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Comparison of measurements of peroxyacetyl nitrates and primary carbonaceous aerosol concentrations in Mexico City determined in 1997 and 2003

N. A. Marley, J. S. Gaffney, R. Ramos-Villegas, and B. Cárdenas González

1Chemistry Department, University of Arkansas at Little Rock, Little Rock, Arkansas, USA
2Secretariat of the Environment, Government of the Federal District, Mexico City, D.F., Mexico
3El Centro Nacional de Investigación y Capitación Ambiental, Instituto Nacional de Ecología, Secretaría de Medio Ambiente y Recursos Naturales, Mexico City, D.F., Mexico

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Abstract. The concentrations of peroxyacetyl nitrate (PAN) in ambient air can be a good indicator of air quality and the effectiveness of control strategies for reducing ozone levels in urban areas. As PAN is formed by the oxidation of reactive hydrocarbons in the presence of nitrogen dioxide (NO$_2$), it is a direct measure of the peroxyacetyl radical levels produced from reactive organic emissions in the urban airshed. Carbon soot, known as black carbon (BC) or elemental carbon (EC), is a primary atmospheric aerosol species and is a good indicator of the levels of combustion emissions, particularly from diesel engines, in major cities. Mexico City is the second largest megacity in the world and has long suffered from poor air quality. Reported here are atmospheric measurements of PAN and BC obtained in Mexico City during the Mexico Megacity 2003 field study. These results are compared with measurements obtained earlier during the Investigación sobre Materia Particulada y Deterioro Atmosférico – Aerosol and Visibility Research (IMADA-AVER) campaign in 1997 to obtain an estimate of the changes in emissions in Mexico City and the effectiveness of control strategies adopted during that time. Concentrations of PAN in 1997 reached a maximum of 34 ppb with an average daily maximum of 15 ppb. The PAN levels recorded in 2003 were quite different, with an average daily maximum of 3 ppb. This dramatic reduction in PAN levels observed in 2003 indicate that reactive hydrocarbon emissions have been reduced in the city due to controls on olefins in liquefied petroleum gas (LPG) and also due to the significant number of newer vehicles with catalytic converters that have replaced older higher emission vehicles. In contrast, black/elemental carbon levels were similar in 1997 and 2003 indicating little improvement likely due to the lack of controls on diesel vehicles in the city. Thus, while air quality and ozone production have improved, Mexico City and other megacities continue to be a major source of black carbon aerosols, which can be an important species in determining regional radiative balance and climate.

1 Introduction

Megacities are large urban and suburban complexes whose populations are in the tens of millions of inhabitants (Lynn, 1999). With the rapid growth of the world’s population and the continuing industrialization and migration of the populace towards major urban centers, the numbers of these megacities are increasing. Although New York City was the only megacity in the world in 1950, they now number 14 and their distribution is growing most rapidly in the tropical areas of South America and Asia. Today, the largest metropolitan complexes are centered at Tokyo, Japan, and Mexico City, Mexico (Molina and Molina, 2002a). Within the next 10 to 15 years it is predicted that there will be more than 30 megacities worldwide.

Megacities have become important global sources of air pollutants from the associated mobile and stationary sources and the emissions from megacities are leading to regional and global increases in many key trace gases as well as primary and secondary aerosols. The Mexico City metropolitan area (MCMA), which occupies ∼1300 km$^2$ with a population of ∼18 M, is one of the largest megacities in the world and is well known for its high levels of air pollution and visibility reduction (Molina and Molina, 2002b). It has been estimated that emissions from the Mexico City basin contribute 15 Mega-tons of fine aerosol (PM$_{2.5}$) per year to the surrounding regions (Gaffney et al., 1999). This fine aerosol is composed of approximately 32% organic carbon, 15% elemental
carbon, 10% ammonium nitrate and 20% ammonium sulfate (Chow et al., 2002). The emissions of sulfate aerosols alone from the MCMA are estimated to amount to 1% of the total global burden (Barth and Church, 1999). While sulfate is an important light scattering aerosol species, black carbon is an important light absorbing aerosol species. The aerosol emissions from megacities such as Mexico City therefore play potentially important roles in regional radiative balance and climate (Jacobson, 2002; Gaffney and Marley, 2005).

Elevated levels of ozone have been known for some time in Mexico City (Bravo et al., 1989; MARI, 1994; Streit and Guzman, 1996; Fast and Zhong, 1998). Mexico City is located at an altitude of 7200 ft in an air basin surrounded by mountains (Fast and Zhong, 1998; Doran et al., 1998) and at a latitude of 19° of North, which implies high levels of incoming solar radiation all year long. In the past, peak ozone levels exceeding 300 ppb were not uncommon, particularly during the late dry winter months of February and March (The one-hour standard is 0.11 ppm (Molina and Molina, 2002a)). These very high levels of ozone require high levels of reactive hydrocarbons as well as elevated levels of nitrogen oxide (NO) emissions to produce the coupled OH and peroxyradical chemistry that is key to the formation of urban ozone (Finlayson-Pitts and Pitts, 2000). Hydrocarbon measurements taken in Mexico City (Blake and Rowland, 1995) showed that the heavy domestic and commercial use of LPG has led to very high levels of butane and propane in that megacity’s air, exceeding parts-per-million (ppm) of carbon in many cases. Indeed, LPG was proposed as an important source of the reactive volatile organic carbon compounds (VOCs) propene and butenes, which could account for an appreciable portion of the observed urban ozone in Mexico City (Blake and Rowland, 1995). Volatile organic carbon samples collected before and during a Mexican national holiday with reduced automobile traffic clearly showed that mobile emissions are equally important as LPG as sources of reactive olefins such as the butenes (Gaffney et al., 1999). In addition, earlier work suggested that mobile sources contribute 75% of the total hydrocarbons to the Mexico City air (Riveros et al., 1998) and that NO₂ was probably the most important contributor to ozone production in Mexico City (Raga et al., 2001a).

Peroxyacetyl nitrates (PANs) are important indicator compounds of peroxyradical activity in an urban air shed (Finlayson-Pitts and Pitts, 2000; Gaffney et al., 1989). Formed by the reaction of peroxyacetyl radicals with NO₂, they exist in equilibrium with the peroxy radical species according to:

$$\text{RC} = \text{O} - \text{O} + \text{NO}_2 \rightarrow \text{RC} = \text{O} - \text{O} - \text{O} - \text{NO}_2 \quad (1)$$

where R is typically (in order of importance): CH₃ - (peroxyacetyl nitrate, PAN), CH₃CH₂ - (peroxypropionyl nitrate, PPN), and CH₂CH₃CH₂O - (peroxybutylnitrate, PBN) in an urban environment. The peroxyacetyl radicals are formed from the reactive olefins directly or via formation of aldehydes that can react with OH to form the peroxyacetyl species. While the PANs are thermally labile, the reverse reaction to reform the PANs is relatively fast leading to no net loss at elevated temperatures in areas of high NO₂ concentrations (Finlayson-Pitts and Pitts, 2000; Gaffney et al., 1989).

Thus, measurements of the temporal variability of the PANs are useful in determining the oxidative reactions involved in the formation of ozone as well as other secondary air pollutants and aerosol species such as nitric acid and ammonium nitrate as they are a direct measure of the peroxyacetyl radical formation activity. The PANs therefore serve as a measure of the peroxy radical concentrations in the atmosphere. Since the PANs have low aqueous solubilities, low reactivity with OH, and are slow to photolyze, they can lead to the transport of NO₂ over long distances and can simultaneously act as a reservoir for NO₂ during transport of the urban plume. This can have regional scale impacts on ozone, nitrate aerosols, and other pollutants associated with megacity plumes (Gaffney et al., 1989).

Carbonaceous particulate matter or “soot” particles are produced from the partial combustion of hydrocarbons, particularly from diesel fuels, and are therefore a measure of the combustion emissions in an urban area. Although the elemental composition of these particles is dominated by carbon (>90%), soot particles may be regarded as a complex organic polymer, rather than an amorphous form of elemental carbon (Andreae and Gelencser, 2006). The absolute identification of carbon soot is difficult and the techniques commonly used, such as Raman spectroscopy, electron microscopy, and mass spectrometry are impractical for routine monitoring of this material. Various measurement methods have been developed for the routine quantitation of aerosol soot content that make use of some of its characteristic properties. These methods have created operational definitions such as “black carbon” or “elemental carbon” depending on the key property being measured (Gaffney and Marley, 2006). The term “black carbon” arises from the use of optical attenuation methods and refers to the highly absorbing nature of carbon soot aerosols. In contrast, the term “elemental carbon” is used when thermal combustion methods are employed for detection and refers to the refractory nature of the aerosols at temperatures up to 350–400°C. Comparisons between the optical and thermal combustion methods in different environments have yielded correlation coefficients of 0.99 (Hansen and McMurry, 1990), 0.97 (Allen et al., 1999), and 0.94 (Babich et al., 2000) and the various terms used to identify combustion derived aerosols are commonly used interchangeably.

Although originally identified as a pollutant and a tracer for combustion emissions, carbon soot has more recently gained attention as a major light absorbing species with significant impacts on the radiative balance of the atmosphere (Ramanathan et al., 2005). Some model calculations suggest that the contribution of carbon soot aerosols to global warming may be as much as 0.3–0.4°C, rivaling the contributions from atmospheric methane (Jacobson, 2004; Chung and
Seinfeld, 2005). The ultimate climate effects from carbon aerosols will depend on their physical and chemical properties, as well as their residence times and distributions in the atmosphere (Jacobson, 2001). In order to adequately assess the aerosol impacts on global and regional climate, a better understanding of black carbon aerosol emission rates and atmospheric distributions will be essential.

In an attempt to better understand the Mexico City air chemistry with regard to oxidants and aerosols, a comprehensive collaborative study was carried out during February-March 1997 that included a wide variety of chemistry, aerosol and meteorological measurements (Edgerton et al., 1999; Doran et al., 1998; Fast and Zhong, 1998). This field campaign (IMADA-AVER) was jointly sponsored by the U.S. Department of Energy’s Office of Biological and Environmental Research Atmospheric Science Program (DOE/ASP) and Petroleos Mexicanos (PEMEX) through the Mexican Petroleum Institute (Instituto Mexicano de Petroleo, or IMP).

As part of IMADA-AVER, measurements of near-surface tropospheric PANs were made at the IMP laboratories to determine the concentrations and temporal variability of these species in the megacity (Gaffney et al., 1999). Levels of the PANs were found to approach 40 ppb in the central metropolitan area. Peroxyacetyl nitrate was the major species, although PPN and PBN were also observed in the low ppb ranges. These are the highest values of the PANs seen in any urban area since 50 ppb values were reported for PAN in the late 1970s downwind of Los Angeles in the south coast air basin in southern California (Tuazon et al., 1981). Volatile organic carbon measurements obtained at IMP during the same time indicated that automobiles were responsible for much of the reactive hydrocarbons in the atmosphere (Gaffney et al., 1999). The meteorological measurements demonstrated that the air basin was subject to a strong advection of the boundary layer in the afternoon leading to a clearing out of the pollutants emitted and formed during the day. This regional meteorology leads to very little carryover of the pollutants in Mexico City from day to day and atmospheric chemistry that is dominated by reactions that took place typically over a one day period (Gaffney et al., 1999; Fast and Zhong, 1998). The application of a simple box model indicated that considerable amounts of air pollutants, both oxidants and aerosols, were being transported out of the Mexico City air basin (Gaffney et al., 1999; Elliott et al., 1997).

In April 2003, the DOE/ASP again conducted a field study in Mexico City (Mexico Megacity 2003) in collaboration with the MCMA 2003 air quality study organized by M. J. Molina and L. T. Molina of the Massachusetts Institute of Technology. Continuous measurements of the PANs and black carbon aerosols were obtained during April 2003 at the National Center for Environmental Research and Training (Centro Nacional de Investigación y Capacitación Ambiental, or CENICA), on the Iztapalapa campus of the Universidad Autónoma Metropolitana (UAM). Data were collected before and during the Easter holiday to assess changes in PAN and BC loadings as a function of vehicle traffic levels.

Results are presented here for PAN and BC concentrations obtained in Mexico City during the Mexico Megacity 2003 field study. These results are compared with those obtained 6 years earlier during the IMADA-AVER study. Changes in PAN and carbonaceous aerosol levels are presented as an indication of changes in emission levels during that time and the effectiveness of ozone control strategies that have been adopted in Mexico City since 1997. The black carbon aerosol comparisons are of particular importance in the evaluation of the regional impacts of the changing megacity emissions of this key absorbing aerosol species and its importance for regional climate considerations.

2 Experimental methods

Measurements were obtained from February 20 to March 23, 1997 at Building No. 24 (Refinación y Petroquímica) of the IMP laboratories (Eje Central Lazaro Cardenas No. 152, Delegacion Gustavo A. Madero, México, Distrito Federal). This site is located in the north central part of Mexico City (19°29′19.392″N, 99°08′50.258″W). The PANs were determined using an automated gas chromatograph equipped with an electron capture detection system (GC/ECD). This system has been described in detail elsewhere (Gaffney et al., 1993, 1997, 1998, 1999). A 2-cm$^3$ sample was injected automatically onto a packed GC Carbowax 400 column every 30 min. Data were collected using a recording integrator and processed manually for each of 1380 samples. Calibration of the instrument for the PANs was accomplished by manual injection of standards synthesized by strong acid nitration of the corresponding peracids (Gaffney et al., 1984; Gaffney and Marley, 2005a).

Data for ozone and NO$_2$ concentrations were obtained from the IMP monitoring station of the Mexico City ambient air monitoring network (la Red Automática de Monitoreo Atmosférico, or RAMA), operated at that time by the Dirección General de Prevención y Control de la Contaminación de la Ciudad de México. Nineteen of the 33 urban RAMA stations measure ozone by UV absorption and nitrogen oxide (NO) and total nitrogen oxides (NO$_x$) by chemiluminescence among other criteria pollutants, which are reported as hourly averages. As the chemiluminescent nitrogen oxides analyzer in the NO$_x$ mode measures the sum of NO$_2$, NO and PAN concentrations, the 1997 NO$_2$ concentrations were estimated by subtraction of the NO and PAN concentrations from NO$_x$ results (NO$_2$=NO$_x$-NO-PAN).

Measurements of the PANs, NO$_2$ and fine aerosol black carbon content were obtained from 3 April to 1 May 2003 on the rooftop of the CENICA laboratory building (19°21′44.541″N, 99°04′16.425″W) on the UAM Iztapalapa
Fig. 1. (A) Concentration profiles for peroxyacetyl nitrate (PAN) measured in Mexico City from 20 February to 23 March 1997 and (B) from 3 March to 1 April 2003.

campus (Calle “Sur 10” No. 230, Colonia La Vicentina, Delegación Iztapalapa, México, D.F.). This site is approximately 16.1 km (10.0 mi) south-southwest of the site at the IMP site that was used in 1997. Measurements of the PANs and NO$_2$ were obtained by fast gas chromatography with luminol detection. This system has been described in detail elsewhere (Marley et al., 2004) but will be briefly reviewed here. A 2-cm$^2$ sample loop was used to automatically inject samples onto a 30-ft. capillary DB-1 column at 1-min intervals. Both NO$_2$ and PAN were detected by the chemiluminescent reaction with luminol and the intensity of the emission at 425 nm was measured with a photon counting module. The instrument was controlled by a 1.8-GHz Notebook computer with Windows 2000 operating system and a custom software application programmed in LabVIEW. The instrument was calibrated for NO$_2$ by dilution of a 2.8-ppm NO$_2$ tank standard in air with a gas calibrator (Dasibi; Model 1009-CP) and for the PANs with the synthetic standards described above. This instrument has been compared with the GC/ECD method in previous studies and found to give good agreement both in field studies and with synthetic PAN standards (Gaffney et al., 1998, 2005a; Marley et al., 2004).

The black carbon content of fine aerosols was measured by using a seven-channel aethalometer (Andersen) with a sample inlet designed to collect aerosols in the 0.1 to 2 micron size range. The aerosols in the air sample are collected within the instrument by continuous filtration through a paper tape strip. The optical transmission of the deposited aerosol particles is then measured sequentially at seven wavelengths (370, 450, 520, 590, 660, 880, and 950 nm). Since black carbon is a strongly absorbing aerosol species with an absorption coefficient relatively constant over a broad spectral region (Marley et al., 2001) the instrument can automatically calculate the black carbon content from the transmission measurements by assuming black carbon to be the main absorbing aerosol species in the samples with a 1/λ dependence typical of broadband absorbers and a mass specific absorption coefficient of 16.6 m$^2/g$ at 880 nm (Hansen et al., 1982). The instrument is operated by an embedded computer with a display screen and keypad that controls all instrument functions and automatically records the data to a built-in 3.5” floppy diskette. Data were recorded for each of the seven channels at a two-minute time resolution. In addition, the analog output of the 520 nm channel was monitored continuously and one minute averages of this channel were recorded separately.

Measurements of total PM$_{2.5}$ mass concentrations were also obtained at the CENICA site by a Tapered Element Oscillating Microbalance (TEOM, Ruppert & Pastashnick) operated at 35°C. This instrument measures the total fine aerosol mass concentration by using a vibrating element whose frequency is dependent on the particle mass collected on a filter located at the end of the element (Hinds, 1999). Hourly averages of ozone for this time period were obtained from the RAMA station at Cerro de la Estrella (Calzada San Lorenzo, Colonia Paraje San Juan, Delegación Iztapalapa, México D.F.). This site (19° 20’ 09.184’’ N, 99° 04’ 28.829’’ W) is located 3.0 km (1.8 mi) from the main sampling site at CENICA.

3 Results and discussion

The 30-min concentrations of PAN measured in 1997 are shown in Fig. 1a. The strong diurnal concentration pattern observed is evidence of the regional afternoon clearing of the pollutants from the MCMA basin (see Fig. 3 for expansion of diurnal detail). The PAN concentrations reached a maximum of 34 ppb with an average daily maximum of 15 ppb. The higher PAN analogs were also observed in 1997. Levels of PPN and PBN reached 5 and 1.1 ppb respectively, giving a maximum of only 8 ppb early in the study with an average daily maximum for the time period of 3 ppb. In addition, neither PPN nor PBN were observed above the detection limit.
of 0.02 ppb at any time during the 2003 study period. The typical diurnal concentration pattern is evident in the first 14 days of the study, indicating the same daily meteorological patterns observed in 1997 leading to a daily clearing of the basin.

18 April 2003 (Julian day 108) was Good Friday marking the beginning of the holiday period accompanied with a decrease in traffic levels and a decrease in mobile and stationary emissions. Average carbon monoxide concentrations dropped by a factor of 2 and peak concentrations dropped by a factor of 4 from the levels reported on the previous Friday. This resulted in even lower PAN levels after that day. The average daily maximum before 18 April was 4 ppb while the average daily maximum afterwards was 1.7 ppb. The IMADA-AVER study in 1997 also incorporated a Mexican national holiday (Benito Juarez’s Birthday) on 21 March (Julian Day 80). The maximum PAN concentration observed on that day was 12 ppb.

The hourly average ozone concentrations reached a maximum of 242 ppb during the 1997 study period but only reached 135 ppb in 2003. The ratio of PAN to ozone daily maximum average concentrations is shown in Fig. 2a for 1997 and Fig. 2b for 2003. Past measurements of PAN and ozone concentration in polluted air masses have yielded a ratio of 0.1 (Tuazon et al., 1981; Finlayson-Pitts and Pitts, 2000; Gaffney et al., 1989). The observed ratio of PAN/ozone in 1997 generally agrees with the value of 0.1. However, many days exceeded 0.1 and 6 out of the 32 study days exceeded a ratio of 0.2. This has been attributed to the significant presence of PAN precursors, including higher aldehydes and olefins, in the Mexico City air during that time and also to the fact than the higher PAN levels act to tie up NO2 thus reducing the ozone production rate and the subsequent atmospheric ozone levels (Gaffney et al, 1999). The PAN/ozone values observed in 2003 only reached the value of 0.1 the first two days of the study. The PAN/ozone ratios after 5 April (Julian Day 95) were significantly below 0.1 with an average daily value of 0.02 for the rest of the study period and an average of 0.03 for the period before the Easter holiday.

As expected, NO2 levels were also lower in 2003 than in 1997. The maximum NO2 concentrations reached 205 ppb in 1997 and 137 ppb in 2003. Figure 3a shows the concentration profiles of ozone, NO2, and PAN for two representative days in 1997. Figure 3b shows similar concentration profiles for 2003. Although the overall concentration levels are lower in 2003 than in 1997, the concentration profiles appear to follow similar patterns. There is a rapid conversion of NO to NO2 in the Mexico City air in both 1997 and 2003.
as indicated by the fact that NO₂ reaches a maximum before noon. At this time both ozone and PAN begin to be produced with PAN reaching a maximum earlier than ozone. Shortly after midday the boundary layer height increases leading in a reduction of NO₂ atmospheric concentrations. In late afternoon PAN levels begin to drop faster than ozone. As this rapid decrease in PAN is not likely due to reaction with NO, because this would cause ozone to be lost faster than PAN, it was proposed in 1997 that this behavior might be due to heterogeneous loss of PAN on carbonaceous aerosol surfaces (Gaffney et al., 1999). It has been shown in laboratory studies that PAN can be lost on contact with soot surfaces at low ppb levels during relatively short contact times (Gaffney et al., 1998). Figure 3b also includes the black carbon aerosol profiles obtained in 2003. The NO₂ concentration levels correspond well with the black carbon concentrations as both black carbon and NO₂, the NO₂ precursor, are produced from combustion. The PAN levels in Fig. 3b begin to drop after the black carbon concentrations reach their maximum supporting the suggestion of heterogeneous PAN loss. This same loss pattern has also been observed in Santiago, Chile where PAN levels have been seen to exceed 20 ppb (Rappenglück et al., 1998, 2000). This city has a large diesel bus fleet and the black carbon levels are likely to be even higher than those observed in Mexico City.

Daily average PM₂.₅ concentrations have been reported elsewhere for La Merced site during the AMADA-AVER campaign in 1997 (Edgerton et al., 1999; Chow et al., 2002). These fine aerosol levels ranged from 21–60 μg/m³ in 1997 with an average of 36 μg/m³. The daily average PM₂.₅ during the 2003 study ranged from 26–69 μg/m³ with an average of 45 μg/m³. This is not a significant difference in fine atmospheric aerosols from 1997 to 2003. The samples collected in 1997 were also analyzed for elemental carbon content by thermal evolution analysis (Chow et al., 1993, 2001). Figure 4a shows the elemental carbon content of the 1997 PM₂.₅ aerosol fraction as calculated from the previously reported results (Edgerton et al., 1999; Chow et al., 2002). The percent elemental carbon in the fine aerosol samples collected in 1997 varied from 6–23% with an average during the campaign of 15(%σ=5%). Figure 4b shows the daily average black carbon content of fine aerosols as determined from light absorption and TEOM measurements made in 2003 (Salcedo et al., 2006). The daily average percent black carbon in fine aerosols measured in 2003 ranged from 5–18% with an average of 10(%σ=3%). This may represent a slight decrease in black/elemental carbon aerosol content in 2003, especially at higher PM₂.₅ concentrations. However, the thermal evolution method used in 1997 to determine elemental carbon may sometimes result in high values if corrections are not adequately made for charring of the sample during analysis (Chow et al., 2004).

The decreasing trend in PAN levels observed in Mexico City from 1997 to 2003 is similar to that observed in southern California where the maximum PAN concentrations have steadily decreased from 50–70 ppb in the 1960s to 3–10 ppb in the 1990s (Grosjean, 2003). The PPN concentrations in southern California have also decreased from 5–6 ppb in the 1960s to less than 1 ppb in the 1990s. Peak ozone concentrations were about 540 ppb in 1960 and 210 in 1997 (Grosjean, 2003; South Coast Air Quality Management District, 1995; www.aqmd.org). This trend in decreasing atmospheric oxidants reflects the increasingly stringent controls on VOC and NOₓ emissions in the state of California. Although the trend for both PAN and ozone in southern California is downward, the ambient levels of PAN have decreased faster than those of ozone yielding a decrease in the PAN/ozone ratio from 0.13 in the 1960s to 0.02 in 1997. Since California’s emission controls have focused on reducing the most reactive VOCs thus reducing the overall photochemical reactivity of the emissions (Carter, 1994) it has been suggested that this may have resulted in a larger reduction of the PAN precursors compared to the VOCs that produce ozone but do not produce PAN (Grosjean, 2003).

As the Mexico City and Los Angeles areas have many similarities in their air pollution problems including, a high density of mobile emissions resulting in similar atmospheric
chemistries, the Mexican government utilized a similar approach to improving the air quality in the MCMA as that used by the state of California (Molina and Molina, 2002b). This included the removal of reactive olefins from LPG (Blake and Rowland, 1995; DDF et al., 1996), re-strengthening motor vehicle emission standards with limits on the most reactive VOCs and the introduction of two-way catalytic converters in new vehicles starting with the model year 1991 and three-way catalytic converters starting with 1993 models (Molina and Molina, 2002b). Fuel-based motor vehicle emission inventories for Mexico City have subsequently estimated a decrease in NOx and hydrocarbon emissions of 26% and 39%, respectively, from 1998 to 2000 (Schiffer et al., 2003, 2005). Vehicle remote sensing studies made in 2000 showed that tailpipe hydrocarbon emissions (measured in pounds of HC per pound of fuel) were lowered to one ninth of those emitted by the 1991 fleet (Beaton et al., 1992; CAM/IMP, 2000). Atmospheric measurements have also reported a decrease in ambient hydrocarbon levels from 1992 to 1996 (Arriaga et al., 1997). It is therefore not surprising that results from the recent studies in Mexico City should yield results similar to those observed in the Los Angeles area with respect to atmospheric oxidants, particularly the PANs. However, fine atmospheric aerosols, including black carbon aerosols remain a problem in this megacity. This is primarily due to the lack of controls on diesel particulate emissions.

The presence of the highly absorbing black carbon aerosols in Mexico City leads to a reduction in solar flux of 17.6% locally (Raga et al., 2001b). The mass of these absorbing aerosols exported from this megacity into the surrounding region is estimated to be 6000 metric tons per day or 2 mega-tons per year of black carbon (Gaffney et al., 1999). Since freshly formed black carbon aerosols are hydrophobic, they are expected to be more resistant to washout and have longer lifetimes than more hygroscopic aerosols such as sulfate and nitrate (Gaffney and Marley, 2005b). Since these aerosols are also transported out of the area. This may lead to regional impacts that were not explored in this study. It is recommended that measurements of PANs in the outflow regions should be undertaken to evaluate the potential impacts of ozone and PAN on ecosystems in the region.

The results for black carbon aerosol content indicate that this pollutant is not being reduced in Mexico City, consistent with no control strategies being put into place. It is clear that this and other megacities continue to be major sources of this key aerosol species on a global scale. As the lifetimes for black carbon aerosols are anticipated to be longer than that for the more hygroscopic aerosol species, the global impact of the emissions from megacities such as Mexico City must continue to be evaluated (Gaffney and Marley, 2005b). Considering the importance of black carbon aerosols in radiative balance, these major sources of absorbing aerosols cannot be ignored if we are to adequately assess their role in climate change.

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4 Conclusions and recommendations

A comparison of results for PAN in Mexico City in 2003 with those obtained in 1997 indicate that the overall reactivity of the urban air chemistry has changed consistent with the control strategies that have been put into place in Mexico City, i.e., reduction of olefin content in LPG sources and the replacement of motor vehicles with those that employ the use of catalytic converters to reduce the reactive VOC emissions. Although the levels of PAN are still fairly high for an urban center, PAN and ozone levels are gradually dropping in Mexico City with similar trends to that observed over a number of decades in the Los Angeles air basin (Grosjean, 2003).

The Mexico City urban air is generally transported downwind of the city on a daily basis leading to the strong diurnal cycles observed for PAN at the urban sites. It is apparent that both NOx and reactive VOC emissions in Mexico City, although still high, are being reduced. Although the reduction of reactive VOCs in the urban emissions leads to the reduction of photochemical oxidants in the MCMA basin, the lower reactivity of the VOCs will certainly lead to the production of PANs downwind of Mexico City as the emissions are transported out of the area. This may lead to regional impacts that were not explored in this study. It is recommended that measurements of PANs in the outflow regions should be undertaken to evaluate the potential impacts of ozone and PAN on ecosystems in the region.
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