at the continental site (by average 299 16.3 and 9.3 pg m⁻² h⁻¹ for FLT and PYR, respectively; Degrendele et al., 2016) or 300 occasionally from surface seawater at the marine site (during at least 1 day-time interval out of 301 302 in total 3 of this data subset; Lammel et al., 2016). In the data set from the continental site, we study day/night (D/N) effects (subsets listed in Tables 1-3, too): PAH concentrations (ctot) 303 were \approx 60% higher during the day than during the night, while c_{tot} of NPAH were by average 304 \approx 5% lower during the day (Table 2). NPAHs are subject to photolysis, while PAHs are not. At 305 306 the site, the PAH concentrations were driven by re-volatilisation from soil, determined by temperature variation (Degrendele et al., 2016). For NPAHs (partly primary emitted) this 307 indicates that the higher emissions during the day (due to re-volatilisation and road traffic) 308 309 were compensated by shorter lifetime. NPAH lifetimes may be limited by heterogeneous photolysis, but available kinetic data are scarce and limited to few aerosol types (Fan et al., 310 1996; Feilberg and Nielsen, 2000, 2001; García-Berríos et al., 2017). Also, different 311 NPAH/PAH ratios (the potential NPAH yields), which were 5.6% and 8.9% at the marine and 312 313 continental sites, respectively, reflect the combination of emission sources and photochemical sinks. The NPAH/PAH ratios at the two sites were influenced by similar substance patterns 314 upon emission, similar irradiation (summer, no or almost no clouds) and deposition velocities 315 (θ in the range 0.05-0.20 for Σ_{11} 3-4rNPAH and Σ_{6} 4rPAH, no precipitation), but different re-316

volatilisation fluxes and different characteristic transport times elapsed. The distance to major urban source areas was 300- >1000 km at the marine and 100-500 km at the continental site. The NPAH/PAH ratios being lower at the more distant receptor site, the marine site, may suggest that photochemical degradation of NPAHs along transport was on average faster than degradation of the precursors. xxx

The NPAH levels observed in marine background air are the lowest ever reported. Remarkably, the concentrations are much lower, by more than one order of magnitude, than one decade before at the same site during the same season (Tsapakis and Stephanou, 2007). The concentrations observed now are a factor of 4-10 lower than in a forest site in Amazonia two decades before (which might have been influenced by biomass burning emissions), a factor of 3 lower (for 2NPYR) than observed at an extremely remote site in the Himalayas two decades before (Ciccioli et al., 1996), and comparable to a high altitude site in the Alps (with the exception of 2NPYR which was observed one order of magnitude higher there in winter; Albinet et al., 2008a; Table 3). The NPAH levels observed at the marine site with influence of pollution and at the continental site are comparable, but also at the lower end of the range spanned by previous observations at rural and remote sites (Table 3).

Gas-particle partitioning

The time-weighted mean NPAH phase distributions (Σ_{11} 3-4rNPAH) differ, corresponding to θ = 0.05 and 0.17 at the marine and continental sites, respectively, – despite similar temperatures (Table 1). In contrast and despite of similar temperature ranges, the 4-ring PAHs' (Σ_6 4rPAH) particulate mass fraction was higher at the marine than at the continental site (θ = 0.42 and 0.20, respectively). Both 4-ring PAHs and 3-4 ring NPAHs were more associated with PM in polluted air than in clean air. This trend is weak for PAHs with θ = 0.02 for Σ_6 4rPAH in marine background but 0.07 in background with urban influence (and θ = 0.09 and 0.20 for CHR; Table 2), but is obviously strong for NPAHs, namely θ = 0.19 for 2NPYR in marine background but 0.69 in background with urban influence, ≈0.93 in polluted continental air, and θ = 0.01 for Σ_{11} 3-4rNPAH in marine background but 0.22 in background with urban influence (Table 2). The urban influenced air at the marine site is also reflected in a much higher OC (a factor of 3 higher than the all-campaign mean) and elevated EC, (less prominent, ≈50% above mean). This confirms the understanding that gas-particle partitioning

of both PAHs (Lohmann and Lammel, 2004; Shahpoury et al., 2016) and NPAHs (Tomaz et 349 al., 2016) is mostly determined by absorption in POM and adsorption to soot. When 350 comparing polluted air at the continental site and background with urban influence at the 351 marine site, a strong shift of Σ_6 4rPAH towards the particulate phase, $\theta \approx 0.21$ vs. 0.07, 352 respectively, is found, while for Σ_{11} 3-4rNPAH θ are similar i.e., ≈ 0.16 vs. 0.22, respectively. 353 This phase partitioning trend of the 4rPAHs could be explained by sorption to EC, which is a 354 factor of \approx 2 higher, but not by OC (only \approx 20% higher). In conclusion, these observations 355 consistently indicate that sorption to soot is less significant for gas-particle partitioning of 356 NPAHs than for PAHs. 357 While NPAHs were significantly phase-shifted ($\theta = 0.24$ during day-time but $\theta = 0.58$ during 358 night-time), this was not the case for 4rPAHs ($\theta = 0.18$ during day-time and $\theta = 0.23$ during 359 360 night-time). This is in line with the perception that the temperature sensitivity of phase change is stronger for the substance class with stronger molecular interactions in the condensed phase, 361 NPAHs. E.g., the enthalpies of phase change between air and OC of FLT and NFLT are -98 362 and -75 kJ mol⁻¹, respectively (OC represented by DMSO; ACD, 2015). 363 Good agreement is found for the prediction of NPAH partitioning using the multi-phase (3-364 phase) ppLFER with most values predicted within one order of magnitude of the observed 365 366 values (Fig. 2; quantification of deviations in S2.3.1). While the sensitivity of assumptions regarding PM phase composition, made in the model do not contribute significantly to the 367 deviations (<< 1, log Kp units), a significant part can be attributed to the usage of estimated 368 solute-specific Abraham solvation parameters (taken from ACD, 2015), in lack of 369 experimentally based descriptors. E.g., for an urban site (Tomaz et al., 2016) it was found that 370 experimentally based descriptors used for 9NPAH lead to better predictions than the estimated 371 372 descriptors i.e., RMSEs differed by 0.43 log units. The agreement of the ppLFER prediction is 373 better than assuming absorption (into OM) to be the only relevant process (K_{0a} model; see S2.3.2, Fig. S5). The same was found when studying gas-particle partitioning of NPAHs in 374 urban air (Tomaz et al., 2016). This supports the perception that gas-particle partitioning of 375 NPAHs is governed by various molecular interactions with OM, with its polarity being well 376 represented by DMSO, better than by octanol. Earlier, it had been found for eight 3-4rNPAHs 377 at urban and rural sites (Li et al., 2016) that the dual model, assuming adsorption (to soot) and 378 absorption (into OM) predicts better than single adsorption (to the total aerosol surface i.e., 379 Junge-Pankow) or single absorption (K_{oa}) models do. 380

381 The interactions with the aerosol matrix of 9NPHE (continental site) and 5NACE, 2NFLN, 2NFLT and 1NPYR (marine site) are less well represented than other NPAHs by the model as 382 suggested by low slopes of their log K_p experimental/log K_p predicted relationships. The reason is 383 384 unknown. Moreover, sampling or sample handling artefacts cannot be excluded, even so same temperature range, sampler and sampling protocols applied across sites with both satisfactory 385 and deficient agreement between predicted and observed K_p. Further conclusions are not 386 supported by the limited amount of data and uncertainties on both the model (estimated ppLFER parameters) and experimental (concentrations close to LOQ) sides.

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387 388 Mass size distribution 390 391 The NPAH mass size distribution had its maximum in the <0.49 µm size range at both sites. The 4-ring PAHs mass size distribution had 2 maxima, <0.49 µm and between 0.95 and 1.5 392 μm, at the marine site, but one at <0.49 μm at the continental site (Table 1). This is probably 393 394 related to the presence of aged aerosol at the marine site vs. a larger contribution of fresh 395 aerosols at the continental site. This is, furthermore, supported by the analysis of air mass 396 origins that shows significant influence of urban areas for only few samples at the marine and 397 for all samples at the continental site (SM S2). Sums of NPAHs' and PAHs' mass size distributions are found unimodal with the maximum 398 in particles <0.49 µm, except PAHs at the marine site, which shows a second maximum 399 400 between 1.5 and 3.0 µm (Fig. 3). At the marine site, 50 and 69% of 1NPYR and 2NFLT, respectively, were found associated with particles <0.45 µm and 68 and 86%, respectively, 401 with particles <0.95 µm, and even more, 83% and 100%, respectively, with particles <0.45µm 402 403 at the continental site. Σ_{c} 4rPAH mass size distributions are shifted to larger particles in background with urban 404 405 influence as compared to marine background air (both collected at the marine site) i.e., MMD = 0.19 and 0.28, respectively. However, such a trend is not apparent for NPAHs (Table 2). The 406 407 size shift of PAHs is not corresponding to the PM₁₀ mass size distribution: The MMD of PM₁₀ 408

for all samples collected at the marine site was 0.58 µm, while it was 1.13 and 0.62 µm in the marine background and background with urban influence data subsets, respectively. The PM₁₀ as well as the OC mass size distributions were bimodal with maxima corresponding to < 0.49 μm and 3.0-7.2 μm particles (MMDs listed in Table 2), while the EC mass size distribution was unimodal, with the maximum concentration in the finest fraction. At the continental site, the Σ_{11} 3-4rNPAH mass size distribution was bimodal with maxima corresponding to < 0.49

 μm and 7.2-10 μm particles, while the Σ_6 4rPAH mass size distribution was unimodal, with the 414 415

maximum concentration in the finest fraction (for all samples as well as for day and night data

subsets; Table 1). 416

The formation of a second maximum, at larger particles than emitted, reflects the 417 418 redistribution of semivolatile organics in an aged aerosol, hence, is expected at receptor sites such as the marine site. This was also observed in polluted air, at rural and suburban sites, but 419 420 not at traffic sites or in winter at a rural site, when primary emissions dominated (unimodal;

421 Albinet et al., 2008b; Ringuet et al., 2012b).

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Substance patterns and NPAH formation during long-range atmospheric transport

Among the targeted NPAHs and apart from NNAPs, which were highest concentrated, 2NFLT and 3NPHE prevailed at the marine site (accounting together for ≈60% of the NPAH mass, excluding the NNAPs), while at the continental site 9NANT and 2NFLT prevailed (accounting for ≈65% together) (Fig. 1, summarised in Fig. S4). The analytical method did not separate the isomers 2NFLT and 3NFLT, but at receptor sites, far from diesel emissions it appears justified to assume $c_{2NFLT} >> c_{3NFLT}$ (Finlayson-Pitts and Pitts, 2000; Zimmermann et al., 2012). The ratio 1NPYR/2NPYR is higher, ≈1, at the continental site than at the marine site (≈ 0.25), which reflects the significance of primary sources for polluted air (Atkinson and Arey, 1994; Finlayson-Pitts and Pitts, 2000; Zimmermann et al., 2012). This ratio was found similarly high or even higher at urban sites (Ringuet et al., 2012c; Tomaz et al., 2016). Similarly, the ratio 2NFLT/1NPYR, the concentration of a secondarily formed over a primary emitted NPAH, has been used as indicator for fresh emissions (if < 5) vs. photochemically aged air mass (Keyte et al., 2013). These values were >> 5 in 21 out of 22 and 7 out of 8 samples at the marine and continental sites, respectively. The only sample collected at the continental site with elevated primary NPAH (2NFLT/1NPYR = 4.3) was possibly influenced by emissions from Budapest, which was passed by the advected air within the last hours before arrival. The only sample collected at the marine site with elevated primary NPAH (2NFLT/1NPYR = 5.9) was indeed directly influenced by emissions into the boundary layer above the Izmir and Istanbul metropolitan areas (urban fractional dose $D_u = 5.0\%$ for samples no. 1 and 2 in Fig. S3). In conclusion, these results from receptor / background sites confirm the existing knowledge about primary emitted and secondarily formed NPAHs. The ratio of two secondarily formed NPAHs, 2NFLT/2NPYR, indicative for day- vs. night-

time formation paths (Atkinson and Arey, 1994; Ciccioli et al., 1996), is found ≈2 at the

marine and ≈ 8 at the continental site (normalised to the precursor ratio i.e., 447 2NFLT/2NPYR/(FLT/PYR); Table 4). Such low values point to day-time (OH initiated) 448 formation, while night-time (NO₃ initiated) formation was negligible, practically excluded at 449 the marine site. This is in line with the perception that NO₃ must have been very low in this 450 remote environment. (NO_x levels at the marine site were in the range 0.2-0.6 ppbv). A similar 451 conclusion had been drawn in a semi-rural environment (Feilberg et al., 2001). 452 For 2NFLT and 2NPYR (secondary sources only) and for 1NPYR, which has mostly primary 453 sources (Finlayson-Pitts and Pitts, 2000; Ringuet et al., 2012a; Jariyasopit et al., 2014a, 454 2014b) we infer the potential yields (Table 4). Here, yield is defined as c_{NPAH}/c_{PAH} (total 455 concentrations). This yield is called 'potential' as it reflects an upper estimate, as other PAH 456 photochemical sinks, such as formation of oxy-PAHs, are neglected. The yield of 2NFLT in 457 polluted air exceeds the one in background air only slightly, while the yield of 2NPYR in 458 polluted air exceeds the one in background air much more (a factor of 3 higher). 459 As expected, the highest potential yield of 1NPYR is found in polluted air (both sites), 460 461 reflecting the dominance of primary emissions of 1NPYR. Similarly, higher yields of secondary NPAHs are found for marine background air compared to background air with 462 urban influence (marine site), reflecting the longer reaction times elapsed since PAH 463 464 emission. The yield for 2NFLT, c_{2NFLT}/c_{FLT} , $\approx 2-4\%$ at both sites ranges higher than the one 465 for 2NPYR, c_{2NPYR}/c_{PYR} , which is found $\approx 0.5-2\%$. Note that because of the co-elution of 2NFLT and 3NFLT, and neglect of 3NFLT, the so derived values of c_{2NFLT}/c_{FLT} represent 466 actually upper estimates. Apart from sites which were immediately influenced by PAH sources 467 (road traffic, power plant, biomass burning), only very few studies reported NPAH together 468 with precursor data in both phases of ambient air. $c_{2NPYR}/c_{PYR} = 1.0\%$, similar to our finding at 469 470 remote sites, but a very high $c_{2NFLT}/c_{FLT} = 12.9\%$ were reported from a suburban site in France in summer during day-time (corresponding values for night-time were 2.0 and 9.4%, 471 respectively; Ringuet et al., 2012c). 2NFLT was not separated from 3NFLT (similar to our 472 data set). A suburban site will be influenced by direct 3NFLT emissions, such that c_{2NFLT}/c_{FLT} 473 is an upper estimate. Much lower ratios, $c_{2NFLT}/c_{FLT} = 0.20\%$ and $c_{2NPYR}/c_{PYR} = 0.08\%$ were 474 reported as the median values for 90 sites of various categories, rural and urban, in northern 475 476 China in summer (Lin et al., 2015). These yields are somewhat higher for the subset of the rural sites. The potential yields found at the marine site in our study are close to the yields for 477 OH-initiated photochemistry observed in laboratory experiments under high NO_x conditions 478

i.e., 3% for c_{2NFLT}/c_{FLT} and 0.5% for c_{2NPYR}/c_{PYR} (Atkinson and Arey, 1994).

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4. Conclusions

For the first time pollution contained in individual background air samples was quantified, by means of a fractional dose. The fractional dose indicated how much the collected volume of air had been exposed to an urban boundary layer within a given time horizon. This is found suitable to discriminate among samples and discuss results, clearly beyond qualitative reasoning on back trajectories alone. The concept could be applied to any type of georeferenced origin and might be useful to track the influence of land use of various kind, or ship and aircraft routes.

Our measurements confirmed the occurrence of mutagenic NPAHs, earlier reported from polluted atmospheric environments of America, Europe and Asia, also for the European background atmosphere and the outflow of Europe. These substances are obviously subject to intercontinental transport and might indeed be distributed ubiquitously. The mass size distribution is determined by the particle size upon emission (primary NPAHs) and condensation and redistribution in the aerosol along transport, hence, does not include the short-lived coarse mass fraction. This indicates a high long-range transport potential. However, the observation of 3.8 and 0.92 pg m⁻³ of 2NFLT and 2NPYR, respectively, measured at the southeastern outflow of Europe (the lowest ever reported concentrations; Table 3), may indicate that their abundance in the remote global environment could be less than anticipated. Earlier, this was based on a single measurement of 2NPYR, 3 pg m⁻³, at an extremely remote site in central Asia two decades before (Ciccioli et al., 1996). Moreover, this air, classified as marine background, was not completely clean, but had been exposed to a non-zero fractional urban pollution dose (0.4% of the total, time horizon of 2 days). More measurements at remote sites should verify NPAH levels globally. PAHs have been abated significantly in Europe during the last decades (EEA, 2014), which should also be reflected in long-term trends of their derivatives. However, a temporal trend for the Aegean or the southeastern outflow of Europe in summer cannot be inferred based on the current and the earlier (2002; Tsapakis and Stephanou, 2007) campaign data. NPAHs should be included in monitoring programmes to better assess the exposure of human health hazards of atmospheric pollution, even in remote areas.

Our understanding of NPAH formation in ambient air is still rudimentary. Both, the kinetics of NPAH formation and photolysis remain to be quantitatively studied under conditions of the

- background atmosphere i.e., low NO_x and on various aerosol matrices, including sea salt.
- 513 More studies into NPAH atmospheric fate, both field observations and kinetic data, are
- 514 needed in order to assess and quantify spatial and temporal trends, the long-range transport
- 515 potential and persistence.

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 Technol., 47, 8434-8442, 2013.

Site	Phase	Σ_{11} 3-4rNPAH (pg m ⁻³)	$\Sigma_6 4 \text{rPAH (pg m}^{-3})$		
Marine	particulate	4.1 (3.5/0.6/0.2/0.03/0.03/0.00)	43 (28/8.1/1.2/6.2/4.3/1.7)		
	(n=8)	(B: 0.2/P: 8.7)	(B: 7.9/P: 42.4)		
	gas	18.4 (B: 13.2/P:31.1)	403 (B:321/P:580)		
	(n = 21)				
	T(°C)	25.6 (B: 27.1/P: 22.0)			
Continental	particulate (n = 22)	24.3 (20.5/2.9/0.7/0.04/0.06/0.15)	129 (87/28/12/0.6/0.0/0.0)		
		(D:13.9/N:34.6)	(D:146/N:116)		
	gas	34.2 (D:42.9/N:25.5)	517 (D:649/N:384)		
	(n = 22)				
	T(°C)	23.1 (D:28.8/N:17.5)			

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	1NPYR (pg/m³)	2NFLT (pg/m ³)	2NPYR (pg/m³)	References
background CEu summer 2013	1.1	15 ^a	1.3	this work (continental)
E Mediterranean summer 2012	0.74	8.6 a	2.5	this work (marine)
E Mediterranean clean summer 2012	0.21	3.8 ^a	0.92	this work (marine background b)
E Mediterranean clean summer 2002	-	29	21	Tsapakis and Stephanou, 2007
Ross Sea coast, Antarctica	<0.02 °	<0.03 °		Vincenti et al., 2001
Himalayas, Nepal 1991	-	-	3	Ciccioli et al., 1996
Forest Amazonia 1993	2	15	8	
Rural Northern Germany 1991	-	-	3	
Rural Denmark winter-spring 1982	9±5 °	-	-	Nielsen et al., 1984
Semi-rural Denmark all year 1998-99	40	97	6.3	Feilberg et al., 2001
Remote Alps 2002	2.2	-	-	Schauer et al., 2004
Rural Alps 2002	6.6	-	-	
Rural Alps ^d winter 2002-03	21	96 ^a	81	Albinet et al., 2008a
Rural Alps ^d summer 2003	4.2	28 ^a	5.7	
Remote Alps ^e winter 2002-03	2.4	1.3 ^a	14.8	
Remote Alps ^e summer 2003	0.6	1.8 ^a	0.7	
Rural Southern France 2004	0.6	2.6 ^a	1.6	Albinet et al., 2007

^a co-eluted with 3NFLT, assuming c_{3NFLT} = 0
^b samples No. 9, 10, 19 and 22 in Fig. S3
^c particulate phase concentration only
^d Val de Maurienne sites (Albinet et al., 2008a)

^e Plan de l'Aiguille site (Albinet et al., 2008a)

Site			Continental		
	Data	all	marine	background	all
	subset	$(n=8^a)$	background	with urban	(n = 22)
			$(n=2^b)$	influence	
				$(n=2^c)$	
Primary	FLT	213±161	161	259	342±215
	PYR	146±130	103	188	226±131
Primary and secondary (potential	2NFLN d	0.038±0.12	<0.18	0.15	0.034±0.044
	1NPYR	0.62±1.1	0.21	1.4	1.1±0.6
yield)		(0.4±0.2%)	(0.2%)	(0.7%)	(0.6±0.3%)
Secondary (yield)	2NFLT e	7.7±8.5	1.68	11.0	15±10
		(3.6±2.0%)	(1.0%)	(4.2%)	(6.5±7.5%)
	2NPYR	2.2±2.6	0.92	3.3	1.3±1.7
		(1.5±0.7%)	(0.9%)	(1.8%)	(0.74±1.09%)

^{736 &}lt;sup>a</sup> 8 filter and 21 PUF samples

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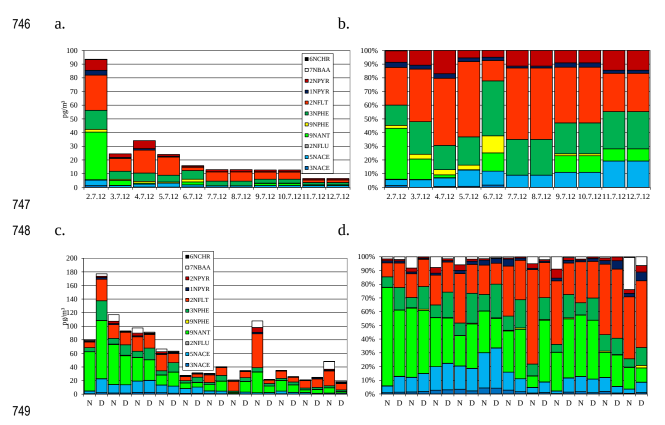
⁷³⁷ b 2 filter and 6 PUF samples i.e., No. 9-10 and 19-22 in Fig. S3 (urban fractional dose $D_u = 0.4\%$)

 $^{^{\}rm c}$ 2 filter and 5 PUF sample i.e., No. 1-2 and 15-18 in Fig. S3 ($D_{\rm u}=3.1\%$)

⁷⁴⁰ d no yield given as c_{FLN} not quantified

⁷⁴¹ e co-eluted with 3NFLT, assuming $c_{3NFLT} = 0$

Fig. 1: Time series of absolute (a, c; pg m⁻³) and relative (b, d) total (gas + particulate) NPAH concentrations at the (a, b) marine (24 h means shown ^a) and (c, d) continental site (day / night means)



^a gas-phase (PUF) sampled day / night, particulate phase (filter) sampled 1-4 subsequent days / nights, 4 during the period 7.-12.7.12

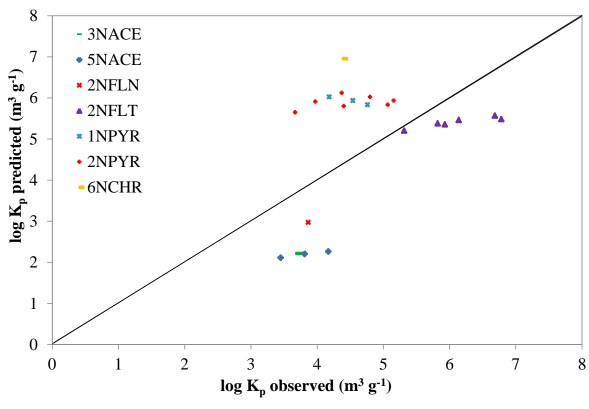
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Fig. 2: Predicted versus experimental log K_p (m^3 air g^{-1} PM) for NPAHs using the multi-phase ppLFER model at the (a) marine and (b) continental site

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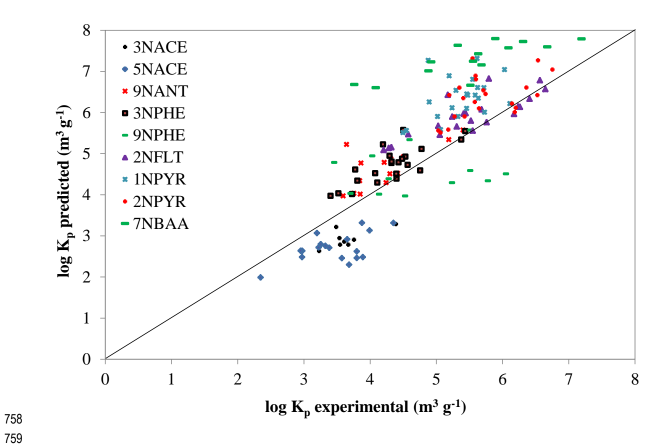


Fig. 3. Time-weighted mean Σ_6 4rPAH and Σ_{11} 3-4rNPAH mass size distributions (pg m⁻³) at the marine and continental sites. The error bars show the standard deviation from the campaign mean.

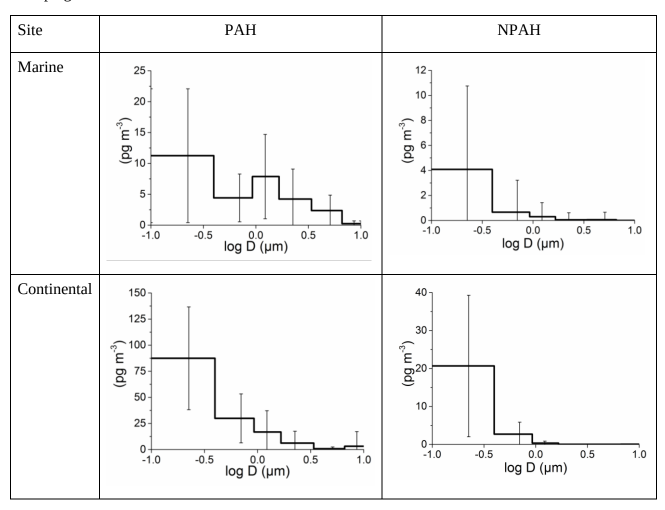


Table 2. Total (g + p) time-weighted concentrations, c_{tot} $(pg \, m^{-3})$, particulate mass fraction, $\theta = c_p / c_{tot}$, and mass median diameter (MMD, μm), of of 2-4-ring NPAHs and 4-ring PAHs at the marine (as 'mean (background mean/ urban influence mean)', $n = 8(2^a/2^b)$) and continental (as 'mean (day mean/ night mean)', n = 22(11/11)) sites, together with temperature and supporting aerosol parameters (PM₁₀ and carbonaceous mass fractions). LOQ = limit of quantification, nd = no data.

	Marine			Continental			
	c _{tot} (pg m ⁻³)	Θ	MMD (µm)	c _{tot} (pg m ⁻³)	Θ	MMD (µm)	
FLT	226 (161/259)	0.07 (0.03/0.07)	0.58 (0.43/0.52)	342 (432/251)	0.11 (0.11/0.12)	0.062 (0.101/0.034)	
PYR	158 (103/188)	0.04 (0.01/0.05)	0.21 (0.022/0.22)	226 (276/176)	0.18 (0.18/0.19)	0.075 (0.105/0.055)	
BBN	4.1 (2.0/5.5)	0.01 (nd/0.05)	0.022 (nd/0.022)	15 (16/13)	0.61 (0.58/0.65)	0.079 (0.127/0.053)	
BAA	2.8 (<loq 3.4)<="" td=""><td>0.28 (nd/0.29)</td><td>0.022 (nd/0.022)</td><td>16 (14/18)</td><td>0.91 (0.90/0.92)</td><td>0.070 (0.090/0.060)</td></loq>	0.28 (nd/0.29)	0.022 (nd/0.022)	16 (14/18)	0.91 (0.90/0.92)	0.070 (0.090/0.060)	
ТРН	12 (8.5/14)	0.02 (nd/0.05)	0.022 (nd/0.022)	23 (26/21)	0.51 (0.41/0.63)	0.070 (0.090/0.057)	
CHR	23 (10/29)	0.22 (0.09/0.20)	0.15 (0.022/0.15)	41 (44/38)	0.75 (0.71/0.80)	0.074 (0.105/0.055)	
$\Sigma_6 4rPAH$	426 (284/499)	0.07 (0.02/0.07)	0.31 (0.19/0.28)	663 (808/517)	0.21 (0.19/0.25)	0.071 (0.10/0.051)	
3NACE	0.21 (0.17/0.39)	0.05 (0.00/0.14)	0.022 (nd/0.022)	1.0(1.0/1.0)	0.05 (0.01/0.11)	0.022 (nd/0.022)	
5NACE	1.8 (1.5/2.0)	0.07 (0.00/0.00)	0.022 (nd/nd)	6.8 (7.6/6.0)	0.03 (0.01/0.05)	0.022 (0.022/0.022)	
2NFLN	0.01 (<loq 0.15)<="" td=""><td>0.02 (nd/0.00)</td><td>1.19 (nd/nd)</td><td>0.035 (0.035/0.034)</td><td>0.00 (0.00/0.00)</td><td>nd</td></loq>	0.02 (nd/0.00)	1.19 (nd/nd)	0.035 (0.035/0.034)	0.00 (0.00/0.00)	nd	
9NPHE	0.73 (0.84/0.55)	0.00 (0.00/0.00)	nd	0.21 (0.28/0.13)	0.36 (0.43/0.20)	0.022 (0.022/nd)	

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3NPHE	4.8 (3.4/5.0)	0.00 (nd/nd)	nd	7.4 (10.0/4.8)	0.24 (0.15/0.44)	0.109 (0.067/0.116)
9NANT	4.2 (1.1/8.2)	0.00 (0.00/0.00)	nd	22 (22/22)	0.23 (0.14/0.33)	0.022 (0.022/0.022)
2NFLT ^c	8.6 (3.8/11)	0.32 (nd/0.45)	0.040 (nd/0.080)	15 (13/18)	0.78 (0.54/0.95)	0.054 (0.035/0.050)
1NPYR	0.75 (0.21/1.4)	0.33 (0.00/0.58)	0.061 (nd/0.14)	1.1 (1.1/1.2)	0.82 (0.76/0.88)	0.030 (0.031/0.029)
2NPYR	2.5 (0.92/3.3)	0.53 (0.19/0.69)	0.058 (0.060/0.055)	1.3 (0.73/2.0)	0.93 (0.83/0.97)	0.070 (0.040/0.061)
7NBAA	<loq< td=""><td>nd</td><td>nd</td><td>2.5 (0.77/4.2)</td><td>0.91 (0.56/0.97)</td><td>0.074 (0.038/0.057)</td></loq<>	nd	nd	2.5 (0.77/4.2)	0.91 (0.56/0.97)	0.074 (0.038/0.057)
6NCHR	0.02 (<loq 0.07)<="" td=""><td>1.00 (nd/1.00)</td><td>2.12 (nd/2.12)</td><td>0.01 (<loq 0.02)<="" td=""><td>1.00 (nd/1.00)</td><td>0.022 (nd/0.022)</td></loq></td></loq>	1.00 (nd/1.00)	2.12 (nd/2.12)	0.01 (<loq 0.02)<="" td=""><td>1.00 (nd/1.00)</td><td>0.022 (nd/0.022)</td></loq>	1.00 (nd/1.00)	0.022 (nd/0.022)
Σ ₁₁ 3-	23.7 (11.8/32.0)	0.22 (0.01/0.22)	0.34(0.33/0.34)	58 (56/59)	0.16 (0.13/0.17)	0.039 (0.036/0.040)
4rNPAH						
$PM_{10} (\mu g/m^3)$	34.9 (21.0/55.5)		0.58 (1.13/0.62)	22.1 (19.5/24.5)		nd
EC (µg/m³)	0.11 (0.09/0.17)		0.03(0.05/0.03)	0.31 (0.28/0.33)		0.21(0.19/0.22)
OC (µg/m³)	1.9 (1.5/3.0)		0.17(0.16/0.15)	3.6 (3.3/3.9)		0.16(0.13/0.18)
T (°C)	25.6 (27.0/22.2)			23.1 (28.8/17.5)		
	l .					

^a 2 filter and 4 PUF samples i.e., No. 9, 10, 19 and 22 in Fig. S3 ^b 1 filter and 1 PUF sample i.e., No. 1 and 2 in Fig. S3 ^c co-eluted with 3NFLT, assuming c_{3NFLT} = 0