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# Composition and mass size distribution of nitrated and oxygenated aromatic compounds in ambient particulate matter from southern and central Europe – implications for the origin

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**Abstract.** Nitro-monoaromatic hydrocarbons (NMAHs), such as nitrocatechols, nitrophenols and nitrosalicylic acids, are important constituents of atmospheric particulate matter (PM) water-soluble organic carbon (WSOC) and humic-like substances (HULIS). Nitrated and oxygenated derivatives of polycyclic aromatic hydrocarbons (NPAHs and OPAHs) are toxic and ubiquitous in the ambient air; due to their light absorption properties, together with NMAHs, they are part of aerosol brown carbon (BrC). We investigated the winter concentrations of these substance classes in size-resolved PM from two urban sites in central and southern Europe, i.e. Mainz (MZ), Germany, and Thessaloniki (TK), Greece. The total concentration of 11 NMAHs ( $\sum_{11}$ NMAH concentrations) measured in  $PM_{10}$  and total  $\overline{PM}$  were 0.51-8.38 and 12.1–72.1 ng m<sup>-3</sup> at the MZ and TK sites, respectively, whereas  $\sum_{7}$  OPAHs were 47–1636 and 858–4306 pg m<sup>-3</sup>, and  $\sum_{8}$  NPAHs were  $\leq 90$  and 76–578 pg m<sup>-3</sup>, respectively. NMAHs contributed 0.4% and 1.8% to the HULIS mass at MZ and TK, respectively. The mass size distributions of the individual substances generally peaked in the smallest or second smallest size fraction i.e. < 0.49 or  $0.49-0.95 \,\mu m$ . The mass median diameter (MMD) of NMAHs was 0.10 and 0.27 µm at MZ and TK, respectively, while the MMDs of NPAHs and OPAHs were both 0.06 µm at MZ and 0.12 and 0.10 µm at TK. Correlation analysis between NMAHs, NPAHs, and OPAHs from one side and WSOC, HULIS, sulfate, and potassium from the other suggested that fresh biomass burning (BB) and fossil fuel combustion emissions dominated at the TK site, while aged air masses were predominant at the MZ site.

# 1 Introduction

Atmospheric humic-like substances (HULIS) represent a complex mixture of aliphatic and aromatic compounds with multiple functional groups, such as hydroxyl, carbonyl, carboxyl, nitro, nitrooxy, and sulfate groups (Havers et al., 1998; Graber and Rudich, 2006; Hallquist et al., 2009; Claevs et al., 2012). They are a major constituent of aerosol water-soluble organic carbon (WSOC), contributing between 9 % and 72 % to WSOC mass (Decesari et al., 2000; Graber and Rudich, 2006; Lin et al., 2010; Zheng et al., 2013). The distribution of HULIS molecular weights (MWs) is unimodal and ranges between 100 and 500 Da, with most of the compounds grouping around 200 Da (Graber and Rudich, 2006; Claeys et al., 2012; Song et al., 2018), unlike soil humic and fulvic acids with MW distributions extending well beyond 1000 Da. Due to the presence of light-absorbing polyconjugated and aromatic compounds (Duarte et al., 2005; Graber and Rudich,

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2006; Claeys et al., 2012; Zheng et al., 2013), HULIS are an important constituent of aerosol water-soluble brown carbon (BrC; Laskin et al., 2015, and references therein). The intense light absorption of HULIS in the ultraviolet, violet, and blue visible regions, between 200 and 500 nm, can affect aerosol optical properties and atmospheric photochemical processes (Andreae and Gelencsér, 2006). Owing to the presence of highly polar polyfunctional material, HULIS have surface-active properties and can make aerosols act as cloud condensation nuclei (CCN). In the aerosol aqueous phase, HULIS can increase the solubility of hydrophobic organic compounds and change the reactivity and solubility of metal aerosols, owing to metal-complexation properties (Graber and Rudich, 2006). Finally, due to the presence of redox-active moieties, HULIS can catalyse electron transfer reactions and formation of reactive oxygen species (ROS), which could pose oxidative stress in humans upon inhalation (Verma et al., 2015).

Biomass burning (BB) is considered as one of the main sources of HULIS in the atmosphere (Lin et al., 2010; Claeys et al., 2012; Pavlovic and Hopke, 2012; Zheng et al., 2013) and an important source of aerosol nitroaromatic compounds (Claeys et al., 2012; Song et al., 2018). Recent studies found that nitro-monoaromatic hydrocarbons (NMAHs), such as 4-nitrocatechol (4-NC; MW: 155 Da) and isomeric methylnitrocatechols (MNCs; MW: 169 Da), are abundant constituents of particulate matter (PM) HULIS, originating from BB (Claeys et al., 2012; Song et al., 2018).

NMAHs are emitted into the atmosphere by primary and secondary processes. 4-NC, MNCs, nitroguaiacols (NGs) and nitrosalicylic acids (NSAs) are predominantly formed by secondary oxidation of lignin thermal decomposition products (e.g. m-cresol, phenols, methoxyphenols, catechols, salicylic acid) in the gas and aqueous phases (Iinuma et al., 2010; Kelly et al., 2010; Kroflič et al., 2015; Frka et al., 2016; Teich et al., 2017; Finewax et al., 2018; Xie et al., 2017; Wang et al., 2019). Therefore, the yellow-coloured watersoluble 4-NC and MNCs have been proposed as suitable tracers for highly oxidized secondary BB aerosols (Iinuma et al., 2010; Kitanovski et al., 2012b; Kahnt et al., 2013; Caumo et al., 2016; Chow et al., 2016). In the past decade, the ambient PM nitrocatechols (NCs) have been measured in several studies worldwide, i.e. in Europe (Iinuma et al., 2010; Zhang et al., 2010; Kitanovski et al., 2012b; Kahnt et al., 2013; Mohr et al., 2013; Teich et al., 2014; Frka et al., 2016), South America (Claeys et al., 2012; Caumo et al., 2016), North America (al-Naiema and Stone, 2017), Asia (Chow et al., 2016; Li et al., 2016; Wang et al., 2019), and Australia (Iinuma et al., 2016). They represent a significant fraction of the PM organic carbon (OC), e.g. 0.8 % in winter PM<sub>10</sub> collected at an urban background location in Slovenia (range 0.4 %-1.3 %; Kitanovski et al., 2012b), 0.75 % in winter PM<sub>10</sub> collected at rural site in Belgium (Kahnt et al., 2013), and  $\approx 0.3 \%$  in PM<sub>10</sub> collected in Brazil during the BB season (Caumo et al., 2016). Nitrosalicylic acids (2hydroxy-nitrobenzoic acids) have been reported in PM samples collected at rural (van Pinxteren and Herrmann, 2007; van Pinxteren et al., 2012; Teich et al., 2017; Wang et al., 2018), urban (Kitanovski et al., 2012a, b; Teich et al., 2017; Wang et al., 2018) and remote (Wang et al., 2018) sites. Similar to NCs, they are mainly associated with secondary BB aerosols (Kitanovski et al., 2012b; Teich at el., 2017; Wang et al., 2018). Nitrophenols (NPs), compounds structurally related to NCs, are emitted from primary sources (e.g. traffic, coal, and wood combustion, industry, and the agricultural use of pesticides), which predominate their secondary formation in urban areas (Harrison et al., 2005; Cecinato et al., 2005; Hoffmann et al., 2007; Iinuma et al., 2007; Zhang et al., 2010; Ganranoo et al., 2010; Özel et al., 2011; Mohr et al., 2013; Kitanovski et al., 2012a, b; Inomata et al., 2015; Teich et al., 2017; Wang et al., 2018; Lu et al., 2019a, b).

Polycyclic aromatic hydrocarbons (PAHs) and their nitrated and oxygenated derivatives (NPAHs and OPAHs), as well as hydroxy derivatives (OH-PAHs), are ubiquitous in the atmosphere (Walgraeve et al., 2010; Lammel, 2015; Bandowe and Meusel, 2017; Shahpoury et al., 2018). They are primarily emitted from the incomplete combustion of fossil fuels (Zielinska et al., 2004; Karavalakis et al., 2010; Pham et al., 2013; Inomata et al., 2015) and wood, coal, and biomass burning (Ding et al., 2012; Shen et al., 2012, 2013a, b; Huang et al., 2014; Vicente et al., 2016). The PAH derivatives are secondarily formed by the reaction of parent PAHs with atmospheric oxidants such as OH, NO<sub>x</sub> and O<sub>3</sub>. Some NPAHs have distinct sources; for instance, 3nitrofluoranthene (3-NFLT) and 1-nitropyrene (1-NPYR) are specifically associated with combustion sources, whereas 2nitrofluoranthene (2-NFLT) and 2-nitropyrene (2-NPYR) are produced through the oxidation of their parent species in the atmosphere (Bandowe and Meusel, 2017). Similarly, OPAHs benzanthrone (OBAT), benz(a)fluorenone (BaOFLN), and benz(b)fluorenone (BbOFLN) have been associated with primary sources, whereas 9,10-anthraquinone (9,10-O<sub>2</sub>ANT), 1,2-benzanthraquinone (1,2-O<sub>2</sub>BAA), and 9-fluorenone (9-OFLN) have been attributed to both source types (Kojima et al., 2010; Souza et al., 2014; Lin et al., 2015; Zhuo et al., 2017). The primary sources dominate in wintertime with residential heating surpassing traffic emission (Lin et al., 2015). It is anticipated that functionalized two- and three-ring PAHs (e.g. two- and three-ring OPAHs) would exhibit the highest hydrophilicity among their analogues and could also be part of PM HULIS (Vione et al., 2014; Fan et al., 2016; Haynes et al., 2019). The water-soluble OPAHs, in particular quinones, were suggested to contribute to light absorption properties of brown carbon (Laskin et al., 2015; Haynes et al., 2019). Moreover, the ROS activity of HULIS from PM2.5 was associated with OPAHs, i.e. quinones and hydroxy-quinones (Verma et al., 2015). It has been shown in controlled experiments that the chemical ageing of PM from various origins would increase its ROS activity, and this effect is enhanced in the presence of O<sub>3</sub> (Li et al., 2009; McWhinney et al., 2011; Stevanovic et al., 2013; Verma et al., 2015; Antiñolo et al., 2015). This process has been attributed to the oxidation of PAHs and formation of water-soluble derivatives.

NMAHs, PAHs, NPAHs and OPAHs significantly contribute to the aerosol BrC due to their light absorption capacity in the ultraviolet (UV) and visible range (Mohr et al., 2013; Samburova et al., 2016; Teich et al., 2017; Xie et al., 2017; Huang et al., 2018). Determining the size-resolved mass distribution of the PM molecular tracers is important for assessing the particle emission sources, atmospheric transport, and health effects (Neusüss et al., 2000). In particular, there is limited knowledge about the size-resolved characteristics of NMAHs, NPAHs and OPAHs and their relation to atmospheric HULIS (Claeys et al., 2012; Song et al., 2018). Therefore, the aim of the present work is to fill this gap by studying the size-resolved PM from polluted urban air at two locations in central and southern Europe, i.e. Mainz (MZ), Germany, and Thessaloniki (TK), Greece, and to apply these data to determine the possible emission sources. These sites were selected to reflect the dominant emission sources in the study areas – while TK is a biomass burning hotspot in southeastern Europe (Saffari et al., 2013; Velali et al., 2019), MZ in central Europe is dominated by traffic emission and long-range transport (Winkler and Junge, 1972; Wesp et al., 2000; Dusek et al., 2006).

# 2 Experimental

## 2.1 Collection of samples

Size-segregated wintertime (season of 2015-2016) PM samples were collected at MZ and TK. In this period, the emissions influencing the sample sites are very different, and, in terms of temperature changes and synoptically, the sampling period is characterized by southwesterly advection with moderate winds at MZ and weak southerly or northeasterly winds at TK (Saffari et al., 2013; Voliotis et al., 2017). At MZ ( $\approx$  200 000 inhabitants), the sampling was done on the rooftop (12 m a.g.l.) of the Max Planck Institute for Chemistry, in a suburb about 2 km away from the city centre. This area was influenced by air masses which consisted of urban and rural continental boundary layer air. At TK (> 1 000 000 inhabitants), the sampling was performed on a rooftop (25 m a.g.l.) at the Aristotle University campus, in a residential area substantially influenced by wood burning for domestic purposes in winter (Voliotis et al., 2017). For each sample set, the air was collected for the duration of 70 and 48 h at MZ and TK, respectively.

All PM samples were collected using a five-stage high-volume cascade impactor with effective cutoff diameters: 0.49, 0.95, 1.5, 3, and 7.2  $\mu$ m of aerodynamic particle size,  $D_{\rm p}$ , and a backup filter collecting particles < 0.49  $\mu$ m (Table 1). The sampling in MZ was done using a high-volume air sampler Baghirra HV-100P (Baghirra, Prague, Czech Re-

public) equipped with a multi-stage cascade impactor (Tisch Environmental Inc., Cleves, Ohio, USA; series 230, model 235) and a PM<sub>10</sub> head. The PM was sampled on slotted quartz fibre filters (QFFs, TE-230-QZ, Tisch Environmental Inc.;  $14.3 \times 13.7$  cm) and a QFF backup filter (Whatman;  $20.3 \times 25.4$  cm). Four sets of samples were collected at MZ between November and December 2015, each over the period of 70 h (flow rate:  $60 \, \text{m}^3 \, \text{h}^{-1}$ ; Table 1). The impactor used in TK was a Sierra Instruments model 235; the PM samples were collected on QFFs (Tisch Environmental Inc.; TE-230-QZ; slotted  $5.7 \times 5.7$  cm) and on QFF backup filters (Pall Corporation, Port Washington, New York, USA; 2500 QAT-UP), without a PM<sub>10</sub> head, as described in Voliotis et al. (2017). Five sample sets were collected at TK between January and March 2016 (Table 1).

# 2.2 Sample preparation and analytical methods

# 2.2.1 Chemical analysis of NMAHs

Extraction of the filter samples for NMAH analysis was done using a validated procedure (Kitanovski et al., 2012b) with small modifications. Briefly, a 1.5 cm<sup>2</sup> section of the filter was spiked with 2,4,6-trinitrophenol and 4-nitrophenold<sub>4</sub> (internal standards – IS; spiked mass: 100 ng; Sect. S1) and subsequently extracted three times (5 min each) with a 10 mL methanolic solution of ethylenediaminetetraacetic acid (EDTA; 3.4 nmol mL<sup>-1</sup>) in an ultrasonic bath. The combined extracts were concentrated to 0.5 mL using a TurboVap II (bath temperature: 40 °C; nitrogen gas pressure: 15 psi; Biotage, Uppsala, Sweden). The concentrated extract was filtered through a 0.2 µm PTFE syringe filter (4 mm; Whatman, GE Healthcare, Little Chalfont, UK) into a 2 mL vial and was evaporated to near dryness under the gentle stream of nitrogen (99.999 %; Westfalen AG, Münster, Germany). Finally, the extract was dissolved in a methanol/water mixture (3/7,v/v) containing a 5 mM ammonium formate buffer with a pH of 3 and 400 µM of EDTA for liquid chromatography/mass spectrometry (LC/MS) analysis.

The NMAHs were determined using an Agilent 1200 Series high-performance liquid chromatography (HPLC) system (Agilent Technologies, Waldbronn, Germany) coupled to an Agilent 6130B Series single quadrupole mass spectrometer equipped with an electrospray ionization (ESI) source. High-purity nitrogen was used as a nebulizer and drying gas. The separation of the targeted analytes was done on an Atlantis T3 column (150 mm × 2.1 mm ID; 3 µm particle size; Waters, Milford, Massachusetts, USA) connected to an Atlantis T3 VanGuard pre-column (5 mm  $\times$  2.1 mm ID; 3 µm particle size; Waters), using isocratic elution with a mobile phase consisting of a methanol/tetrahydrofuran/water (30/15/55, v/v/v) mixture containing 5 mM ammonium formate buffer with a pH of 3 (Sect. S1 in the Supplement). The mobile-phase flow rate, column temperature and injection volume were 0.2 mL min<sup>-1</sup>, 30 °C and 10 µL, respec-

	Cutoff diameters (µm)	Sampling date	Sample volume (m <sup>3</sup> )
Mainz	10–7.2		
49.99° N	7.2–3	17-20 November 2015	3402
8.23° E	3-1.5	26-29 November 2015	4124
	1.5-0.95	1-4 December 2015	4088
	0.95-0.49	4-7 December 2015	4197
	< 0.49		
Thessaloniki		27–29 January 2016	3228
40.63° N	10-3*	8-10 February 2016	3228
22.96° E	3-0.95*	16-18 February 2016	3228
	0.95-0.49	22-24 February 2016	3172
	< 0.49	17-19 March 2016	3175

<sup>\*</sup> Pooled from two impactor stages.

tively (Kitanovski et al., 2012b). The detection and quantification of NMAHs was done in single-ion monitoring and negative ESI mode (Table 2). The optimized ESI-MS parameters were as follows:  $-1000\,\mathrm{V}$  for the ESI capillary voltage, 30 psi for the nebulizer pressure, and  $12\,\mathrm{L}$  min $^{-1}$  and  $340\,^{\circ}\mathrm{C}$  for the drying gas flow and temperature, respectively. Due to the lack of a reference standard for 3-methyl-4-nitrocatechol (3-M-4-NC), its concentrations were calculated based on the calibration curve of 4-M-5-NC. This is justified based on the structural similarity of the two substances and therefore similar ionization efficiency under ESI-MS conditions. LC/MSD ChemStation (Agilent Technologies) was used for data acquisition and analysis. The mean recovery of target NMAHs at  $100\,\mathrm{pg}\,\mu\mathrm{L}^{-1}$  was  $91\pm3\,\%$ .

# 2.2.2 Chemical analysis of NPAHs and OPAHs

NPAHs and OPAHs were extracted from PM samples following a QuEChERS (quick, easy, cheap, effective, rugged, and safe) method with slight modifications (Albinet et al., 2014; Shahpoury et al., 2018). Briefly, two strips of each filter paper were placed inside a glass centrifuge tube (Duran, Schott, Mainz, Germany) and spiked with a mixture of internal standards containing 60 ng of 1-nitronaphthalene-d<sub>7</sub>, 2-nitrofluorene-d<sub>9</sub>, 9-nitroanthracene-d<sub>9</sub>, 3-nitrofluoranthene-d<sub>9</sub>, 1-nitropyrened<sub>9</sub>, 6-nitrochrysene-d<sub>11</sub>, 9,10-anthraquinone-d<sub>8</sub>, and 9fluorenone-d<sub>8</sub> each. A total of 7 mL of dichloromethane (DCM) was then added to each tube; the tubes were capped, and the samples were extracted by vortexing for 1.5 min. The extracts were passed through a glass funnel plugged with deactivated glass wool and concentrated to 0.5 mL using a TurboVap II. The concentrated extracts were loaded on preconditioned SiO<sub>2</sub> solid-phase extraction cartridges (500 mg; Macherey-Nagel, Weilmünster, Germany), and the target analytes were eluted with 9 mL of 65 : 35 *n*-hexane-DCM.

The purified extracts containing the analytes were concentrated to 0.5 mL, and the solvent was exchanged by adding 5 mL of ethyl acetate, concentrating the solution to 0.5 mL, and the process was repeated three times. The sample volumes were adjusted to 0.3 mL and transferred to 2 mL vials containing 0.4 mL glass inserts. All solvents used for NPAH and OPAH analysis were high-purity (Suprasolv, GC-MS grade; Merck, Darmstadt, Germany). All glassware used for analysis was pre-washed with lab-grade detergent, tap water and deionized water and baked at 310 °C for 12 h.

The samples were analysed using a Trace 1310 gas chromatograph (GC; Thermo Scientific, Waltham, Massachusetts, USA) interfaced to a TSQ8000 Evo triple quadrupole mass selective detector (MS/MS; Thermo Scientific). The analysis was performed in negative chemical ionization with methane used as an ionization gas  $(1.5 \,\mathrm{mL\,min^{-1}})$  flow rate; > 99.99 %; Messer, Bad Soden, Germany). The analytes were separated on a 30 m DB-5ms capillary column (0.25 mm ID; 0.25 µm film thickness; J&W, Santa Clara, California, USA) with helium (99.99 %; Westfalen AG, Münster, Germany) as a carrier gas at a 1 mL min<sup>-1</sup> flow rate. The GC inlet temperature was set to 250 °C and operated in the pulsed splitless mode (30 psi pulsed pressure for 1.5 min; splitless time of 1.8 min). The GC oven temperature was held at 60 °C for 2 min at the start of the analysis, and it was then increased to 180°C at 15 °C min<sup>-1</sup> and to 280 °C at 5 °C min<sup>-1</sup>, followed by a final hold time of 15 min. The MS transfer line and ion source temperature were set to 290 and 230 °C, respectively. Emission current and electron energy were set to  $100 \,\mu\text{A}$  and  $-70 \,\text{eV}$ , respectively. The target analytes were detected in the selected-ion monitoring mode, identified using their retention times and quantification ions (Table 2). The quantification was performed using the internal calibration method and 11-point calibration curves ranging from 0.25 to  $1000 \text{ pg } \mu \text{L}^{-1}$ . Trace Finder (Thermo Scientific, Waltham, Massachusetts, USA) was used for chromato-

Table 2. Analytes targeted in this study.

Analyte	Abbreviation	Q1
3-Nitrosalicylic acid	3-NSA	182
5-Nitrosalicylic acid	5-NSA	182
4-Nitrocatechol	4-NC	154
4-Nitroguaiacol	4-NG	168
4-Methyl-5-nitrocatechol	4-M-5-NC	168
4-Nitrophenol	4-NP	138
2,4-Dinitrophenol	2,4-DNP	183
3-Methyl-4-nitrophenol	3-M-4-NP	152
3-Methyl-5-nitrocatechol	3-M-5-NC	168
3-Methyl-4-nitrocatechol	3-M-4-NC	168
2-Methyl-4-nitrophenol	2-M-4-NP	152
2-Methyl-3,5-dinitrophenol	DNOC	197
(Dinitro-ortho-cresol)		
1-Nitronaphthalene	1-NNAP	173.1
2-Nitronaphthalene	2-NNAP	173.1
5-Nitroacenaphthene	5-NACE	199.1
2-Nitrofluorene	2-NFLN	211.1
9-Nitroanthracene	9-NANT	223.1
9-Nitrophenanthrene	9-NPHE	223.1
3-Nitrophenanthrene	3-NPHE	223.1
2-Nitrofluoranthene	2-NFLT	247.1
3-Nitrofluoranthene	3-NFLT	247.1
1-Nitropyrene	1-NPYR	247.1
2-Nitropyrene	2-NPYR	247.1
7-Nitrobenz(a)anthracene	7-NBAA	273.1
6-Nitrochrysene	6-NCHR	273.1
1,3-Dinitropyrene	1,3-N <sub>2</sub> PYR	292.1
1,6-Dinitropyrene	1,6-N <sub>2</sub> PYR	292.1
1,8-Dinitropyrene	1,8-N <sub>2</sub> PYR	292.1
6-Nitrobenz(a)pyrene	6-NBAP	297.1
1,4-Naphthoquinone	1,4-O <sub>2</sub> NAP	158.1
9-Fluorenone	9-OFLN	180.1
9,10-Anthraquinone	9,10-O <sub>2</sub> ANT	208.1
2-Nitro-9-fluorenone	2-N-9-OFLN	225.1
Benz(a)fluorenone	BaOFLN	230.1
Benz(b)fluorenone	BbOFLN	230.1
Benzanthrone	OBAT	230.1
1,2-Benzanthraquinone	1,2-O <sub>2</sub> BAA	258.1

Q1: m/z (mass-to-charge ratio) of ions used for quantification in ESI(–)MS for NMAHs and NCI-MS (negative chemical ionization mass spectrometry) for NPAHs and OPAHs.

graphic data acquisition and analysis. The mean recoveries of target NPAHs and OPAHs at 200 pg  $\mu L^{-1}$  were 73  $\pm$  15 % and 72  $\pm$  18 %, respectively.

# 2.2.3 Quality control and data analysis

Field blanks (n = 3) were prepared during sample collection by mounting the pre-baked filters on the sampler without switching it on. These filters were subsequently retrieved and processed along with the rest of the samples. Limits of quantification (LOQs) for analytes were calculated as the mean concentration of each analyte in blanks +3 stan-

dard deviations. When analyte concentrations in the samples exceeded the LOQ, mean blank concentrations were subtracted from those in the corresponding samples. Microsoft Office Excel 2013 (Microsoft Corp., Redmond, Washington, USA) and OriginPro 9.0 (OriginLab Corp., Northampton, Massachusetts, USA) were used for statistical analysis and data visualization. Mass size distributions (MSDs) of NMAHs, NPAHs and OPAHs were additionally characterized by the mass median diameter (MMD), defined as  $\log \text{MMD} = \sum (c_i \log D_i) / \sum c_i$ , with  $c_i$  and  $D_i$  being the concentration (ng m<sup>-3</sup>) and geometric mean diameter, respectively, of six impactor stages. For consistency across the samples, 0.001 µm was adopted as the lower cutoff of the lowermost stage (backup filter), and 10 µm was the upper cutoff of the uppermost stage, even in the absence of a PM<sub>10</sub> inlet in the case of TK samples. Although this may introduce a small underestimation of MMDs for TK samples, during the sampling at TK, wind velocities did not favour resuspension of large particles and sea spray; hence, we expect that the contribution of PM  $> 10 \,\mu m$  would be negligible. The concentrations of ions, organic acids, HULIS, and HULIS-C in the samples used in this study can be found in a companion paper (Voliotis et al., 2017).

#### 3 Results and discussion

# 3.1 Sources of NMAHs, NPAHs and OPAHs at Thessaloniki and Mainz

# 3.1.1 Concentrations and sources of NMAHs

From the 11 targeted NMAHs, 8 were consistently detected in size-segregated PM from MZ and TK. 4-NG and DNOC were not detected in MZ samples, while they were sporadically detected in the coarse PM (> 3 µm) from TK. 2,4-DNP was detected more frequently in TK (three sample sets) than in MZ samples (one sample set). The concentrations of NMAHs associated to PM<sub>10</sub> (MZ) and total PM (TK) are given in Table S3 in the Supplement. The  $\sum_{11}$ NMAH concentrations in PM from MZ and TK were 0.51-8.38 and 12.1–72.1 ng m<sup>-3</sup>, respectively. In all sample sets, 4-NC was the most abundant NMAH with concentrations ranging within  $0.05-3.90 \,\mathrm{ng} \,\mathrm{m}^{-3}$  (mean:  $2.46 \,\mathrm{ng} \,\mathrm{m}^{-3}$ ; Table S3) in MZ samples and concentrations that were 10 times higher in TK samples  $(5.89-36.33 \text{ ng m}^{-3})$ ; mean:  $22.11 \text{ ng m}^{-3}$ ; Table S3). 4-NP was the second-most abundant NMAH in MZ with concentrations between 0.24 and  $1.27 \,\mathrm{ng}\,\mathrm{m}^{-3}$ (mean:  $0.83 \,\mathrm{ng}\,\mathrm{m}^{-3}$ ; Table S3), while 4-M-5-NC was the second-most abundant NMAH in TK samples (2.54- $16.05 \,\mathrm{ng}\,\mathrm{m}^{-3}$ ; mean:  $9.79 \,\mathrm{ng}\,\mathrm{m}^{-3}$ ; Table S3). In general, the concentration trends of NMAHs were 4-NC > MNCs > 4-NP > NPs > NSAs > DNP (dinitrophenols) for MZ samples and 4-NC > MNCs > 4-NP > NSAs > NPs > DNP for TK samples. These trends are in good agreement with other studies, where 4-NC, MNCs, and 4-NP were the most abundant NMAHs (Kitanovski et al., 2012b; Chow et al., 2016). However, we previously found different concentration trends in snow-scavenged atmospheric particles collected in MZ, where 4-NC and MNCs were the second-most abundant NMAH species following NPs (Shahpoury et al., 2018). NMAH winter concentrations at TK were higher than those found in winter PM2.5 and PM10 from Hong Kong (Chow et al., 2016) and rural Belgium (Kahnt et al., 2013), respectively, but they were lower than NMAH concentrations in winter PM<sub>10</sub> samples from Ljubljana, Slovenia (Kitanovski et al., 2012b), and Shanghai, China (Li et al., 2016). The concentrations of individual NMAHs in winter PM<sub>10</sub> from MZ were among the lowest values reported so far (Iinuma et al., 2010; Kitanovski et al., 2012b; Kahnt et al., 2013; Mohr et al., 2013; Chow et al., 2016; Li et al., 2016; Teich et al., 2017; Wang et al., 2019).

In Table S3, one can easily notice the consistently higher  $(\approx 10 \text{ times})$  total PM concentrations of 4-NC, MNCs, and NSAs in TK samples compared to those found in PM<sub>10</sub> samples from MZ. Smaller concentration discrepancies among the sites were observed for 4-NP and methyl-nitrophenols (MNPs; up to 3 times higher concentrations in TK samples). Since 4-NC, MNCs, and NSAs are considered as suitable tracers for secondary BB aerosols (Iinuma et al., 2010; Kitanovski et al., 2012b; Kahnt et al., 2013; Caumo et al., 2016; Chow et al., 2016; Teich et al., 2017), this suggests that the air masses over TK during sample collection were most likely influenced by BB emissions. To test this hypothesis, a correlation analysis was done for NMAHs. Initially, we did the correlation analysis on PM<sub>10</sub> (MZ) and total PM (TK) samples (Tables S4 and S7). Although the correlation analysis was performed using a limited number of sample sets per location (five for TK and four for MZ), it showed several interesting features. Based on these results, we propose the most probable sources for NMAHs at both sampling sites. Overall, except for NPs in TK samples, high correlations were observed within the NMAH sub-groups (NSAs, NCs, and NPs;  $R_{\text{adj}}^2 > 0.8$ ; Tables S4 and S7). In TK samples, 5-NSA was highly correlated ( $R_{\text{adj}}^2$  0.81–0.83) with 4-NP and potassium cation (K<sup>+</sup>), but it showed insignificant correlations with 4-NC, MNCs, and nitrate (Table S4). 3-NSA showed significant (p < 0.05) correlation with K<sup>+</sup>, but it had a moderate correlation with 4-NP, whereas 4-NP was highly correlated with  $K^+$  and nitrate ( $R_{adj}^2$  0.94 and 0.81, respectively). Secondly, 4-NC and 4-M-5-NC showed low correlations with K<sup>+</sup> and nitrate, but it was highly correlated with 3-M-4-NP ( $R_{\rm adj}^2$  0.74 and 0.78, respectively). Finally, the high correlations between  $K^+$  and WSOC or HULIS ( $R_{\mathrm{adj}}^2 \sim$ 0.9) suggest a strong influence of primary BB emissions on WSOC concentrations at TK. Based on the correlation analysis, our results from TK indicate distinct emission sources for NSAs and 4-NP on the one hand and for 4-NC and MNCs on the other hand. NSAs and 4-NP most likely had the same emission source, i.e. BB (correlate with  $K^+$ ;  $0.81 < R_{adi}^2 <$ 0.94). Additionally, they also showed moderate-to-high correlations with WSOC and HULIS (0.59  $< R_{adi}^2 < 0.78$ ; Table S4). Within specific PM size ranges, PM<sub>0.97</sub> (Table S5) and PM<sub>3-0.97</sub> (Table S6), we found significant correlations (p < 0.05) between NSAs and K<sup>+</sup>, WSOC, HULIS, and 4-NP in the sub-micrometre particles ( $PM_{0.97}$ ). This could indicate either fresh emissions of NSAs during BB (for instance, 3-NSA could also be primarily emitted by BB; Wang et al., 2017) or secondary formation (nitration of salicylic acid, primarily emitted by BB; Iinuma et al., 2007) and subsequent gas-to-particle conversion. 4-NP correlated significantly with K<sup>+</sup>, WSOC, and HULIS in PM<sub>0.97</sub> and PM<sub>3-0.97</sub> but also with NSAs and sulfates in these size ranges, respectively. These results imply that BB is a predominant source of 4-NP in sub-micrometre particles, while additional anthropogenic sources (e.g. coal burning and industry) might also contribute to its concentrations in PM<sub>3-0.97</sub> (high correlation with sulfate, Table S6; Lu et al., 2019a). In contrast, low correlations of 4-NC and MNCs with K<sup>+</sup>, WSOC and HULIS (Tables S5 and S6) suggest that BB might not be a significant emission source for NCs and that their possible source could be fossil fuel combustion (e.g. gas-phase nitration of NC precursors emitted by coal combustion; Kourtchev et al., 2014; Xie et al., 2017; Finewax et al., 2018; Wang et al., 2019; Lu et al., 2019a). Significant correlations of 3-M-4-NP with 4-NC, 4-M-5-NC, and 3-M-5-NC were observed ( $R_{\text{adi}}^2 \ge 0.71$ , p < 0.05; Table S5), in contrast to the insignificant correlations with 4-NP. MNP isomers (2-M-4-NP and 3-M-4-NP) in PM<sub>0.97</sub> and PM<sub>3-0.97</sub> showed strong inter-correlations (0.75  $\leq R_{\rm adj}^2 \leq$  0.92; Tables S5 and S6) but low correlations with K<sup>+</sup>, WSOC, and HULIS. This suggests that, similar to 4-NC and MNCs, MNPs' predominating source at TK was fossil fuel combustion (Noguchi et al., 2007; Lu et al., 2019a). Regardless, fresh BB emission remains a major contributor to PM WSOC at TK, as observed by the significant correlations of WSOC and HULIS with K<sup>+</sup>  $(R_{\rm adi}^2 = 0.9;$  Tables S4 and S5). In conclusion, the emission profile and correlation analysis for NMAHs at TK suggest a complex interplay of different emission sources, particularly dominated by fresh BB and fossil fuel combustion emissions.

Correlation analysis for NMAHs in MZ samples presents a different picture (Table S7). Significant correlations (p < 0.05) were observed among different NMAH compound groups (i.e. NSAs, NCs, and NPs), with a majority of  $R_{\rm adj}^2$  being higher than 0.8. In our previous work, high correlations ( $R_{\rm adj}^2 > 0.8$ ) between NSAs and 4-NC or MNCs were also observed, which indicated the presence of BB secondary organic aerosol (SOA) (Kitanovski et al., 2012b). The correlations between K<sup>+</sup> and NSAs, 4-NC, and MNCs in PM<sub>0.95</sub> were moderate to high ( $0.4 < R_{\rm adj}^2 < 0.8$ ; Table S8), inferring that BB could have contributed to the formation of these species in the sub-micrometre particles. In the same PM size range, nitrate showed moderate-to-high correla-

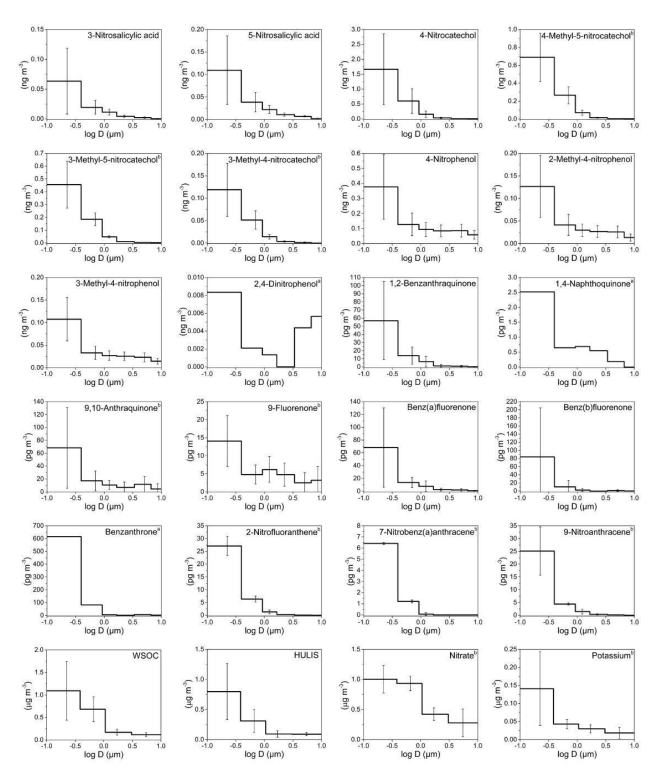
tions  $(0.5 \le R_{\text{adj}}^2 < 0.9)$  with NSAs, 4-NP, MNPs, 4-NC, and MNCs (Table S8), which are much higher than the corresponding ones in PM<sub>0.97</sub> samples from TK (Table S5). In PM<sub>3–0.95</sub>, HULIS showed significant correlations with K<sup>+</sup>  $(R_{\text{adi}}^2 = 0.99)$  and all NMAH species  $(0.66 < R_{\text{adi}}^2 < 0.98;$ Table S9), except for 2,4-DNP, suggesting that NMAHs and PM HULIS had similar sources (i.e. BB). The significant correlations of 4-NC and MNCs with 4-NP and MNPs in PM<sub>0.95</sub> and PM<sub>3-0.95</sub> suggest similar sources for NCs and NPs over MZ. Moreover, high correlations of 4-NC and MNC with K<sup>+</sup> in PM<sub>0.95</sub> indicate that BB was a significant emission source over MZ (Chow et al., 2016; Voliotis et al., 2017; Wang et al., 2018), whereas their high correlations with sulfate in  $PM_{3-0.95}$  (0.66 <  $R_{adj}^2$  < 1.00; Table S9) could infer possible anthropogenic emissions, i.e. coal combustion (Lu et al., 2019a).

# 3.1.2 Concentrations and sources of NPAHs and OPAHs

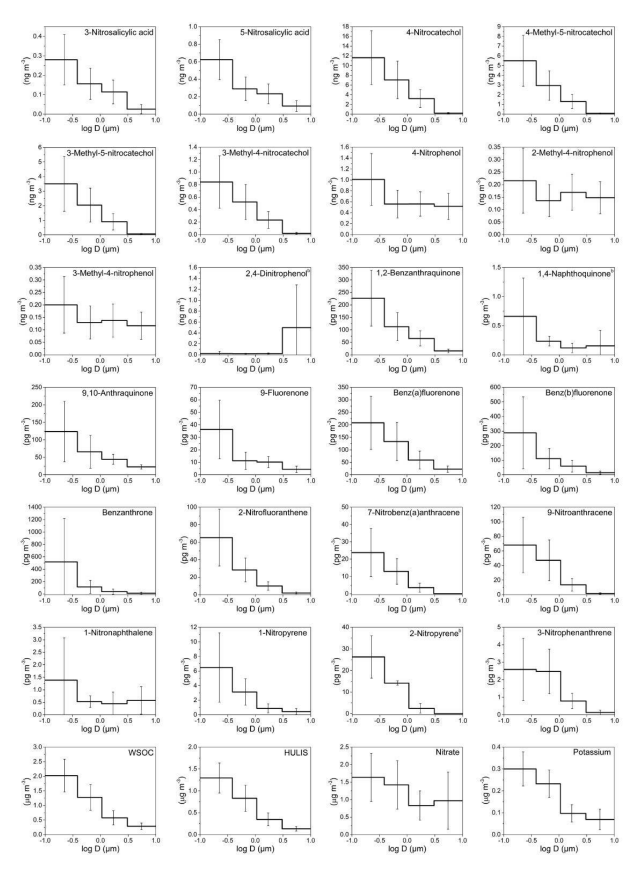
NPAHs and OPAHs were studied in size-resolved PM at both the MZ and TK sites. At both sites, particle-phase OPAHs were detected more frequently than NPAHs: seven out of eight OPAHs targeted for analysis were detected in nearly all MZ and TK samples (Table S3; Figs. S3 and S4 in the Supplement). In contrast, only 8 out of 17 targeted NPAHs were found in the PM samples, of which only 1-nitronaphthalene (1-NNAP), 9-nitroanthracene (9-NANT), 2-NFLT, and 7-nitrobenz(a)anthracene (7-NBAA) were detected in both MZ and TK samples. Interestingly, 3nitrophenanthrene (3-NPHE), 3-NFLT, and 1- and 2-NPYR were only found in TK samples. This was not due to differences in individual LOQs between the two sites (see Table S2). The mean concentrations of NPAHs in PM were dominated by 9-NANT followed by 2-NFLT and 7-NBAA at both sites (Figs. 1 and 2, Table S3), with concentrations reaching to 225, 154, and 71 pg  $m^{-3}$ , respectively. This pattern closely resembles those previously reported for PM from several locations in central Europe (Tomaz et al., 2016, and references therein), including NPAHs found in snowscavenged atmospheric particles from the MZ sample site (Shahpoury et al., 2018). As for OPAHs, the mean analyte concentrations in PM were dominated by OBAT, followed closely by BbOFLN, BaOFLN, 9,10-O<sub>2</sub>ANT, and 1,2-O<sub>2</sub>BAA. The latter two quinones could be of high importance due to their redox activity and their potential to catalyse the formation of reactive oxygen species within the human respiratory system (Ayres et al., 2008; Bates et al., 2019). The two substances were found to dominate two out of four MZ samples with concentrations up to 221 and 137 pg m $^{-3}$ , respectively. These concentrations were higher at the TK site and reached 354 and 514 pg  $m^{-3}$ , respectively.

Overall, all NPAHs and OPAHs showed considerably higher concentrations in TK than in MZ samples. ∑NPAH concentrations in PM<sub>10</sub> from MZ and in total PM from TK were < LOQ-90 and 76–578 pg m<sup>-3</sup>, respectively, whereas ∑OPAHs demonstrated much higher levels ranging from 47 to 1636 and from 858 to  $4306 \,\mathrm{pg}\,\mathrm{m}^{-3}$ , respectively. The sum of three quinones, 1,4-naphthoquinone (1,4-O<sub>2</sub>NAP), 9,10- $O_2$ ANT, and 1,2- $O_2$ BAA, were 30–363 and 428–873 pg m<sup>-3</sup> at these sites, respectively. The levels of particle-phase NPAHs found in MZ fall in the lower end of the range  $(50-500 \,\mathrm{pg}\,\mathrm{m}^{-3})$  observed for various types of sites in Europe (Tomaz et al., 2016, and references therein). The levels at TK represent the upper end of this range, while they are within the concentration range previously found at other sites in Thessaloniki (1204  $\pm$  249 pg m<sup>-3</sup> at a traffic site and  $383 \pm 77 \,\mathrm{pg}\,\mathrm{m}^{-3}$  at an urban background site; Besis et al., 2017). The total OPAH concentrations at both sites fall in the lower end of the range previously observed in Europe (0.5–  $50 \,\mathrm{ng}\,\mathrm{m}^{-3}$ ; Tomaz et al., 2016 and references therein).

NPAHs and OPAHs were predominant in the submicrometre PM fraction (PM<sub>0.95</sub>; 85 %–91 % of PM<sub>10</sub> at MZ and 78 %-85 % of total PM at the TK site; Figs. 1, 2, S3, S4 and S5), with relatively more enrichment in PM<sub>0.49</sub> compared to  $PM_{0.49-0.95}$  across the two sites. The mean concentrations of  $\sum$ NPAHs in PM<sub>0.49</sub> from MZ and TK were 101  $\pm$ 73 and  $417 \pm 134 \text{ pg m}^{-3}$ , whereas in PM<sub>0.49-0.95</sub> they were  $22.8 \pm 15.9$  and  $222 \pm 95$  pg m<sup>-3</sup>, respectively.  $\Sigma$ OPAHs showed similar patterns at the MZ and TK sites – they were  $460 \pm 566$  and  $1426 \pm 1210$  pg m<sup>-3</sup> in PM<sub>0.49</sub>, respectively, and  $81.6 \pm 78.8$  and  $555 \pm 209 \,\mathrm{pg} \,\mathrm{m}^{-3}$  in  $PM_{0.49-0.95}$ . The targeted NPAHs did not show a second mode in any sample, whereas for 9-OFLN and 9,10-O<sub>2</sub>ANT a second mode was found in MZ samples. Such differences between size distributions indicate that 9-OFLN and 9,10-O<sub>2</sub>ANT are subject to different atmospheric processes compared to all other NPAHs and OPAHs that we studied in the present work. This could point at different emission and formation pathways in the atmosphere. Some of the OPAHs with a single O atom, namely OBAT, BaOFLN, and BbOFLN, originate from primary sources (i.e. combustion of fossil fuels and biomass; Albinet et al., 2007; Karavalakis et al., 2010; Shen et al., 2013b; Souza et al., 2014; Huang et al., 2014; Tomaz et al., 2016; Vicente et al., 2016), whereas some quinones, such as 9,10-O<sub>2</sub>ANT and 1,2-O<sub>2</sub>BAA, are associated with both primary and secondary sources (Kojima et al., 2010; Souza et al., 2014; Lin et al., 2015; Zhuo et al., 2017). The presence of 3-NFLT and 1-NPYR at TK indicates the influence of primary sources at that site (Bandowe and Meusel, 2017); notably, these two NPAHs were not found in MZ samples. In order to better understand the potential sources of the target substances, we performed correlation analysis between the measured levels of NPAHs and OPAHs and other PM constituents, namely, WSOC, HULIS, nitrate, sulfate, and K<sup>+</sup>. For this analysis, we considered the compositions of PM<sub>10</sub> (at MZ) and total PM (at TK), as well as the constituents



**Figure 1.** Mass size distributions (MSDs) of PM-bound NMAHs, NPAHs, OPAHs, WSOC, HULIS, and ions in Mainz (Germany). The error bars represent standard deviations. <sup>a</sup> Compound MSD calculated from one (out of four) sample set (detected and quantified in one sample set only); <sup>b</sup> compound MSD calculated from three (out of four) sample sets (detected and quantified in three sample sets only).



**Figure 2.** Mass size distributions (MSDs) of PM-bound NMAHs, NPAHs, OPAHs, WSOC, HULIS, and ions in Thessaloniki (Greece). The error bars represent standard deviations. <sup>b</sup> Compound MSD calculated from three (out of five) sample sets (detected and quantified in three sample sets only).

of PM<sub>0.97</sub> and PM<sub>3-0.97</sub> at both sites. We found a significant correlation (n = 5; p < 0.05) between 9,10-O<sub>2</sub>ANT and 1,2-O<sub>2</sub>BAA at the TK site, regardless of the considered PM size range, which suggests a common emission source (Table S10). The data shown in Table S10 also indicate significant correlations (p < 0.05) between the levels of BaOFLN and 1-NPYR (produced by primary sources) and WSOC, HULIS, and K<sup>+</sup> (BB tracer) in the TK total PM samples. Considering the PM<sub>0.97</sub> size fraction (Table S11), the correlation with K<sup>+</sup> was only significant for 1-NPYR at TK, whereas both BaOFLN and 1-NPYR correlated with WSOC and HULIS in this size fraction. 1-NPYR is the predominant congener among NPAHs found in diesel engine exhaust particles and was proposed as a tracer for diesel emission (Bamford et al., 2003; IARC, 2013), but it may also be emitted with relatively small quantities from biomass-fuelled combustion (Shen et al., 2012; Orakij et al., 2017). These findings suggest the importance of primary emission sources including BB and diesel exhaust in the TK study area. For MZ PM<sub>10</sub> samples, we found significant correlations (n = 4; p < 0.05) of 9-OFLN, BaOFLN, and 9-NANT with WSOC and HULIS (Table S13) but no significant correlations to K<sup>+</sup>. We found similar correlations in PM<sub>0.97</sub>, which suggest the predominance of chemically aged air masses that were advected during the MZ campaign. This is further supported by the absence of NPYR isomers in MZ samples, which are indicative of road traffic and industrial emissions (IARC, 1989; Finlayson-Pitts and Pitts, 2000; Lammel et al., 2017; Voliotis et al., 2017). Finally, K<sup>+</sup>, WSOC, and HULIS correlated significantly at TK (p < 0.05,  $R_{\text{adi}}^2$  0.89–0.90), whereas such correlations were not found at MZ. In summary, while NPAHs and OPAHs from TK samples were influenced by primary emissions related to BB and fossil fuel combustion, those from MZ samples were dominated by aged air masses.

## 3.2 Mass size distributions of target compounds

# 3.2.1 Mass size distributions of NMAHs

MSDs of NMAHs over the two sampling locations are given in Figs. 1 and 2. NSAs (3-NSA and 5-NSA) and NCs (4-NC, 4-M-5-NC, 3-M-5-NC, and 3-M-4-NC) showed unimodal distributions with MSDs generally peaking in the finest PM fraction (PM<sub>0.49</sub>) in both MZ and TK samples. Overall, NMAHs were prominent in smaller size fractions (PM<sub>0.95</sub>) in MZ compared to TK (Figs. 1 and 2). In one of the four samples collected at MZ, NSA MSDs peaked in PM<sub>1.5-0.95</sub>, while the PM<sub>0.95</sub> mass fractions of 3-NSA and 5-NSA were 22 % and 44 %, respectively (Fig. S1a). In this sample only, 5-NSA showed a bimodal distribution (dominant peaks in PM<sub>0.49</sub> and PM<sub>1.5-0.95</sub>). Moreover, 4-NP and MNPs were the most abundant NMAHs (Fig. S1a); the abundance of 4-NP and MNPs could indicate the influence of primary traffic emissions (vehicle exhaust; Seki et al., 2010; Inomata et al., 2015; Lu et al., 2019b) at the beginning of the sampling campaign in MZ. During the next sampling periods at the MZ site (Fig. S1b, c and d), 75 %–86 % of NSAs' PM<sub>10</sub> mass was associated with PM<sub>0.95</sub>, which is in line with the observations at TK (66 %–82 % of total PM mass belongs to PM<sub>0.95</sub>; Fig. S2). At both sites, usually more than 90 % of the compound total mass was associated with PM<sub>3</sub> (range: 83 %– 99 %). We found that 87 %–93 % and 82 %–88 % of NCs at MZ and TK were associated with PM<sub>0.95</sub> (Figs. S1, S2, and S5). The coarse mode ( $> 3 \mu m$ ) accounted for only 1 % (MZ) or 2.5 % (TK). The unimodal distributions of NCs peaking in the fine PM fraction are in line with the only report on MSDs of 4-NC (Li et al., 2016). The MSDs of HULIS in MZ and TK closely followed the MSDs of NCs and NSAs (Figs. 1 and 2), suggesting that they may have undergone similar atmospheric processes. The accumulation of the NCs' and NSAs' mass in the sub-micrometre (< 0.95 µm) PM fractions could indicate fresh combustion emissions (e.g. BB) and/or gas-to-particle conversion processes of their precursors over MZ and TK (Li et al., 2016).

Nitrophenols (i.e. 4-NP, 2-M-4-NP, and 3-M-4-NP) showed bimodal distributions with a dominant peak in the finest fraction (PM<sub>0.49</sub>) and a smaller peak in PM<sub>3-0.95</sub> (Figs. 1, 2, S1, S2 and S5). Bimodal distribution of NPs (i.e. 4-NP, 4-NG, 2,6-dimethyl-4-nitrophenol, and 2,6-dinitrophenol), with peaks in the fine and coarse PM fractions, was recently reported during winter haze episodes over Shanghai, China (Li et al., 2016). For 4-NP at both sites, nearly 80 % of the PM<sub>10</sub> mass (or of the total PM mass at TK) was associated with PM<sub>3</sub>, while  $\approx$  60 % was associated with PM<sub>0.95</sub> (Figs. S1 and S2). Similarly, for methyl-nitrophenols 83 %–88 % of PM<sub>10</sub> mass at MZ and 75 %–83 % of total PM mass at the TK site were associated with PM<sub>3</sub>, while 58 %–65 % of PM<sub>10</sub> at MZ and 48 %–61 % of total PM mass at the TK site were associated with PM<sub>0.95</sub> (Figs. S1 and S2).

The MMD of NMAHs was  $0.10\,\mu m$  (0.24 for NPs, 0.07 for NCs, and 0.11  $\mu m$  for NSAs) at MZ vs. 0.27  $\mu m$  (0.60 for NPs, 0.24 for NCs, and 0.31  $\mu m$  for NSAs) at TK. The larger MMDs at TK could be indicative of aerosol ageing. In aged aerosols, semivolatile organic species are expected to be redistributed with the MMD approaching the surface mean diameter, which for urban and continental aerosol peaks around 0.2  $\mu m$  (Jaenicke, 1988), a shift which could not be resolved by the sampling technique applied here. Note that the low size resolution (six stages) may hide modes, which in particular applies to the so-called accumulation mode, which adds mostly to PM<sub>0.49</sub> but also to the size fraction between 0.49 and 0.95  $\mu m$ .

# 3.2.2 Mass size distributions of NPAHs and OPAHs

NPAH and OPAH MSDs are shown in Figs. 1 and 2. On average, the MMDs of NPAHs were 0.06  $\mu m$  at MZ and 0.12  $\mu m$  at TK, while those for OPAHs were 0.06  $\mu m$  at MZ and 0.10  $\mu m$  at TK. The MMDs for quinones were 0.07 and 0.15 at the two sites, respectively. We found two distinct MSD pat-

terns among the samples: the first pattern observed in three samples across the two sites (one sample set from MZ and two sets from TK; Figs. S3c, S4a, d, and e) was dominated by OBAT followed by BbOFLN. The MMD of OPAHs in these three samples was on average 0.06 µm (ranging within 0.05–0.09 µm). The unique analyte distribution in these samples was accompanied by a noticeably higher enrichment in PM<sub>0.49</sub> as well as relatively high concentrations compared to the rest of samples. The preferential enrichment of OBAT, BaOFLN, and BbOFLN in sub-micrometre PM was previously reported from locations in Europe, Asia, and the USA (Allen et al., 1997; Albinet et al., 2008; Ladji et al., 2009; Ringuet et al., 2012; Shen et al., 2016; Gao et al., 2019). The second pattern, which was seen in the remaining six sample sets, was considerably different: the target substances were more evenly distributed across different PM size ranges, and they were often dominated by relatively high abundance of quinones 9,10-O<sub>2</sub>ANT and 1,2-O<sub>2</sub>BAA – the two quinones were previously reported with preferential enrichment in ultrafine PM (Ringuet et al., 2012; Shen et al., 2016). The MMD of OPAHs in these five sample sets was on average  $0.25 \,\mu m$  (ranging within  $0.08-0.49 \,\mu m$ ).

In terms of the inter-site variability of the target substance MSDs, the size fraction PM<sub>0.49</sub> was more prominent in MZ than in TK, i.e. on average 74 % for NPAHs, 75 % for OPAHs, and 69 % for quinones at MZ, compared to 55 %, 60 %, and 52 %, respectively, at the TK site (Figs. 1-2 and S3–S5). The largest differences found among each substance group were for 9-NANT (28 % higher at MZ), BbOFLN (25 % higher), and 1,2-O<sub>2</sub>BAA (17 % higher). The values for NPAHs from TK were lower than those previously found for wintertime PM at this site (59 % and 71 % for a traffic and urban background site, respectively; Besis et al., 2017). The higher enrichment of predominant NPAHs (9-NANT and 2-NFLT; Figs. S3-S4) in PM<sub>0.49</sub> in the present study is in agreement with the MSDs reported for these compounds from several other locations in Europe and Asia (Ringuet et al., 2012; Lan et al., 2014; Lammel et al., 2017). The preferential enrichment of NPAHs and OPAHs in sub-micrometre PM, especially  $PM_{0.49}$ , raises concerns with respect to the inhalation toxicity of airborne PM; this is due to the fact that PM<sub>0.49</sub> is capable of reaching deeper regions in the lung (Oberdörster et al., 2005). This is exacerbated by the ability of quinones to catalyse redox reactions and the formation of ROS in the respiratory system (Ayres et al., 2008).

# 3.3 NMAHs as part of HULIS

Because of their water solubility, NMAHs are constituents of PM HULIS and WSOC (Claeys et al., 2012; Teich et al., 2017). This substance class contributed  $\approx 0.4\,\%$  and 1.8 % to HULIS mass at the MZ and TK sites, respectively (Table 3). This contribution was fairly even across the size fractions addressed, while showing a maximum for particles size 0.95–3  $\mu$ m, namely  $\approx 0.7\,\%$  and 2.0 % by mass at the MZ

**Table 3.** Mean absolute concentrations and mass mixing ratios (in brackets) of HULIS<sup>a</sup> in WSOC<sup>a</sup> as well as of NMAHs in HULIS in (a) Mainz and (b) Thessaloniki PM.

(a)			
Particle size	WSOC	HULIS $\mu$ g m <sup>-3</sup>	NMAHs ng m $^{-3}$
(µm)	$\mu \mathrm{gC}\mathrm{m}^{-3}$	(% C/C)b	(%)
< 0.49	1.14	0.80 (39)	3.41 (0.43)
0.49 - 0.95	0.68	0.31 (25)	1.24 (0.40)
0.95-3	0.18	0.09 (28)	0.65 (0.73)
3–10	0.12	0.09 (42)	0.27 (0.30)
Total	2.07	1.29 (33)	5.58 (0.43)
(b)			
Particle size	WSOC	HULIS $\mu$ g m <sup>-3</sup>	NMAHs $ng m^{-3}$
(µm)	$\mu \mathrm{gC}\mathrm{m}^{-3}$	(% C/C)	(%)
< 0.49	2.02	1.29 (34)	24.0 (1.9)
0.49 - 0.95	1.28	0.83 (34)	13.9 (1.7)
0.95-3	0.57	0.35 (32)	6.89 (2.0)
> 3	0.33	0.11 (18)	1.87 (1.7)
Total	4.20	2.58 (32)	46.6 (1.8)

<sup>&</sup>lt;sup>a</sup> Voliotis et al. (2017). <sup>b</sup> Fraction (%) of HULIS carbon in organic carbon.

and TK sites, respectively. The large particle size,  $0.95-3\,\mu m$ , points to the significance of aqueous-phase processes and in general the slower formation of NMAHs (Voliotis et al., 2017). The activation of condensation-mode particles (under high humidity) into cloud droplets, as well as the subsequent possible aqueous-phase reactions, leads to the formation of larger particles in aged and cloud-processed aerosols. In central Europe, characteristic times of formation of coarse-mode secondary inorganic aerosols and OC peak around 60–72 h (Lammel et al., 2003).

Our reported NMAH contribution to HULIS mass is in good agreement with the results of previous reports from urban sites in Europe (Kitanovski et al., 2012b; Claeys et al., 2012) and Brazil (Caumo et al., 2016). Specifically, Kitanovski et al. (2012b) found that NMAHs contributed 0.4 %-1.3 % to the OC mass in winter PM<sub>10</sub> from Ljubljana (Slovenia), while in another study 4-NC alone contributed 0.46% and 0.04% to the HULIS mass in urban spring and summer PM<sub>2.5</sub> from Budapest, Hungary, respectively (Claeys et al., 2012). Moreover, NMAHs (4-NP, 4-NC, MNCs, and dimethyl-nitrocatechols; DMNCs) contributed 0.28 % and 0.35 % to the OC mass in winter PM<sub>10</sub> samples from São Paulo, Brazil (Caumo et al., 2016). Lower NMAH contribution to HULIS (or OC) mass were reported for rural sites in Europe. For example, 4-NC contributed 0.03 % to the HULIS mass in summer PM2.5 from K-puszta, Hungary (Claeys et al., 2012), while total NMAHs (NPs, 4-NC, MNCs, and DMNCs) represented 0.75 % of OC mass in winter PM<sub>10</sub> sampled at a rural background site in Belgium (Kahnt et al., 2013).

In Sect. 3.2.1, we emphasized the similar MSDs at both locations between HULIS on one side and NCs and NSAs on the other. These two NMAH subclasses on average contributed to  $\approx 83\,\%$  and  $\approx 94\,\%$  of total NMAHs in PM<sub>0.95</sub> and  $\approx 55\,\%$  and 87 % of total NMAHs in PM<sub>3-0.95</sub> at the MZ and TK sites, respectively (Table S16). At both sites, NCs were the dominant NMAH species. It is also interesting to note that HULIS showed higher correlations with NSAs and NCs in MZ (0.68 <  $R_{\rm adj}^2$  < 0.98; Table S7) than in TK (0.24 <  $R_{\rm adj}^2$  < 0.59; Table S4).

With mass mixing ratios of the order of 1 %, NMAHs are constituents of HULIS with limited significance by mass, but their relevance is more significant due to their optical properties (Mohr et al., 2013; Laskin et al., 2015; Teich et al., 2017; Xie et al., 2017). Teich et al. (2017) found that the mass contributions of total NMAHs (NPs and NSAs) to WSOC on average was five times lower than their contribution to the light absorption of the aqueous PM extract at 370 nm (Teich et al., 2017). This implies that even small fractions of chromophoric HULIS compounds such as NMAHs can have an excessive influence on the aerosol light absorption (Mohr et al., 2013; Teich et al., 2017) and the atmospheric photochemical processes, especially in polluted areas (Laskin et al., 2015; Teich et al., 2017).

#### 4 Final remarks

We studied the composition and MSDs of NMAHs, NPAHs, and OPAHs in PM from urban locations in Germany and Greece, with some of the target substances (i.e. NSAs, MNCs, and MNPs) studied in size-resolved PM for the first time. At both locations, NCs were the most abundant NMAH species, and OPAHs were more abundant and more frequently detected than NPAHs. The total concentrations of the most abundant NMAHs, NCs, NPAHs, and OPAHs were up to 10 times higher in TK than in MZ. Correlation analysis of NMAHs revealed distinct features among the sites, suggesting mixed air masses influenced by fresh BB and fossil fuel combustion emissions at TK and aged advected air influenced by combustion emissions (i.e. BB and coal combustion) at MZ.

The MSDs of NMAHs, OPAHs, and NPAHs were rather similar, but they exhibited temporal and spatial variations due to daily changes in atmospheric conditions and different sources. In general, NCs, NSAs, OPAHs, and NPAHs showed unimodal MSDs peaking in the finest PM fraction, PM<sub>0.49</sub>, which was more prominent in MZ than in TK. NPs exhibited bimodal MSDs with the dominant peak in PM<sub>0.49</sub>. The MMDs of all chemical classes were lower at MZ than at TK; the larger MMDs at TK could be an indication of aerosol ageing. On average, NMAHs contributed up to 1.8 % to the HULIS mass in the study areas. Although NMAHs represent a small fraction of PM HULIS (and WSOC), due to their light absorption properties, their impact on the total aerosol

light absorption could be disproportionally large. This is particularly important for atmospheric photochemical processes in polluted areas.

Data availability. The dataset used in this paper is included in the Supplement, and further information is available from the corresponding authors (z.kitanovski@mpic.de; p.shahpoury@mpic.de).

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/acp-20-2471-2020-supplement.

Author contributions. GL and CS conceived the study. PS and AV conducted the air sampling and field measurements. ZK and PS did the chemical analysis of samples. ZK, PS, and GL did the data analysis. ZK, PS, and GL discussed the results and wrote the paper with input from all co-authors.

Competing interests. The authors declare that they have no conflict of interest.

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