

# Insights into the sources of ultrafine particle numbers at six European urban sites obtained by investigating COVID–19 lockdowns

Alex Rowell<sup>1</sup>, James Brean<sup>1</sup>, David C.S. Beddows<sup>1</sup>, Tuukka Petäjä<sup>2</sup>, Máté Vörösmarty<sup>3</sup>, Imre Salma<sup>4</sup>, Jarkko V. Niemi<sup>5</sup>, Hanna E. Manninen<sup>5</sup>, Dominik van Pinxteren<sup>6</sup>, Thomas Tuch<sup>6</sup>, Kay Weinhold<sup>6</sup>, Zongbo Shi<sup>1</sup>, Roy M. Harrison<sup>1,7</sup>

<sup>1</sup>School of Geography, Earth & Environmental Sciences, University of Birmingham, Birmingham B15 2TT, United Kingdom

<sup>2</sup>Institute for Atmospheric and Earth System Research (INAR) / Physics, Faculty of Science, University of Helsinki, Finland

<sup>3</sup>Hevesy György Ph. D. School of Chemistry, Eötvös Loránd University, Budapest, Hungary

<sup>4</sup>Institute of Chemistry, Eötvös Loránd University, Budapest, Hungary

<sup>5</sup>Helsinki Region Environmental Services Authority (HSY), Helsinki, Finland

<sup>6</sup>Leibniz Institute for Tropospheric Research (TROPOS), Atmospheric Chemistry Department (ACD), Permoserstr. 15, 04318 Leipzig, Germany

<sup>7</sup>Department of Environmental Sciences, Faculty of Meteorology, Environment and Arid Land Agriculture, King Abdulaziz University, Jeddah 21589, Saudi Arabia

*Correspondence to:* Roy M. Harrison ([r.m.harrison@bham.ac.uk](mailto:r.m.harrison@bham.ac.uk))

## ABSTRACT

Lockdown restrictions in response to the COVID–19 pandemic led to the curtailment of many activities and reduced emissions of primary air pollutants. Here, we applied Positive Matrix Factorization to particle size distribution (PSD) data from six monitoring sites (three urban background and three roadside) between four European cities (Helsinki, Leipzig, Budapest, and London) to evaluate how particle number concentrations (PNCs) and their sources changed during the respective 2020 lockdown periods compared to the reference years 2014–2019. A number of common factors were resolved between sites, including nucleation, road traffic semi–volatile fraction (road traffic<sub>svf</sub>), road traffic solid fraction (road traffic<sub>sf</sub>), diffuse urban (woodsmoke + aged traffic), ozone–associated secondary aerosol (O<sub>3</sub>–associated SA), and secondary inorganic aerosol (SIA). Nucleation, road traffic, and diffuse urban factors were the largest contributors to mean PNCs during the reference years and respective lockdown periods. However, SIA factors were the largest contributors to particle mass concentrations, irrespective of environment type. Total mean PNCs were lower at two of the urban background and all roadside sites during lockdown. Nucleation and road traffic<sub>svf</sub> factors response to lockdown restrictions were highly variable, although road traffic<sub>sf</sub> factors were consistently lower at roadside sites. The responses of diffuse urban factors were largely consistent and were mostly lower at urban background sites. Secondary aerosols (O<sub>3</sub>–associated SA and SIA) exhibited extensive reductions to their mean PNCs at all sites. These variegated responses to lockdowns across Europe point to a complex network of sources and aerosol sinks contributing to PSDs.

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## 1. INTRODUCTION

The COVID–19 pandemic and the resultant curtailment of human activities had profound impacts on global atmospheric chemistry, reflected in the concentrations of greenhouse gases (Le Quéré et al., 2020), gas phase pollutants (Shi et al., 2021), and particulate matter (Hammer et al., 2021; Putaud et al., 2023; Torkmahalleh et al., 2021). Ambient atmospheric aerosol particles are of scientific concern due to their detrimental effects on human health (Cohen et al., 2017) and the uncertainties they cause in models of global radiative forcing (Storelvmo et al., 2016). These health and climatic effects depend on particle size, as their ability to enter the lung, reflect or refract solar radiation, and form clouds, are size dependent processes. For their parameterisation in air quality and climate models, it is therefore important to understand the sources of different sized aerosol particles. Particle mass concentrations (PMCs) are typically dominated by larger particles, such as those with mobility diameters  $>100$  nm, whereas, particle number concentrations (PNCs) are typically dominated by smaller particles, such as those  $\leq 100$  nm, commonly referred to as ultrafine particles (UFP). The human health effects of UFP are less clear, however; there is epidemiological evidence for the adverse health effects of UFP exposure when weighted by number (Ohlwein et al., 2019), and particle count in different size ranges is an important metric for understanding the climatic effects of aerosols (Jiang et al., 2021).

Number size distributions are typically comprised of a series of lognormal modes. Each mode represents a different source, or aggregate of sources, (referred to as a factor) as modified by aerosol microphysical processes, such as particle shrinkage. Different sources of particles produce different modal diameters (Vu et al., 2015). In urban environments, primary and secondary particle production ( $<1,000$  nm) arises from factors of natural and anthropogenic origin. In the existing literature, a number of common source–related factors have been identified around the world, including nucleation, traffic (multiple), heating, ozone–associated secondary aerosol, and secondary inorganic aerosol, as well as biomass burning, and various unidentified factors (Hopke et al., 2022). However, the identification of such factors is not straightforward. The shape of the particle size distribution from primary traffic emissions, for example, depends on a host of variables, including fuel type, and driving conditions (Rönkkö and Timonen, 2019). Secondary aerosols arise in the smallest diameters from new particle formation (NPF) processes, where vapours such as sulphuric acid and amines cluster in the atmosphere to form new thermodynamically stable aerosol particles at  $\sim 1.5$  nm, before growing to larger sizes due to the condensation of oxidised organic molecules (OOMs), acids, and other suitably involatile vapours (Kulmala et al., 2014; Lee et al., 2019). Vapour nucleation and the subsequent formation of new particles occurs most commonly during photochemically active periods. Secondary accumulation mode aerosols are typically comprised largely of nitrate, sulphate, organic matter, and

ammonium, and the modal diameter depends on precursor concentrations and rates of oxidation chemistry. Whilst size distribution data does not provide any compositional information, sources and mechanisms can be inferred through their modal diameters, as well as their daily, weekly, and monthly cycles.

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The COVID–19 pandemic, and the associated lockdown periods, will have impacted these factors and their respective contributions to total PNCs. In order to better understand these changes, a mathematical receptor model (Positive Matrix Factorisation, PMF) was applied to particle size distribution data from six monitoring sites between four European cities (Helsinki, Leipzig, Budapest, and London) to identify the major factors contributing to total PNCs during lockdown compared to the associated periods in the reference years 2014–2019, depending on data coverage.

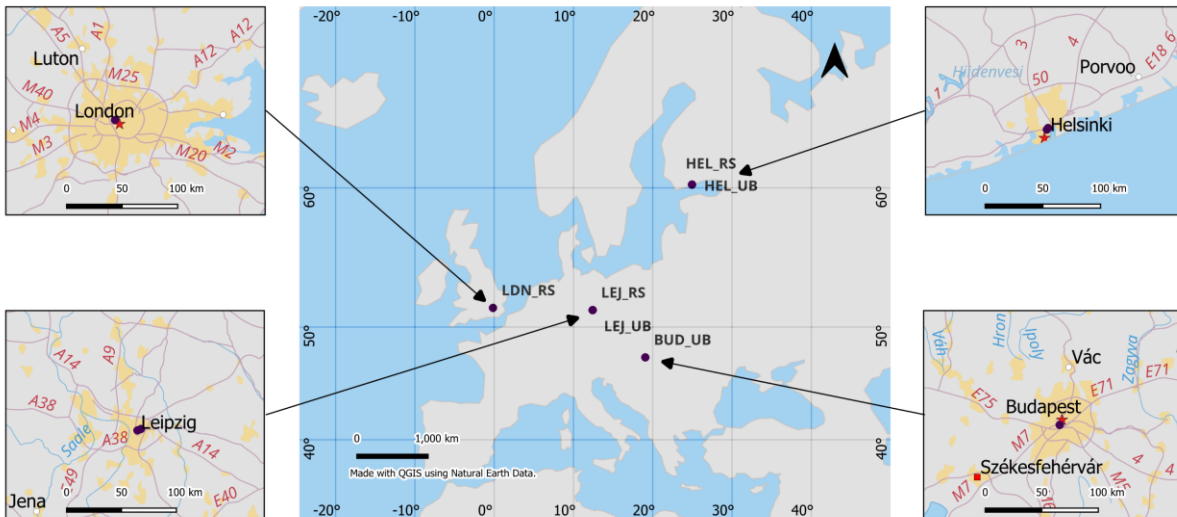
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## 2. METHODOLOGY

### 2.1. Monitoring Stations

This study is based on data from six monitoring sites spanning four European countries including Finland, Germany, Hungary, and the United Kingdom (**Figure 1**). Each site was assigned a unique identifier denoting its location and environment type (**Table 1**). The monitoring sites incorporate two different environment types, including urban background (denoted as UB) and roadside (denoted as RS) locations. Three urban background and three roadside sites were included in this study. The urban background sites are not dominated by any single source or street and are deemed representative of a well-mixed, average atmospheric environment within their respective cities. Their surroundings are similarly varied and broadly consist of commercial and residential property, transport infrastructure, and greenspace. The roadside sites are located such that their pollution levels are heavily influenced by the emissions from nearby traffic. Their surroundings are also similarly varied; however, they differ in terms of their aspect ratios (average building height divided by the most frequent width of the street canyon) and daily traffic volumes. As the COVID lockdown was different in each country, we chose the start point as where a substantial limitation on human movement was imposed around mid-March, and the end of the period as when some large change was made to human movement in mid-May. Using any single metric would lead to vastly different sampling periods where differences in meteorology would dominate our results, rather than differences in emissions.

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**Figure 1:** Monitoring site locations. *Above left*, London (United Kingdom) roadside; *above right*, Helsinki (Finland) urban background & roadside; *below left*, Leipzig (Germany) urban background & roadside; *below right*, Budapest (Hungary) urban background.

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**Table 1:** Monitoring site information. An Asterisk (\*) in the Site ID column denotes if a site is participating in the Aerosol, Clouds and Trace Gases Research Infrastructure (ACTRiS) Infrastructure Consortium.

City (country)	Monitoring site	Site ID	Environment Type	Coordinates and height above sea level	Lockdown pe- riods	Peak traffic hours (local times)
Helsinki (Finland)	Kumpula (SMEAR III)	HEL_UB*	Urban back- ground	N60°12'10" E24°57'40" 29 m	March 17th – May 15th	A.M. 08:00 P.M. 16:00
Leipzig (Germany)	TROPOS rooftop	LEJ_UB*	Urban back- ground	N51°21'09" E12°26'04" 127 m	March 23rd – May 5th	A.M. 07:00– 08:00 P.M. 15:00–16:00
Budapest (Hungary)	Lágymányos (BpART)	BUD_UB	Urban back- ground	N47°28'30" E19°3'45" 115 m	March 17th – May 19th	A.M. 07:00– 09:00 P.M. 15:00–18:00
Helsinki (Finland)	Mäkeläkatu	HEL_RS	Roadside	N60°11'48" E24°57'06" 33 m	March 17th – May 15th	As previously stated.
Leipzig (Germany)	Eisenbahnstrasse	LEJ_RS*	Roadside	N51°20'44" E12°24'23" 120 m	March 23rd – May 5th	As previously stated.
London (United Kingdom)	Marylebone Road	LDN_RS	Roadside	N51°31'21" W000°09'17" 35 m	March 27th – May 16th	A.M.–P.M. 07:00–20:00

Peak traffic hours were obtained from TomTom traffic index <https://www.tomtom.com/traffic-index/>

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### 2.1.1. Helsinki, Finland

Helsinki is located in Southern Finland, at the shore of the Gulf of Finland in the Baltic Sea. The capital is Finland's largest city, with 0.66 million inhabitants or 1.59 million inhabitants inclusive of its sub-regional units. Finland has 497 passenger cars per 1,000 inhabitants, which is below the average motorisation rate in the European Union (EU). However, Helsinki's car density is ~1,035 cars per km<sup>2</sup> which is comparable with densities observed in other cities across Europe.

Urban background aerosol data was obtained from the SMEAR III research station (Jarvi et al., 2009), located in the Kumpula campus of the University of Helsinki. Measurements are taken at 4 m above ground level and at distances >125 m from highly-trafficked roads bordering the site. The busy roadways experience 44,000 vehicles per workday (Järvi et al., 2012). The stations immediate surroundings also include multi-storey buildings, access roads, allotted parking bays, and greenspace.

Roadside aerosol data was obtained from a supersite monitoring station along one of Helsinki's main thoroughfares known as Mäkelänkatu. The street is ~42 m in width and is flanked by three- and four-storey buildings, yielding an aspect ratio of 0.40 (often referred to as an avenue canyon) (Rönkkö et al., 2017) and experiences 28,000 vehicles per day (Helin et al., 2018; Kuula et al., 2020). The stations immediate surroundings also include six lanes of traffic (three in each direction of travel), a central tramline bordered by tall vegetation, two footpaths, and on-street parking.

### 2.1.2. Leipzig, Germany

Leipzig is located in the German State of Saxony in east Germany. Leipzig is the 8<sup>th</sup> most populated city in Germany, with 0.6 million inhabitants. Germany has 574 passenger cars per 1,000 inhabitants which is comparable with the average motorisation rate in the EU. However, Leipzig's vehicle density is ~913 vehicles per km<sup>2</sup> which is comparable with densities observed in other cities across Europe.

Urban background aerosol data was obtained from an atmospheric research station operated by the Leibniz Institute for Tropospheric Research (TROPOS) within the Leipzig Science Park. Measurements are taken on the roof of an institute building at 16 m above ground level and at distances >100 m from highly-trafficked roads bordering the site (Klose et al., 2009; Birmili et al., 2016). The Science Park contains other research institutes and related companies, greenspace, and allotted parking bays, including a multi-storey carpark. The Park perimeter includes transport infrastructure (road, rail and tramways), commercial property (restaurants, hotels, a petrol station etc.), residential property, on-street parking, and greenspace.

Roadside aerosol data was obtained from a permanent observation site located along an important connecting road in the east of the city known as Eisenbahnstrasse. Measurements are taken from an apartment window at 6 m above ground level on the northern side of the street. The street is ~20 m in width and is flanked by multi-storey period buildings, yielding an aspect ratio of 0.90, and experiences 12,000 vehicles per working day (Birmili et al., 2016). The stations immediate surroundings also include two-lanes of traffic (one in each direction of travel), an integrated tramline, on-street parking, two bicycle lanes (one in each direction of travel), two footpaths, and scant vegetation.

### 2.1.3. Budapest, Hungary

Budapest is located in the Carpathian Basin in central Hungary. It is the capital and the largest city of the country, with 1.72 million inhabitants. Hungary has 395 passenger cars per 1,000 inhabitants which is one of the lowest motorisation rates in the EU. However, Budapest's car density is ~1,315 cars per km<sup>2</sup> which is comparable with densities commonly observed in other European cities.

Urban background aerosol data was obtained from the Budapest platform for Aerosol Research and Training (BpART) Laboratory, located on the second-floor balcony of the Northern block in the Lágymányos Campus of the Eötvös Loránd University. The balcony is 11 m above the street level of the closest road and is situated 85 m from the right bank of the River Danube (Salma et al., 2016). Sampling inlets and sensors are set up at heights of between 80 and 150 cm above the rooftop level of the measurement container.

### 2.1.4. London, United Kingdom

London is located in southeast England (United Kingdom, UK). The capital is the UK's largest city, with 9 million people occupying greater London. The UK has 589 passenger cars per 1,000 inhabitants which is comparable with the average motorisation rate in the EU. However, Greater London's car density is ~1,938 cars per km<sup>2</sup> which is well above average densities commonly observed in other cities across Europe.

Roadside aerosol data was obtained from a supersite monitoring station, located along Marylebone Road opposite one of London's top attractions, Madame Tussauds. Measurements are taken on the roof of a cabin positioned kerbside at 4 m above ground level (Harrison et al., 2019). The street is ~34 m in width and is flanked by multi-storey buildings, yielding an aspect ratio of 1.00 (often referred to as a regular street canyon), and experiences 80,000 vehicles per day (Harrison et al., 2019). The monitoring stations surroundings also include six lanes of traffic (three in each direction of

travel), two footpaths, and scant vegetation. Analyses of air quality data from this site have been reported by Kamara and Harrison (2021).

## 2.2. Instrumentation and Data Coverage

Instrumentation used to sample aerosol size distributions, as well as gaseous pollutants, particle mass concentrations, black carbon concentrations, and vehicle counts (referred to as auxiliary data) at the different monitoring sites are stated in **Table 2**, their sampling methodologies are outlined in **SI section 1.1**, and the applicable data coverage is plotted in **Figure S1**. The particle sizers covered different size ranges therefore a reduced common range (10–600 nm) was selected (although the whole range was utilised in the model; see **SI section 1.2**). The years of study varied depending on the availability of the data but each dataset covered the associated spring 2020 lockdown period, as well as an equivalent time period between 2014 and 2019, for comparison purposes.

**Table 2:** Sampling equipment used at each monitoring site.

Site ID	Particle sizer (size range, nm)	Black carbon (BC)	Nitrogen oxides (NO <sub>x</sub> )	Ozone (O <sub>3</sub> )	Sulphur dioxide (SO <sub>2</sub> )	Particulate matter (PM <sub>2.5</sub> )	Vehicle count (VC)
HEL_UB	DMPS (3–800)	–	Thermo TEI42	Thermo TEI49	Horiba APSA 360	–	–
LEJ_UB	TDMPS (3–800)	–	Horiba APNA 370	Horiba APOA–350E	Thermo Electron TE 43C–TL	–	–
BUD_UB	DMPS (6–1,000)	–	Thermo 42C	Ysselbach 43C	–	Environment MP101M	Traffic counter
HEL_RS	DMPS (6–800)	MAAP	Horiba APNA–370	Thermo Electron Model 49i/Horiba APOA–370	–	Thermo TEOM 1405	Traffic counter
LEJ_RS	DMPS / TDMPS (5/10– 800)	–	–	–	–	–	–
LDN_RS	SMPS (16.6–604)	Magee AE33	Teledyne API 200E	Teledyne API 400E	–	Palas Fidas 200	–

## 2.3. Positive Matrix Factorisation

The application of PMF is similar to the previous work of Rivas et al. (2020) and implemented using PMF2 (Positive Matrix Factorization (PMF2) (Sep 25, 2012) vers. 4.2, Copyright 1993, 2004 Pentti Paatero, Helsinki, Finland). PMF is a well-established receptor model (Paatero and Tapper, 1994) used to solve functional mixing models when the source profiles are unknown and presumed to be constant. PMF solutions are constrained to be non-negative, and a least squares algorithm is applied which accounts for uncertainties in the dataset. PMF is therefore quantitative and identifies a user-



specified number of sources depending on how well the outputs describe the monitoring site. These  
200 make it suitable for the source apportionment of size distribution data. Details of the PMF methodol-  
ogy can be found in **SI section 1.2**.

## 2.4. Condensation Sink

The condensation sink (CS) represents the rate at which a vapour phase molecule will collide with  
pre-existing particle surface, and was calculated from the size distribution data as follows:

$$205 \quad CS = 2\pi D \sum_{d_p} \beta_{m,d_p} d_p N_{d_p} \quad (1)$$

where  $D$  ( $\text{m}^2 \text{s}^{-1}$ ) is the diffusion coefficient of the diffusing vapour (assumed sulphuric acid),  $\beta_m$  is a  
transition regime correction,  $d_p$  (m) is particle diameter, and  $N_{d_p}$  ( $\text{m}^{-3}$ ) is the number of particles at  
diameter  $d_p$ .

## 2.5. Statistical Analysis

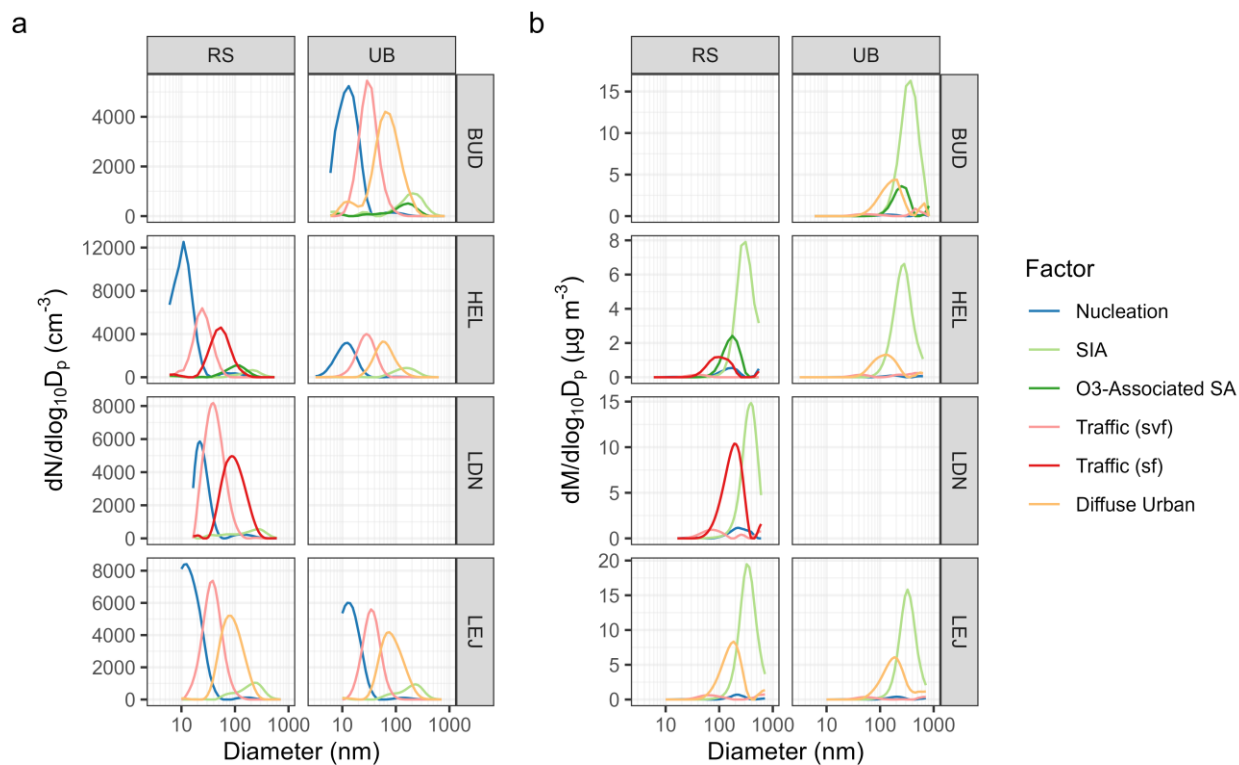
210 An Independent Samples T-test was performed to compare the means of the resolved factors from  
the PMF model (from the source apportionment of size distribution data; **section 2.3**) for the respec-  
tive lockdown periods at each site to the equivalent periods in the reference years (depending on data  
availability). The purpose of this test was to determine if the resolved factors were different between  
the associated periods at each location. The significance level was set at 0.05 (i.e. a p value <0.05  
215 was deemed statistically significant).

# 3. RESULTS AND DISCUSSION

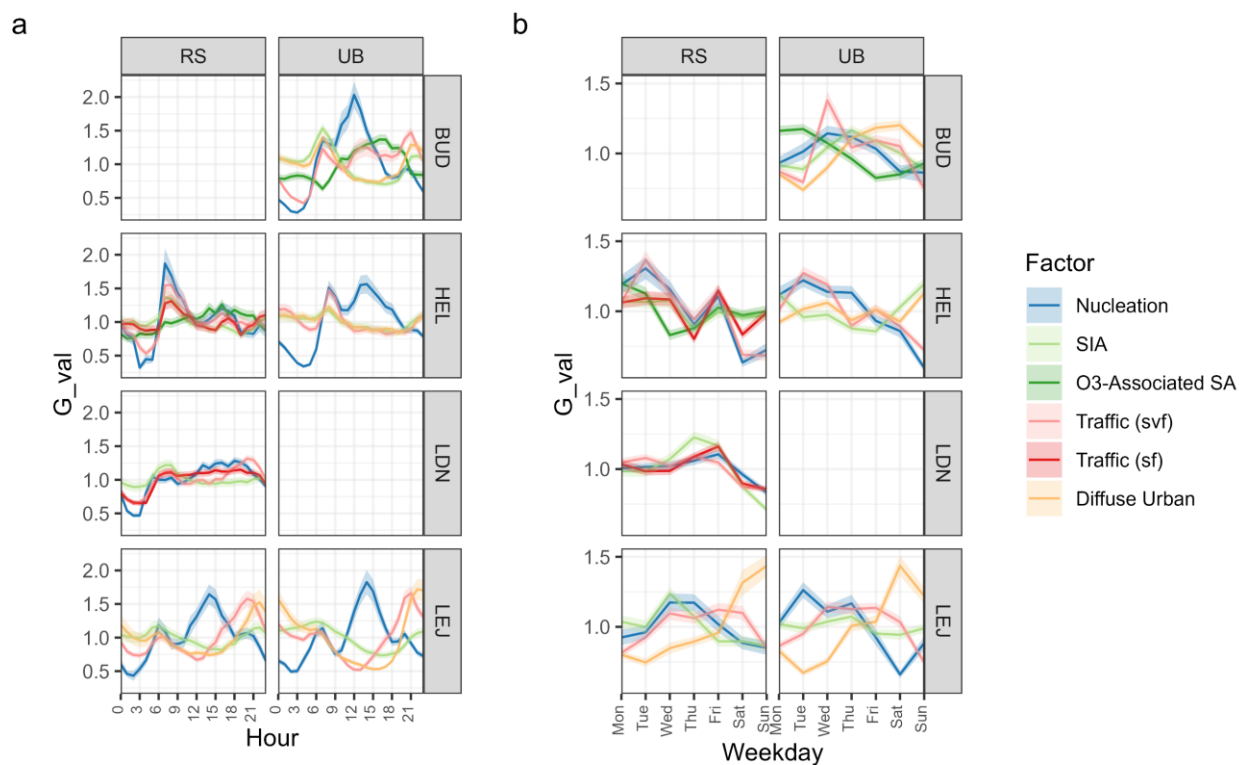
## 3.1. Identification of Major Factors

Data covering the spring 2020 lockdown period for each monitoring site were individually analysed  
using PMF. Multiple outputs were compared and a solution was selected based on the cogency and  
220 spatiotemporal behaviour of each factor. A number of common factors were resolved between the  
sites including nucleation, road traffic semi-volatile fraction (road traffic<sub>svf</sub>), road traffic solid frac-  
tion (road traffic<sub>sf</sub>), diffuse urban (woodsmoke + aged traffic), ozone-associated secondary aerosol  
(O<sub>3</sub>-associated SA), and secondary inorganic aerosol (SIA). We do not expect a substantial contribu-  
tion from dust and sea salt in the particle number size distribution at diameters <600 nm. Despite their  
225 commonalities, factors exhibited varying profiles at each site. Their particle number (**Figure 2a**) and  
mass (**Figure 2b**) size distributions, mean daily (**Figure 3a, diurnals during and outside of lock-  
downs in Figure S5**) and weekly (**Figure 3b**) cycles, relative associations with available auxiliary  
data (**Figure 4**), and polar plots showing wind directions and speeds coincidental with the top 25<sup>th</sup>

percentile of factor intensity (**Figure S6**) were tabled and/or plotted for analysis. The factors are  
 230 broadly summarised below and site-by-site descriptions can be found in **SI section 2.1**.

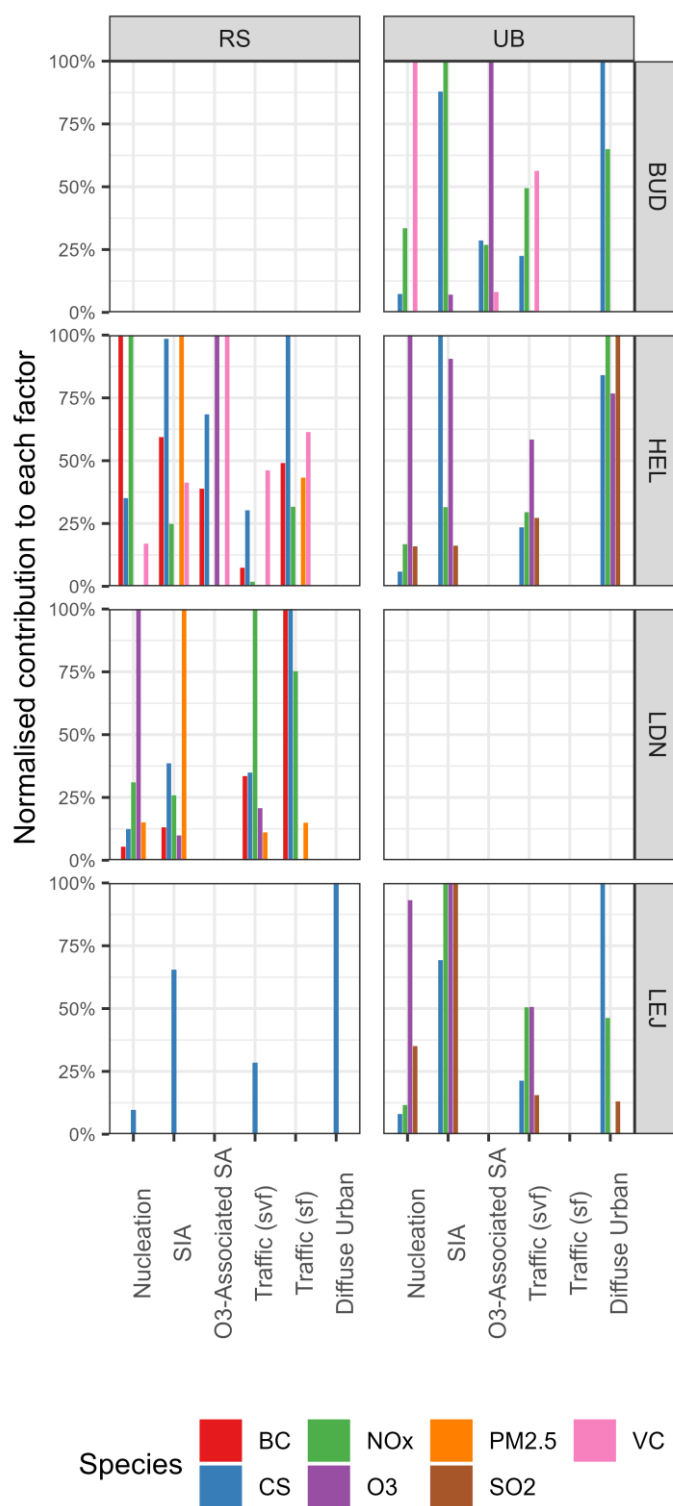


**Figure 2:** Number (a) and mass (b) size distribution data for each factor at each monitoring site. Each panel represents a factor and each colour represents a site.



235 **Figure 3:** Mean G values from the PMF model for each factor, showing (a) daily and (b) weekly cycles. Each panel represents a factor and each colour represents a monitoring site. The shaded region

represents the standard error of the mean. The G value is the time-series component of the PMF solution (see supplement).



240 **Figure 4:** Normalised contribution of downweighted auxiliary variables to each factor for each monitoring site. The mean G values for each variable for each site are normalised to a maximum of 1 and expressed as a percentage. The normalisation is performed so each variable has a maximum value of 1 for easy comparison. The G value is the time-series component of the PMF solution (see supplement).

### 245 3.1.1. Nucleation

Factors attributed to nucleation were resolved at all sites. The factors had a major mode in the size distribution which peaked at ~11–22 nm (**Figure 2a**) and a mass distribution dominated by particles in the accumulation mode (**Figure 2b**). Factor contributions typically peaked in the afternoon and with busy traffic periods (**Figure 3a**), and were higher on weekdays compared to weekends (**Figure 3b**). The factors were associated to varying degrees with BC, NO<sub>x</sub>, SO<sub>2</sub>, O<sub>3</sub>, and PM<sub>2.5</sub> (**Figure 4**). Additionally, the factors were consistently associated with low CS (**Figure 4**) which is a key determinant for NPF in urban environments (Deng et al., 2021). CS is the largest sink for vapours such as sulphuric acid, as well as other low volatility molecules, and occurs synchronously with high coagulation losses of small particles. We inferred that this factor is likely the sum of particles produced by means of photochemically induced nucleation processes, as well as some combustion-related processes, such as ultrafine vehicle emissions and the formation of new particles through the dilution and cooling of vehicle exhaust (Charron and Harrison, 2003; Pérez et al., 2010). The diurnal profiles at most sites suggest a predominantly non-traffic formation source. However, the nucleation factor at HEL\_RS is mainly related to the traffic source, since the diurnal variation shows clear morning and afternoon rush hour peaks. and the contribution of NO<sub>x</sub> and BC are very high.

### 3.1.2. Road traffic semi-volatile fraction (road traffic<sub>svf</sub>)

Factors attributed to road traffic<sub>svf</sub> were resolved at all sites. The factors had a major mode in the size distribution which peaked at ~25–40 nm (**Figure 2a**) and a mass distribution dominated by Aitken and/or accumulation mode particles (**Figure 2b**). Factor contributions typically peaked in the morning and afternoon/evening (**Figure 3a**), were higher on weekdays compared to weekends (**Figure 3b**), and were associated to varying degrees with combustion-related pollutants including BC, NO<sub>x</sub>, SO<sub>2</sub>, and PM<sub>2.5</sub> (**Figure 4**). In Europe, diesel vehicles are responsible for much of the exhaust PM from road traffic (Damayanti et al., 2023). Particles emitted in diesel exhaust fall into two main categories: semi-volatile and solid graphitic (black carbon) particles (Harrison et al., 2018; Kittelson et al., 2006). We inferred that these factors likely represent the former particle type, as well as a likely contribution from other mobile and/or stationary combustion-related activities (i.e. cooking and heating emissions). The factors closely resemble those referred to as ‘traffic 1’ in the literature, typically in reference to spark-ignition vehicle emissions or freshly emitted traffic particles (Hopke et al., 2022) and may include gasoline vehicle emissions.

### 275 3.1.3. Road traffic solid fraction (road traffic<sub>sf</sub>)

Factors attributed to road traffic<sub>sf</sub> were resolved at HEL\_RS and LDN\_RS. The factors had a major mode in the size distribution which peaked at ~55–90 nm (**Figure 2a**) and a mass distribution

dominated by particles in the Aitken and/or accumulation mode (**Figure 2b**). Factor contributions typically peaked in the morning and afternoon/evening (**Figure 3a**), were higher on weekdays compared to weekends (**Figure 3b**), and were associated to varying degrees with combustion-related pollutants including BC, NO<sub>x</sub>, SO<sub>2</sub>, and PM<sub>2.5</sub> (**Figure 4**). We inferred that these factors predominantly represent the solid particle mode arising from diesel road traffic (Harrison et al., 2018; Kittelson et al., 2006). The factors closely resemble those referred to as ‘traffic 2’ in the literature, typically in reference to diesel vehicle emissions or distant traffic particles (Hopke et al., 2022).

#### 285 **3.1.4. Diffuse urban**

Factors attributed to diffuse urban were mainly resolved at urban background sites. The factors had a major mode in the size distribution which peaked at ~75–90 nm (**Figure 2a**) and a mass distribution dominated by accumulation mode particles (**Figure 2b**). Factor contributions typically peaked in the morning and evening (**Figure 3a**), were higher on weekends compared to weekdays (**Figure 3b**), and were associated to varying degrees with combustion-related pollutants, including NO<sub>x</sub> and SO<sub>2</sub> (**Figure 4**). The factors closely resemble those referred to as ‘urban background’ by Beddows et al. (2015), and later ‘diffuse urban’ by Beddows and Harrison (2019), representing aged woodsmoke and road traffic emissions. A contribution of aged natural aerosol may also be present in this factor. ‘Urban background’ has other connotations and therefore we opted to use ‘diffuse urban’ as a factor name as it better categorised the emission source, with less confusion amongst the literature.

#### 3.1.5. Ozone-associated secondary aerosol (O<sub>3</sub>-associated SA)

Factors attributed to O<sub>3</sub>-associated SA were resolved at BUD\_UB and HEL\_RS. The factors had a major mode in the size distribution which peaked at ~125–175 nm (**Figure 2a**) and a mass distribution dominated by particles in the accumulation mode (**Figure 2b**). Factor contributions peaked in the daytime (**Figure 3a**), were higher on weekdays compared to weekends (**Figure 3b**), and were strongly associated with O<sub>3</sub> (**Figure 4**). The exact nature of these factors are uncertain; however, they are consistent with other such observations (often referred to as O<sub>3</sub>-rich SA) in the available literature (Ogulei et al., 2007; Liu et al., 2014; Squizzato et al., 2019) and likely represent particles which have grown through the condensation of secondary material (Hopke et al., 2022). In an urban environment it is reasonable to presume that most SA precursors are anthropogenic, but an influence of natural SA precursors will also contribute some fraction of total SA.

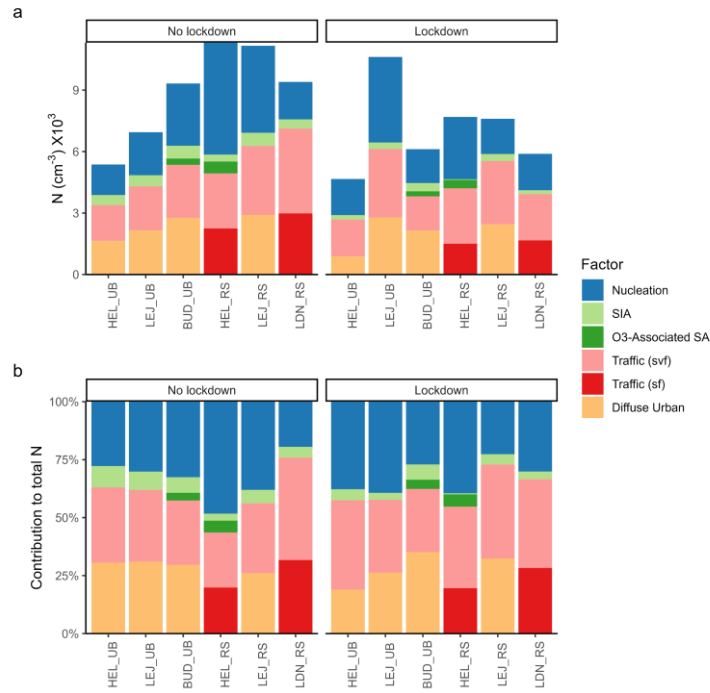
#### 3.1.6. Secondary inorganic aerosol (SIA)

Factors attributed to SIA were resolved at all sites. The factors had a major mode in the size distribution which peaked at ~175–265 nm (**Figure 2a**) and a mass distribution dominated by accumulation

310 mode particles (**Figure 2b**). Factor contributions typically peaked in the morning and evening/night  
(**Figure 3a**), were higher on weekdays compared to weekends (**Figure 3b**), and were associated with  
a host of auxiliary variables (**Figure 4**). Multimodal number size distributions have also been ob-  
served at other locations and suggest the presence of both local and distant particles thought to have  
315 been formed through the atmospheric processing of NO<sub>x</sub> and other gaseous precursor emissions  
(Ogulei et al., 2007; Kasumba et al., 2009).

### 3.2. Changes in Factor Contributions Under Lockdown Restrictions

To evaluate the effects of lockdown restrictions on PNCs, the applicable 2020 lockdown periods at  
each monitoring site were compared to the equivalent days of the year in the reference years 2014–  
2019, depending on data coverage. In examining temporal changes affecting pollutant concentrations,  
320 it is now common practice to remove the influences of changes in weather variables which affect  
primary pollutants concentrations (Vu et al., 2019; Shi et al., 2021) primarily by accounting for dilu-  
tion effects. Such a treatment was not applied in this study as weather variables affect both photo-  
chemical nucleation, with elevated temperatures accelerating the evaporation rate of clusters, and  
intense solar radiation accelerating the generation of NPF precursors (Lee et al., 2019; Bousiotis et  
325 al., 2021), and the semi-volatile fraction of both traffic and secondary aerosols through evaporation  
(Charron et al., 2003). Similarly, emissions of organic aerosol precursors are temperature-dependent  
(Lee et al., 2019). The dynamics of the number size distribution are therefore more complex than can  
be accounted for by de-weathering methods, and such results could not be interpreted with confi-  
dence. Rather, the trends in meteorological variables are presented in **Figure S8**, and commented  
330 upon in the text, where considered relevant.



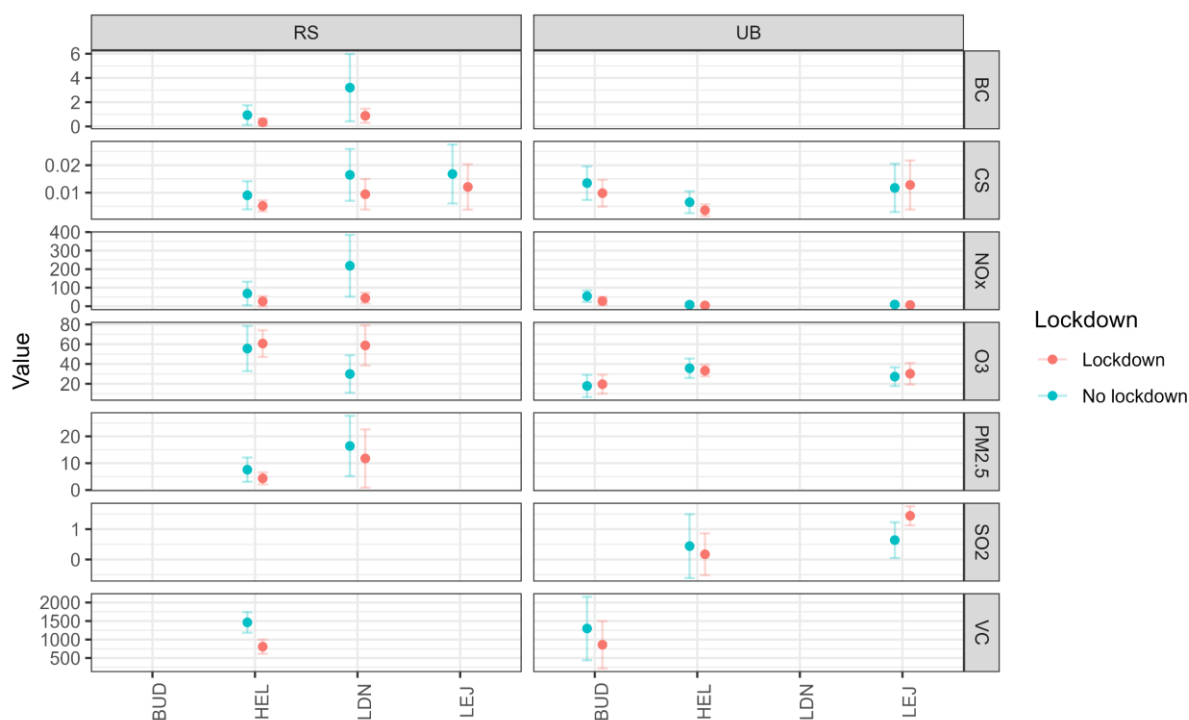
**Figure 5:** Mean total particle number concentrations (PNCs) (a) and mean contributions to total PNCs (b) for the 2020 lockdown period for each factor at each monitoring site and the equivalent periods in the reference years 2014–2019, depending on data availability. Also see Tables 3 and 4.

335 **Table 3:** Mean and min/max particle number concentrations ( $\text{cm}^{-3}$ ) during lockdown compared to the equivalent periods in the reference years 2014–2019 (depending on data availability) for urban background sites. An Asterisk (\*) in the Lockdown column denotes if a factor is significantly different ( $p$  value is less than 0.05) between the reference years and lockdown period by way of an Independent Samples T-test.

Factors	HEL_UB				LEJ_UB				BUD_UB			
	Reference		Lockdown		Reference		Lockdown		Reference		Lockdown	
	Mean	Min/Max	Mean	Min/Max	Mean	Min/Max	Mean	Min/Max	Mean	Min/Max	Mean	Min/Max
Nucleation	1,490	0/22,495	1,754*	0/14,053	2,178	0/50,641	4,176*	0/38,805	3,037	0/42,887	1,656*	0/26,723
Road traffic <sub>svf</sub>	1,746	0/19,577	1,789	79/9,659	2,211	0/19,586	3,334*	0/14,803	2,589	0/30,302	1,665*	0/14,886
Road traffic <sub>sf</sub>	–	–	–	–	–	–	–	–	–	–	–	–
Diffuse urban	1,642	0/10,943	889*	0/4,969	2,172	0/36,819	2,789*	0/33,532	2,762	0/13,977	2,150*	0/8,891
O <sub>3</sub> -associated SA	–	–	–	–	–	–	–	–	307	0/1,560	245*	0/1,155
SIA	492	0/3,893	221*	14/1,192	500	0/2,216	318*	5/1,316	626	0/2,452	402*	0/1,870
Total	5,370	–	4,653	–	6,944	–	10,617	–	9,321	–	6,118	–

340 **Table 4:** Mean and min/max particle number concentrations ( $\text{cm}^{-3}$ ) during lockdown compared to the equivalent periods in the reference years 2014–2019 (depending on data availability) for roadside sites. An Asterisk (\*) in the Lockdown column denotes if a factor is significantly different ( $p$  value is less than 0.05) between the reference years and lockdown period by way of an Independent Samples T-test.

Factors	HEL_RS				LEJ_RS				LDN_RS			
	Reference		Lockdown		Reference		Lockdown		Reference		Lockdown	
	Mean	Min/Max	Mean	Min/Max	Mean	Min/Max	Mean	Min/Max	Mean	Min/Max	Mean	Min/Max
Nucleation	5,464	0/65,412	3,041*	0/23,259	4,243	0/45,090	1,721*	0/19,449	1,830	0/11,119	1,775	0/8,393
Road traffic <sub>svf</sub>	2,684	0/26,689	2,704	391/12,156	3,362	0/24,168	3,083*	8/12,747	4,145	43/16,990	2,257*	0/7,807
Road traffic <sub>sf</sub>	2,247	0/16,009	1,503*	370/3,718	–	–	–	–	2,981	0/12,525	1,664*	0/8,318
Diffuse urban	–	–	–	–	2,907	0/38,104	2,456*	88/24,864	–	–	–	–
O <sub>3</sub> -associated SA	579	0/2,758	409*	0/1,349	–	–	–	–	–	–	–	–
SIA	342	16/2,702	34*	0/194	648	0/2,572	340*	11/1,411	444	16/2,422	194*	0/1,973
Total	11,320	–	7,692	–	11,140	–	7,600	–	9,400	–	5,890	–



350 **Figure 6:** Means of auxiliary variables during lockdown compared to the equivalent periods in the reference years 2014–2019 (depending on data availability) for each monitoring site included in this study. Black carbon (BC) =  $\mu\text{g m}^{-3}$ , condensation sink (CS) =  $\text{s}^{-1}$ , nitrogen oxides ( $\text{NO}_x$ ), ozone ( $\text{O}_3$ ), and sulphur dioxide ( $\text{SO}_2$ ) = ppb, and vehicle count (VC) =  $\text{h}^{-1}$ .

### 3.2.1. Comparison of total particle number concentrations during lockdowns to the reference years

355 During lockdown, total mean PNCs were lower at most urban background (Figure 5 and Table 3) and all roadside sites (Figure 5 and Table 4) included in this study, in comparison to the reference years. Regression estimates also showed that measured PNCs were lower than predicted at the majority of monitoring sites (Figure S7a), and the diurnal cycle of each source-related factor is similar for both the lockdown and reference years (Figure S5).

360 Among urban background sites, total mean PNCs were higher at LEJ\_UB, and lower at HEL\_UB and BUD\_UB, during lockdown compared to the equivalent periods in the reference years (Figure 5 and Table 3). Nucleation, road traffic<sub>svf</sub>, and diffuse urban factors were primarily responsible for the changes in total mean PNCs at LEJ\_UB.

365 Mean PNCs were lower at all of the roadside sites during lockdown compared to the equivalent periods in the reference years (Figure 5 and Table 4). Changes to nucleation and road traffic factors were primarily responsible for the fall in total mean PNCs at roadside sites.



### 3.2.2. Comparison of road traffic–related particle number concentrations during lockdowns to the reference years

370 Mean PNCs from factors attributed to road traffic<sub>svf</sub>, during reference and lockdown periods, were amongst the largest of the resolved factors, at both roadside and urban background sites (**Figure 5** and **Table 3/****Table 4**). The response of traffic<sub>svf</sub> to lockdown restrictions were varied. Mean PNCs from factors attributed to road traffic<sub>sf</sub> factors were also substantial, regardless of lockdown measures (**Figure 5** and **Table 4**). However, their decrease during lockdown restrictions were consistent across  
375 all sites.

Among urban background sites, mean PNCs from road traffic<sub>svf</sub> factors were higher at LEJ\_UB, comparable at HEL\_UB, and lower at BUD\_UB during lockdown compared to the equivalent periods in the reference years, with a mean change of +3.7% across the three sites (**Figure 5** and **Table 3**). Large  
380 reductions in traffic volumes were observed on major roads across Finland (Riuttanen et al., 2021) and the German state of Saxony (Jaekel and Muley, 2022). However, the associated emissions reductions may have been offset by increased emissions from private households. Fuel oils are regularly used in urban households for a variety of domestic activities and predominantly generate Aitken and nucleation mode particles (Tiwari et al., 2014). Lockdown restrictions had immediate and varied im-  
385 pacts on energy use, with increased residential demand due to people being confined to their homes. This points to a systematic misattribution of this factor throughout the literature, where this traffic factor contains some contribution from domestic heating. Road traffic volumes in central Budapest were also substantially reduced (Salma et al., 2020). These reductions were reflected in the traffic<sub>svf</sub> factor at BUD\_UB.

390 At roadside sites, mean PNCs from road traffic<sub>svf</sub> factors were comparable at HEL\_RS, and lower at LEJ\_RS and LDN\_RS during lockdown compared to the equivalent periods in the reference years, with a mean decrease of -21% (**Figure 5** and **Table 4**). Mean PNCs from road traffic<sub>sf</sub> factors were also lower at HEL\_RS and LDN\_RS during lockdown, with a mean decrease of -39% (**Figure 5** and  
395 **Table 4**). Changes to mean PNCs from road traffic factors were notably more pronounced at LDN\_RS than at other roadside sites included in this study. This is in some measure due to the enormous volumes of traffic typically present along Marylebone Road which were significantly reduced during lockdown (Hicks et al., 2021). However, BC concentrations have reduced considerably in recent years (**Figure S7c**), likely associated with the increased proportion of Euro 6/VI-compliant  
400 vehicles (Damayanti et al., 2023; Luoma et al., 2021). The adoption of Euro 6/VI vehicle emissions standards and the compulsory emission technologies (such as diesel particle filters, DPFs) will have

impacted road traffic factors over time, especially the road traffic<sub>sf</sub> factors resolved at HEL\_RS and LDN\_RS. DPFs do not suppress the semi-volatile mode of the emissions with high efficiency (Damayanti et al., 2023), explaining the smaller reduction in traffic<sub>svf</sub>.

### 405 **3.2.3. Comparison of diffuse urban particle number concentrations during lockdowns to the reference years**

Mean PNCs from factors attributed to diffuse urban, during reference and lockdown periods, were amongst the largest of the resolved factors (**Figure 5** and **Table 3/Table 4**). The response of these factors to lockdown measures were largely consistent, and the diurnal cycle in of each source-related  
410 factor is similar for both the lockdown and reference years (Figure S5).

Among urban background sites, mean PNCs from diffuse urban factors were higher at LEJ\_UB, and lower at HEL\_UB and BUD\_UB, during lockdown compared to the equivalent periods in the reference years, with a mean decrease of -11% (**Figure 5** and **Table 3**). Mean PNCs from the factor were  
415 also lower at LEJ\_RS during lockdown by -16% (**Figure 5** and **Table 4**). Changes to mean PNCs from diffuse urban factors likely reflect variations in residential wood combustion and reductions to traffic volumes. As previously mentioned, large reductions to traffic volumes were reported across Helsinki (Riuttanen et al., 2021), the German state of Saxony (Jaekel and Muley, 2022), and Central Budapest (Salma et al., 2020). Variations in residential wood combustion, on the other hand, may  
420 reflect responses to outdoor temperature fluctuations (**Figure S8**), as well as the more social and cultural aspects of urban air pollution. It is commonplace in Northern countries to use firewood in sauna stoves and various fireplaces as supplementary heating (Kukkonen et al., 2020). Reductions in BC concentrations in Northern Helsinki may be related to decreased wood-burning (Harni et al., 2023) and/or weather conditions during lockdown. Nevertheless, biomass burning has previously  
425 been identified as a major contributor to PM concentrations in Helsinki (PM<sub>2.5</sub>; Saarnio et al., 2012), Leipzig (PM<sub>10</sub>; van Pinxteren et al., 2016), and Budapest (PM<sub>2.5</sub>; Salma et al., 2017) during the heating period.

### **3.2.4. Comparison of nucleation particle number concentrations during lockdowns to the reference years**

430 Mean PNCs from factors attributed to nucleation, during reference and lockdown periods, were amongst the largest of the resolved factors, irrespective of environment type (**Figure 5** and **Table 3/Table 4**). The response of these factors to lockdown restrictions were varied.

Among urban background sites, mean PNCs from nucleation factors were higher at HEL\_UB and LEJ\_UB, and lower at BUD\_UB, during lockdown compared to the equivalent periods in the reference years with a mean increase of 13% (**Figure 5** and **Table 3**). Changes to mean PNCs from nucleation factors likely reflect variations to NPF, as well as reductions to primary and delayed primary emissions (Rönkkö et al., 2017). NPF in these urban areas is likely driven by sulphuric acid, bases such as dimethylamine, and OOMs, likely from AVOC oxidation (Lee et al., 2019), meaning our NPF is mostly driven by anthropogenic emissions. Variations in solar radiation (**Figure S8**) could also influence NPF via photochemical processes (Shen et al., 2021), but do not explain the trends we see in our data. The insolation was markedly higher in London during the lockdown period than in previous years, but the other cities show only a small increase. Strong solar radiation favours OH production which through various formation and oxidation processes produces sulphuric acid and other low volatility vapours in the atmosphere (Wang et al., 2023). NPF is inhibited by high particle surface area, which was lower during lockdowns at all sites except LEJ\_UB (**Figure 6**) (Du et al., 2022). Primary and delayed nucleation particles are linked to the number of vehicles on the road, as well as the associated emissions technologies, which can significantly impact the formation mechanisms and composition of emitted nanocluster aerosol (Rönkkö et al., 2017). The mixed response of nucleation factors at urban background sites to lockdown restrictions likely represents the interplay between these complex variables.

Across all urban background sites, the largest increase to nucleation was at LEJ\_UB, 92% increase, where the greatest increase to traffic factors was also seen (51% increase to  $\text{traffic}_{\text{svf}}$ ), while CS also increased (**Figure 6**). The largest decrease in nucleation was at BUD\_UB, 45% decrease, where the greatest reduction in traffic factors was also seen (-36% decrease to  $\text{traffic}_{\text{svf}}$ ), while CS decreased. Primary NCA will have a lifetime on the order of tens of minutes, and will need to grow to 10 nm and also undergo dilution while being transported from the roadside to the urban background measurement stations. Primary and delayed primary particle emissions will be less important here than at the roadside, and we infer some substantial role of primary gaseous traffic emissions in accelerating NPF (Brean et al., 2023).

465 **3.2.5. At roadside sites, mean PNCs from nucleation factors were comparable at LDN\_RS, and lower at HEL\_RS and LEJ\_RS, during lockdown compared to the equivalent periods in the reference years, with a mean decrease of -43% (Figure 5 and Table 4). It is important to acknowledge that instrumentation used to sample particle size distribution data at LDN\_RS covered a narrower diameter range compared to instruments used at other monitoring sites included in this study (Table 2). A common size range was selected between sites (10–600 nm) to help streamline the analysis except at LDN\_RS (17 to 600 nm). This discrepancy will have led to lower PNCs at LDN\_RS and likely disproportionately impacted the nucleation factor at this location. However, all particle sizers were equipped with an aerosol dryer to limit relative humidity in the sampled air (SI section 1.1). Relative humidity was controlled to minimise diameter changes due to hygroscopic growth and the resultant particle losses in the dryer were characterised and accounted for in the data analysis as recommended by Wiedensohler et al. (2012). The harmonised approach aided in the comparability between measurements, particularly in the lower size cut. Nevertheless, nucleation factors at roadside locations revealed some wide-ranging reductions to their mean PNCs during lockdown. Moreover, nucleation factors at monitoring sites in close proximity to one and other, but of different environment type, responded differently to lockdown measures. Again, this points to the importance of traffic and its multifaceted contribution to nucleation mode particles. Road traffic leads to an increased condensation sink which has a negative effect on nucleation, but is a source of organic vapours which can rapidly oxidise forming molecules of low volatility which enhance particle growth rates and survivability (Brean et al., 2023). The large decrease to nucleation at the roadside can be attributed either to a decrease in primary nanocluster aerosol emissions, or to a reduction in NPF precursor emissions (Brean et al., 2023). Comparison of secondary aerosol particle number concentrations during lockdowns to the reference years**

490 Mean PNCs from factors attributed to secondary aerosols ( $O_3$ -associated SA and SIA), during the reference and lockdown periods, were the smallest of the resolved factors, irrespective of environment type (Figure 5 and Table 3/Table 4). Though, SIA factors were the largest contributors to PMCs (Figure 2b) due to the large median diameters of the modes. The response of these factors to the lockdown periods were consistent across all of the monitoring sites included in this study.

495 Mean PNCs from  $O_3$ -associated SA and SIA factors were lower at urban background and roadside sites during lockdown compared to the equivalent periods in the reference years, with mean decreases

of -42% in SIA at the urban background sites, and -42% and -60% for O<sub>3</sub>-associated SA and SIA at the roadside (**Figure 5** and **Table 3/Table 4**). The response of SIA factors to lockdown restrictions may reflect reductions in gaseous precursor pollutants (**Figure 4**). Mechanisms of secondary aerosol formation changed under lockdown conditions in Beijing, for example, when NO<sub>x</sub> levels substantially declined (Yan et al., 2023). SIA is generated by the transfer of inorganic material from the vapour to the aerosol phase following the chemical processing of emitted gas-phase precursor emissions (McFiggans et al., 2015). Both precursors and particles may be emitted locally or transported long distances from adjacent source regions. Road traffic is typically the largest source of NO<sub>x</sub> in an urban area, as well as an under-recognised source of ammonia (Cao et al., 2022). Ammonia reacts with acid pollutants such as oxidation products of NO<sub>x</sub> and SO<sub>2</sub> to form ammonium nitrate and ammonium sulphate which is essential for the generation of SIA in PM (Duan et al., 2021). These interactions may help to explain why changes to mean PNCs from SIA factors were typically more pronounced at roadside sites. Similarly, if O<sub>3</sub>-associated SA is generated through ozonolysis of VOCs, VOC concentrations will have declined substantially during lockdown periods, although, O<sub>3</sub> concentrations increased during lockdown periods (Shi et al., 2021). This decline to secondary aerosol is large, consistent across all sites, and will result in a substantial reduction to PM mass.

#### 4. CONCLUSION

A multivariate factor analysis technique, PMF, was applied to particle number size distribution data to better understand how PNCs and their sources changed during the respective spring 2020 lockdown periods, compared to the equivalent days of the year in the reference years 2014–2019, depending on data coverage. The analysis involved six monitoring sites (three urban background and three roadside) between four European cities, including Helsinki, Leipzig, Budapest, and London. A number of common factors were resolved between the different sites, including nucleation, road traffic<sub>svf</sub>, road traffic<sub>sf</sub>, diffuse urban, O<sub>3</sub>-associated SA, and SIA. Despite their commonalities, factors exhibited varying profiles between sites, illustrative of the complex network of aerosol sources and sinks contributing to particle size distributions in urban areas.

The factors attributed to nucleation, road traffic, and diffuse urban were the largest contributors to mean PNCs, during the reference years and the respective lockdown periods. Total mean PNCs were lower at two of the three urban background sites and at all of the roadside sites during lockdown compared to the reference years. Nucleation factors showed highly variable behaviour. This perhaps demonstrates the important contribution from traffic to nucleation mode particles – either through the direct emission of primary aerosol or via key precursor compounds, such as amines and organic molecules. Road traffic<sub>svf</sub> factors were also highly variable. This likely reflects the complex interplay

between decreased precursor emissions and a lower condensation/coagulation sink giving variable outcomes. It is also possible that reduced traffic volumes and economic activities were partly counterbalanced by increased domestic emissions. Mean PNCs from road traffic<sub>sf</sub>, on the other hand, were notably lower at roadside locations. The response of diffuse urban factors to lockdown measures were largely consistent and perhaps reflect the more social and cultural aspects of urban air pollutant emissions. Secondary aerosols (O<sub>3</sub>-associated SA and SIA) exhibited extensive reductions to their mean PNCs during lockdown at all sites. However, SIA remained the largest contributor to PMCs.

The analyses reveal a complex and varied response in the particle size distributions to the curtailment of human mobility during the COVID-19 lockdown periods. The analyses also offer a glimpse into the future, where the electrification of road transport, together with traffic reduction schemes, may reduce mean PNCs from traffic, as well as potentially shift the relative importance of other sources in urban areas, driving the need for further air quality interventions and policy changes.

Furthermore, as alluded to in this study, the literature encompasses a wide range of named factors, often characterised by substantial overlap, particularly when it comes to factors related to road traffic. We argue that the named factors introduced in this study describe the reality better than variants present in other works and should be used going forward.

## DATA AND MATERIALS AVAILABILITY

Data supporting this publication are openly available from the UBIRA eData repository at

<https://doi.org/10.25500/edata.bham.00001040>

## AUTHOR CONTRIBUTIONS

Conceptualisation – JB; data curation and/or resources – TP, MV, IS, JN, HM and DvP; formal analysis – AR and JB; funding acquisition – RH; investigation – AR and JB; methodology – JB; project administration – RH; software – JB and DB; supervision – RH and ZS; visualisation – AR, JB, and DB; writing (original draft preparation) – AR; writing (review & editing) – JB, DB, ZS, RH, TP, IS, JN, and DvP.

## COMPETING INTERESTS

At least one of the (co-)authors is a member of the editorial board of Atmospheric Chemistry and Physics.

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## REFERENCES

- 585 Beddows, D. C. S. and Harrison, R. M.: Receptor modelling of both particle composition and size distribution from a background site in London, UK -- a two-step approach, *Atmos. Chem. Phys.*, 19(7), 4863–4876, doi:10.5194/acp-19-4863-2019, 2019.
- 590 Beddows, D. C. S., Harrison, R. M., Green, D. C. and Fuller, G. W.: Receptor modelling of both particle composition and size distribution from a background site in London, UK, *Atmos. Chem. Phys.*, 15(17), 10107–10125, doi:10.5194/acp-15-10107-2015, 2015.
- 595 Birmili, W., Weinhold, K., Rasch, F., Sonntag, A., Sun, J., Merkel, M., Wiedensohler, A., Bastian, S., Schladitz, A., Löschau, G., Cyrus, J., Pitz, M., Gu, J., Kusch, T., Flentje, H., Quass, U., Kaminski, H., Kuhlbusch, T. A. J., Meinhardt, F., Schwerin, A., Bath, O., Ries, L., Gerwig, H., Wirtz, K. and Fiebig, M.: Long-term observations of tropospheric particle number size distributions and equivalent black carbon mass concentrations in the German Ultrafine Aerosol Network (GUAN), *Earth Syst. Sci. Data*, 8(2), 355–382, doi:10.5194/essd-8-355-2016, 2016.
- 600 Bousiotis, D., Brean, J., Pope, F. D., Dall’Osto, M., Querol, X., Alastuey, A., Perez, N., Petäjä, T., Massling, A., Klenø Nøjgaard, J., Nordstrøm, C., Kouvarakis, G., Vratolis, S., Eleftheriadis, K., Niemi, J. V., Portin, H., Wiedensohler, A., Weinhold, K., Merkel, M., Tuch, T. and Harrison, R. M.: The effect of meteorological conditions and atmospheric composition in the occurrence and development of new particle formation (NPF) events in Europe, *Atmos. Chem. Phys.*, 21(5), 3345–3370, doi:10.5194/acp-21-3345-2021, 2021.
- 605 Brean, J., Rowell, A., D. C. S., Weinhold, K., Mettke, P., Merkel, M., Tuch, T., Rissanen, M., Dal Maso, M., Kumar, A., Barua, S., Iyer, S., Karppinen, A., Wiedensohler, A., Shi, Z., Harrison, R. M.: Road traffic emissions lead to much enhanced new particle formation through increased growth rates, submitted, 2023.
- 610 Cao, H., Henze, D. K., Cady-Pereira, K., McDonald, B. C., Harkins, C., Sun, K., Bowman, K. W., Fu, T.-M. and Nawaz, M. O.: COVID-19 Lockdowns Afford the First Satellite-Based Confirmation That Vehicles Are an Under-recognized Source of Urban NH<sub>3</sub> Pollution in Los Angeles, *Environ. Sci. Technol. Lett.*, 9(1), 3–9, doi:10.1021/acs.estlett.1c00730, 2022.
- 615 Charron, A. and Harrison, R. M.: Primary particle formation from vehicle emissions during exhaust dilution in the roadside atmosphere, *Atmos. Environ.*, 37(29), 4109–4119, doi:10.1016/S1352-2310(03)00510-7, 2003.
- 620 Cohen, A. J., Brauer, M., Burnett, R., Anderson, H. R., Frostad, J., Estep, K., Balakrishnan, K., Brunekreef, B., Dandona, L., Dandona, R., Feigin, V., Freedman, G., Hubbell, B., Jobling, A., Kan, H., Knibbs, L., Liu, Y., Martin, R., Morawska, L., Pope, C. A., Shin, H., Straif, K., Shaddick, G., Thomas, M., van Dingenen, R., van Donkelaar, A., Vos, T., Murray, C. J. L. and Forouzanfar, M. H.: Estimates and 25-year trends of the global burden of disease attributable to ambient air pollution: an analysis of data from the Global Burden of Diseases Study 2015, *Lancet*, 389(10082), 1907–1918, doi:10.1016/S0140-6736(17)30505-6, 2017.
- 625 Damayanti, S., Harrison, R. M., Pope, F. and Beddows, D. C. S.: Limited impact of diesel particle filters on road traffic emissions of ultrafine particles, *Environ. Int.*, 174, 107888, doi:10.1016/j.envint.2023.107888, 2023.
- Deng, C., Cai, R. and Yan, C.: Formation and growth of sub-3 nm particles in megacities : impact of background aerosols, *Faraday Discuss.*, 348–363, doi:10.1039/d0fd00083c, 2021.



- 635 Du, W., Cai, J., Zheng, F., Yan, C., Zhou, Y., Guo, Y., Chu, B., Yao, L., Heikkinen, L. M., Fan, X.,  
Wang, Y., Cai, R., Hakala, S., Chan, T., Kontkanen, J., Tuovinen, S., Petäjä, T., Kangasluoma, J.,  
Bianchi, F., Paasonen, P., Sun, Y., Kerminen, V.-M., Liu, Y., Daellenbach, K. R., Dada, L. and  
Kulmala, M.: Influence of Aerosol Chemical Composition on Condensation Sink Efficiency and New  
Particle Formation in Beijing, *Environ. Sci. Technol. Lett.*, 9(5), 375–382,  
doi:10.1021/acs.estlett.2c00159, 2022.
- 640 Duan, X., Yan, Y., Peng, L., Xie, K., Hu, D., Li, R. and Wang, C.: Role of ammonia in secondary  
inorganic aerosols formation at an ammonia-rich city in winter in north China: A comparative study  
among industry, urban, and rural sites, *Environ. Pollut.*, 291, 118151,  
doi:10.1016/j.envpol.2021.118151, 2021.
- 645 Hammer, M. S., Donkelaar, A. Van, Martin, R. V, McDuffie, E. E., Lyapustin, A., Sayer, A. M., Hsu,  
N. C., Levy, R. C., Garay, M. J., Kalashnikova, O. V and Kahn, R. A.: Effects of COVID-19  
lockdowns on fine particulate matter concentrations, *Sci. Adv.*, 7(26), 1–11,  
doi:10.1126/sciadv.abg767, 2021.
- 650 Harni, S. D., Saarikoski, S., Kuula, J., Helin, A., Aurela, M., Niemi, J. V, Kousa, A., Rönkkö, T. and  
Timonen, H.: Effects of emission sources on the particle number size distribution of ambient air in  
the residential area, *Atmos. Environ.*, 293, 119419, doi:10.1016/j.atmosenv.2022.119419, 2023.
- 655 Harrison, R. M., Beddows, D. C. S. and Dall’Osto, M.: PMF analysis of wide-range particle size  
spectra collected on a major highway, *Environ. Sci. Technol.*, 45(13), 5522–5528,  
doi:10.1021/es2006622, 2011.
- 660 Harrison, R. M., Rob MacKenzie, A., Xu, H., Alam, M. S., Nikolova, I., Zhong, J., Singh, A., Zeraati-  
Rezaei, S., Stark, C., Beddows, D. C. S., Liang, Z., Xu, R. and Cai, X.: Diesel exhaust nanoparticles  
and their behaviour in the atmosphere, *Proc. R. Soc. A Math. Phys. Eng. Sci.*, 474(2220), 20180492,  
doi:10.1098/rspa.2018.0492, 2018.
- 665 Harrison, R. M., Beddows, D. C. S., Alam, M. S., Singh, A., Brean, J., Xu, R., Kotthaus, S. and  
Grimmond, S.: Interpretation of particle number size distributions measured across an urban area  
during the FASTER campaign, *Atmos. Chem. Phys.*, 19(1), 39–55, doi:10.5194/acp-19-39-2019,  
2019.
- 670 Helin, A., Niemi, J. V, Virkkula, A., Pirjola, L., Teinilä, K., Backman, J., Aurela, M., Saarikoski, S.,  
Rönkkö, T., Asmi, E. and Timonen, H.: Characteristics and source apportionment of black carbon in  
the Helsinki metropolitan area, Finland, *Atmos. Environ.*, 190, 87–98,  
doi:10.1016/j.atmosenv.2018.07.022, 2018.
- 675 Hicks, W., Beevers, S., Tremper, A. H., Stewart, G., Priestman, M., Kelly, F. J., Lanoisellé, M.,  
Lowry, D. and Green, D. C.: Quantification of Non-Exhaust Particulate Matter Traffic Emissions  
And The Impact Of COVID-19 lockdown at London Marylebone Road, *Atmosphere (Basel)*, 12(2),  
doi:10.3390/atmos12020190, 2021.
- 680 Hopke, P. K., Feng, Y. and Dai, Q.: Source apportionment of particle number concentrations: A  
global review, *Sci. Total Environ.*, 819, 153104, doi:10.1016/j.scitotenv.2022.153104, 2022.
- 685 Jaekel, B. and Muley, D.: Transport impacts in Germany and State of Qatar: An assessment during  
the first wave of COVID-19, *Transp. Res. Interdiscip. Perspect.*, 13, 100540,  
doi:10.1016/j.trip.2022.100540, 2022.

- Jarvi, L., Hannuniemi, H., Hussein, T., Junninen, H., Aalto, P. P., Hillamo, R., Makela, T., Keronen, P., Siivola, E., Vesala, T. and Kulmala, M.: The urban measurement station SMEAR III: Continuous monitoring of air pollution and surface-atmosphere interactions in Helsinki, Finland, *Boreal Environ. Res.*, 14, 86–109, 2009.
- 690 Järvi, L., Nordbo, A., Junninen, H., Riikonen, A., Moilanen, J., Nikinmaa, E. and Vesala, T.: Seasonal and annual variation of carbon dioxide surface fluxes in Helsinki, Finland, in 2006–2010, *Atmos. Chem. Phys.*, 12(18), 8475–8489, doi:10.5194/acp-12-8475-2012, 2012.
- 695 Jiang, S., Zhang, F., Ren, J., Chen, L., Yan, X., Liu, J., Sun, Y. and Li, Z.: Evaluation of the contribution of new particle formation to cloud droplet number concentration in the urban atmosphere, *Atmos. Chem. Phys.*, 21, 14293–14308, doi:10.5194/acp-21-14293-2021, 2021.
- 700 Kamara, A. A. and Harrison, R. M.: Analysis of the air pollution climate of a central urban roadside supersite: London, Marylebone Road, *Atmos. Environ.*, 258, 118479, doi:10.1016/j.atmosenv.2021.118479, 2021.
- 705 Kasumba, J., Hopke, P. K., Chalupa, D. C. and Utell, M. J.: Comparison of sources of submicron particle number concentrations measured at two sites in Rochester, NY, *Sci. Total Environ.*, 407(18), 5071–5084, doi:10.1016/j.scitotenv.2009.05.040, 2009.
- Kittelson, D. B., Watts, W. F. and Johnson, J. P.: On-road and laboratory evaluation of combustion aerosols—Part 1: Summary of diesel engine results, *J. Aerosol Sci.*, 37(8), 913–930, doi:10.1016/j.jaerosci.2005.08.005, 2006.
- 710 Klose, S., Birmili, W., Voigtländer, J., Tuch, T., Wehner, B., Wiedensohler, A. and Ketzel, M.: Particle number emissions of motor traffic derived from street canyon measurements in a Central European city, *Atmos. Chem. Phys. Discuss.*, 9, 3763–3809, doi:10.5194/acpd-9-3763-2009, 2009.
- 715 Kukkonen, J., López-Aparicio, S., Segersson, D., Geels, C., Kangas, L., Kauhaniemi, M., Maragkidou, A., Jensen, A., Assmuth, T., Karppinen, A., Sofiev, M., Hellén, H., Riikonen, K., Nikmo, J., Kousa, A., Niemi, J. V, Karvosenoja, N., Santos, G. S., Sundvor, I., Im, U., Christensen, J. H., Nielsen, O.-K., Plejdrup, M. S., Nøjgaard, J. K., Omstedt, G., Andersson, C., Forsberg, B. and Brandt, J.: The influence of residential wood combustion on the concentrations of PM<sub>2.5</sub> in four  
720 Nordic cities, *Atmos. Chem. Phys.*, 20(7), 4333–4365, doi:10.5194/acp-20-4333-2020, 2020.
- 725 Kulmala, M., Petäjä, T., Ehn, M., Thornton, J., Sipilä, M., Worsnop, D. R. and Kerminen, V.-M.: Chemistry of atmospheric nucleation: on the recent advances on precursor characterization and atmospheric cluster composition in connection with atmospheric new particle formation., *Annu. Rev. Phys. Chem.*, 65, 21–37, doi:10.1146/annurev-physchem-040412-110014, 2014.
- 730 Kuula, J., Kuuluvainen, H., Niemi, J. V, Saukko, E., Portin, H., Kousa, A., Aurela, M., Rönkkö, T. and Timonen, H.: Long-term sensor measurements of lung deposited surface area of particulate matter emitted from local vehicular and residential wood combustion sources, *Aerosol Sci. Technol.*, 54(2), 190–202, doi:10.1080/02786826.2019.1668909, 2020.
- 735 Lee, S. H., Gordon, H., Yu, H., Lehtipalo, K., Haley, R., Li, Y. and Zhang, R.: New particle formation in the atmosphere: From molecular clusters to global climate, *J. Geophys. Res. Atmos.*, doi:10.1029/2018JD029356, 2019.
- Liu, Z. R., Hu, B., Liu, Q., Sun, Y. and Wang, Y. S.: Source apportionment of urban fine particle number concentration during summertime in Beijing, *Atmos. Environ.*, 96, 359–369,

- doi:10.1016/j.atmosenv.2014.06.055, 2014.
- 740 Luoma, K., Niemi, J. V, Aurela, M., Fung, P. L., Helin, A., Hussein, T., Kangas, L., Koussa, A., Rönkkö, T., Timonen, H., Virkkula, A. and Petäjä, T.: Spatiotemporal variation and trends in equivalent black carbon in the Helsinki metropolitan area in Finland, *Atmos. Chem. Phys.*, 21(2), 1173–1189, doi:10.5194/acp-21-1173-2021, 2021.
- 745 McFiggans, G., Alfarra, M., Allan, J., Coe, H., Hamilton, J., Harrison, R., Jenkin, M., Lewis, A., Moller, S. and Williams, P.: A review of the state-of-the-science relating to secondary particulate matter of relevance to the composition of the UK atmosphere: Full technical report to Defra, project AQ0732., 2015.
- 750 Ogulei, D., Hopke, P. K., Chalupa, D. C. and Utell, M. J.: Modeling source contributions to submicron particle number concentrations measured in Rochester, New York, *Aerosol Sci. Technol.*, 41(2), 179–201, doi:10.1080/02786820601116012, 2007.
- 755 Ohlwein, S., Kappeler, R., Kutlar, M. and Nino, J.: Health effects of ultrafine particles: a systematic literature review update of epidemiological evidence, *Int. J. Public Health*, 7, doi:10.1007/s00038-019-01202-7, 2019.
- 760 Okuljar, M., Kuuluvainen, H., Kontkanen, J., Garmash, O., Olin, M. and Niemi, J. V: Measurement report: The influence of traffic and new particle formation on the size distribution of 1-800 nm particles in Helsinki: a street canyon and an urban background station comparison, *Atmos. Chem. Phys.*, (January), 1–29, doi:10.5194/acp-21-9931-2021, 2021.
- 765 Paatero, P. and Tapper, U.: Positive matrix factorization: A non-negative factor model with optimal utilization of error estimates of data values, *Environmetrics*, 5(2), 111–126, doi:10.1002/env.3170050203, 1994.
- 770 Pérez, N., Pey, J., Cusack, M., Reche, C., Querol, X., Alastuey, A. and Viana, M.: Variability of particle number, black carbon, and PM<sub>10</sub>, PM<sub>2.5</sub>, and PM<sub>1</sub> levels and speciation: Influence of road traffic emissions on urban air quality, *Aerosol Sci. Technol.*, 44(7), 487–499, doi:10.1080/02786821003758286, 2010.
- 775 van Pinxteren, D., Fomba, K. W., Spindler, G., Müller, K., Poulain, L., Iinuma, Y., Löschau, G., Hausmann, A. and Herrmann, H.: Regional air quality in Leipzig Germany: detailed source apportionment of size-resolved aerosol particles and comparison with the year 2000, *Faraday Discuss.*, 189(0), 291–315, doi:10.1039/C5FD00228A, 2016.
- 780 Putaud, J.-P., Pisoni, E., Mangold, A., Hueglin, C., Sciare, J., Pikridas, M., Savvides, C., Ondracek, J., Mbengue, S., Wiedensohler, A., Weinhold, K., Merkel, M., Poulain, L., van Pinxteren, D., Herrmann, H., Massling, A., Nordstroem, C., Alastuey, A., Reche, C., Pérez, N., Castillo, S., Sorribas, M., Adame, J. A., Petaja, T., Lehtipalo, K., Niemi, J., Riffault, V., de Brito, J. F., Colette, A., Favez, O., Petit, J.-E., Gros, V., Gini, M. I., Vratolis, S., Eleftheriadis, K., Diapouli, E., van der Gon, H., Yttri, K. E. and Aas, W.: Impact of 2020 COVID-19 lockdowns on particulate air pollution across Europe, *EGUsphere*, 2023, 1–21, doi:10.5194/egusphere-2023-434, 2023.
- 785 Le Quéré, C., Jackson, R. B., Jones, M. W., Smith, A. J. P., Abernethy, S., Andrew, R. M., De-gol, A. J., Willis, D. R., Shan, Y., Canadell, J. G., Friedlingstein, P., Creutzig, F. and Peters, G. P.: Temporary reduction in daily global CO<sub>2</sub> emissions during the COVID-19 forced confinement, *Nat. Clim. Chang.*, 10, 647–654, doi:10.1038/s41558-020-0797-x, 2020.

790

Riuttanen, A., Ponkilainen, V., Kuitunen, I., Reito, A., Sirola, J. and Mattila, V. M.: Severely injured patients do not disappear in a pandemic: Incidence and characteristics of severe injuries during COVID-19 lockdown in Finland, *Acta Orthop.*, 92(3), 249–253, doi:10.1080/17453674.2021.1881241, 2021.

795

Rivas, I., Beddows, D. C. S., Amato, F., Green, D. C., Järvi, L., Hueglin, C., Reche, C., Timonen, H., Fuller, G. W., Niemi, J. V., Pérez, N., Aurela, M., Hopke, P. K., Alastuey, A., Kulmala, M., Harrison, R. M., Querol, X. and Kelly, F. J.: Source apportionment of particle number size distribution in urban background and traffic stations in four European cities, *Environ. Int.*, 135, 105345, doi:10.1016/j.envint.2019.105345, 2020.

800

Rönkkö, T. and Timonen, H.: Overview of Sources and Characteristics of Nanoparticles in Urban Traffic-Influenced Areas, *J. Alzheimer's Dis.*, 72, 15–28, doi:10.3233/JAD-190170, 2019.

805

Rönkkö, T., Kuuluvainen, H., Karjalainen, P., Keskinen, J., Hillamo, R., Niemi, J. V., Pirjola, L., Timonen, H. J., Saarikoski, S., Saukko, E., Järvinen, A., Silvennoinen, H., Rostedt, A., Olin, M., Yli-Ojanperä, J., Nousiainen, P., Kousa, A. and Dal Maso, M.: Traffic is a major source of atmospheric nanocluster aerosol, *Proc. Natl. Acad. Sci. U. S. A.*, 114(29), 7549–7554, doi:10.1073/pnas.1700830114, 2017.

810

Saarnio, K., Niemi, J., Saarikoski, S., Aurela, M., Timonen, H., Teinila, K., Myllynen, M., Frey, A., Lamberg, H., Jokiniemi, J. and Hillamo, R.: Using monosaccharide anhydrides to estimate the impact of wood combustion on fine particles in the Helsinki Metropolitan Area, *Boreal Environ. Res.*, 17(3–4), 163–183, 2012.

815

Salma, I., Németh, Z., Kerminen, V., Aalto, P., Nieminen, T. and Weidinger, T.: Regional effect on urban atmospheric nucleation, *Atmos. Chem. Phys.*, 16(14), 8715–8728, doi:10.5194/acp-16-8715-2016, 2016.

820

Salma, I., Németh, Z., Weidinger, T., Maenhaut, W., Claeys, M., Molnár, M., Major, I., Ajtai, T., Utry, N. and Bozóki, Z.: Source apportionment of carbonaceous chemical species to fossil fuel combustion, biomass burning and biogenic emissions by a coupled radiocarbon–levoglucosan marker method, *Atmos. Chem. Phys.*, 17(22), 13767–13781, doi:10.5194/acp-17-13767-2017, 2017. Salma, I., Vörösmarty, M., Gyöngyösi, A. Z., Thén, W. and Weidinger, T.: What can we learn about urban air quality with regard to the first outbreak of the COVID-19 pandemic? A case study from central Europe, *Atmos. Chem. Phys.*, 15725–15742, doi:10.5194/acp-20-15725-2020, 2020.

825

Shen, X., Sun, J., Yu, F., Wang, Y., Zhong, J., Zhang, Y., Hu, X., Xia, C., Zhang, S. and Zhang, X.: Enhancement of nanoparticle formation and growth during the COVID-19 lockdown period in urban Beijing, *Atmos. Chem. Phys.*, 21(9), 7039–7052, doi:10.5194/acp-21-7039-2021, 2021.

830

Shi, Z., Song, C., Liu, B., Lu, G., Xu, J., Vu, T. Van, Elliott, R. J. R., Li, W., Bloss, W. J. and Harrison, R. M.: Abrupt but smaller than expected changes in surface air quality attributable to COVID-19 lockdowns, *Science (80-. )*, 7(3), doi:10.1126/sciadv.abd6696, 2021.

835

Squizzato, S., Masiol, M., Emami, F., Chalupa, D. C., Utell, M. J., Rich, D. Q. and Hopke, P. K.: Long-Term Changes of Source Apportioned Particle Number Concentrations in a Metropolitan Area of the Northeastern United States, *Atmosphere (Basel)*, 10(1), 27, doi:10.3390/atmos10010027, 2019.

840

Storelvmo, T., Leirvik, T., Lohmann, U., Phillips, P. C. B. and Wild, M.: Disentangling greenhouse

- warming and aerosol cooling to reveal Earth's climate sensitivity, *Nat. Geosci.*, 9(4), 286–289, doi:10.1038/ngeo2670, 2016.
- 845 Tiwari, M., Sahu, S. K., Bhangare, R. C., Yousaf, A. and Pandit, G. G.: Particle size distributions of ultrafine combustion aerosols generated from household fuels, *Atmos. Pollut. Res.*, 5(1), 145–150, doi:10.5094/APR.2014.018, 2014.
- 850 Torkmahalleh, M. A., Akhmetvaliyeva, Z., Omran, A. D., Omran, F. F. D., Kazemitabar, M., Naseri, M., Naseri, M., Sharifi, H., Malekipirbazari, M., Adotey, E. K., Gorjinezhad, S., Eghtesadi, N., Sabanov, S., Alastuey, A., de Fátima Andrade, M., Buonanno, G., Carbone, S., Cárdenas-Fuentes, D. E., Cassee, F. R., Dai, Q., Henríquez, A., Hopke, P. K., Keronen, P., Khwaja, H. A., Kim, J., Kulmala, M., Kumar, P., Kushta, J., Kuula, J., Massagué, J., Mitchell, T., Mooibroek, D., Morawska, L., Niemi, J. V., Ngagine, S. H., Norman, M., Oyama, B., Oyola, P., Öztürk, F., Petäjä, T., Querol, X., Rashidi, Y., Reyes, F., Ross-Jones, M., Salthammer, T., Savvides, C., Stabile, L., Sjöberg, K., Söderlund, K., Raman, R. S., Timonen, H., Umezawa, M., Viana, M. and Xie, S.: Global air quality and COVID-19 pandemic: Do we breathe cleaner air?, *Aerosol Air Qual. Res.*, 21(4), 200567, doi:10.4209/aaqr.200567, 2021.
- 860 Vu, T. V., Delgado-Saborit, J. M. and Harrison, R. M.: Review: Particle number size distributions from seven major sources and implications for source apportionment studies, *Atmos. Environ.*, 122, 114–132, doi:10.1016/j.atmosenv.2015.09.027, 2015.
- 865 Vu, T. V., Shi, Z., Cheng, J., Zhang, Q., He, K., Wang, S. and Harrison, R. M.: Assessing the impact of clean air action on air quality trends in Beijing using a machine learning technique, *Atmos. Chem. Phys.*, 19(17), 11303–11314, doi:10.5194/acp-19-11303-2019, 2019.
- 870 Wang, K., Ma, X., Tian, R. and Yu, F.: Analysis of new particle formation events and comparisons to simulations of particle number concentrations based on GEOS-Chem–advanced particle microphysics in Beijing, China, *Atmos. Chem. Phys.*, 23(7), 4091–4104, doi:10.5194/acp-23-4091-2023, 2023.
- 875 Wiedensohler, A., Birmili, W., Nowak, A., Sonntag, A., Weinhold, K., Merkel, M., Wehner, B., Tuch, T., Pfeifer, S., Fiebig, M., Fjåraa, A. M., Asmi, E., Sellegri, K., Depuy, R., Venzac, H., Villani, P., Laj, P., Aalto, P., Ogren, J. A., Swietlicki, E., Williams, P., Roldin, P., Quincey, P., Hüglin, C., Fierz-Schmidhauser, R., Gysel, M., Weingartner, E., Riccobono, F., Santos, S., Gruning, C., Faloon, K., Beddows, D., Harrison, R., Monahan, C., Jennings, S. G., O'Dowd, C. D., Marinoni, A., Horn, H. G., Keck, L., Jiang, J., Scheckman, J., McMurry, P. H., Deng, Z., Zhao, C. S., Moerman, M., Henzing, B., De Leeuw, G., Löschau, G. and Bastian, S.: Mobility particle size spectrometers: Harmonization of technical standards and data structure to facilitate high quality long-term observations of atmospheric particle number size distributions, *Atmos. Meas. Tech.*, 5(3), 657–685, doi:10.5194/amt-5-657-2012, 2012.
- 880 Yan, C., Tham, Y. J., Nie, W., Xia, M., Wang, H., Guo, Y., Ma, W., Zhan, J., Hua, C., Li, Y., Deng, C., Li, Y., Zheng, F., Chen, X., Li, Q., Zhang, G., Mahajan, A. S., Cuevas, C. A., Huang, D. D., Wang, Z., Sun, Y., Saiz-Lopez, A., Bianchi, F., Kerminen, V.-M., Worsnop, D. R., Donahue, N. M., Jiang, J., Liu, Y., Ding, A. and Kulmala, M.: Increasing contribution of nighttime nitrogen chemistry to wintertime haze formation in Beijing observed during COVID-19 lockdowns, *Nat. Geosci.*, doi:10.1038/s41561-023-01285-1, 2023.
- 890