Particle Concentration and Characteristics near a Major Freeway with Heavy-Duty Diesel Traffic

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This study presents the number, surface and volume concentrations, and size distribution of particles next to the I-710 freeway during February through April 2006. I-710 has the highest ratio (up to 25%) of heavy-duty diesel vehicles in the Los Angeles highway network. Particle concentration measurements were accompanied by measurements of black carbon, elemental and organic carbon, and gaseous species (CO, CO₂). Using the incremental increase of CO₂ over the background to calculate the dilution ratio, this study makes it possible to compare particle concentrations measured next to the freeway to concentrations measured in roadway tunnels and in vehicle exhaust. In addition to the effect of the dilution ratio on the measured particle concentrations, multivariate linear regressions showed that light and heavy organic carbon concentrations are positively correlated with the particle volume in the nucleation and accumulation modes, respectively. Solar radiation was also positively correlated with the particle surface concentration and the particle volume in the accumulation (40–638 nm) mode, presumably as a result of secondary particle formation. The methods developed in this study may be used to decouple the effect of sampling position, meteorology, and fleet operation on particle concentrations in the proximity of freeways, roadway tunnels, and in street canyons.

Introduction

Various epidemiological studies have concluded that a positive correlation exists between airborne particulate matter (PM) and adverse effects on human health (1, 2). Recent toxicological studies corroborate these findings both in vitro and in vivo (3, 4). Although this compelling evidence exists, the causal mechanisms behind these toxic responses remain uncertain. Consequently, which metric of PM—mass, surface area, or number of particles—may be responsible for toxicity remains unknown. Recent studies suggest that ultrafine particles (dₚ < 100 nm) may elicit a higher adverse response per unit mass than finer (dₚ < 2.5 μm) and coarse (2.5 μm < dₚ < 10 μm) particles (5, 6).

Dynamometer experiments have shown motor vehicles to emit high concentrations of particles by number, and heavy-duty diesel vehicles (HDVs) without particle traps, in particular, emit high PM mass concentrations (7, 8). Recently, sampling conducted in freeway microenvironments has also confirmed high concentrations of PM due to motor vehicles (9, 10). Zhu et al. (9) found that particle number concentration was highly influenced by distance from the freeway, with a large spike in particle number next to the roadway that decays to background levels within 100–150 m. PM₂.₅ (particulate matter less than 2.5 μm in aerodynamic diameter) mass concentration was nearly unaffected over the same distances. Westerdahl et al. (10) reported concentrations of PM and various co-pollutants while driving on various roads and highways in Los Angeles. Concentrations on some of the freeways sampled in that study peaked as high as one million particles per cm³. Thus, it has been suggested that exposure to motor vehicle exhaust emissions during commute constitutes a large fraction of daily personal PM exposure, especially to ultrafine PM.

The vast freeway network in Los Angeles accommodates over six million registered vehicles. Therefore, it is reasonable that most PM₂.₅ originates from vehicles, which has been proven in other urban areas (11, 12). PM is both directly emitted, via combustion and mechanical wear, and formed via secondary reactions of direct emissions, where primary and inorganic vapors undergo gas-to-particle conversion in the atmosphere (13, 14). Roadways with high HDV traffic have been shown to have elevated particle number and mass concentrations, and especially high levels of black carbon (10).

In this study, detailed information is provided on the physical and chemical characteristics of particles originating from a major freeway during the winter season, with a significant HDV fraction. This was accomplished by measuring semi-continuous and time-integrated particle number, surface area, mass, and chemistry adjacent to I-710 from February to April 2006. The results are compared to those of previous studies, after correcting for the dilution ratio in different sampling locations, and a regression analysis is conducted to reveal the effect of meteorology and traffic conditions on the observed particle concentrations.

Materials and Methods

The I-710 freeway is a 26 m wide eight-lane highway connecting the ports complex of Long Beach and San Pedro to the shipping yards in East Los Angeles. For this reason, as much as 25% diesel traffic has been reported on this freeway (http://traffic-counts.dot.ca.gov). Total traffic counts are also very high, with between 150,000 and 200,000 vehicles per day passing the sampling location. The sampling site was located in a paved property run by the Imperial Flood Control Yard in South Gate, CA. This location, shown in Figure 1, is directly adjacent to the roadway with no other immediate sources either upwind or downwind. Just downwind of the site are a bike path and the Los Angeles River, a concrete-lined drainage channel.

To ensure that sample probes were within the plume of the freeway, wind speed and direction were monitored with a portable weather station (Monitor II, Weather Systems Company, San Jose, CA). This instrument also recorded local temperature and relative humidity. Data were logged over 5 min intervals during the entire study. The monitor was located 2 m above the ground, and direction was calibrated daily by aligning the weather vane with a compass.

Traffic data were collected by two methods that were used to create a comprehensive traffic profile. Total traffic volume (including all vehicle types) and average vehicle speed were obtained from the California Department of Transportation.
Particle number and mass (by assuming an average density of 1.0 g cm$^{-2}$) concentrations in the range of 0.7–2.5 μm (aerodynamic diameter) were measured in 5 min intervals by means of the Aerodynamic Particle Sizer (APS). Particle number concentrations by both SMPS and APS were determined by integrating over the particle number size distribution. Continuous particle surface concentration was measured with the recently developed Nanoparticle Surface Area Monitor (NSAM). This measures the total particle surface area available for diffusion of ions. A dual beam aethalometer measured black carbon (BC) concentrations every 5 min. In addition, an organic carbon (OC)/elemental carbon (EC) field instrument (OC/EC) measured hourly PM$_{2.5}$ elemental and organic carbon concentrations. This instrument allows for the separation of particulate OC into four response peaks representing different volatility fractions of OC (J5). For the purposes of this study, peaks 2–4 were summed and considered as the least volatile or heavy OC fraction (HOC), while peak 1 represented the lighter, most volatile OC fraction (LOC).

**Results and Discussion**

**Particle and Co-Pollutant Concentrations Next to the Freeway.** Meteorological and traffic parameters were measured during the course of sampling. The average wind direction was 215 ± 39 degrees, which placed the sampling site downwind of freeway traffic emissions, and the mean wind speed was 1.8 ± 0.9 ms$^{-1}$ (all ranges given as mean ± standard deviation). The temperature and relative humidity were 19.6 ± 5.0 °C and 46.2 ± 16.3%, respectively, with a diurnal profile shown later in the paper. Caltrans reported total volumes of north- and south-bound lanes in the range of 5180 ± 640 h$^{-1}$ and 5640 ± 800 h$^{-1}$, respectively. The average traffic speeds of north and south bound lanes were 78 ± 28 km h$^{-1}$ and 85 ± 12 km h$^{-1}$, respectively. Video recordings revealed that 11–21% of vehicles on the freeway were HDVs. However, when expressed as an absolute hourly density on the highway, the HDV number did not significantly differ during the sampling hours (1700–2100 h$^{-1}$). This range is slightly lower than previously reported in studies conducted in 2001 (16) on the same freeway, possibly due to the opening of a railway line (Alameda Corridor) in 2002, which is used for the transport of goods from the Los Angeles port.

Table 2 presents the mean particle and copollutant levels measured near the freeway over the 7-week sampling period (120–140 h of data, depending on the magnitude considered). The average CPC particle number concentration was 86 000 cm$^{-3}$ during the course of the whole campaign whereas the average SMPS concentration was 19 000 cm$^{-3}$. The difference between the two instruments can be attributed to the different size ranges (CPC > 6 nm, SMPS 16–638 nm) and particle losses in the SMPS, and it is a strong indication that the particle concentration near the freeway is dominated by particles less than 16 nm in diameter. The high particle number concentration below 16 nm is consistent with the results of Zhu et al. (9), conducted in the winter of 2002 next to the same freeway. The particle surface concentration measured with the NSAM was 153 ± 55 μm$^2$ cm$^{-3}$. With regard
to carbon species, BC concentration measured 4.4 ± 2.1 µg m⁻³, which is at similar levels with EC (3.2 ± 2.2 µg m⁻³) and total OC (4.4 ± 1.6 µg m⁻³). The fraction of EC and BC to PM₁₀⁻².₅ was 50% and 68%, respectively. In a previous study at the Caldecott tunnel (17) with 3–6% diesel vehicles, EC accounted for 30–38% of PM₁₀. It was also reported that, on average, BC accounted for 50% of PM₁ at the entrance and 30% at the exit in Gubrist tunnel with 16% diesel vehicles (18). EC is considered a marker for diesel exhaust aerosol production rate increased mildly due to the acceleration and deceleration driving pattern. However, the increase is not substantial due to the relative reduction of the total traffic road activity. This has been calculated on the basis of activity data provided by Caltrans for driving speeds and traffic volume, the split in diesel HDV and gasoline cars, and appropriate CO₂ emission functions for each vehicle category (21). The total CO₂ production rate increased very little during the study, due to the relatively constant speeds and traffic volumes during the day. During congestion, the CO₂ production rate increased mildly due to the acceleration–deceleration driving pattern. However, the increase is not substantial due to the relative reduction of the total traffic volume.

Both EC and BC average concentrations display a bimodal pattern and agree with each other very well, indicating identical sources of EC and BC (Figure 2c). Two peaks were observed at 11:00 and later in the afternoon (15:00–16:00) corresponding to the diurnal pattern of the traffic rush hours. The diurnal patterns of OC species do not follow traffic as closely and this is a further indication that these may come from both direct traffic emissions as well as secondary formation mechanisms through photochemical reactions.

With regard to particle concentrations, SMPS total number concentration, surface concentration, and volume of nucleation and accumulation modes all peaked at 16:00 when traffic was congested. These associations strongly indicate the link between particle concentrations and traffic emission on the freeway. Harrison et al. (22) have also reported that roadside particle number and surface concentrations have the same general pattern of behavior as the vehicle emission source. On the other hand, the CPC number concentration showed a monotonically increasing trend from 50,000 cm⁻³ to 110,000 cm⁻³ as the day progressed. This can be explained by the nucleation process of small particles with decreasing temperature in the evening. Similar results have also been observed by Kuhn et al. (23). These results indicate that the

### TABLE 2. Summary of Particle and Copollutant Levels Next to the Freeway for the Three Cases: Campaign, a Rainy Day, and a Day When the Freeway Was Closed

<table>
<thead>
<tr>
<th></th>
<th>Average</th>
<th>Range</th>
<th>SD*</th>
<th>Average</th>
<th>Range</th>
<th>SD*</th>
<th>Average</th>
<th>Range</th>
<th>SD*</th>
</tr>
</thead>
<tbody>
<tr>
<td>N_{CPC} (cm⁻³)</td>
<td>8.6 x 10⁴</td>
<td>1.1 x 10⁻⁴–5.4 x 10⁻⁴</td>
<td></td>
<td>3.4 x 10⁴</td>
<td>1.3 x 10⁻⁴–2.2 x 10⁻⁴</td>
<td></td>
<td>3.1 x 10⁴</td>
<td>2.5 x 10⁻⁴–6.8 x 10⁻³</td>
<td></td>
</tr>
<tr>
<td>N_{SMPS} (cm⁻³)</td>
<td>1.9 x 10⁴</td>
<td>3.7 x 10⁻⁴–9.4 x 10⁻³</td>
<td></td>
<td>5.5 x 10⁴</td>
<td>2.3 x 10⁻³–3.0 x 10⁻³</td>
<td></td>
<td>6.5 x 10⁴</td>
<td>5.1 x 10⁻³–1.4 x 10⁻³</td>
<td></td>
</tr>
<tr>
<td>PMₐ⁻³₀⁻².₅ (µg/m³)</td>
<td>6.4</td>
<td>1.7–30</td>
<td>3.6</td>
<td>8.2</td>
<td>6.7–8.0</td>
<td>0.85</td>
<td>2.1</td>
<td>1.7–2.3</td>
<td>0.27</td>
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<td>surface (µm/cm³)</td>
<td>153</td>
<td>36–83</td>
<td>55</td>
<td>41</td>
<td>20–64</td>
<td>19</td>
<td>57</td>
<td>46–71</td>
<td>12</td>
</tr>
<tr>
<td>V_{NM} (µm³/cm³)</td>
<td>0.094</td>
<td>0.011–0.22</td>
<td>0.047</td>
<td>0.031</td>
<td>0.011–0.061</td>
<td>0.02</td>
<td>0.049</td>
<td>0.035–0.058</td>
<td>0.13</td>
</tr>
<tr>
<td>V_{AM} (µm³/cm³)</td>
<td>16.3</td>
<td>3.66–36.5</td>
<td>6.96</td>
<td>7.1</td>
<td>4.73–9.79</td>
<td>1.86</td>
<td>4.3</td>
<td>3.7–5.6</td>
<td>1.1</td>
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<tr>
<td>Geomean dp (nm)</td>
<td>50</td>
<td>28–96</td>
<td>11</td>
<td>53</td>
<td>40–60</td>
<td>7</td>
<td>46</td>
<td>42–52</td>
<td>4</td>
</tr>
<tr>
<td>EC (µg/m³)</td>
<td>3.2</td>
<td>0.3–11.2</td>
<td>2.2</td>
<td>0.7</td>
<td>0.3–11.5</td>
<td>0.3</td>
<td>0.40</td>
<td>0.3–0.5</td>
<td>0.2</td>
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<tr>
<td>BC (µg/m³)</td>
<td>4.4</td>
<td>0.5–10.1</td>
<td>2.1</td>
<td>2.6</td>
<td>0.9–9.6</td>
<td>2.6</td>
<td>0.8</td>
<td>0.8–1.8</td>
<td>0.8</td>
</tr>
<tr>
<td>LOC (µg/m³)</td>
<td>2.4</td>
<td>0.9–5.3</td>
<td>0.8</td>
<td>1.3</td>
<td>1.1–1.7</td>
<td>0.2</td>
<td>3.0</td>
<td>2.8–3.2</td>
<td>0.3</td>
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<tr>
<td>HOC (µg/m³)</td>
<td>2.0</td>
<td>0.1–1.2</td>
<td>1.4</td>
<td>1.2</td>
<td>0.9–1.8</td>
<td>0.3</td>
<td>1.6</td>
<td>1.5–1.8</td>
<td>0.2</td>
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<td>CO (ppmv)</td>
<td>0.23</td>
<td>0.10–3.6</td>
<td>0.35</td>
<td>0.01 &lt;0.09</td>
<td>0.02</td>
<td>0.29 &lt;1.2</td>
<td>0.53</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CO₂ (ppmv)</td>
<td>426</td>
<td>359–567</td>
<td>30</td>
<td>402</td>
<td>384–430</td>
<td>14</td>
<td>405</td>
<td>401–410</td>
<td>3.6</td>
</tr>
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</table>

* Standard deviation.
Particle concentration and characteristics next to the freeway are influenced by a combination of the traffic activity and the meteorological conditions. The statistical analysis presented in a following section will help to further identify the underlying associations.

Comparison with Previous Studies. Particle characterization studies in the vicinity of freeways or tunnels with diesel HDV traffic have been conducted in the past. Westerdahl et al. (10) measured particle characteristics by driving a mobile lab on fixed routes in the Los Angeles road network, including a stretch of the I-710 freeway where they chased different vehicle plumes. In another study, Geller et al. (24) measured size distributions in the Caldecott tunnel (Bore 1), located in Orinda, CA. Table 3 compares measured particle

![Graph showing diurnal profile of measured variables normalized over the maximum hourly value recorded (given in the legend within parentheses): (a) meteorological conditions, (b) traffic data, (c) particle chemical parameters and CO2, and (d) particle physical parameters.](image)

**FIGURE 2.** Diurnal profile of measured variables normalized over the maximum hourly value recorded (given in the legend within parentheses): (a) meteorological conditions, (b) traffic data, (c) particle chemical parameters and CO2, and (d) particle physical parameters.

**TABLE 3.** Comparison of Concentrations in Different Studies Involving Heavy-Duty Diesel Vehicle Traffic (Italized Values Have Been Corrected for the Different Dilution Ratios, Calculated on the Basis of the CO2 Concentrations)

<table>
<thead>
<tr>
<th></th>
<th>2006 study</th>
<th>Westerdahl et al. (10)</th>
<th>Geller et al. (24)</th>
</tr>
</thead>
<tbody>
<tr>
<td>location</td>
<td>I-710</td>
<td>I-710</td>
<td>Caldecott Tunnel (Bore 1)</td>
</tr>
<tr>
<td>sampling site</td>
<td>20 m from freeway median strip</td>
<td>mobile lab following traffic</td>
<td>tunnel exit</td>
</tr>
<tr>
<td>period</td>
<td>Feb–Apr 2006</td>
<td>Feb–Apr 2003</td>
<td>Aug 2004</td>
</tr>
<tr>
<td>sampling hours</td>
<td>12 pm–4 pm</td>
<td>5 pm–7 pm</td>
<td>12 pm–6 pm</td>
</tr>
<tr>
<td>passenger cars (h⁻¹)</td>
<td>8359</td>
<td>10250</td>
<td>7580</td>
</tr>
<tr>
<td>light-duty trucks (h⁻¹)</td>
<td>600</td>
<td>360</td>
<td>91</td>
</tr>
<tr>
<td>heavy-duty trucks (h⁻¹)</td>
<td>1630</td>
<td>1225</td>
<td>1040</td>
</tr>
<tr>
<td>temperature (°C)</td>
<td>21.4</td>
<td>14.4</td>
<td>21</td>
</tr>
<tr>
<td>RH (%)</td>
<td>42</td>
<td>60</td>
<td>23.3</td>
</tr>
<tr>
<td>wind speed (m/s)</td>
<td>2.2</td>
<td>1.2</td>
<td>59</td>
</tr>
<tr>
<td>measured CPC (cm⁻³)</td>
<td>75000</td>
<td>98500</td>
<td>366000 (54000)</td>
</tr>
<tr>
<td>measured CO (ppm)</td>
<td>0.27</td>
<td>0.11</td>
<td>0.28 (0.05)</td>
</tr>
<tr>
<td>measured BC (µg/m³)</td>
<td>4.6</td>
<td>2.8</td>
<td>3.4 (0.6)</td>
</tr>
<tr>
<td>corrected for DR⁵</td>
<td>measured</td>
<td>corrected for DR⁵</td>
<td>637500</td>
</tr>
<tr>
<td>CPC (cm⁻³)</td>
<td>75000</td>
<td>98500</td>
<td>366000 (54000)</td>
</tr>
<tr>
<td>CO (ppm)</td>
<td>0.27</td>
<td>0.11</td>
<td>0.28 (0.05)</td>
</tr>
<tr>
<td>BC (µg/m³)</td>
<td>4.6</td>
<td>2.8</td>
<td>3.4 (0.6)</td>
</tr>
<tr>
<td>relative dilution ratio</td>
<td>1:1</td>
<td>1:10.2 (1.22)</td>
<td>1:7.3 (0.82)</td>
</tr>
</tbody>
</table>

* Daily averages from CalTrans. ** Values in parentheses correspond to calculation uncertainty (see Supporting Information). ^ Elemental carbon.
number and copollutant concentrations from these two studies to the current one. All studies were conducted under similar temperature and relative humidity conditions and the wind-speed was also very similar in the two non-tunnel studies. The diesel truck fraction was ~3.7% in the Caldecott tunnel, and the road had a positive gradient of +4.2%.

It is important to correct for the dilution ratio (DR) in the various sampling locations before comparing pollutant concentrations between these studies. This correction enables us to decouple the effects of fleet operation and ambient conditions on pollutant levels from the influence of the exact position of the monitoring station. The DR calculation is based on the ratio of the fleet-average exhaust CO\(_2\) concentration over the incremental ambient CO\(_2\) increase, and it is given as Supporting Information. Carbon dioxide has been used in several studies as a tracer to derive fuel-specific emissions from ambient pollutants concentrations (25) or to deduce emission factors in tunnel measurements (26). While the fundamentals of those approaches are similar to our study, we use the incremental increase in ambient CO\(_2\) to deduce dilution conditions in different environments. The calculations show that the DR by the freeway (present study) is 10.2 ± 1.22 times higher than in-freeway, and 7.3 ± 0.83 times higher than in the tunnel. These ratios are consistent with the conceptual model of Zhang et al. (27), who estimated that the road-to-ambient process is associated with an additional DR of ~10.

Concentrations varied in the range 75 to 637.5 thousand particles per cm\(^3\) in the different studies before correcting for the DR. After correction (italicized values in Table 3), concentrations reached rather similar levels, which can lead to interesting conclusions. The in-freeway concentrations of particle number and BC measured by Westerdahl et al. (10) are lower than the values of this study while CO is at the same levels. This may be attributed to the different mixing of the exhaust when sampling in the freeway as opposed to next to the freeway. When following traffic, there is a larger probability to follow a car than a truck, due to the larger density of passenger cars. The different mixing would have a limited effect on the DR calculated, since the CO\(_2\) concentration in the exhaust of the two vehicle types is different (gasoline, 12.5%; diesel, 7.1%). However, it is not possible to calculate a more precise DR than the one corresponding to mean fleet operation (Table S2), unless the probabilities of following a car vs a truck plume are known. Nevertheless, the effect of different mixing on DR is negligible compared to the orders of magnitude difference expected for the concentration of BC and particle number in the exhaust of the two vehicle types. Therefore, the higher CO/BC ratio of the Westerdahl et al. (10) study, compared to the present results, is suggestive of the larger CO/BC ratio from gasoline vehicles. This is consistent with the much higher ratio of CO over light extinction by the exhaust plume of gasoline compared to diesel vehicles (28). For the same reason, the DR-corrected particle number concentration is lower than that measured next to the freeway, which is due to the lower on-road particle number emission factors of gasoline cars vs diesel trucks (24, 29).

On the other hand, the particle number and BC concentration in the Caldecott tunnel are at the same or slightly higher levels than next to the freeway after DR correction, despite the much lower fraction of diesel vehicles in the tunnel. This is most probably due to the uphill driving in the tunnel, which has a significant effect on the engine load. Using typical values for road resistance and solving the equation of motion for LGVs and HDDVs (see Supporting Information), 4.2% positive road gradient corresponds to ~1.6 times increase in engine load for a light duty gasoline engine and a ~2.9 increase for a diesel truck engine. The high particle number concentrations in this case are consistent with the findings of Kittelson et al. (29) and Maricq et al. (8), who show a large increase in particle number from gasoline vehicles as the engine load increases. The effect of engine load on particle number concentration is not as significant in diesel engines (7) but does increase BC emissions (30), which leads to the relatively high ambient concentration of BC, despite the small fraction of diesel vehicles.

**Particle Size Distributions.** Figure 3 shows particle size distributions measured next to the freeway with typical exhaust size distributions of two heavy-duty vehicles measured in the laboratory, after correction for the dilution ratio. (a) Mean geometric hourly size distributions measured during the 7-week campaign, (b) effect of individual events on the particle size distributions, and (c) comparison of the grand average size distribution measured next to the freeway with typical exhaust size distributions of two heavy-duty vehicles measured in the laboratory, after correction for the dilution ratio.
b), and, finally, in comparison to HDV particle exhaust distributions measured in the laboratory (panel c). The ambient particle distributions (campaign grand average and hourly averages) are the geometric mean of the corresponding size distributions recorded over the 7-week sampling period.

Figure 3a shows that the particle size distribution next to the diesel freeway was bimodal, with a nucleation mode in the size range below 40–50 nm and an accumulation mode peaking at 70–80 nm during all sampling hours. This bimodal distribution has been typically encountered in the proximity of freeways with diesel traffic (9) or when following a diesel vehicle on the road (31–33) but has not been present when sampling in the proximity of a gasoline freeway (23). During freeway closure (Figure 3b), there was no traffic-induced emission source and the particle concentration dropped significantly compared to the average. In addition, the size distribution became monomodal within the range 16–638 nm studied, with a peak concentration of 12,000 cm$^{-3}$ at 30 nm. During the rainy day, the concentration of the whole size distribution dropped by 70%, similar to the observations of Mozurkewich et al. (34).

Figure 3c attempts to explain the bimodal size distribution by comparing the average size distribution recorded next to the 710 freeway to typical size distributions obtained by older technology (Euro I) and more recent technology (Euro III) HDVs (35). These distributions correspond to mode 5 of the European heavy-duty Stationary Cycle (ESC 5), which stands for moderate speed (~60% of maximum speed) and 50% load, both typical of freeway driving. The exhaust concentrations have been corrected for the DR considered at the sampling location (1917 ± 219; Supporting Information) and the contribution of HDV exhaust flowrate to the mean fleet exhaust flowrate. The comparison shows excellent agreement in the size range of both particle modes, between freeway and laboratory sampling. In addition, the concentration of particles larger than 40–50 nm (accumulation mode) measured in the laboratory is very close to the levels measured next to the freeway. However, the accumulation mode distribution is wider next to the freeway than in the laboratory studies, which may be due to the averaging effect on the size distributions of several vehicles, and/or to the contribution of other particle sources (secondary particles and non-exhaust vehicle particles).

With regard to the nucleation mode particles, the size range determined in the laboratory from the Euro I engine (employing a primary DR of 12.5:1, a dilution temperature of 32 ℃, and a residence time of 2.5 s) is similar to the size distribution measured next to the freeway. However, the actual number concentration depends on the vehicle technology and the sampling conditions. In addition, the nucleation mode formed next to the freeway also depends on species emitted by gasoline vehicles.

The comparison of the tunnel, on-road, dynamometer, and present studies demonstrates that the measurement of the incremental increase in ambient CO$_2$ due to traffic is a good indicator of the DR at the sampling location and can be used to infer concentrations at other locations with similar conditions. This can be used to decouple the effect of sampling position from the difference in the measured pollutant levels.

### Statistical Processing of the Sampled Variables

A multivariate linear regression analysis was performed on a dataset of 86 hourly average values of traffic, ambient, and pollutant concentration data, in order to associate particle concentrations next to the freeway with traffic and ambient conditions as well as co-pollutant concentrations. The dataset size was limited to 86 h because it was necessary to have a complete set of values for all 22 variables recorded during sampling. The variables included all magnitudes shown in Figure 2, with the addition of the PM$_{10-7.5}$ mass and the time of sampling. “CO$_2$ Production” was excluded because it was not possible to have a precise estimate for each hour of the sampling period. The dependent variables considered in the regressions were the CO$_2$ concentration, the CPC particle number, the surface concentration, and the volumes of particles in the nucleation (16–40 nm) and accumulation (40–638 nm) modes. All other recorded variables were considered independent. For the CO$_2$ regression in particular, only traffic data and ambient conditions were considered as independent variables. All regressions were carried out with SPSS 12.0 and only final linear combinations at a confidence level of 99.5% were considered.

The results of the multivariate linear regression analysis are shown in Table 4. The values in the table correspond to the multiplication factors for the standardized independent variables (input). The linear combination of the multiplication factor and the standardized input variable leads to a reconstructed value of the dependent variable. The correlation of the dependent (measured) variables and the reconstructed variables is satisfactory with regression slopes on the order of 0.91–1.0 and $R^2$ above 0.6 (Figure S2 in Supporting Information). However, it was not possible to derive a reconstructed variable that would satisfactorily correlate with the CPC particle number. This is probably because the nucleation of new particles is a highly sensitive, nonlinear function of the input variables, and it is therefore not possible to linearly associate it with the independent variables considered.

Given the satisfactory correlations between the reconstructed and the dependent variables, the values in Table 4 may be used to establish the associations of the dependent with the independent variables. The multiplication coefficients in this table correspond to standardized variables. For example, a decrease in the wind speed by one standard deviation (which corresponds to a variation of 0.9 m/s) would increase the CO$_2$ concentration by 0.24 standard deviations (which corresponds to an increase of $0.24 \times 24$ ppm = $-5.8$ ppm). Based on these values, it is evident that CO$_2$ concentration increases as wind speed decreases, as wind direction changes toward the west (moving the sampling point closer...
to the freeway), and in the afternoon and evening hours as the sun sets and the mixing height decreases. These relationships confirm that CO2 is a very good indicator of the DR. Interestingly, CO2 concentration does not depend on the traffic volume and conditions because there was little variation of the CO2 emitted per hour on the highway, as shown in Figure 2b (standard deviation over mean was 13% for the whole measurement campaign). Presumably, this variation is not high enough to cause a measurable difference in the CO2 concentration levels.

The CO2 concentration was then used as an independent variable to reconstruct the concentration of the particle surface and volume concentrations. Table 4 shows that all three particle concentration expressions are positively affected by the CO2 concentration, which is consistent with an increase in the concentration of these variables with decreasing DR. All concentrations are also positively associated with BC. Interestingly, LOC is positively associated with the nucleation mode volume, while HOC is associated with the accumulation mode. There is evidence (30, 36) that the gasoline OC profile is shifted to heavier species than the diesel one. This would imply that the nucleation mode volume in the size range 16–40 nm is mainly associated with the diesel activity on the freeway. This is consistent both with the laboratory measurements (Figure 3c) and with the findings of Zhu et al. (9) who found that next to the diesel freeway (I-710) a distinct particle mode appears in the 20–30 nm range. This mode is clearly separated by a higher nucleation mode forming below 10 nm and was not evident when sampling next to a (mainly) gasoline freeway (I-405). It needs to be noted that a high particle concentration below 16 nm is also evident in our study when comparing the CPC and SMPS recordings. Unfortunately, due to the lack of linear correlation of the CPC recordings with any of the other measured variables, it was not possible to link the origin of this nanoparticle (6–16 nm) mode to either diesel or gasoline vehicles.

The regression analysis also shows that the NM volume concentration is negatively correlated to the wind-speed. This is in addition to the effect of wind on dilution, expressed by the CO2 correlation, and can be an indication of NM evaporation as the wind-speed increases. Finally, the volume of accumulation mode is positively correlated with the solar radiation. This indicates that there may be other processes, unrelated to traffic, that contribute to a significant portion of the aerosol in that size range, such as secondary formation by photochemical activity.

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Supporting Information Available
Dilution ratio calculation scheme, concentrations calculation and uncertainty, engine load calculation, tables of data, figures, and references. This material is available free of charge via the Internet at http://pubs.acs.org.

Literature Cited


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