

# Emission factors of black carbon and co-pollutants from diesel vehicles in Mexico City

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Abstract. Diesel-powered vehicles are intensively used in urban areas for transporting goods and people but can substantially contribute to high emissions of black carbon (BC), organic carbon (OC), and other gaseous pollutants. Strategies aimed at controlling mobile emissions sources thus have the potential to improve air quality and help mitigate the impacts of air pollutants on climate, ecosystems, and human health. However, in developing countries there are limited data on the BC and OC emission characteristics of diesel-powered vehicles, and thus there are large uncertainties in the estimation of the emission contributions from these sources. We measured BC, OC, and other inorganic components of fine particulate matter (PM), as well as carbon monoxide (CO), nitrogen oxides  $(NO_x)$ , sulfur dioxide  $(SO_2)$ , ethane, acetylene, benzene, toluene, and C2-benzenes under real-world driving conditions for 20 diesel-powered vehicles encompassing multiple emission level technologies in Mexico City with the chasing technique using the Aerodyne mobile laboratory. Average BC emission factors ranged from 0.41- $2.48 \,\mathrm{g \, kg^{-1}}$  of fuel depending on vehicle type. The vehicles were also simultaneously measured using the cross-road remote sensing technique to obtain the emission factors of nitrogen oxide (NO), CO, total hydrocarbons, and fine PM, thus allowing for the intercomparison of the results from the two techniques. There is overall good agreement between the two techniques and both can identify high and low emitters, but substantial differences were found in some of the vehicles, probably due to the ability of the chasing technique to capture a larger diversity of driving conditions in comparison to the remote sensing technique. A comparison of the results with the US EPA MOVES2014b model showed that the model underestimates CO, OC, and selected VOC species, whereas there is better agreement for NO<sub>x</sub> and BC. Larger OC / BC ratios were found in comparison to ratios measured in California using the same technique, further demonstrating the need for using locally obtained diesel-powered vehicle emission factor database in developing countries in order to reduce the uncertainty in the emissions estimates and to improve the evaluation of the effectiveness of emissions reduction measures.

### 1 Introduction

On-road mobile sources can substantially contribute to high emissions of black carbon (BC), organic carbon (OC), and other particulate matter (PM) components in urban areas. Although both gasoline and diesel-powered vehicles are emitters of primary fine particulate matter, the available evidence indicates that when normalized to fuel consumption, PM emission factors are more than an order of magnitude higher for heavy-duty diesel vehicles compared to light-duty gasoline vehicles (e.g., Ban-Weiss et al., 2008; Dallmann et al., 2014). Freight tractor trailers, public transport buses, and heavy-duty trucks are typically powered by diesel fuel due to their high requirements of power, durability, and fuel efficiency. However, diesel-power vehicles can also contribute to high levels of nitrogen oxides (NO<sub>x</sub>), carbon monoxide (CO), volatile organic compounds (VOCs), and other harmful co-pollutants. Thus, controlling diesel-powered mobile emissions has the potential to improve air quality and help mitigate the impacts of air pollutants on climate, ecosystems, and human health.

Compared to gaseous pollutant emissions, measurementbased estimations of emission factors for PM components from diesel-powered vehicles are less abundant. Until recently most of the measurements of PM from diesel-powered vehicles have been obtained using either semi-quantitative opacity-based techniques or by using time-integrated gravimetric measurements that are subsequently analyzed in the laboratory to estimate mass fractions of BC, OC, and other chemical PM components. In many of these studies, results are obtained using dynamometers to achieve preestablished engine loads, standardized driving cycles, and controlled sampling conditions (e.g., Zhen et al., 2009; Cadle et al., 2009; Khalek et al., 2015). Recent technological advancements have allowed for the direct measurement of BC emissions from diesel-powered vehicles under real-world driving conditions using mobile laboratories (e.g., Thornhill et al., 2010; Wang et al., 2012; Lau et al., 2015; Ježek et al., 2015) and tunnel studies (e.g., Geller et al., 2005; Ban-Weiss et al., 2008, 2009; Brimblecombe et al., 2015). Cross-road remote sensing studies and measurements obtained with an onboard portable emission measurement systems (PEMS) have also been used to characterize  $NO_x$ , CO, hydrocarbons (HC), and other gaseous emissions from heavy-duty diesel vehicles (e.g., Burgard et al., 2006; Frey at al., 2008; He et al., 2010; Carslaw and Rhys-Thyler, 2013).

Exhaust emission measurements obtained using on-road or roadside mobile laboratories, traffic tunnel sampling, cross-road remote sensing, and PEMS sampling techniques vary substantially in their sampling size, sampling time, captured driving modes, and pollutants sampled. For example, traffic tunnel sampling and cross-road remote sensing studies can sample hundreds of vehicles in relatively short periods but are limited in the range of driving conditions captured. In contrast, on-road exhaust plume interception studies with mobile laboratories and PEMS can provide large amounts of information on emissions under diverse driving conditions but are often limited in their sample size. Nevertheless, the overall results from the available studies have shown that there are important differences in the emission factors obtained under real-world driving conditions when compared to dynamometer-based studies. Furthermore, recent research suggests that in-use emissions of  $NO_x$  are routinely underestimated relative to certification standards (Anenberg et al., 2017; Franco et al., 2014). The differences arise because in real-world driving conditions there are multiple parameters (e.g., driving behavior, fuel quality, engine mechanical conditions, road conditions, etc.) that simultaneously affect the emission characteristics of on-road vehicles. These effects may not be properly captured under controlled tests (Ropkins et al., 2009). There have been some efforts to incorporate emissions and activity data obtained with PEMS into dynamometer-based tests to improve the representation of real-world driving conditions for heavy-duty diesel trucks, but there are still substantial challenges for standardizing the certification and compliance testing procedures (e.g., Zhen et al., 2009; Giechaskiel et al., 2016; Maricq et al., 2016). As mobile emission inventories should aim to accurately represent real-world driving conditions, there is a continuing need to better characterize on-road emission factors using realworld sampling techniques.

Current estimates suggest that on-road diesel vehicles are a major source of BC and other submicron carbonaceous particles in many parts of the world (Bond et al., 2013). However, the estimates are highly uncertain due to different assumptions about emission factors and the fraction of highemitting vehicles in developing countries' fleets. In Mexico, the most recent BC emissions estimates from the 2013 greenhouse gases and black carbon emission inventory (2013 GHG-BC MNEI) suggest that on-road vehicles contribute about 25 % of the total 125 Gg annual BC emissions (SE-MARNAT, 2015). However, due to a lack of locally obtained data, Mexico's BC and co-pollutants estimates for the diesel vehicle fleet were obtained using the default databases in the MOVES2014 EPA model (EPA, 2015) without adjusting emission factors or ancillary data. Therefore, there is a strong need to better characterize fine PM and gaseous pollutants emitted from diesel-powered vehicles in Mexico. In particular, the development of accurate emission factors and activity data for on-road vehicles is a critical step towards reducing uncertainties in Mexico's on-road emissions inventories.

In this pilot study we have estimated the fuel-based emission factors for BC, OC, CO,  $NO_x$ , and selected VOCs under real-world driving conditions for 20 on-road diesel vehicles in Mexico using the Aerodyne Research Inc. mobile laboratory (AML). The emission factors of NO, CO, HC, and fine PM were simultaneously obtained using the crossroad remote sensing technique, thus allowing for the intercomparison of the results obtained by the two techniques. The sampled vehicles included service trucks, Metrobuses, Turibuses, and intercity urban buses encompassing EPA98, EPA03, EPA04, and EURO3-5 emission level technologies. The results of this pilot study are useful to better understand the emission characteristics of the diesel vehicle fleet, to evaluate the emission factors used for the development of emissions inventories in Mexico, and to provide insights into diesel vehicle fleet emissions in other developing countries.

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Measurements were performed at Modulo 23, a large facility that is part of the Mexico City public transportation service (Red de Transporte de Pasajeros, or RTP), in collaboration with the Mexico City Secretariat of Environment (Secretaría del Medio Ambiente, or SEDEMA) during 25– 28 February 2013 as part of the field measurement campaign to characterize the emissions from key sources of short-lived climate forcers (SLCF-2013 Mexico). The Modulo 23 is typically used by RTP as a parking and maintenance facility for their public transport buses (see Fig. S1 in the Supplement). For this pilot project SEDEMA authorities redirected all of their scheduled RTP buses during the measurement period so that the parking area was empty and free of buses, except those selected for this study.

#### 2.1 Sampling techniques

#### 2.1.1 ARI mobile laboratory

The measurements were obtained using the AML by targeting on-road vehicles in "chase" and stationary roadside "exhaust plume sampling" techniques following the procedures described in Zavala et al. (2006). Tested vehicles were driven on prescribed routes inside and outside the Modulo 23 parking facility while the AML followed behind target diesel vehicles to continuously sample their exhaust emissions with fast-response onboard instrumentation (see the Supplement for a detailed description of the driving conditions). For these on-road chase measurements the AML's velocity and acceleration were also recorded continuously as the AML trailed target vehicles sampling their exhaust plumes for a variety of driving conditions. Emissions ratios are obtained by correlating the sampled exhaust plume gaseous and particle signals with above-background CO<sub>2</sub> and CO, which contain nearly all of the precombustion fuel carbon. Respective amounts of exhaust plume and background pollutant concentrations are determined by comparing background levels measured just before and after each plume encounter with those inside the exhaust plumes, effectively correcting for background and providing an exhaust emission ratio that can be used to obtain fuel-based emission factors (Zavala et al., 2006).

In addition to the on-road chase technique, the AML also employed the stationary roadside exhaust plume technique consisting of positioning the mobile laboratory downwind of the sampled target vehicles' exhaust. For instance, in collaboration with SEDEMA authorities, the mobile laboratory was strategically parked in one of the city's main bus rapid transit (Metrobus) passenger stations to measure the emission plumes of incoming and departing Metrobuses. Only lowspeed decelerating and accelerating plumes were sampled at this location. A total of 101 Metrobuses were sampled at the passenger station, encompassing multiple model years, manufacturers, and engine emissions tiers.

The measurement of vehicle emissions with the mobile laboratory is possible due to the use of high-time-resolution onboard instrumentation that is capable of capturing the highly transient conditions of the sampled plumes. BC and OC were measured using a soot particle aerosol mass spectrometer (SP-AMS) developed by ARI (Onasch et al., 2012). The application of the SP-AMS for the characterization of real-world vehicle emissions has been described in detail by Dallmann et al. (2014). The SP-AMS uses the laser-induced incandescence of absorbing soot particles to vaporize both the coatings and black carbon cores of exhaust soot particles within the ionization region of the AMS, thus providing a high-sensitivity measurement of the refractory BC (rBC) mass and the particle organic and inorganic coating materials (see Petzold et al., 2013). For simplicity, we refer to rBC as BC in this paper. The detection limits in the mass spectrum mode of the SP-AMS for BC and OC were 30 and  $60 \text{ ng m}^{-3}$ , respectively, with a nominal time resolution of 1 s. In addition to BC and OC, the SP-AMS measures other inorganic PM components including nitrates, sulfates, ammonium, and chlorides corresponding to a particle size range of 35 nm-1 µm. In this paper, we refer to PM emission factors obtained with the mobile laboratory as the sum of the BC, OC, and inorganic components simultaneously measured with the SP-AMS for each sampled vehicle.

Additional instruments were deployed in the AML to characterize the gaseous pollutants of the sampled vehicles. Quantum cascade tunable infrared laser differential absorption spectrometers (QC-TILDASs) were used to measure CO, SO<sub>2</sub>, ethane (C<sub>2</sub>H<sub>6</sub>), and acetylene (C<sub>2</sub>H<sub>2</sub>) (Dallmann et al., 2013). A proton transfer reaction mass spectrometer (PTR-MS) operated using H<sub>3</sub>O<sup>+</sup> as the ionization reagent (Rogers et al., 2006) was run in multiple ion detection mode to measure selected VOCs. Species measured with the PTR-MS included acetaldehyde, benzene, toluene, and C2-benzenes (sum of C<sub>8</sub>H<sub>10</sub> isomers: xylenes + ethylbenzene and benzaldehyde). Two Thermo Electron 42i chemiluminescent detectors modified for fast-response

measurements of NO and NO<sub>y</sub> and a Li-Cor 6262 nondispersive infrared (NDIR) instrument for CO<sub>2</sub> and water vapor were also used. Calibrations of these instruments were checked using certified gas standards. Other instruments onboard the mobile laboratory included a global positioning system (GPS), a sonic anemometer, and a video camera. Further details on the typical detection limits of the ARI instruments are presented in Table S1 of the Supplement.

#### 2.1.2 Remote sensing

A remote sensing (RS) unit model 4650 developed by Environmental Systems Products was deployed in a location near the start of the prescribed routes inside the Modulo 23 (see Fig. S1 in the Supplement). The RS unit included both low-level (0.15 m) and high-level (3.9 m) sampling elevations for measuring exhaust emissions, which is an important advan-

					Sampling	g size*
Vehicle type	Vehicle ID	Make	Model year	Tier	AML	RS
Service truck	ST1	Freightliner	1998	EPA98	22	3
	ST2	Freightliner	1998	EPA98	21	7
	ST3	International	2011	EPA04	14	6
	ST4	Freightliner	2006	EURO3	15	4
	ST5	HINO	2011	EURO4	14	4
	ST6	Kenworth	2010	EPA04	15	4
	ST7	Mercedes-Benz	1995	EPA94	4	4
	ST8	Freightliner	1999	EPA98	9	3
	ST9	Freightliner	1999	EPA98	8	3
Urban bus	UB1	International	2002	EPA98	38	8
	UB2	International	2009	EPA04	21	6
	UB3	Mercedes Benz	2002	EPA98	29	6
	UB4	Mercedes Benz	2009	EPA04	9	6
	UB5	Dina	2013	EURO5	15	15
Metrobus	MT1	Scania	2005	EURO3	9	5
	MT2	Volvo	2009	EURO4	9	8
	MT3	Mercedes Benz	2011	EURO5	10	13
	MT4	Volvo	2012	EURO5	6	8
Turibus	TU1	Scania	2006	EURO3	6	5
	TU2	Scania	2002	EURO3	8	5

 Table 1. Summary of sampled vehicle characteristics.

\* Sampling size refers to the number of emission factors obtained with the mobile laboratory (AML) and remote

sensing (RS) unit from the prescribed routes. See text for an explanation of the sampling periods for each technique.

tage when characterizing emissions from diesel-powered vehicles that have elevated tailpipes. The grade at the Modulo 23 is  $0^{\circ}$ . The location of the RS close to the start of the prescribed route was selected on the basis of obtaining an accelerating plume from the tested vehicle. Speed bar detectors were used to obtain vehicle speed and acceleration at the moment of passing through the RS unit. A video camera was placed down the road from the RS unit to take pictures of license plates when triggered.

In the RS, an NDIR exhaust gas analyzer with an optical filter of a wavelength known to be uniquely absorbed by the molecule of interest is placed in front of each detector, determining its specificity. The light source is shone across the road and reflected back. Reduction in the signal caused by the absorption of light by the molecules of interest reduces the detector's signal, and thus the number of molecules of the pollutant can be inferred. The RS instrument measures CO<sub>2</sub>, CO, and total HC (as propane equivalents) using infrared light, whereas ultraviolet spectrometers are used for NO and NO<sub>2</sub>. PM levels are not directly measured but inferred from a "smoke factor" estimated by the manufacturer from the ultraviolet and infrared absorption (Schuchmann et al., 2010). The target gas analyzers were calibrated daily with a mixture of certified gases. Technical specifications on the accuracy of the RS unit instruments are included in Table S2 of the Supplement.

#### 2.2 Vehicles sampled

Vehicles sampled in this pilot study included nine service trucks, four Metrobuses, two Turibuses, and five urban (RTP) buses encompassing models years 1995 to 2011 and EPA98, EPA04, and EURO3-5 technologies (Table 1 and Fig. S2 in the Supplement). RTP urban buses are single two-axle vehicles used for public transport with typical capacities of 60-90 passengers. RTP urban buses typically start operations very early in the morning and are continuously driven using designated intracity routes until nighttime when they are returned to Modulo 23 for regular maintenance, refueling, and overnight parking. Thus, RTP urban buses are continuously used throughout the day and often driven in low-speed but intense urban traffic conditions. Metrobuses are buses of one or two (merged) units that are used for transporting a large number of passengers (typical capacity is about 170 passengers) and have a dedicated (exclusive) driving lane on their route roads. The intracity routes of Metrobuses are selected for connecting highly populated but largely separated areas in Mexico City using in-between passenger stations. Since no other vehicles are allowed to travel in the designated lanes, Metrobuses often are driven at higher speeds than the rest of the surrounding fleet and are less affected by traffic.

Turibuses are double-decker buses that take passengers on guided tours through the main touristic landmarks of the city. Turibuses are usually driven at lower speeds than the Metrobuses, with gentler driving modes, and are well maintained. The service trucks tested were medium class 7 diesel trucks used for transporting goods for Coca Cola FEMSA. The sampled service trucks are typically driven in urban roads and are subject to intense traffic conditions. All vehicles sampled used ultra-low sulfur diesel (ULSD; <15 ppm in S) except the Turibuses, which used biodiesel with a B20 blend. All tested vehicles were ballasted in normal load operating conditions either with actual goods (for service trucks) or volunteers (for RTP buses, Metrobuses, and Turibuses) during the measurements.

#### 2.3 Data processing

Data from the two sampling techniques were processed to obtain fuel-based emission factors using established analytical protocols for the ARI mobile laboratory as described in detail in Zavala et al. (2006) and for the RS measurements as described in Bishop et al. (2008). In essence, the estimation of fuel-based emission factors for the two techniques is based on obtaining the mass ratio of the species of interest,  $m_i$ , to the total carbon mass found in above-background CO<sub>2</sub> and CO,  $m_{CO_2}$ , and  $m_{CO}$  (and HC in the case of RS) normalized by the mass fraction carbon content of the diesel fuel  $w_c$  (assumed as 0.87) as shown in Eq. (1):

$$\mathrm{EF}_{i} = \frac{\int m_{i} \mathrm{d}t}{\int \left(m_{\mathrm{CO}_{2}} + m_{\mathrm{CO}}\right) \mathrm{d}t} w_{\mathrm{c}}.$$
(1)

For the AML technique the gaseous species mass ratio is obtained using the moles of the pollutant and the total moles of emitted carbon by multiplying with their respective molecular weights, whereas the PM component measurements are directly obtained in  $\mu gm^{-3}$ , which is therefore the denominator unit for the total carbon content to be converted accordingly to  $\mu g Cm^{-3}$  using the respective CO<sub>2</sub> and CO molecular weights. The RS technique measures the difference in the number of molecules of the pollutant *i* before and after the sampled vehicle passes through the detector; thus the fuelbased emission factor is estimated from the ratio of emitted *n* moles of *i* to *n* moles of CO<sub>2</sub>  $(n_i/n_{CO_2})$  by dividing it by the sum of carbon moles in the  $CO_2$ , CO, and HC ratios to CO<sub>2</sub> (1,  $n_{\rm CO}/n_{\rm CO_2}$ , and  $n_{\rm HC}/n_{\rm CO_2}$ , respectively) and multiplying by the corresponding molecular weights and  $w_c$ . As described above, the PM levels in the RS technique are estimated from opacity measurements using a smoke factor to scale the absorption reading to grams of PM / grams of fuel and are thus semi-quantitative in nature, but they are useful for intercomparing vehicle emissions within the same experiment. For all the analyses, standard temperature and pressure conditions were used.

The important differences in the data analysis for the two techniques arise with the sampling frequency and thus with the integration periods ( $\Delta t$ ) used to calculate the emission factors. In the mobile laboratory technique, an emission fac-



**Figure 1.** Comparison of average fuel-based emission factors  $(g kg^{-1} fuel)$  between the mobile laboratory and remote sensing techniques by vehicle type. Variability bars represent 1 standard deviation of the observed values. \* Turibuses are fueled by biodiesel B20.

tor is typically obtained from multiple individual plume periods of 5–20 s depending on the truck velocity and wind conditions following the procedures described in Zavala et al. (2006). In this pilot study, each vehicle was sampled multiple times on prescribed routes with the mobile laboratory (see sampling size in Table 1) for about 3–10 min each time, therefore capturing hundreds of individual plume measurements for each vehicle. In the RS technique, the light source travels multiple times back and forth between the detectors during the short time of the passing plume of the targeted truck and the integration period is close to 1 s. Thus, the resulting estimated emission factor represents a snapshot for the driving condition at the time when the vehicle is passing through the detector.

#### 3 Results

Table 2 shows the average and 1 standard deviation of gaseous and PM fuel-based emission factors measured with the AML and the RS techniques for each of the sampled vehicles. The table also includes the results of the emission factors measured for the Metrobuses in stationary sampling mode as described above. Since the measurements were obtained under similar prescribed driving routes, differences in the results mainly reflect variability among vehicle engines and emission control characteristics. The large standard deviations reported in Table 2 indicate that most vehicles presented high variability in their emission factors under the prescribed driving routes. The urban bus UB1 was visually identified as a high emitter during the experiments due to its intense black smoke exhaust plumes, and this is confirmed by the much higher emission factors for this vehicle. Additionally, the relatively newer Dina urban bus with EURO5 technology had a malfunctioning selective catalytic reduction (SCR) device.

The average of the measured emission factors by vehicle type are shown in Figs. 1 and 2. Figure 1 shows a compar-

TU2       22.1 (11)       36.1 (12) $ $ 16.1 (4)       15.5 (2)       26.7 (6)       3.0 (0.7)       0.17 (0.08)       0.06 (0.04)       0.23 (0.1)       0.08 (0.01)       0.12 (0.04)       0.11 (0.03)       0.76 (0.8)       0.78 (0.3)       0.04 (0.01)       5.03 (1.8)         Numbers in parenthesis represent standard deviations. AML and RS stand for mobile laboratory and remote sensing techniques. Vehicle identification codes can be found in Table 1. <sup>2</sup> C2-B corresponds to the sum of $C_8H_{10}$ isomers. "Inorg," represents the sum of ammonium, chloride, sulfates, and incate from the AML are expressed as MO <sub>2</sub> equivalents.         routine equivalents. ''O <sub>2</sub> , emission factors from the AML are obvioued from 0.5 epitied to absorption measurements. <sup>4</sup> MTs represent the measurements obtained in stationary mode for 101 Metrobases. See text for further details. <sup>5</sup> Total HC emission factors expressed as more activated as the stationary mode for 101 Metrobases. See text for further details. <sup>5</sup> Total HC emission factors expressed as more activated as the stationary mode for 101 Metrobases. See text for further details. <sup>5</sup> Total HC emission factors expressed as the stationary mode for 101 Metrobases. See text for further details. <sup>5</sup> Total HC emission factors expressed as the stationary mode for 101 Metrobases. See text for further details. <sup>5</sup> Total HC emission factors from the AML are expressed as the stationary mode for 101 Metrobases.	0.8) $0.78$ $(0.3)norg" represents the sum ofor further details. 5 Total$	0.8) norg" rej for furth	0.76 (0.8) ) isomers. "Inorg" lses. See text for fr	0.11 (0.03) the sum of $C_8H_{10}$ de for 101 Metrobu	0.12 (0.04) 2-B corresponds to d in stationary mo	0.08 (0.01) und in Table 1. <sup>2</sup> ( tsurements obtain	0.23 (0.1) on codes can be for s represent the mea	0.06 (0.04) Vehicle identificatio asurements. <sup>4</sup> MT	0,17 (0.08) nsing techniques. <sup>v</sup> d to absorption me	3.0 (0.7) ry and remote ser ke factor" applie	26.7 (6) mobile laborato ned from a "smo O <sub>2</sub> equivalents.	15.5 (2) nd RS stand for expressed as N	16.1 (4) viations. AML a s from remote so om the AML are	36.1 (12) eent standard dev MS. <sup>3</sup> PM value ission factors fro	TU2 22.1 (11) 36.1 (12) $ $ 16.1 (4) 15.5 (2) 26.7 (6) 3.0 (0.7) 0.17 (0.08) 0.06 (0.04) 0.23 (0.1) 0.08 (0.01) 0.12 (0.04) <sup>1</sup> Numbers in parenthesis represent standard deviations. AML and RS stand for mobile laboratory and remote sensing techniques. Vehicle identification codes can be found in Table 1. <sup>2</sup> C2-B corresponding the standard deviation factor from tenote sensing are obtained from a "snoke factor" applied to absorption measurements. <sup>4</sup> MTs represent the measurements obtained in stationary propane equivalents.	TU2 <sup>1</sup> Numbers nitrate meas propane equ
(0.1) 0.16 (0.1) 0.07 (0.02) 0.50 (0.05) 0.04 (0.01)	) 0.16 (0.1) 0.07 (0.02)	) 0.16 (0.1) 0.07 (0.02)		(0.1)	0.16 (0.1	0.09 (0.01)	0.27 (0.1)	0.16 (0.1)	0.06 (0.03)	12.7 (4.0)	29.1 (6)	21.9 (3)	19.0 (5)	33.2 (17)	24.1 (7)	TUI
0.50 (0.5) 0.40 (0.4) 0.99 (1.2) 0.84 (0.7) 0.14 (0.04)	) 0.40 (0.4) 0.99 (1.2)	) 0.40 (0.4)	Ŭ	0 (0.5)	0.5	0.27 (0.3)	0.51 (0.7)	0.20 (0.2)	0.18 (0.2)		21.2 (8)		13.5 (5)		53.9 (44)	$\mathrm{MTs}^4$
0.15 (0.04) 0.12 (0.04) 0.23 (0.1) 0.40 (0.1) 0.03 (0.01)	) 0.12 (0.04) 0.23 (0.1)	) 0.12 (0.04)	)	5 (0.04)	0.1:	0.07 (0.02)	0.16 (0.04)	0.06 (0.02)	0.13 (0.03)	6.0 (7.3)	22.6 (4)	12.7 (4)	14.3 (3)	28.4 (25)	7.8 (5)	MT4
0.90 (0.3) 0.29 (0.1) 0.27 (0.2) 0.44 (0.1) 0.03 (0.01)	0.29 (0.1) 0.27 (0.2)	0.29 (0.1)	-	.90 (0.3)	0	0.14 (0.05)	0.26 (0.1)	0.17 (0.1)	0.22 (0.1)	0.3(0.1)	37.0 (6)	21.5 (5)	21.1 (4)	24.9 (16)	43.6 (23)	MT3
0.37 (0.2) 0.33 (0.1) 0.82 (0.4) 0.91 (0.3) 0.06 (0.01)	0.33(0.1) $0.82(0.4)$	0.33 (0.1)	-	.37 (0.2)	c	0.17 (0.1)	0.26 (0.1)	0.08 (0.03)	0.25 (0.1)	2.4 (1.3)	31.8 (5)	24.6 (10)	18.3 (4)	32.9 (40)	67.5 (57)	MT2
0.19 (0.1) 0.18 (0.1) 3.64 (1.7) 4.15 (1.8) 0.03 (0.01)	) 0.18 (0.1) 3.64 (1.7)	) 0.18 (0.1)	)	0.19 (0.1)	(	0.13 (0.03)	0.18 (0.04)	0.09 (0.03)	0.16 (0.1)	10.7 (6.2)	29.0 (2)	17.6 (3)	15.4 (2)	30.0 (11)	35.6 (14)	MT1
0.27 (0.2) 0.17 (0.1) 0.12 (0.1) 0.57 (0.4) 0.06 (0.02)	$0.17\ (0.1) \qquad 0.12\ (0.1)$	0.17 (0.1)	Ũ	0.27 (0.2)		0.11 (0.1)	0.17 (0.1)	0.05 (0.02)	0.41 (0.1)	6.0 (4.3)	58.9 (7)	35.8 (6)	32.8 (7)	15.3 (6)	28.6 (14)	UB5
$0.13\ (0.1) \qquad 0.13\ (0.1) \qquad 0.58\ (0.2) \qquad 0.88\ (0.3) \qquad 0.04\ (0.01)$	) 0.13 (0.1) 0.58 (0.2)	) 0.13 (0.1)	)	0.13(0.1)		$0.10\ (0.03)$	0.15 (0.04)	0.05 (0.03)	0.12 (0.1)	0.5 (0.1)	31.2 (8)	21.9 (6)	18.2 (5)	19.4 (9)	23.4 (10)	UB4
0.23 (0.2) 0.28 (0.3) 1.01 (1.4) 0.85 (0.6) 0.05 (0.02)	0.28 (0.3) 1.01 (1.4)	0.28 (0.3)	-	0.23 (0.2)		0.14 (0.1)	0.36 (0.4)	0.08 (0.03)	0.24 (0.4)	4.4 (2.8)	28.8 (10)	17.1 (4)	18.0 (7)	17.2 (4)	33.6 (15)	UB3
0.23 (0.1) 0.21 (0.1) 0.30 (0.9) 0.55 (0.7) 0.04 (0.01)	0.21 (0.1) 0.30 (0.9)	0.21 (0.1)	Ũ	0.23 (0.1)		0.13 (0.1)	0.23 (0.1)	0.08 (0.04)	0.21 (0.1)	5.4 (6.5)	39.9 (8)	21.1 (4)	20.3 (8)	17.4 (14)	23.5 (13)	UB2
0.60 (0.7) 0.58 (0.7) 10.37 (11.6) 4.50 (2.8) 0.10 (0.03)	) 0.58 (0.7) 10.37 (11.6)	) 0.58 (0.7)	Ŭ	0.60 (0.7)		0.30 (0.2)	0.34 (0.2)	0.39 (0.3)	0.47 (0.4)	5.2 (3.9)	21.4 (7)	10.1 (2)	11.9 (5)	34.3 (15)	140.3 (131)	UB1
0.65 (0.9) 0.39 (0.4) 2.19 (1.6) 1.50 (0.8) 0.08 (0.01)	0.39 (0.4) 2.19 (1.6)	0.39 (0.4)	-	0.65 (0.9)		0.21 (0.2)	0.38 (0.3)	0.12 (0.1)	0.51 (0.2)	3.5 (0.6)	21.0 (6)	14.8 (5)	14.9 (8)	14.3 (5)	33.8 (13)	ST9
$0.22\;(0.1) \qquad 0.18\;(0.1) \qquad 0.40\;(0.2) \qquad 0.80\;(0.2) \qquad 0.03\;(0.01)$	0.18 (0.1) 0.40 (0.2)	0.18 (0.1)	Ū	0.22 (0.1)		0.12 (0.03)	0.26 (0.1)	0.11 (0.03)	0.42 (0.2)	2.4 (1.7)	21.9 (6)	13.1 (1)	11.8 (2)	26.2 (3)	26.7 (4)	ST8
0.18 (0.1) 0.20 (0.02) 0.14 (0.1) 2.07 (0.2) 0.03 (0.01)	$)  0.20 \ (0.02) \qquad 0.14 \ (0.1)$	) 0.20 (0.02)		0.18 (0.1)		0.13 (0.04)	0.50 (0.2)	0.11 (0.03)	0.35 (0.1)	21.4 (3.1)	28.9 (6)	18.0 (2)	18.4 (4)	23.6 (7)	23.1 (3)	ST7
0.38 (0.3) 0.30 (0.2) 0.74 (0.7) 0.92 (0.6) 0.06 (0.01)	) 0.30 (0.2) 0.74 (0.7)	) 0.30 (0.2)	Ŭ	0.38 (0.3)		0.18 (0.1)	0.37 (0.3)	0.08 (0.1)	0.43 (0.1)	2.5 (0.6)	29.2 (6)	18.5 (7)	16.1 (3)	15.4 (5)	22.0 (24)	ST6
0.30 (0.2) 0.25 (0.1) 0.72 (0.6) 0.85 (0.4) 0.06 (0.01)	0.25(0.1) $0.72(0.6)$	0.25 (0.1)	Ũ	0.30 (0.2)		0.18 (0.1)	0.54 (0.1)	0.18 (0.06)	0.63 (0.1)	2.5 (0.6)	17.6 (3)	13.8 (3)	11.5 (3)	78.8 (7)	61.9 (17)	ST5
0.58 (0.8) 0.52 (0.8) 0.96 (0.5) 1.11 (0.6) 0.11 (0.03)	) 0.52 (0.8) 0.96 (0.5)	) 0.52 (0.8)	Ŭ	0.58 (0.8)		0.26 (0.3)	0.49 (0.7)	0.12 (0.1)	0.72 (0.4)	8.0 (1.3)	45.5 (14)	35.8 (5)	33.8 (25)	23.4 (4)	32.8 (18)	ST4
0.23 (0.2) 0.31 (0.4) 0.71 (0.6) 1.05 (1.3) 0.09 (0.05)	) 0.31 (0.4) 0.71 (0.6)	) 0.31 (0.4)	Ŭ	0.23 (0.2)		0.12 (0.1)	0.24 (0.1)	0.05 (0.04)	0.40 (0.2)	2.5 (0.9)	36.1 (8)	14.2 (1)	24.9 (20)	11.1 (1)	17.0 (9)	ST3
0.21 (0.1) 0.19 (0.1) 2.01 (1.4) 1.49 (0.7) 0.04 (0.01)	) 0.19 (0.1) 2.01 (1.4)	) 0.19 (0.1)	)	0.21 (0.1)		0.10 (0.04)	0.17 (0.1)	0.06 (0.03)	0.16 (0.1)	2.0 (1.1)	19.7 (3)	9.8 (1)	11.4 (3)	15.2 (5)	20.3 (10)	ST2
0.24 (0.2) 0.23 (0.2) 0.59 (0.5) 0.88 (0.8) 0.14 (0.1)	0.23 (0.2)  0.59 (0.5)	0.23 (0.2)	-	0.24 (0.2)		0.16 (0.2)	0.15 (0.1)	0.12 (0.1)	0.16 (0.1)	0.8 (0.2)	23.4 (5)	12.5 (1)	13.5 (3)	17.1 (1)	24.3 (15)	ST1
AML AML AML AML AML	AML AML	AML		AML		AML	AML	AML	AML	RS	AML	RS	AML	RS	AML	Ð
Toluene $C2-B^2$ BC OC $Inorg^2$	C2-B <sup>2</sup> BC	$C2-B^2$	,	Toluene		Benzene	$C_2H_4O$	$C_2H_2$	$SO_2$	HC <sub>2</sub>	$NO_x^6$	0	NO		СО	

 Table 2. Measured on-road average fuel-based emission factors (g kg<sup>-1</sup> fuel)\*.



**Figure 2.** Comparison of average VOCs, PM components, and SO<sub>2</sub> fuel-based emission factors ( $g kg^{-1}$  fuel) measured with the mobile laboratory by vehicle type. PM inorganics refer to the sum of ammonium, chloride, sulfates, and nitrate measured with the SP-AMS. \* Turibuses are fueled by biodiesel B20.

ison of the CO and NO average emission factors obtained with the AML and the RS techniques, whereas Fig. 2 shows the average emission factors of BC, OC, PM inorganics, toluene, C2-benzenes, benzene, acetaldehyde, acetylene, and SO<sub>2</sub> measured with AML instruments. The inorganic component of PM was estimated as the sum of nitrate, sulfate, chloride, and ammonium measured with the SP-AMS. For consistency in the comparisons, the emission factors shown for Metrobuses in Figs. 1 and 2 do not include data obtained in stationary sampling mode but only those obtained during the co-sampling of the two techniques. Non-paired Wilcoxon rank tests indicate that there is statistically significant difference (at the 0.05 significance level) between emission factors from service trucks, urban buses, and Metrobuses for the CO,  $NO_x$ , and SO<sub>2</sub> pollutants, whereas their corresponding VOCs, BC, OC, and PM inorganic emission factors were not significantly different. VOCs, BC, and PM inorganic emission factors from biodiesel-fueled Turibuses were significantly different from the corresponding emission factors from service trucks, urban buses, and Metrobuses.

The results show that the urban RTP buses produced the highest gaseous and PM emission factors. Conversely, the two sampled Turibuses running on biodiesel produced the lowest emission factors, particularly for BC, OC, and aromatics. The results also show that overall the emission factors measured with the remote sensing technique presented smaller variability with respect to those measured with the mobile laboratory. The higher variability observed with the mobile laboratory is likely the result of the larger range of driving conditions captured with this technique along the sampling routes, whereas the emission factors measured with the RS technique capture only the driving conditions at the time when the vehicle passes the detectors.



**Figure 3.** Box plots of PM emission factors measured by control technology. The numbers in parenthesis represent the number of sampled vehicles. Upper vertical central lines, upper level box lines, middle horizontal lines, lower box lines, and lower vertical central lines represent 90, 75, 50, 25, and 10% of the data. The first box plot of the EPA98 technology category includes the high emitter vehicle UB1 (see Table 2), whereas the adjacent box plot does not include this vehicle. PM was obtained as the sum of BC, OC, chlorides,

ammonium, sulfates, and nitrate components measured by the SP-

AMS.

Figure 3 shows a comparison of PM emission factors classified by vehicle control technology. Despite the small sampling size, the results indicate that there are marked differences between PM emissions depending on the vehicle's emissions control technology. Overall, the comparisons indicate lower PM emission factors due to improved control technology. The average PM emission factor decreases from 4.3 to  $0.72 \text{ g kg}^{-1}$  for vehicles with EURO3 to EURO5 technologies, respectively. However, the results also show that there is a large effect on the average PM emission factors when the data from the high-polluting vehicle UB1 (EPA98) are included in the comparison. The average PM emission factor for the EPA98 category is reduced from 5.7 to 2.0 g kg<sup>-1</sup> when the large emissions from this high-polluting vehicle are not included. Data from the single older 1995 ST7 vehicle with EPA94 technology were not included in Fig. 3 due to the relatively smaller sampling size (see Table 1). However, its average PM emission factor of  $2.24 \text{ g kg}^{-1}$  is slightly higher than the 2.0 g kg<sup>-1</sup> average of the EPA98 technology excluding the high-emitting vehicle, which is consistent with the observed higher PM emission factors for vehicles with older technologies. Non-paired Wilcoxon rank tests indicate that there is statistically significant difference (at the 0.05 significance level) between the PM2.5 emission factors obtained for the EPA98 and EPA04 control technologies and among the EURO3, EURO4, and EURO5 technologies. However, the results for the EPA98 and the EURO3 technologies were not significantly different.



Figure 4. Comparison of (a) CO and (b) NO fuel-based emission factors measured with the mobile laboratory and remote sensing techniques. Dashed lines represent 1 : 1.5, 1 : 1, and 1 : 0.25 lines.

As shown in Fig. 1, the comparison of average CO and NO emission factors by vehicle type suggests an overall good agreement between the mobile laboratory and the remote sensing techniques, particularly for NO. However, rather than comparing the averages of emission factors, a proper comparison accounting for the actual co-sampling periods between the two techniques is required. Figure 4 shows the comparison of the individual CO and NO emission factors measured for each co-sampled vehicle. Since the remote sensing technique measures the emission factor of the sampled vehicle only while it passes through the detectors, only the emission factors obtained with the mobile laboratory  $\sim$  10 s before and up to the corresponding actual moment of co-sampling with the remote sensing detector were considered for the comparison between the two techniques. Thus, we assume that a time period of 10s is sufficient to capture a large portion of the emission plume sampled by the mobile laboratory.

Figure 4 shows a linear but disperse correlation of the NO emission factors and a poor linear correlation of CO emission factors between the two techniques. Paired t tests indicate that there is no statistically significant difference (at the 0.05 significance level) between the two measurement techniques for both cases of CO and NO<sub>x</sub> emission factors. Arguably, the results show an overall good agreement between the two techniques in terms of their ability to distinguish low and high CO and NO emitters; however, there is some indication that the agreement varies substantially by vehicle type. Whereas the overall Pearson linear correlation coefficient (R) between the two techniques is only 0.60 (slope 0.25:1) for CO, the coefficient increases to 0.96 and 0.92 for Metrobuses and service trucks, respectively. The lower CO emission factors for the UB1 high emitter measured by the RS in comparison with the AML contributed significantly to the overall small linear correlation coefficient: R increases to 0.78 if the CO data for the UB1 are not included in the comparison. Similarly, the overall R for NO emission factors between the two techniques is only 0.75 (slope 0.77 : 1), but it increases to 0.86 and 0.84 for service trucks and urban buses, respectively. Paired t tests indicate that there is no statistically significant difference (at the 0.05 significance level) between the two measurement techniques for both cases of CO and NO emission factors. Although the sampling size may be too small to provide a more precise quantification of the agreement between the two techniques, the results suggest that overall both techniques can be used to adequately distinguish between high and low emitters but that distinction should consider the sampling efficiency by vehicle type.

#### 4 Discussion

#### 4.1 Comparison with MOVES2014 Mexico model

MOVES2014 is currently the most advanced model for estimating on-road emissions in the US at the national, state, county, and project level as it incorporates emissions data obtained from field studies over a wide range of vehicle types, pollutants, emission processes, fuel types, and operating modes (EPA, 2015). A number of studies indicate that the use of the model can improve the emissions estimates of inventories in Mexico with respect to older emission models (Zavala et al., 2013; Guevara et al., 2017). However, its efficient application to other countries requires the adjustment of multiple internal parameters, among which the emission factor databases are of key importance. A recent project was developed by the Eastern Research Group (ERG) to adjust the model's default emission factors and deterioration rates for the gasoline fleet using remote sensing data obtained in major Mexican cities (Koupal et al., 2016). The resulting model, MOVES2014 Mexico, also considers Mexican vehicle emissions and fuel quality standards, vehicle population by age and state, fuel properties, and fuel consumption. However, emission factors for the diesel fleet in the model were not adjusted using field measurement data.

The heavy-duty emission exhaust database for the MOVES2014 model's previous version (MOVES2010) was originally constructed using the results of several real-world in-use dedicated studies for gaseous pollutants, including (1) measurements of 124 trucks and buses with model years 1999 through 2007 using the Real-time On-Vehicle Emissions Reporter (ROVER) PEMS system developed by EPA and (2) measurements of 188 trucks with model years 1994 through 2003 using the Mobile Emissions Measurement System (MEMS) by West Virginia University (WVU). The current version of MOVES2014 builds upon these studies using two additional real-world studies: (1) the Heavy-Duty Diesel In-Use Testing (HDIU) program in which data were collected by manufacturers during normal operation for 243 diesel trucks of model years 2003-2009 and (2) the Houston Drayage Data (HDD) study in which the EPA collected emissions data from 27 trucks with model years 1991-2006 in drayage service using PEMS in the Houston-Galveston Area. Among other changes resulting from the emissions database updates, MOVES2014 estimates higher  $NO_x$  emission factors than MOVES2010 (EPA, 2015).

Databases of PM emission factors in MOVES2014 were constructed from the CRC E-55/59 research program that consisted of sampling 71 diesel vehicles with model years 1974–2004 (Clark et al., 2007). However, the PM speciation data were collected from only nine different vehicles using the WVU's Transportable Heavy-Duty Vehicle Emissions Testing Laboratory (EPA, 2014). Importantly, the measurements did not include transit buses, and thus the PM emission factors for the urban bus vehicle category were proportionated using data from other measured vehicle types.

Figure 5 shows a comparison between emission factors measured with the AML and those from the MOVES2014 Mexico model in the "exhaust" emission process category. The figure compares the measured emission factors of urban buses and Metrobuses with those estimated for the transit bus vehicle category in the model. Measured emission factors of service trucks are compared against those estimated for the single-unit short-haul truck vehicle category in the model. Turibuses are not included in this comparison. In addition, in the comparison only the vehicle age groups in the MOVES2014 Mexico model corresponding to the model years of the sampled vehicles are included.

The results indicate very good agreement between the modeled and measurement-based  $NO_x$  emission factors for both buses and service trucks, but suggest a significant model underestimation of CO emission factors. Model-based BC emission factors are well within the observed values for service trucks, but the results show higher variability in the measurements for the urban buses and Metrobuses compared to the model. The results also suggest a large underestimation of



Figure 5. Comparison between AML measurements and MOVES2014 Mexico emission factors.

OC emission factors in the model for both buses and service trucks. Interestingly, despite the underestimation of OC there is a better agreement between the model and measurements for the total PM emission factor that results from a compensating effect of overestimation of the inorganic PM components in the model. The measured emission factors for acetaldehyde, benzene, and toluene were all much higher than those obtained from the MOVES2014 Mexico model, which is consistent with the observed underestimation of CO emission factors.

Overall, the model underestimated the CO, OC, and selected VOCs but had better agreement for NO<sub>x</sub> and BC emission factors. Due to the small sampling size in this pilot study, caution should be exercised when attempting to extrapolate the results from this comparison to other vehicle categories and model years. Nevertheless, the results demonstrate the need for locally adjusting the emission factors database for the diesel vehicle fleet in the MOVES2014 Mexico model using real-world driving conditions to improve the emission estimates during inventory development.

Location and sampling year	Source type	Sampling technique	Mean and SD $(g kg^{-1} fuel)$	Reference
Mexico City, 2013	Turibus	Chasing	$0.41 \pm 0.7$	This study
Mexico City, 2013	Metrobus	Chasing	$1.24\pm1.8$	This study
Mexico City, 2013	Urban bus	Chasing	$2.48\pm7.3$	This study
Mexico City, 2013	Service trucks	Chasing	$0.94 \pm 1.1$	This study
Los Angeles, 1997	HDDV	Tunnel	1.3	Kirchstetter et al. (1999)
San Francisco, 2005	MDDT, HDDV	Tunnel	$0.78\pm0.09$	Geller et al. (2005)
San Francisco, 1997, 2006	MDDT, HDDV	Tunnel	$0.92\pm0.07$	Ban-Weiss et al (2008)
Mexico City, 2006	Diesel fleet	Chasing <sup>c</sup>	1.4 (1.3–1.6) <sup>a</sup>	Thornhill et al. (2010)
San Francisco, 2006	HDDV	Tunnel	$1.7\pm2.3$	Ban-Weiss et al. (2009)
Wilmington, CA, 2007	HDDT	Chasing <sup>c</sup>	0.5 (0.07–0.1) <sup>b</sup>	Park et al. (2011)
Beijing, 2009	HDDT	Chasing	2.2 (0.4-1.7) <sup>a</sup>	Wang et al. (2012)
Chongqing, 2010	HDDT	Chasing	1.6 (0.7–1.6) <sup>a</sup>	Wang et al. (2012)
Beijing, 2010	HDDT	Chasing	1.1 (0.2–0.8) <sup>a</sup>	Wang et al. (2012)
San Francisco, 2010	HDDT	Tunnel	$0.62\pm0.17$	Dallmann et al. (2014)
Los Angeles, 2011	HDDV freeways	Chasing <sup>c</sup>	$1.33\pm0.33$	Hudda et al. (2013)
Slovenia, 2011	Buses	Chasing	0.4 (0.24–0.65) <sup>b</sup>	Ježek et al. (2015)
Oakland, CA, 2011–2013	HDDT	Tunnel	$1.15 \pm 0.19$ $0.09 \pm 0.04$	Preble et al. $(2015)^d$
Hong Kong, 2013–2014	HDDV	Chasing	$2.2 \pm 0.3$	Lau et al. (2015)
Hong Kong, 2014	Diesel fleet	Tunnel	$1.28\pm0.76$	Brimblecombe et al. (2015)

Table 3. Comparison of measurements of BC emission factors from diesel-powered sources
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<sup>a</sup> Represent average and first and third quartiles of data. <sup>b</sup> Mean and 95 % confidence interval. HDDT: heavy-duty diesel vehicle; HDDT: heavy-duty diesel truck; MDDT: medium-duty diesel truck. <sup>c</sup> Includes chasing and fleet average values. <sup>d</sup> Sampling of individual plumes on an overpass. High BC value represents 2009 fleet with 2 % diesel particle filters (DPFs) and 0 % selective catalytic reduction (SCR) installed, whereas low BC value represents 2010–2013 trucks with full DPF and SCR systems installed.

#### 4.2 Comparison with other studies

A recent study by Sheinbaum et al. (2015) investigated the impacts on PM and NO<sub>x</sub> emission levels when using B10 and B20 biodiesel blends for three EPA98 and three EPA04 urban RTP buses in Mexico. They found mixed results on the emission benefits depending on the technology and blend composition. The average reductions of PM for the three EPA04 buses were 66 and 36 % using B10 and B20 blends, whereas the corresponding reductions for  $NO_x$  were 4 and 8 %. For the EPA98 buses PM increased 59 and 15 % when using B10 and B20 blends, respectively, whereas  $NO_x$  correspondingly increased 8 and 3 %. The two biodiesel TU1 and TU2 vehicles sampled in this study have EURO3 technology similar to the ST4 and MT1 vehicles. The results in Table 2 show that the CO and  $NO_x$  emissions factors of the TU1 and TU2 vehicles had no distinguishable differences with respect to the ST4 and MT1 vehicles. The major difference between the biodiesel-fueled vehicles and other vehicles is observed in the much smaller BC and OC emission factors. Therefore, these results also suggest PM emission reduction benefits when using the biodiesel in vehicles with newer technology.

Table 3 compares the measured BC emission factors in this study with those reported in other parts of the world obtained with various sampling techniques. In 2006, the AML measured emissions from mobile sources in Mexico City using the chase technique and applied the positive matrix factorization (PMF) method to obtain an average fleet-wide emission factor of  $1.4 \text{ g kg}^{-1}$  for the diesel fleet (Thornhill et al., 2010). This value is well within the ranges of the measured emission factors in this study and is similar to the values obtained in Beijing and Chongqing by Wang et al. (2012). Preble et al. (2015) found much smaller BC emission factors that corresponded to newer trucks with diesel particle filters (DPFs) and selective catalytic reduction (SCR) control technologies. Our results also indicate that there is a strong effect of control technology on BC emissions. Nevertheless, the results also demonstrate that the information on the fraction of high emitters in the diesel fleet in developing countries is a key parameter for the construction of emissions inventories. In addition, the values in Table 3 indicate that there is large variability in BC emission factors measured worldwide and at different times, highlighting the need for increasing the number of available datasets of emission factors obtained under real-world driving conditions to improve emissions inventory accuracy.

Dallmann et al. (2014) obtained an average BC emission factor of  $0.62 \pm 0.17$  in 2010 in San Francisco also using the SP-AMS instrument and found an OC / BC ratio of  $0.31 \pm 0.1$  for the diesel fleet. The OC / BC emission ratios in this study are much higher: 0.59, 1.19, 1.26, and 1.56 for urban buses, Metrobuses, service trucks, and Turibuses, respec-

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tively. The biodiesel Turibuses presented the larger OC / BC ratio although their BC emission factors were the smallest of all sampled vehicles. Several factors, including driving conditions, vehicle technology, and diesel fuel composition, can contribute to the observed differences, but the quantification of these contributions is beyond the scope of this study. Nevertheless, the higher organic content of the emissions in the sampled Mexican vehicles with respect to those measured in California by Dallmann et al. (2014) illustrates the large emission differences in PM composition that can be found in diesel fleets around the world, thus further indicating the need for locally adjusting the emission factor databases in mobile emission models.

The sampling size in this pilot study is too small to be a representative sample of the entire Mexican fleet. Nevertheless, there are some vehicle age and type characteristics that make the results relevant. According to the 2013 GHG-BC MNEI, the Mexican heavy-duty diesel fleet of about 810,000 vehicles is dominated by diesel trucks with a gross vehicle weight (GVW) of >3 tons ( $\sim 50.8 \%$ ) followed by large trailer trucks (29.4%), urban buses (12.5%), and smaller trucks with GVW < 3 tons (4.5%) (Fig. S3 in the Supplement). Thus, the sampled service trucks corresponding to diesel trucks with GVW > 3 tons, the urban RTP buses, and the Metrobuses belong to large categories of the diesel fleet. In addition, an analysis of the diesel-powered fleet distribution for Mexico City indicates that a large fraction of the inuse diesel vehicle fleet is relatively old and remains in use for longer periods compared to the gasoline vehicle fleet (Fig. S4 in the Supplement). For example, about 61.5 and 64.9 % of the buses and diesel trucks with GVW > 3 tons, respectively, are older than 10 years. The vehicle model years of the sampled service trucks (1995-2011) correspond to about 53.4 % of the diesel trucks with GVW > 3 ton fleet, whereas the model years of the sampled buses are relatively newer (2002-2011) and correspond to only about 36.6 % of the bus fleet.

#### 5 Conclusions

We present the results of measurements of fuel-based emission factors for BC, OC, CO,  $NO_x$ , and selected VOCs for diesel-powered service trucks, urban buses, Metrobuses, and Turibuses in Mexico under real-world driving conditions using the AML and the remote sensing sampling techniques. The results showed higher PM emissions factors for urban buses with older technologies than for the other vehicle types and a marked dependency on vehicle emission control technology. These results further demonstrate the benefits of tighter tier regulations and independent testing to verify the efficacy of reduced emissions standards for diesel vehicles.

The two biodiesel Turibuses presented smaller BC and OC emission factors. Although the effects of using biodiesel fuel could not be quantified in this study, the results suggest substantial emission benefits. Further dedicated studies with larger sampling sizes can help to quantify the benefits.

The comparison between the emission factors obtained with the two sampling techniques suggests that both techniques can be used to identify high and low vehicle emitters, but there are differences in sampling efficiency depending on the vehicle type sampled. In addition, higher variability was observed in the emission factors obtained with the mobile laboratory that likely results from the larger diversity of emission driving conditions captured with respect to the fixed-site remote sensing technique.

A comparison of the measured results with the emission factors estimated in the MOVES2014 Mexico model shows that the model underestimates CO, OC, and selected VOC species but that there is better agreement for  $NO_x$  and BC. The underestimation of organic components in the model is further supported by the larger OC / BC ratios found in comparison to ratios measured elsewhere with the same sampling technique. These results further demonstrate the need to locally adjust the emission factor databases for the diesel vehicle fleet when the MOVES2014 model is applied in countries other than the US in order to reduce the uncertainty in the emissions estimates and to improve the evaluation of the effectiveness of emissions reduction measures.

*Data availability.* The pertinent data described in the manuscript are provided in the Supplement. Additional information not included within the Supplement is available upon request.

# The Supplement related to this article is available online at https://doi.org/10.5194/acp-17-15293-2017-supplement.

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

*Disclaimer.* The use of the MOVES2014 Mexico model in this paper is for illustrative purposes and should not be considered as an evaluation of the model's performance.

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