



Traffic-originated nanocluster emission exceeds H₂SO₄-driven photochemical new particle formation in an urban area

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Abstract. Elevated ambient concentrations of sub-3 nm particles (nanocluster aerosol, NCA) are generally related to atmospheric new particle formation events, usually linked with gaseous sulfuric acid (H₂SO₄) produced via photochemical oxidation of sulfur dioxide. According to our measurement results of H₂SO₄ and NCA concentrations, traffic density, and solar irradiance at an urban traffic site in Helsinki, Finland, the view of aerosol formation in traffic-influenced environments is updated by presenting two separate and independent pathways of traffic affecting the atmospheric NCA concentrations: by acting as a direct nanocluster source, and by influencing the production of H₂SO₄. As traffic density is frequently correlated with solar radiation, it is likely that the influence of traffic-related nanoclusters has been hidden in the diurnal variation, and is thus underestimated because new particle formation events also follow the diurnal cycle of sunlight. Urban aerosol formation studies should, therefore, be updated to include the proposed formation mechanisms. Additionally, the directly emitted NCA poses a potentially elevated health risk, highlighting the need for quantifying the source and loadings in populated areas. The formation of H₂SO₄ in urban environments is here separated in two routes: primary H₂SO₄ is formed in hot vehicle exhaust and is converted rapidly to particle phase; secondary H₂SO₄ results from the combined effect of emitted gaseous precursors and available solar radiation.

1 Introduction

Urban environments exhibit some of the highest aerosol particle concentrations encountered in the Earth's atmosphere. Elevated particle concentrations are related to adverse health effects (Dockery et al., 1993; Pope et al., 2002; Beelen et al., 2014) and various effects on climate (Arneth et al., 2009). Recent studies on urban aerosol particles have focused attention on the formation process of sub-3 nm particles (Zhao et al., 2011; Kulmala et al., 2013; Kontkanen et al., 2017) also called nanocluster aerosol (NCA) (Rönkkö et al., 2017). The importance of photochemical formation mechanisms, involving, e.g., sulfuric acid (H₂SO₄) and ammonia (Yao et al., 2018) or other photochemically produced vapors (Lehtipalo et al., 2018), has



been highlighted. However, these studies omit the important role of direct emission of NCA-sized particles in their analysis, despite recent findings that traffic is a major source of such particles (Rönkkö et al., 2017). The proposed mechanisms also assume that key precursor vapors are formed via photochemical oxidation (Paasonen et al., 2010; Lehtipalo et al., 2018).

The most important gaseous species forming new particles in the atmosphere is H_2SO_4 , the main source of which is usually considered to be sulfur dioxide (SO_2). SO_2 is photochemically oxidized in the atmosphere by an oxidizing agent produced by solar radiation, such as hydroxyl radical (OH) (Kulmala et al., 2014). H_2SO_4 produced via this route is here termed *secondary H_2SO_4* . Sources of regional SO_2 include shipping, power generation, atmospheric oxidation of dimethyl sulfide, and volcanic activity. Additionally, motor vehicles emit SO_2 due to sulfur-containing fuels and lubricant oils; hence, also traffic can contribute to the secondary H_2SO_4 levels. A part of SO_2 formed during combustion is oxidized to H_2SO_4 already in vehicles' oxidative exhaust after-treatment systems (Arnold et al., 2012), which makes vehicles direct H_2SO_4 emitters. In contrast to the secondary H_2SO_4 formed via photochemistry, H_2SO_4 formed in hot exhaust without the need of solar radiation is here termed *primary H_2SO_4* . In principle, primary H_2SO_4 can also contribute to the atmospheric H_2SO_4 concentrations, at least in the vicinity of traffic.

Ambient aerosol particles are either emitted directly into the atmosphere as primary particles or new particles are formed from atmospheric precursor gases in a new particle formation (NPF) process. NPF processes have been shown to occur in a variety of environments, and their occurrence is believed to be controlled by, on one hand, by the availability of particle-forming vapors and, on the other hand, by the reduction of the vapors and fresh cluster-sized nuclei due to pre-existing aerosol surface area acting as a condensation sink (CS) (McMurry and Friedlander, 1979; Kerminen et al., 2018). The observations that many urban areas display high numbers of NCA particles have been puzzling because aerosol in this size range has generally been associated with NPF processes, which are unexpected due to high CS in urban areas.

Simultaneously, an evidence has been mounting that the exhaust of road vehicles often contains high numbers of particles in the nucleation mode size range (5–50 nm) (Kittelson, 1998) and, recently, that traffic is a direct source of NCA-sized particles (Rönkkö et al., 2017). A recent study by Yao et al. (2018) presented data of high NCA concentrations in a highly polluted urban area, with an interpretation that they are formed in a regional NPF process. Here, this view is contrasted and complemented by presenting data from a one-month measurement campaign performed in May 2017 at curbside of a densely trafficked street in an urban area of Helsinki, Finland. NCA concentration data from this curbside measurement have already been analyzed by Hietikko et al. (2018) with the conclusion of traffic inducing a dominant signal on NCA concentrations, according to diurnal variation and wind direction. Here, we extend the analysis with the data of H_2SO_4 concentrations and solar irradiances (SI) to distinguish interfering processes of traffic and regional NPF on NCA concentrations.

Prescribed primary emissions in current regional and global aerosol models do not include particles in the smallest size range (Paasonen et al., 2016). The NCA-sized particle concentrations in models are therefore mainly driven by photochemical NPF processes, omitting the directly emitted NCA-sized particles. Due to the unknown chemical composition of the traffic-originated NCA-sized particles, significant NCA-related health risks cannot be excluded. Especially for solid NCA, their behavior inside the body, such as penetrating directly into brains through the olfactory nerve (Maher et al., 2016), may have hitherto unknown adverse effects.



2 Methods

2.1 Measurement site

The measurement site was located at a street canyon at Mäkelänkatu about 3 km north of the city center of Helsinki, Finland (Fig. 1). The devices for gas and aerosol measurements were installed in two containers next to each other (Fig. 2) located at the curbside of the street canyon.

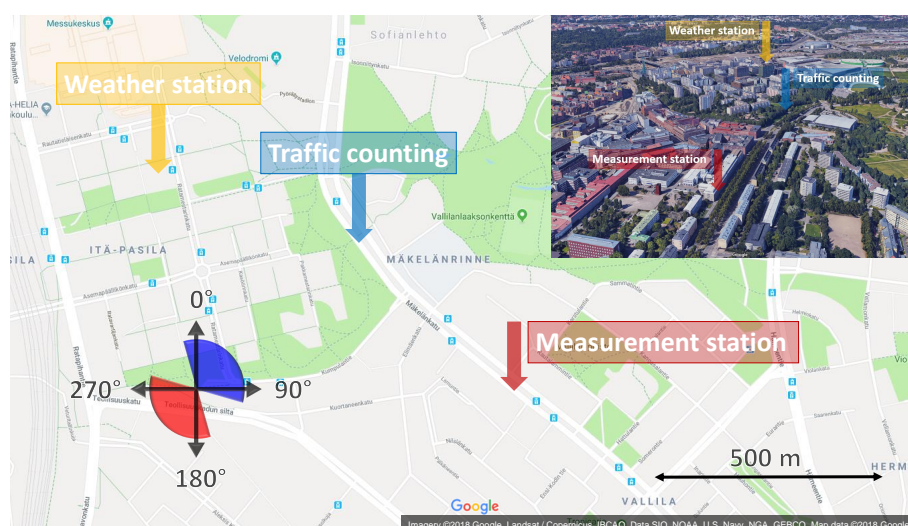


Figure 1. The map of the measurement site. Red and blue sectors denote the wind directions which result in the flow field coming from the street direction and from the background direction, respectively, towards the measurement container.

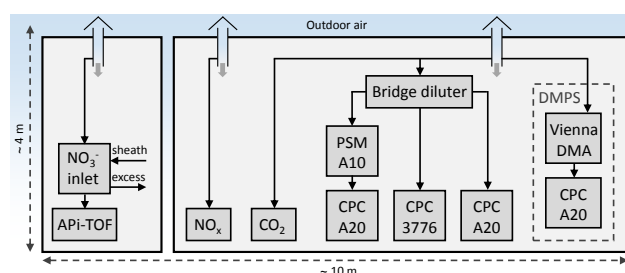


Figure 2. The measurement setup inside the containers at the curbside of the street canyon.

Traffic count was measured by the City of Helsinki at the same street but 600 m north of the measurement containers. Environmental parameters, such as wind velocity, wind direction, SI, air temperature, pressure, relative humidity, and precipitation, were measured at a weather station on the rooftop of a 53 m high building 900 m northwest of the measurement containers.



The location of the weather station provided measurement data which are undisturbed by other buildings but the location was still sufficiently near to the measurement containers to provide representative values.

The street canyon consists of three lanes for cars to both directions, two rows of trees, two tramlines, and two pavements, resulting in total width of 42 m and height of 17 m (Kuuluvainen et al., 2018). Due to the vortex affecting the flow field in a street canyon, the wind direction at the measurement containers was considered opposite to the direction above the roofs (Ahmad et al., 2005). Therefore, the wind direction diagram in Fig. 1 is mirrored by the street canyon axis. However, the street canyon in this case is not a regular street canyon but a wide avenue canyon and it has a displaced building near the measurement location. This can cause some errors to the actual flow field interpreted using the mirrored wind direction, and the effect of wind direction on the measured emissions is not seen as clearly as in an open environment or in a regular street canyon.

The measurement setup inside the containers is shown in Fig. 2. Outdoor air samples to the measurement devices were drawn through the roof of the containers 4 m above the ground, using vertical probes having diameters of 50 mm and flow rates higher than 200 lpm to minimize losses onto the walls of the sampling lines.

2.2 Sulfuric acid measurements

H₂SO₄ was measured in the gas phase using a nitrate-ion-based (NO₃⁻-based) chemical ionization atmospheric pressure interface time-of-flight mass spectrometer (CI-API-TOF; Aerodyne Research Inc.; USA and ToFwerk AG, Switzerland; Jokinen et al., 2012). It consists of a chemical ionization (CI) inlet (Eisele and Tanner, 1993) and an API-TOF mass spectrometer (Junninen et al., 2010).

The CI-inlet was operated by ionizing small concentration of nitric acid (HNO₃) vapor in the sheath air using X-ray to produce NO₃⁻ ions. The sheath air flow rate to the CI-inlet was 20 lpm and it was generated in two ways: during the first two weeks, a small pump followed by an HEPA filter was used; and during the last two weeks, an oil-lubricated compressor followed by a coarse particle, an oil, a water droplet, an HEPA, and an activated carbon filters was used. The excess flow from the CI-inlet using a vacuum pump had a flow rate of 30 lpm, resulting in the sample flow rate of 10 lpm to the inlet.

H₂SO₄ is detected in the CI-API-TOF as bisulfate ions (HSO₄⁻) and as HSO₄⁻ ions clustered with HNO₃ through the following reactions:



where $n = 0, 1, \dots$ The H₂SO₄ concentrations are calculated with the equation:

$$[\text{H}_2\text{SO}_4] = \quad (1)$$

$$\frac{C}{P} \cdot \frac{\{\text{HSO}_4^-\} + \{\text{HSO}_4^- \cdot \text{HNO}_3\} + \{\text{HSO}_4^- \cdot \text{H}_2\text{SO}_4\}}{\{\text{NO}_3^-\} + \{\text{NO}_3^- \cdot \text{HNO}_3\} + \{\text{NO}_3^- \cdot (\text{HNO}_3)_2\}}$$



where C is the calibration coefficient for H_2SO_4 , P is the penetration efficiency of H_2SO_4 in the sampling lines, and the $\{\}$ -braces denote the areas of the peaks at corresponding mass-to-charge ratios in the high-resolution spectra measured by the TOF mass spectrometer. The calibration coefficient was determined by generating known concentrations of H_2SO_4 using the oxidation of SO_2 by OH radical (Kürten et al., 2012). The calibration coefficient was determined for the both sheath air generations: the values are $C = 1.3 \times 10^9 \text{ cm}^{-3}$ for the pump-based sheath air and $C = 9.1 \times 10^9 \text{ cm}^{-3}$ for the compressor-based sheath air. The values differ due to different purities of the sheath air.

The diffusional losses (Brockmann, 2005) of H_2SO_4 in the sampling lines were calculated with the diffusion coefficient of $0.071 \text{ cm}^2/\text{s}$ representing the diffusion coefficient of a hydrated H_2SO_4 molecule in the relative humidity of 60 % and temperature of 283 K (Chapman and Cowling, 1954; Hanson and Eisele, 2000). The calculated penetrations are $P = 0.30$ when pump-based sheath air was used and $P = 0.22$ when compressor-based sheath air was used. These values differ because minor changes to the sampling lines were also done when the compressor was installed.

The H_2SO_4 concentrations from zero measurements are subtracted from the measured H_2SO_4 concentrations. The zero measurements were done by using the sheath air as a sample to obtain the H_2SO_4 concentration originated from the sheath air generation. The H_2SO_4 concentrations during the zero measurements were $3.7 \times 10^5 \text{ cm}^{-3}$ with the pump-based sheath air and $1.8 \times 10^6 \text{ cm}^{-3}$ with the compressor-based sheath air. Due to the limitations of space inside the containers, higher level of purification for the sheath air was not available.

The data from the APi-TOF was recorded with the time resolution of 2 s, but at least 1 min of the raw data is required for averaging to obtain feasible high-resolution spectra.

2.3 Gas measurements

Nitric oxide (NO) and nitrogen dioxide (NO_2) concentrations were measured using Horiba APNA-370 and the data were recorded with a time resolution of 1 min. In this study, only the sum of NO and NO_2 concentrations, denoted as NO_x concentration, is used in the analysis. Carbon dioxide (CO_2) concentration was measured using LI-COR LI-7000 analyzer with a time resolution of 1 s.

The NO_x concentration is used to represent the concentration of the traffic-originated emissions overall, because it correlates well with the traffic density on weekdays (Fig. 3). However, on weekends, much stronger dilution conditions during daytime compared to nighttime are seen. During nighttime, there is a peak in the NO_x concentration though it is nonexistent in traffic density, which suggests stagnant weather conditions coincided at nighttime on weekends for the considered time range. The CO_2 concentration would represent the amount of combusted fuel better than the NO_x concentration; however, because traffic does not cause a clear signal on the measured CO_2 concentration due to a high and varying CO_2 background level, the NO_x concentration was selected to represent the traffic-originated emissions overall. The NO_x concentrations were higher during the morning rush hours than during the afternoon rush hours on weekdays although the traffic density behaved oppositely, which occurs because, during the morning rush hours, traffic was concentrated on the same side of the street as the measurement containers located providing a shorter distance, and thus less dilution, for the emissions to travel to the measurement devices.

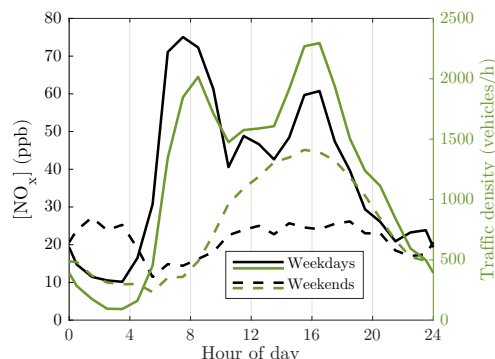


Figure 3. Average diurnal variations of nitrogen oxides (NO_x) concentration, representing the concentration of the traffic-originated emissions overall, and traffic density. The NO_x concentration data are firstly averaged to 1 h time resolution; secondly, the data from different days are averaged geometrically due to the logarithmic nature of the concentrations.

2.4 Particle measurements

The number concentration of particles with the diameters larger than approximately 1.2 nm were measured using an Airmodus A10 Particle Size Magnifier (PSM A10) (Vanhanen et al., 2011) with a diethylene glycol saturator flow rate of 1.3 lpm followed by an Airmodus A20 Condensation Particle Counter (CPC A20). Particles larger than 3 and 7 nm were measured using a TSI 3776 Condensation Particle Counter (CPC 3776) and another Airmodus CPC A20, respectively. The particle size distribution between 6 and 800 nm was measured using a Differential Mobility Particle Sizer (DMPS) consisting of a Vienna-type Differential Mobility Analyzer (DMA) followed by a CPC A20. Due to high particle concentrations at the street canyon, the sample to CPCs was diluted using a bridge diluter having a dilution ratio of 8.2. The dilution ratio for the specific diluter is, however, measured for larger particle sizes only, and because the diluter is based on diffusional losses of the particles, the dilution ratio for NCA-sized particles is higher. Therefore, the NCA concentrations reported here represent the lower limits of the actual concentrations.

The number concentration of nanocluster aerosol (NCA), particles within the diameter range between 1.2 and 3 nm, can be calculated by subtracting the concentration measured by the CPC 3776 from the concentration measured by the PSM. The particle size distribution between 1.2 nm and 800 nm can be calculated with the data from all these aerosol measurement devices by taking the cut diameters of the CPCs and the dilution ratio of the bridge diluter into account. The NCA concentration was measured with a time resolution of 1 s and the size distribution with a time resolution of 9 min.

3 Results and discussion

The data from the off-site measurements of traffic count and environmental parameters are available for the whole four-week measurement campaign starting on 4 May 2017 and ending on 31 May 2017 (see the Supplement for the time series). This



time range provided adequate data for examining NCA and H_2SO_4 formation respect to solar irradiance, because there were sufficient amounts of days both with clear sky and with cloud cover, yet without too many rainy days. There are some gaps in the NCA and H_2SO_4 data during the four weeks due to unavailability of the measurement devices. The data analysis considers only the time ranges for which all the measurement data are available, resulting in three weeks of data.

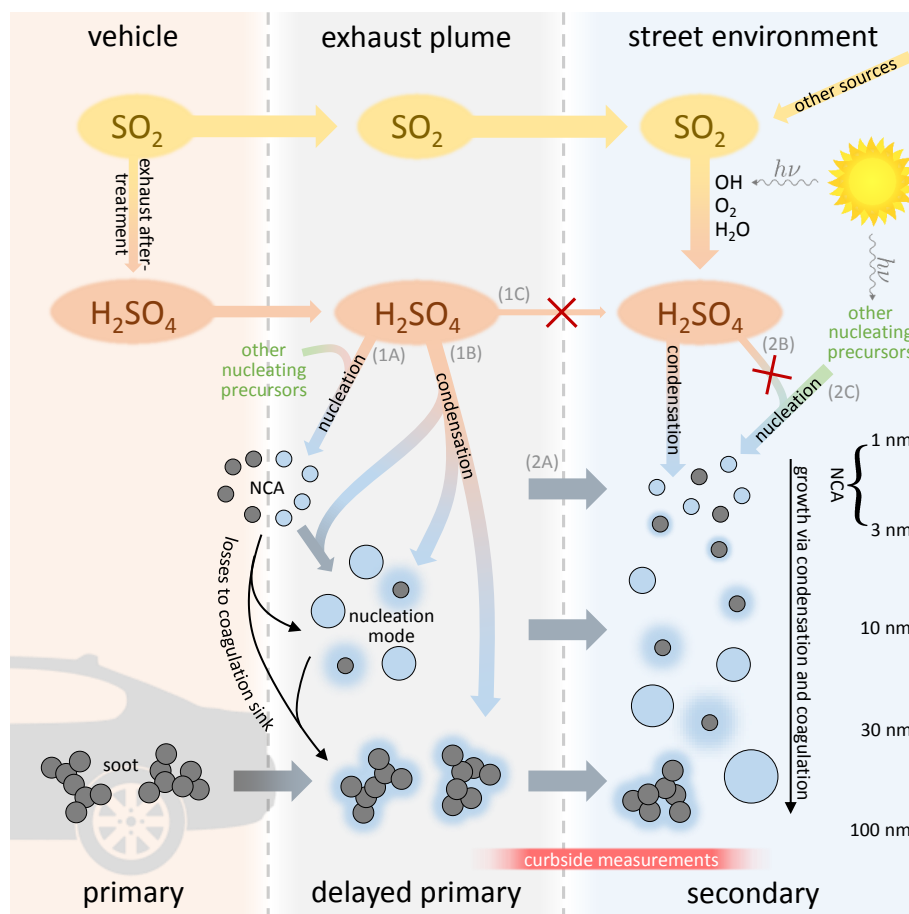


Figure 4. Proposed mechanism of sulfuric acid and particle formation in traffic-influenced areas. The route of primary H_2SO_4 to urban atmosphere (route 1C) is largely terminated to particle phase rapidly after the emission (routes 1A and 1B). Conversely, secondary H_2SO_4 remains in urban atmosphere because it does not experience such a rapid gas-to-particle conversion. NCA presence in urban atmosphere is majorly controlled by traffic emissions (route 2A) and only marginally by nucleation from secondary precursors (routes 2B and 2C), especially the contribution of nucleation from secondary H_2SO_4 (route 2B) is noticeably overridden by traffic emissions (route 2A).

- 5 Figure 4 presents our proposal for the updated mechanism of H_2SO_4 and particle formation in traffic-influenced areas, based on our measurement results. The most noteworthy details are illustrated with red crosses indicating H_2SO_4 routes which were observed to occur barely only, or not at all. As shown later in this section, our measurement at the curbside displays no clear increase in gaseous H_2SO_4 concentrations with increasing traffic volumes, indicating that the majority of primary H_2SO_4 is



- converted to the particle phase via nucleation and condensation rapidly after emission. Conversely, secondary H_2SO_4 remains longer in urban atmosphere because it does not experience conditions favoring such a rapid gas-to-particle conversion, i.e., rapid temperature decrease, high precursor concentrations, and high pre-existing CS. Therefore, the signal of H_2SO_4 measured from the curbside of the street is mainly due to secondary H_2SO_4 only.
- 5 Our results show that both traffic and regional NPF influence NCA concentrations at the urban traffic site, with direct NCA emission from traffic dominating. Comparison of the NCA and H_2SO_4 concentrations with SI and traffic density suggests that while solar radiation favors higher NCA concentrations, the photochemically produced H_2SO_4 may not be the key compound in the presence of NCA in urban areas. Traffic-originated NCA particles may be formed via a delayed primary emission route by rapid nucleation of low-volatile gaseous compounds emitted by vehicles during exhaust cooling after releasing from the
- 10 tailpipe. On the other hand, they may be solid particles emitted directly by engines, via a primary emission route (Sgro et al., 2012; Alanen et al., 2015).

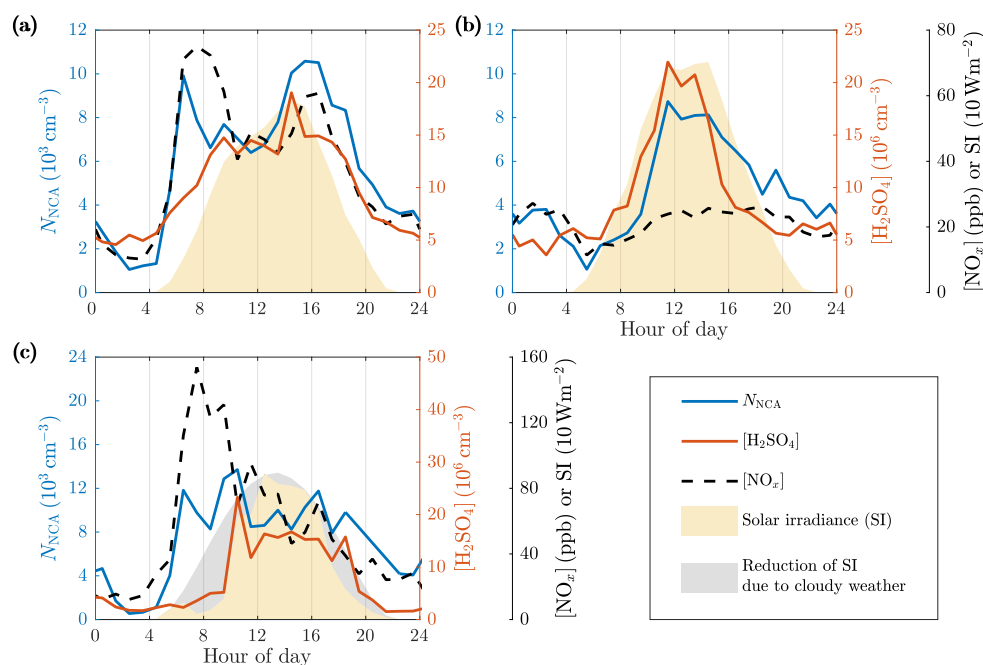


Figure 5. Average diurnal variations of the NCA, H_2SO_4 , and NO_x concentrations and solar irradiance (SI) on (a) weekdays, (b) weekends, and (c) example day 5/22/2017 with cloudy morning and evening. The data are firstly averaged to 1 h time resolution; secondly, the data from different days are averaged geometrically (except SI) due to the logarithmic nature of the concentrations of the emissions.

The first evidence for traffic-contributed concentrations of NCA and H_2SO_4 was found in the diurnal variations of the NCA, H_2SO_4 , and NO_x concentrations and SI (Fig. 5). The diurnal variations on weekdays (Fig. 5a) differ from the diurnal variations



on weekends (Fig. 5b). The main difference between weekdays and weekends are traffic volumes; therefore, such a difference in the concentrations of NCA and H_2SO_4 should only be expected if their formation are in some manner connected to traffic. The connection of NCA to traffic is further strengthened by comparing it to the NO_x concentrations, which are directly linked to traffic densities and traffic-related emissions (Fig. 3). On weekdays, the NCA concentration increased in tandem with the NO_x concentration during the morning rush hours. On weekends, the NCA concentration increased at noontime much more clearly than the NO_x concentration. This can be interpreted as a sign of an ongoing regional NPF process producing NCA particles with high SI. On weekdays, the regional NPF process should only produce higher NCA concentration during afternoon rush hours having higher SI compared to morning. The increased NCA concentrations during the morning rush hours suggest that the traffic-originated NCA does not require solar radiation to form.

We observed that, traffic levels influence the H_2SO_4 concentrations, but they are still mainly controlled by solar radiation. In contrast to the NCA concentration, the H_2SO_4 concentration traced SI much more closely, with a maximum at noontime and minimum at night. On weekdays, a peak in the H_2SO_4 concentration during afternoon rush hours suggests that traffic might also influence the formation of H_2SO_4 . Further evidence for this is found by comparing the diurnal variation of H_2SO_4 between weekdays and weekends. On weekends, the H_2SO_4 concentration increased not until the traffic density and the NO_x concentration were also increased, whereas on weekdays, the traffic density was already high when SI and the H_2SO_4 concentration began to increase. Furthermore, higher irradiances were required on weekends before the rise in the H_2SO_4 concentration and, additionally, the order of the increase in the NCA and H_2SO_4 concentrations was switched.

The time series show that NCA is not similarly controlled by solar radiation but rather by traffic. This is clearly showcased in Fig. 5c which presents data from a day with cloudy weather reducing SI in the morning and in the evening but still with a constant wind direction. The NCA concentration closely traced traffic levels in the morning, whereas the H_2SO_4 concentration only increased when SI increased hours later. This clearly shows that the formation of NCA, in this case, is independent of SI and the H_2SO_4 concentration. It is also noteworthy that no increase in the NCA concentration is observed when SI increased, suggesting that traffic dominated in the NCA formation.

The data suggest that the formation of atmospheric H_2SO_4 is strongly enhanced in the presence of both traffic and sunlight. While a strong correlation between the NCA and NO_x concentrations (Fig. 6a, b: Pearson's $R = 0.84$) confirms the connection between NCA and traffic, a remarkably weaker, but also positive, correlation between the H_2SO_4 and NO_x concentrations ($R = 0.50$) was observed, revealing the connection between H_2SO_4 and traffic. The effect of SI at different traffic densities (data averaged to 17 different SI bins for three different NO_x concentration ranges) shows differing patterns for NCA (Fig. 6c) and H_2SO_4 (Fig. 6d). While high SI is associated with higher NCA and H_2SO_4 levels, traffic density determines the base level for both (the concentrations at zero SI). For H_2SO_4 , the influence of traffic causes a marked increase in the slope of the H_2SO_4 concentration-SI-line. The slope can be interpreted as the production efficiency of H_2SO_4 via photochemistry. It is evident that for NCA, the influence of traffic dominates in comparison to SI, as the traffic-influenced NCA concentration (red data) exceeds the non-traffic concentrations (black data) even during dark times. For H_2SO_4 , the situation is different, as all dark-period H_2SO_4 concentrations are close to equal levels. These differing patterns suggest that the majority of NCA in traffic-influenced areas is formed independently of secondary H_2SO_4 , in contrast to the findings of Yao et al. (2018).

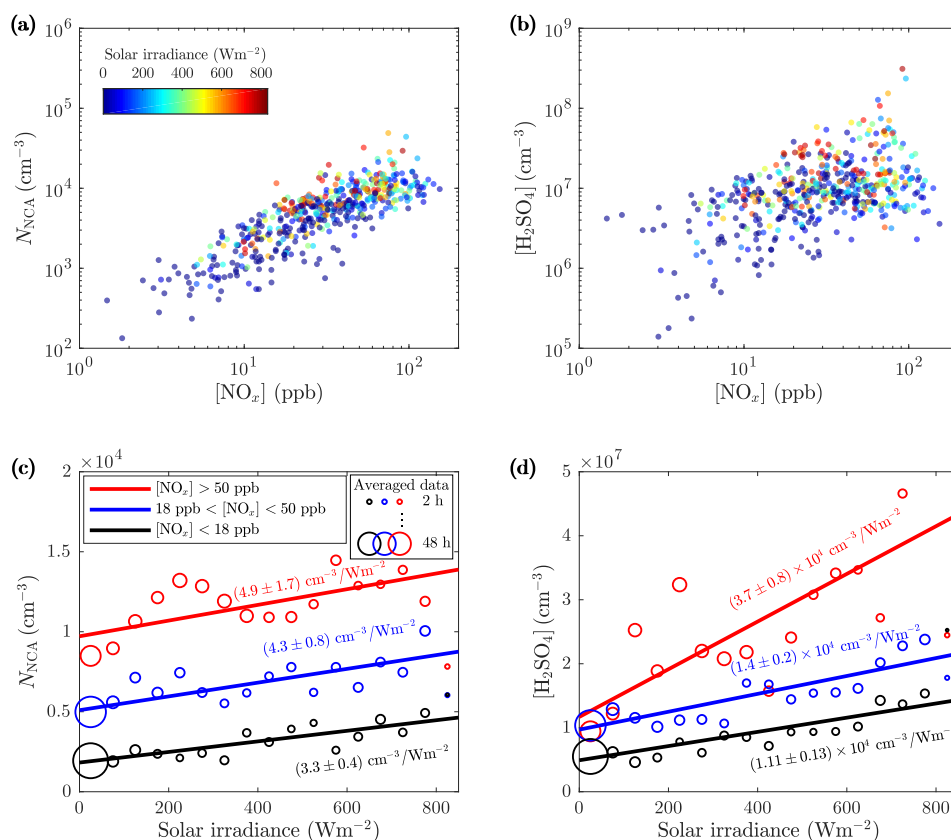


Figure 6. One-hour-averages of the (a) NCA and (b) H_2SO_4 concentrations as a function of the NO_x concentration colored by the solar irradiance (SI) and 10-min-averages of the (c) NCA and (d) H_2SO_4 concentrations further averaged to SI bins for three NO_x concentration ranges. The slopes of the linear fits are marked in the figure. The amount of the averaged data to specific bins are shown with the radii of the circles and are also used as weighting factors for the linear fitting.



Even more compelling evidence for traffic-originated NCA and H_2SO_4 can be found by comparing the observed concentrations to momentary CO_2 concentrations. Such a comparison for measurements near emission sources has been shown to directly represent a fuel-based emission factor using the emission factor of CO_2 , 3.14 kg per 1 kg of fuel combusted (Yli-Tuomi et al., 2005). In Fig. 7, no apparent difference in the emission factors of NCA for different SI are seen; however, in the case of H_2SO_4 , higher SI lead to noticeably higher emission factors of H_2SO_4 . We tested for potential co-correlations between SI and traffic density to examine potential traffic level increase with simultaneous SI increase due to their almost similar diurnal cycles. We found no correlation between CO_2 concentration and SI (Fig. 8) that would have led to the misinterpretation of NCA and H_2SO_4 formation via traffic instead of solar radiation. This result again supports the finding that solar radiation is required for the formation of H_2SO_4 from traffic emissions and demonstrates clearly that both NCA and H_2SO_4 originate from traffic. This is further supported by examining the concentrations in different wind directions (Figs. 9 and 10) which shows that the highest concentrations are measured when the wind blew from the street.

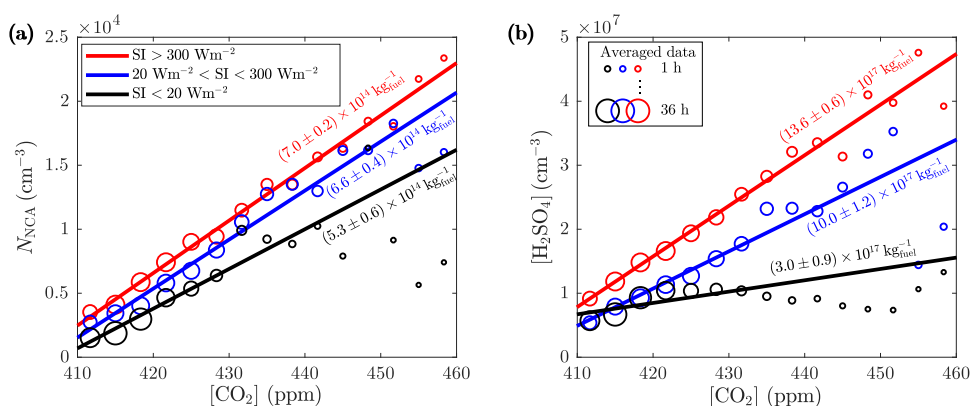


Figure 7. 1-min-averages of the (a) NCA and (b) H_2SO_4 concentrations further averaged to CO_2 concentration bins for three SI ranges. The slopes of the linear fits converted to kilograms of fuel combusted are marked in the figure. The amount of the averaged data to specific bins are shown with the radii of the circles and are also used as weighting factors for the linear fitting.

The annual CO_2 emission rate from traffic in Helsinki in 2017 was $5.38 \times 10^8 \text{ kgCO}_2 \cdot \text{a}^{-1}$ (VTT Technical Research Centre of Finland Ltd, 2017). Using the average NCA emission factor versus CO_2 emission, $2.21 \times 10^{14} \text{ kgCO}_2^{-1}$, the annual NCA emission from traffic in Helsinki becomes $1.19 \times 10^{23} \text{ a}^{-1}$. The annual NCA formation rate via photochemical NPF in Helsinki can be approximated using estimates of nucleation rate, from 1 to $10 \text{ cm}^{-3}\text{s}^{-1}$, NPF event day count per year, from 30 to 120 a^{-1} , NPF duration, from 2 to 4 h, measured in a rural area in Hyytiälä, Finland (Dal Maso et al., 2005; Kulmala and Kerminen, 2008) and in an urban area in Helsinki (Hussein et al., 2008), the total area of Helsinki, 214 km^2 , and a rough estimate for the boundary layer height, 500 m. Multiplying these gives the annual NCA formation rate from 0.23×10^{23} to $18.5 \times 10^{23} \text{ a}^{-1}$. Comparison of these annual rates suggests that in minimum of 6 % but even up to 84 % of NCA particles originates from traffic in Helsinki on an annual basis. Although this range is wide, the contribution of traffic-originated NCA is significant.

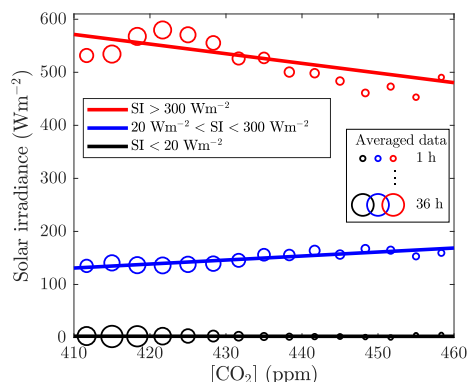


Figure 8. 1-min-averages of the solar irradiances (SI) as a function of CO₂ concentration for the SI ranges used in Fig. 7. The amount of the averaged data to specific bins are shown with the radii of the circles and are also used as weighting factors for the linear fitting.

Considering typical weekday noontime at our measurement location and assuming the annual mean of the daytime maximum SI in Helsinki, 500 Wm⁻², the NCA concentration due to traffic is $\sim 9.5 \times 10^3 \text{ cm}^{-3}$ (the value of the high NO_x line at zero irradiance in Fig. 6) and the increase of the NCA concentration due to photochemistry is $\sim 1.7 \times 10^3 \text{ cm}^{-3}$ (calculated with the slope of the low NO_x line in Fig. 6). These concentrations indicate that $\sim 85\%$ of the total NCA concentration at the street canyon is originated from traffic at noontime. Considering midsummer and midwinter, the daytime maximum SI are 850 Wm⁻² and 100 Wm⁻², giving the contributions of $\sim 78\%$ and $\sim 97\%$, respectively. Therefore, it is evident that the major fraction of NCA originated from traffic at our measurement location, even with the highest available SI values in midsummer.

For H₂SO₄, the concentration due to traffic at our measurement location at typical weekday noontime is $\sim 12 \times 10^6 \text{ cm}^{-3}$ and the increase of the concentration due to photochemistry is $\sim 5.6 \times 10^6 \text{ cm}^{-3}$, indicating 68 % of the total H₂SO₄ concentration at the street canyon originating from local traffic at noontime. For midsummer and midwinter, the contributions become 56 % and 92 %, respectively. These values signify that also the major fraction of H₂SO₄ originated from traffic even though it cannot be seen as clearly from the diurnal variation as is seen in the case of NCA.

Our data clearly demonstrate that NCA-sized particle concentrations in a traffic-influenced environment is controlled by NCA directly emitted by traffic. The data also demonstrate that while generally NCA and photochemically produced nucleating vapor concentrations correlate, this correlation is likely, firstly, due to increased traffic volumes at daytime and, secondly, due to traffic-originated H₂SO₄ and other nucleating vapors. We also showed that H₂SO₄ formation is driven by both solar radiation and a traffic-related source.

4 Conclusions

Our results have several implications on our understanding of aerosol particle formation in traffic-influenced areas. Firstly, because current regional and global air quality models do not include particles in the sub-3 nm size range as primary emissions

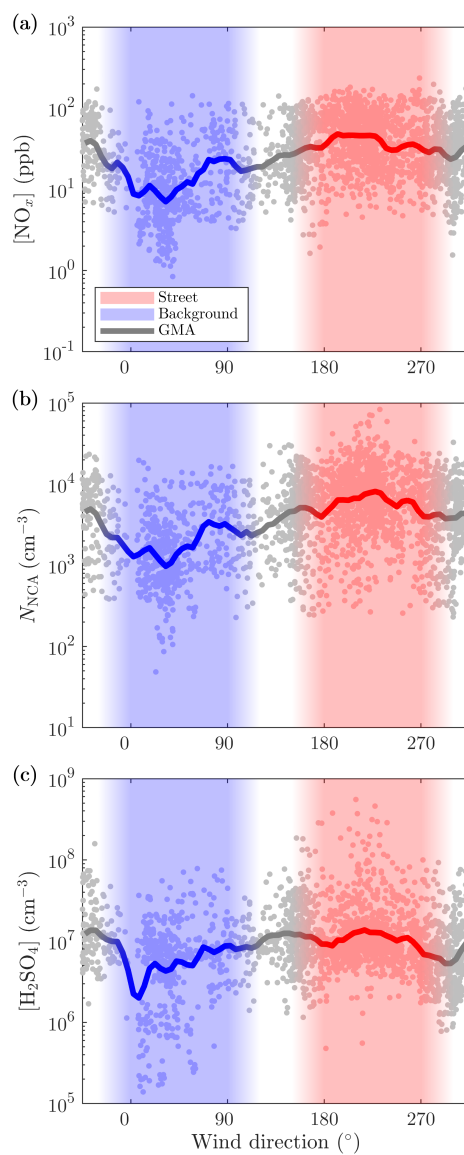


Figure 9. 10-min-averages of the (a) NO_x , (b) NCA, and (c) H_2SO_4 concentrations measured with different wind directions. Wind velocities smaller than 0.5 m/s are excluded. GMA denotes geometric moving average.

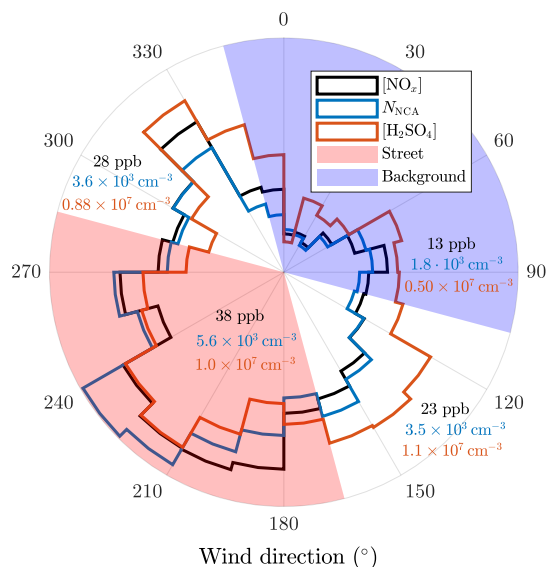


Figure 10. 10-min-averages of the NO_x, NCA, and H₂SO₄ concentrations further averaged to different wind direction sectors. Wind velocities smaller than 0.5 m/s are excluded. The geometric averages of the concentrations across the sectors are shown in the figure.

(Paasonen et al., 2016), the modelled NCA-sized particle concentrations are mainly driven by photochemical NPF processes, neglecting their origin from traffic as primary sources. Thus, our results show an urgent need to update these emissions. In light of our results, it seems evident that there will be areas in which direct emissions dominate the formation of new aerosol. A rough calculation gives that, on an annual basis, up to 84 % of NCA can originate from traffic in Helsinki; and according to the measured NCA concentrations, on typical weekday noontime, ~85 % of the total NCA concentration was contributed by traffic at our studied site. In wintertime, this contribution may reach ~97 % due to lower SI, which highlights the need for updating the annual particle formation cycles in the models. Secondly, our results also show that both traffic emission and regional NCA formation signals can be distinguished for the most of the times, and that traffic also influences the formation of H₂SO₄. Together with the findings of Yao et al. (2018), this presents a significant update on the particle formation mechanisms in urban areas. As illustrated in Fig. 4, the particle concentration is controlled by the interplay of the two processes, with varying importance depending on the proximity of the emission source. Our results call for reconsideration and re-analysis of observations of NPF events observed in traffic-influenced areas. In many cases, there is covariance between traffic volumes and SI, and care should be taken to separate these two variables in the analysis, e.g., by considering CO₂ or NO_x as tracers for traffic volumes. Finally, potential health effects of traffic emissions in urban areas should also be considered more carefully because the composition of the emitted NCA particles is still unknown, especially as some clues for their non-volatility exist.



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