Cyclists’ exposure to air pollution: in situ evaluation with a cargo bike platform

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Received: 22 December 2019 / Accepted: 21 June 2020 / Published online: 29 June 2020
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Abstract Cyclists’ exposure to air pollutants near roadways has been associated with numerous health effects. While the adverse health effects concerning aerosols have traditionally been assessed with data of particle mass concentrations, it appears that the number concentration is also another important indicator of toxicity. Thus, to holistically evaluate one’s exposure to aerosol particles, assessments should be based on mass concentrations and number concentrations. In order to assess individual cyclists’ exposure as they move through space and time, spatiotemporal high-resolution approaches are needed. Therefore, a mobile, fast-response monitoring platform was developed that uses a cargo bicycle as a base. Data of particle mass concentrations (PM₁, PM₂.₅, PM₁₀) and particle number concentrations (PN₁₀) were collected along two different routes, one characterized by high-intensity vehicle traffic and one by low-intensity vehicle traffic. While high spatiotemporal heterogeneity was observed for all measured quantities, the PN₁₀ concentrations fluctuated the most. High concentrations of PN₁₀ could be clearly associated with vehicle traffic. For PM₂.₅, this relation was less pronounced. Mean particle concentrations of all measures were significantly higher along the high-traffic route. Comparing route exposures, the inhalation of PM₂.₅ was similar between both routes, whereas along the high-traffic route, cyclists were exposed to twice the particle number. We conclude that the cargo bike, featuring high-frequency mobile measurements, was useful to characterize the spatial distribution of mass concentrations and number concentrations across an urban environment. Overall, our results suggest that the choice of route is a key factor in reducing cyclists’ exposure to air pollution.

Keywords Cargo bike · Exposure assessment · Particle number concentration · Particle mass concentration

Introduction

Bicycling is—besides walking—widely accepted as the most environmentally friendly means of transport since it does not contribute to air or noise pollution nor to any extra emission of carbon dioxide. Cycling also provides many health benefits such as improving cardiovascular health, muscular strength, and control of blood sugar (Bauman and Rissel, 2009, Kelly et al., 2014, Oja et al., 2011). However, cycling has also been related to many negative health issues due to cyclists’ exposure to high levels of air pollutants and noise (Boogaard et al., 2008) when traveling on roads shared with motor vehicles or on routes adjacent to or near main roads (Badland and Duncan, 2009; De Hartog et al., 2005; Kaur et al., 2007). The typical exposure of cyclists is relatively

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https://doi.org/10.1007/s10661-020-08443-7
short, but the dose is high due to their higher breathing activity compared with those of car or bus passengers (Boogaard et al., 2008, Brand et al., 2019). Consequently, cyclists’ exposure to both gaseous and particulate air pollutants near roadways has been associated with numerous health effects including asthma exacerbation, other respiratory illnesses, and excess risk of mortality from cardiopulmonary disease and stroke (Apparicio et al., 2016; Gordian et al., 2006).

Traditionally, the adverse health effects of particulate matter (PM) have been associated with PM$_{10}$ or PM$_{2.5}$ particles, i.e., the mass concentrations of PM with aero-dynamic diameters below 10 µm and 2.5 µm, respectively. Yet, the specific health effects of ultrafine particles with an aerodynamic diameter ≤0.1 µm (UFP), which do not contribute much to the PM$_{2.5}$ or PM$_{10}$ mass concentrations, are still under discussion. For example, in 2013, the Health Effects Institute concluded that the state of science was inconclusive regarding the toxicity of UFPs due to inconsistencies and limitations in the findings from both short-term and long-term studies (EPA, 2015). While some studies state that the total particle number concentration in air, which is a good indicator of UFP, remains a poor indicator of toxicity markers (Xue et al., 2019), others suggest that UFPs cause a greater lung inflammatory response than is caused by equal mass concentrations of larger particles (Frampton, 2016). Similarly, Brown et al. (2001) and Oberdorster (2001) reported that UFP exhibited greater pro-inflammation activity per mass than the same mass of larger particles. Overall, it appears that the mass concentration of UFP exhibits fewer toxic effects than the number concentration of UFP (Penttinen et al., 2001; von Klot et al., 2002; Ferin et al., 1992, Hosiokangas et al., 1995). Consequently, we hold the view that any evaluation of cyclists’ exposure to aerosol particles should be based both on mass concentration (PM$_{10}$, PM$_{2.5}$) and on number concentration (PNC) data.

Within cities, motor traffic has been identified as a major contributor to the near-road aerosol strain (Karagulian et al., 2015; Manousakas et al., 2017) due to both exhaust and non-exhaust sources (Ketzel et al., 2007; Amato et al., 2014). Raised levels of particle concentrations mostly occur within street canyons (Wurzler et al., 2016). Furthermore, there are diffuse sources of aerosol particles such as resuspension from dry, grassy, and unpaved footpaths containing loose and dry top coating material (Birmili et al., 2013; Paas et al., 2016). Furthermore, Birmili et al. (2013) identified cooking activities and smoking in outdoor restaurant seating areas to be important sources of PM. The combination of various particle sources and the influence of meteorological conditions on the transport of aerosols in the urban boundary layer pose a challenge to determining aerosol concentration patterns (Venkatachari et al., 2006).

An increasing number of studies have used simulation approaches (Baxter et al., 2010) or land use regression and related models (Gryparis et al., 2007) to estimate the spatiotemporal distribution of air pollutants for use in epidemiological studies. While these studies have provided insight into exposure patterns and health effects, they lack sufficient spatial resolution to capture the adverse exposure dose on the micro scale. Furthermore, some of the studies provided significant misclassifications of exposure, which tends to obscure true correlations between air pollutant concentrations and potential health effects (Baxter et al., 2010; Sarnat et al., 2010).

When it comes to individual exposure, the situation is even more complex for subjects moving through space and time (Kousa et al. 2002). With each breath of air, another air mass, with potentially much different concentrations from the earlier or later breath, respectively, is inhaled. In the Netherlands, a tool has been established to assess local cycling conditions, and this tool also includes the evaluation of the current air quality (Boogaard et al., 2008). Commuters can use the spatial information on air pollution to avoid high exposure by choosing less-polluted routes. In addition, urban planners and policymakers can make use of the detailed spatial information to identify local hotspots and to take measures against local air pollution.

However, to evaluate local air pollution, such tools primarily use stationary air quality monitoring networks that only report 30-min or 60-min averages. These networks measure—following legal requirements—particle mass concentrations (PMC, either PM$_{10}$ or PM$_{2.5}$) but not PNC. Furthermore, such stations cannot account for the high variability of air pollutant concentrations in urban environments. For these reasons, no reliable assessment of an individual cyclist’s exposure can be made from the respective data. In fact, the exposure to air pollutants in traffic, which is found to be generally enhanced, is, in many cases, only poorly correlated with simultaneous measurements at stationary monitoring sites (Kaur et al., 2007). Therefore, there is
a clear need for measurements of air pollutants to be taken at much better spatial and temporal resolutions of 1 m and 1 s, respectively, in order to evaluate cyclists’ exposure.

During the last decade, mobile monitoring platforms enabling personalized and high-resolution monitoring have become increasingly popular (Broich et al., 2012; Birmili et al., 2013; Van den Bossche et al., 2015; Dewulf et al., 2016, Gerike et al., 2016). Especially for atmospheric pollutants that experience large spatiotemporal variation in urban environments, mobile monitoring provides valuable insights into the spatial distribution of air pollution that cyclists and pedestrians are exposed to in everyday life. However, to the best of our knowledge, no experimental study has simultaneously characterized the spatiotemporal patterns of both particle mass concentration and particle number concentration in an urban environment. Thus, we used a mobile cargo bike platform to a) assess the potential of real-time PMC and PNC monitoring to measure the exposure of cyclists who commute daily in an urban environment, b) to provide insight into the spatiotemporal variability of fine and ultrafine particle concentrations, and c) to quantify the potential human health risk due to exposure to fine particles depending on the traffic density of the route chosen.

Methods

Site description

This study was carried out in Münster, NW Germany (52.0° N, 7.6° E), a city with approximately 300,000 inhabitants (Fig. 1). It is characterized by flat terrain and hosts some small- to medium-sized industries. The city is well documented in terms of traffic flow, and there is a good understanding of the underlying processes driving air pollution (Gietl and Klemm, 2009). The regional impact of air pollution originates from long-range transport, mainly from the Ruhr industrial area located 60 km to the SW, since the main wind direction is southwest, as determined over the period from 1989 to 2018 (DWD, 2019), and from the intensive agricultural activity of the surrounding area (Gietl et al. 2008; Gietl and Klemm 2009). Local sources are traffic emissions, residential heating, and to a lesser degree industry and power plant emissions (Deventer et al., 2015).

Since almost 40% of daily routes are made by bike in Münster, we selected two routes with different traffic influence, crossing the city from east to west. The high-traffic route (2.45 km) goes along a main thoroughfare that links the central train station to the downtown area and eventually leads to the university campus, while the low-traffic route (2.99 km) goes through a green corridor that surrounds the downtown area, and which may only be used by bicyclists and pedestrians. The two courses were probed in the morning of July 17, 2019, between 09:50 and 10:35 local time (UTC + 2). Average synoptic conditions during data collection were a northwest (290°) wind direction and a wind speed of 1.6 m s⁻¹, as observed at the weather station at the university campus at 34 m above ground, 1800 m beeline from the downtown area (Fig. 1).

Data collection and processing

Data were collected with a cargo bike equipped with a three-dimensional sonic anemometer, type R3-50 (Gill Instruments Ltd., Lymington, Hampshire, UK). The anemometer was installed vertically on the front end of the cargo bike with the center of the sensor head being at a height of 1.47 m above ground (Fig. 2). The north arrow of the anemometer was aligned with the direction of driving to enable correction of wind direction in post-processing. The aerosol particle measurement technology included an optical particle spectrometer (OPS), type 3330, and a condensation particle counter (CPC), type 3025A (both sensors manufactured by TSI Inc., Minnesota, USA). The OPS sampled the PMC in 16 size classes (0.3- to 10-μm diameters), while the CPC sampled the total PNC of particles between 3-nm and 10-μm diameters (PN10). The upper detection limit of the CPC was 53,000 p cm⁻³, meaning that a concentration higher than 53,000 particles per cm³ could not be detected. During post-processing, the respective PMCs of individual size classes were summed up to the most common measures of PM10, PM2.5, and PM1. The air samples were taken at the front end of the cargo bike at a height of 1.1 m above ground and led through individual hoses of 0.96 m (internal diameter 5 mm) and 1.28 m (internal diameter 10 mm) lengths to the OPS and CPC, respectively. To account for the position and ground speed as well as for the orientation of the cargo bike, a GPS device, type 19x HVS (Garmin Ltd., Kansas, USA), and a digital compass module, type HMR3000 (Honeywell International Inc., North Carolina, USA),
were used (Fig. 2). All sensors were operated with 1-Hz data acquisition frequency, centrally controlled by a data logger, type CR3000 (Campbell Scientific Inc., Utah, USA).

Processing of raw data was made with R version 3.5.0 (R Core Team, 2018). The R packages ggplot2 3.0.0 (Wickham, 2016) and OpenStreetMap 0.3.4 (Fellows, 2019) were used for the cartographic display of the data.

Results

Spatiotemporal variability of aerosol particle concentrations

The concentrations of PM$_{2.5}$ and PN$_{10}$ as measured along the two cargo bike routes are shown as cartographic displays and as longitude series in Fig. 3. The two routes that lead from east to west through the city of Münster are primarily different in their influence of motor vehicle traffic. The northern route experienced high vehicle traffic (HT), whereas the southern route was influenced by low traffic intensities (LT).

The second-to-second differences can be as high as 28.5 $\mu$g $m^{-3}$ and 43,500 p $cm^{-3}$ for PM$_{2.5}$ and PN$_{10}$, respectively. Comparing the PM$_{2.5}$ concentration data from the two routes side by side, it becomes apparent that the median values are quite similar for both routes (LT 4.3 $\mu$g $m^{-3}$, HT 4.7 $\mu$g $m^{-3}$). Nevertheless, the HT route exhibited considerably more variation. The mass concentration range was between 2.8 and 7.3 $\mu$g $m^{-3}$ in the LT route and between 2.4 and 48 $\mu$g $m^{-3}$ in the HT route. The PM$_{2.5}$ concentrations did not vary largely along the LT route. However, five hotspots were identified along the HT route. Apart from these few hotspots, the impact of traffic on the observed PM$_{2.5}$ concentrations was detectable yet small. Road traffic is only a minor source of PM$_{2.5}$ compared with the long-range transport of particles.

More pronounced differences between the two routes were observed for PN$_{10}$ compared with PM$_{2.5}$. For the LT route, the particle number concentration ranged between 3400 p $cm^{-3}$ and the value for the maximum detectable concentration, 53,000 p $cm^{-3}$, (median 5400 p $cm^{-3}$), and for the HT route, it ranged between 4400 p $cm^{-3}$ and 53,000 p $cm^{-3}$ (median 17,000 $cm^{-3}$).
The spatial analysis of both the HT and LT routes concerning PN₁₀ unveils that the spatial variations of PN₁₀ concentrations were very high (Fig. 3c). Furthermore, elevated PN₁₀ concentrations could almost always be associated with areas such as street intersections and sectors around traffic lights and, thus, with motorized vehicle traffic.

A cyclist’s exposure dose on individual routes

On average, cyclists were exposed to higher PMC and PNC when biking along the HT route as compared with the LT route. The differences were much larger when comparing PN₁₀ particle number concentrations than PM₁₀ mass concentrations, and they were also larger when comparing larger rather than smaller size fractions of particles (Fig. 4).

The PMCs of PM₁₀, PM₂.₅, and PM₁ showed larger variations for the HT route than for the LT route; the variation coefficients (CV) were larger by factors of 1.34, 3.32, and 2.33, respectively. The spatial analysis revealed that variations of PM₂.₅ are weakly associated with intersections between collector or arterial roads (Fig. 3a). No such difference of CVs between the HT and the LT routes was observed for the particle number concentrations (CV HT route 63.28, CV LT route 75.04). Overall, the high variability of PN₁₀ as compared with PMC is associated with traffic emissions. The more pronounced variability of PN₁₀ along the LT route is caused by the generally lower baseline of PN₁₀ concentrations in combination with the crossing of a few street intersections, where the maximum PN₁₀ concentrations were as high as the maximum concentrations detected along the HT route (Fig. 3c and d).

Our results show much higher concentrations of PM₁₀ and PN₁₀ along the HT route than along the LT route, while levels of PM₂.₅ and PM₁ were also higher for the HT route, but to a lesser degree. In order to arrive at a quantitative estimation of cyclist’s exposition while riding along different route types, we quantify the route exposure (RE) to PM₂.₅ as follows:

\[
\text{RE}_{\mu g} = \left( \frac{\text{concentration (\mu g)}}{\text{m}^3} \right) \times \text{inhalation rate (m}^3/\text{s}) \times \text{time lapse (s)}
\]

and to the particle number p as

\[
\text{RE}_p = \left( \frac{\text{concentration (p)}}{\text{m}^3} \right) \times \text{inhalation rate (m}^3/\text{s}) \times \text{time lapse (s)}
\]

We assumed an average volumetric inhalation rate of 0.065 m³ min⁻¹ for cyclists, based on data from the Exposure Factor Handbook (Environmental Protection Agency, 2011). The overall exposure of each route was calculated with the mean mass of particles < 2.5 μm inhaled as well as the total number of particles < 10 μm inhaled as metrics to estimate the personal exposure to PM₂.₅ and PN₁₀, respectively. We found that cyclists...
Fig. 3 PM$_{2.5}$ concentrations (μg m$^{-3}$) as a) a color-coded cartographic display and as b) a longitude series, PN$_{10}$ concentrations (p cm$^{-3}$) as c) a color-coded cartographic display and as d) a longitude series. Data were collected at 1 Hz along two routes characterized by high traffic influence (northern route, light blue line) and by low traffic influence (southern route, dark blue line) in the city of Münster, Germany.
using the HT route inhaled 4.22 μg of particles < 2.5 μm, while when using the LT route, which lasted longer, they inhaled 4.86 μg of particles < 2.5 μm. Thus, even though the LT route had lower PM$_{2.5}$ concentrations, this was outweighed by its longer distance and, thus, its longer travel time (1010 s vs. 741 s). Note that a cyclist traveling on the HT route also needs to breathe air when not biking (and this air potentially contains PM$_{2.5}$ as well), such that his total exposure during the entire time period of the LT route will be somewhat larger than his calculated HT exposure. For both routes, then, the exposure to PM$_{2.5}$ is quite similar. However, when assessing the cyclists’ exposure to particle number, we found that cyclists using the HT route were exposed to twice the particle number as compared with those using the LT route (14,700 × 10$^6$ vs. 7500 × 10$^6$ particles). Thus, as this particle fraction and metric are more relevant to toxicological effects, the toxicological risk to a cyclist on the HT route is at least twice as high as the toxicological risk to a cyclist on the LT route.

**Discussion**

The overall spatiotemporal variabilities of the measured PM$_{2.5}$ and PN$_{10}$ concentrations at both sampling routes are very large. Boogaard et al. (2008) also found very high variability for 1-min averages of PM$_{2.5}$ and PNC within eleven Dutch cities. They were able to demonstrate that peaks of PNC occurred more often and more suddenly for cyclists than for car drivers with a typical peak length of less than 10 s (Boogaard et al., 2008).

However, the PM$_{2.5}$ concentrations measured at the HT and LT routes are much lower than those measured by Ham et al. (2017), who compared daily commutes using bike trails or arterial trails in Sacramento, CA (6 ± 1.5 μg m$^{-3}$ and 13.5 ± 7 μg m$^{-3}$, respectively). Boogaard et al. (2008) also found higher mean PM$_{2.5}$ concentrations (44.5 μg m$^{-3}$ for cyclists), albeit the variability of concentrations in their 1-min dataset is similar to the range we found in our Münster 1-s dataset (5 to 112 μg m$^{-3}$). The PM$_{2.5}$ hotspots identified along
the LT route can be spatially associated with areas likely to be affected by high traffic intensities at intersections or traffic lights. These areas typically experience stop-and-go vehicle traffic, and therefore, particles are emitted from brake and tire abrasion, which contributes to PM$_{2.5}$ mass concentrations (Ketzel et al., 2007; Amato et al., 2014). Furthermore, as is known from other studies, the particle mass concentration in the size range of PM$_{2.5}$ is heavily influenced by accumulation mode particles that originate from long-range transport (Gidhagen et al., 2004; Ketzel and Berkowicz, 2005, Gietl et al. 2008; Gietl and Klemm 2009). Our findings confirm that compared with the long-range transport sources, local vehicle traffic sources are evident but less important in the case of PM$_{2.5}$.

Regarding differences in PN$_{10}$ concentrations between the sampling routes, our findings agree well with the mean UFP concentrations as measured by Ham et al. (2017) for bike trails and arterial trails in Sacramento, CA (10,000 to 24,000 p cm$^{-3}$), with the results from Vinzent et al. (2005) for data collected in central Copenhagen (geometric mean 32,400 p cm$^{-3}$) and with a study from Thai et al. (2008) that was conducted in Vancouver (UFP concentration range 18,830 to 57,692 p cm$^{-3}$). Our resulting ratio of three to one when comparing the PN$_{10}$ medians of the HT and LT routes agrees well with findings from Ham et al. (2017). The authors of the Sacramento study attributed the differences of mean UFP concentration to the proximity of the bike route to the motorized traffic and to differences in vehicle activity between these two routes. Other authors also published similar ratios despite having different study areas (Jarjour et al., 2013; Zuurbier et al., 2010).

Compared with the spatial concentration variability of PM$_{2.5}$, the measured concentrations of PN$_{10}$ showed a largely different picture. Particle number concentrations are heavily influenced by short-lived particles within the nucleation mode that originate both from primary emissions and from secondary formation from precursor gasses from combustion processes (Ning and Sioutas, 2010; Quiros et al., 2013). Kingham et al. (2013) reported median PN$_{10}$ of 31,414 particles cm$^{-3}$ and 16,641 particles cm$^{-3}$ for on-road and off-road cyclists, respectively, in Christchurch, New Zealand, showing that the overall mean PN$_{10}$ concentrations detected in this study (18,026 and 6700 particles cm$^{-3}$) are rather low in comparison with other cities. Boogaard et al. (2008) also showed that factors like the type of passing vehicles, the time spent waiting at traffic lights and intersections, the type of intersection, and the distance of bike lanes to motorized traffic routes all contributed to the variability of particle number concentration in eleven Dutch cities. Similarly, Pattinson et al. (2017) found 40 to 50% lower ultrafine particle concentrations on a path that was 19 m away from the road than on the main road itself in Christchurch, and Brand et al. (2019) found a rise in black carbon average concentration when the cycle lane inside the Hyde Park approaches to the edge of the park, which was attributed to the proximity of traffic emission sources. Our PN$_{10}$ results confirm that local vehicle traffic sources play a major role for local PN$_{10}$ concentrations.

The analysis of cyclist’s exposure dose on individual routes shows much higher concentrations of PM$_{10}$ and PN$_{10}$ along the HT route than along the LT route. Similarly, MacNaughton et al. (2014) found that bike paths have lower concentrations of traffic-related pollutants than bike lanes along heavy traffic routes. Ham et al. (2017) found that average PM$_{2.5}$ concentrations were lowest for trips taken on a dedicated bike trail. We believe that the health risk for cyclists is better explained by exposure to UFP rather than to PM$_{10}$, because a high number concentration of UFP is of higher toxicological relevance than a high PM$_{10}$ mass concentration (Penttinen et al., 2001; von Klot et al., 2002).

The route exposure metrics reveals that cyclist exposure to PM$_{2.5}$ is quite similar between routes. However, since cyclists’ exposure to particle number at the HT doubled the one at the LT route, the toxicological risk was twice larger at the former route. Similarly, Lonaty et al. (2016) stated that the higher the volume of traffic, the greater the cyclist’s exposure to traffic-related pollutants, in particular UFP. Our results confirm that vehicle traffic intensities play a major role for local UFP concentrations, while the influence of vehicle traffic sources on local PM$_{2.5}$ concentrations is less pronounced.

**Conclusion**

In this study, it was demonstrated that mobile direct measurements of particle mass concentration and particle number concentration are feasible at a high temporal resolution (1 Hz) without requiring any surrogate measures or assumptions. The monitoring platform mounted in the cargo bike was useful for characterizing the spatial
distribution of PMC and PNC simultaneously across an urban environment, clearly showing the contrasting dynamics of PMC and PNC along various routes. Particle mass concentration and particle number concentration differed depending on the proximity to motorized traffic. The temporal as well as the spatial variability of PN_{10} was very high, while the differences were less distinct for the PM_{2.5} mass concentration.

Overall, our results suggest that the choice of route is a key factor for reducing cyclists’ exposure to air pollution, although the duration and distance traveled can—for some metrics—offset the exposure reduction when using low-traffic routes. In our study for example, using the bike path through the city, which was well separated from the street, might result in a higher mass of PM_{2.5} particles inhaled, since this route was longer than the bike lane next to the street. However, the larger amount of UFP particles found on bike lanes adjacent to arterial streets indicates that even if cyclists reduce their exposure time, routes closer to traffic result in much higher levels of particle exposure.

We acknowledge some limitations in the present study. First, not every particle inhaled by a human will indeed reach the pulmonary alveoli; therefore, we might overestimate the particles’ effects. On the other hand, we did not consider the chemical composition of particles by any means, which might have given more insight into specific toxic effects.

Our results stress the importance of studying the levels of UFP, which humans are exposed to. In our view, it is not sufficient to measure the PMC. Legislation and policymakers should acknowledge the need to further improve environmental observation targets such as PNC in order to help mitigate humans’ exposure, specifically cyclists’ exposure, to air pollution.

We performed only a limited number of measurements. Many more trips with the instrumented cargo bike are required to arrive at statistically robust results concerning cyclists’ exposure to air pollution in various cities, climates, traffic situations, commuting distances, and traveling speeds. In that sense, this is a feasibility study that proves the technical and scientific concept to be successful.

Acknowledgments We thank Celeste Brennecka for language editing of the manuscript.

Funding information This study was supported by a Visiting Professor Fellowship of the German Academic Exchange Service (DAAD) to Hebe Carreras, which is gratefully acknowledged.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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