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Can 3-D models explain the observed fractions of fossil and non-fossil carbon in and near Mexico City?

A. Hodzic¹, J. L. Jimenez², A. S. H. Prévôt³, S. Szidat⁴, J. D. Fast⁵, and S. Madronich¹

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Correspondence to: A. Hodzic (alma@ucar.edu)

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¹NCAR Earth System Laboratory, Boulder, CO, USA

²Dept. of Chemistry and Biochemistry, and CIRES, University of Colorado, Boulder, CO, USA

³Laboratory of Atmospheric Chemistry, Paul Scherrer Institut, Switzerland

⁴Dept. of Chemistry and Biochemistry, and Oeschger Centre for Climate Change Research, University of Bern, Switzerland

⁵Pacifc Nortwest National Laboratory, Richland, WA, USA

A 3-D chemistry-transport model has been applied to the Mexico City metropolitan area to investigate the origin of elevated levels of non-fossil (NF) carbonaceous aerosols observed in this highly urbanized region. High time resolution measurements of the fine aerosol concentration and composition, and 12 or 24 h integrated ¹⁴C measurements of aerosol modern carbon have been performed in and near Mexico City during the March 2006 MILAGRO field experiment. The non-fossil carbon fraction (fC_{NF}), which is lower than the measured modern fraction (fC_M) due to the elevated ¹⁴C in the atmosphere caused by nuclear bomb testing, is estimated from the measured fC_M and the available source information. The fC_{NF} contained in PM₁ total carbon (fC_{NF}^{TC}) ranged from 0.37 to 0.67 at the downtown location (T0), and from 0.50 to 0.86 at the suburban site T1. Substantially lower values (i.e. 0.24-0.49) were found for PM₁₀ filters at T0 by an independent set of measurements, which are inconsistent with the modeled and known differences between the size ranges, suggesting higher than expected uncertainties in the measurement techniques of ¹⁴C. An increase in the non-fossil organic carbon (OC) fraction (fC_{NE}^{OC}) by 0.10–0.15 was observed for both sets of filters during periods with enhanced wildfire activity in comparison to periods when fires were suppressed by rain, which is consistent with the wildfire impacts estimated with other methods. Model results show that the relatively high fraction of non-fossil carbon found in Mexico City seems to arise from the combination of regional biogenic SOA, biomass burning OA, as well as non-fossil urban OA. Similar spatial and temporal variations for fC_{NE}^{OC} are predicted between the urban vs. suburban sites, and high-fire vs. low-fire periods. The absolute modeled values of fC_{NF}^{OC} are consistent with the PM₁₀ dataset but lower than the PM₁ filters. Resolving the ¹⁴C measurement discrepancies is necessary for further progress in model evaluation. The model simulations that included secondary organic aerosol (SOA) formation from semi-volatile and intermediate volatility (S/IVOC) vapors showed better skill in explaining both total OA mass and fC_{NE}^{OC} compared to simulations which only included SOA from VOCs. Urban sources of modern carbon are important

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in reducing or closing the gap between model and measurements, even though they are often neglected on the interpretation of 14 C datasets. The fC_{NF} of urban POA and SOA precursors is an important parameter that needs to be better constrained by measurements. Performing faster (≤3 h) ¹⁴C measurements in future campaigns is critical to further progress in this area. To our knowledge this is the first time that radiocarbon measurements are used together with aerosol mass spectrometer (AMS) organic components to assess the performance of a regional model for organic aerosols.

Introduction

Organic aerosols (OA), composed of a complex mixture of primary (POA, emitted in the particle phase) and secondary (SOA, formed due to chemical reactions of organic vapors) compounds, account for a large fraction of the submicron particulate mass over continental regions (20-80%, Jimenez et al., 2009) and in the free troposphere (Murphy et al., 2006). Despite their ubiquity, the sources and formation processes of OA are still largely controversial with major consequences on our ability to predict and regulate OA levels and, therefore, their effects on climate and human health and their changes for future climate and emission scenarios. Of particular interest for pollution regulation is the ability to distinguish between the OA emitted from human activities, of which a major fraction involves fossil fuel burning (and that can be reduced using emission controls or non-combustion renewable energies) and OA generated by non-fossil sources such as biogenic SOA or biomass burning. We note that the fossil/non-fossil distinction does not map to controllable/uncontrollable sources: a fraction of biogenic SOA in polluted regions may be controllable (Carlton et al., 2010), and wildfire emissions can be partially managed through prescribed burning and other measures.

Over the past several decades radiocarbon analyses have provided insights into the relative contribution of fossil and modern sources of carbonaceous aerosols at different locations by measuring the ¹⁴C/¹²C ratio in ambient aerosol samples (Szidat, 2009a). Emissions from fossil fuel combustion do not contain ¹⁴C, as the geological age of the

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fuel is much larger than the half-life of ¹⁴C of ~5730 years. Thus they can be separated from non-fossil carbon sources that have a similar amount of ¹⁴C isotope as atmospheric CO₂ when the carbon in the biomass was photosynthesized (e.g. Hildemann et al., 1994 and references therein). We define the non-fossil carbon fraction (fC_{NF}) ₅ as the fraction of aerosol carbon which arises from sources for which the carbon has been recently fixed, such as forest fires, biogenic SOA, and food cooking. Similarly we can define the fossil carbon fraction (fC_F) as the aerosol carbon arising from fossil sources, such as the combustion of fossil fuels or SOA formed from evaporation of fossil fuels such as gasoline. By definition $fC_{NF}+fC_{F}=1.0$. The traditionally reported modern carbon fraction (fC_M , Stuiver and Polach, 1977) is higher than fC_{NF} because $fC_{\rm M}$ =1.0 for atmospheric CO₂ around 1950, but $fC_{\rm M}$ of atmospheric CO₂ increased greatly due to atomic bomb tests in 1955-1963 (Szidat et al., 2006; Levin et al., 2010). fC_M of atmospheric CO₂ reached almost 2.0 in the Northern Hemisphere in 1963, and has been slowly decreasing since. Biomass photosynthesized 30, 20, 10, and 0 years before the MILAGRO study in 2006 would have fC_M of 1.37, 1.20, 1.12, and 1.06, respectively (Levin et al., 2010). Thus $fC_M + fC_F > 1.0$, and it is important to keep in mind that always fC_{NF} , with their ratio depending on the age of the biomass. We also will use EC, OC, or TC (for elemental, organic, or total carbon, respectively) as a superscript to indicate the carbon fraction whose NF fraction we are referring to (e.g. fC_{NF}^{OC}).

Generally, ¹⁴C measurements are performed on total carbon aerosols (TC) and the resulting modern fraction includes contributions from all carbonaceous aerosol components (i.e. both elemental carbon, EC, and organic carbon, OC). Recently, the quantification of ¹⁴C in sub-fractions of TC, such as EC and OC has been reported (Szidat et al., 2004). The availability of ¹⁴C for these carbon fractions is very useful for source apportionment studies, as OC and EC are unlikely to contain the same fraction of fossil and non-fossil carbon due to the different relative source impacts. OC is emitted from primary sources but also formed as SOA and can partition between the gas and particle phases, and even volatilize back to the gas-phase upon heterogeneous oxidation.

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EC is exclusively generated by combustion of fossil fuels and biomass and is effectively chemically inert on atmospheric residence times of weeks.

The results of aerosol radiocarbon studies have revealed the presence of large amounts of non-fossil carbon in both remote and urban environments of the Northern Hemisphere throughout the year. Rather surprisingly, substantial amounts of non-fossil carbon have been reported even in heavily urbanized areas (e.g. Los Angeles or Mexico City) where large quantities of fossil fuel are being burned. Measurements of fC_{NF}^{OC} reported in recent years are summarized in Fig. 1 and Table 1. One of the largest urban fC_{NIE}^{TC} was reported in Albuquerque, New Mexico (0.76, Klinedinst and Currie, 1999). These levels are comparable to levels found in remote mountain environments (0.62-0.87 of non-fossil contribution in Puy-de-Dome, Gelencser et al., 2007; Legrand and Puxbaum, 2007) or in the Grand Canyon National Park (>0.95) where the anthropogenic influence is expected to be small. Lower, but still substantial $fC_{\rm NF}^{\rm TC}$ were found in the US urban agglomerations ranging from 0.29 to 0.56 (e.g. 0.29-0.38 in Los Angeles, Hildemann et al., 1994; or 0.49-0.56 in Phoenix, Bench et al., 2007), in Zurich (0.54-0.59, Szidat et al., 2006), in Beijing (0.33-0.48, Yang et al., 2005) and in Tokyo (0.30-0.40, Takahashi et al., 2007). Mexico City is no exception. Marley et al. (2009) have reported that modern carbon (fC_M^{TC}) ranged between 0.42–0.74 within the city center, which corresponds to approximately 0.35–0.63 fC_{NF}^{TC} as discussed below. Aiken et al. (2009b) have quantified fC_{NF}^{TC} to range from 0.28 during low biomass burning periods to 0.41 during high biomass burning activity of March 2006 (MILAGRO field project). The presence of substantial levels of non-fossil TC in urban environments during low biomass burning periods poses the question of the influence of sources such as biogenic SOA, primary biological particles (PBAP), and urban sources of modern carbon (such as food cooking, biofuel use, tire wear etc.) on urban air quality. These sources are usually not well captured by current models, and radiocarbon data offers an additional constraint for OA modeling studies.

To estimate the relative contributions of fossil fuel combustion, biomass burning and

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biogenic SOA to OA concentrations, several studies (e.g. Gelencser et al., 2007; Szidat et al., 2009b) have combined radiocarbon measurements with analyses of specific organic tracers. This receptor-modeling approach is exclusively based on data and is sensitive to inaccuracies associated with measurements, missing or below detection limit observations, and to non-uniqueness or atmospheric degradation of tracers. The potential of using the radiocarbon measurements, in combination with other OA measurements, for 3-D model evaluation and source apportionment has not been explored to our knowledge. Such a combined approach may be useful to help interpret the apportionment of modern and fossil carbon between different POA and SOA sources, as the sources of carbonaceous aerosols cannot be directly determined from only 14C measurements because of the contribution of multiple primary and secondary fossil and non-fossil sources.

In this study we combine the results of an air quality model with ¹⁴C and AMS measurements acquired in Mexico City during the MILAGRO field study (March 2006). We have already shown in our previous work (Hodzic et al., 2009, 2010) that the model is able to reproduce the observed concentrations of primary and secondary OC within 30% (a relatively small error given the current state of knowledge) during this campaign. However, there is insufficient evidence that this level of agreement is due to the right mixture of sources. The combined use of AMS and ¹⁴C observations will help determine whether the model captures the concentrations and fractions of OC from modern and fossil carbon sources, or whether error compensation may be occurring among various OA sources. The goal of the paper is twofold: (i) to assess whether the current representation of OA in our model can explain the observed levels of non-fossil carbon in aerosols within Mexico City; and (ii) to use the model results to determine the relative contributions of urban sources, biomass burning (BB), and biogenic emissions to the observed levels of carbon in the vicinity of Mexico City. As SOA formation is one of the major current uncertainties in OA modeling (e.g. Hallquist et al., 2009; de Gouw and Jimenez, 2009), we will examine the sensitivity of the results to the choice of the SOA parameterization. This type of analysis can help determine the strengths

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and weaknesses of current OA and SOA models. The terminology used for organic matter in this paper is summarized in Table 2 for reference.

2 Measurements

Within the framework of the MILAGRO field experiment (Molina et al., 2010), aerosol samples were collected by two groups on quartz fiber filters during March 2006 at both urban (T0) and suburban (T1) locations in Mexico City and analyzed for ¹⁴C content at two different laboratories. Available *f* C_{NF} measurements for MILAGRO are summarized in Table 3.

2.1 Using fC_M vs. fC_{NF} for analysis and model-measurement comparison

The analyses carried out in this paper could use either fC_M or fC_{NF} . Each parameter has some advantages and disadvantages. The measurements report fC_M directly, and to estimate fC_{NF} one needs to make an assumption about the mixture of sources which are responsible for the measured modern carbon. On the other hand, fC_{NF} is the physically meaningful quantity and its use is thus preferable. Also, the model calculates fC_{NF} most directly, and additional assumptions about each source are needed to estimate fC_M . In addition, the use of fC_M causes much confusion, as the large majority of the researchers in the OA field do not appear to be aware of the difference between fC_M and fC_{NF} and thus they tend to interpret reported fC_M values as if they were fC_{NF} , overestimating the importance of non-fossil sources. Considering the uncertainties in both measurements and model as well as the confusion introduced when fC_M is used, we have chosen to only use fC_{NF} in the remainder of this manuscript. The conclusions of this paper would not change if fC_M was used instead.

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The first set of ¹⁴C measurements (Marley et al., 2009, hereafter abbreviated MAR) was collected on submicron filter samples over 12 h periods at the T0 and T1 sites for both daytime (06:00-18:00 Local Time, LT) and nighttime (18:00-06:00 LT). Mar- $_{5}$ ley et al. reported fC_{M}^{TC} which includes modern carbon contained in both EC and OC, and did not attempt to estimate fC_{NE}^{TC} . For the purpose of this study, the fraction of modern carbon in OC was estimated by subtracting the modern carbon contribution from EC (see Table 3). To determine EC concentrations several datasets were available at T0 and T1 locations during MILAGRO as shown in Fig. S1 (supp. Info., see http://www.atmos-chem-phys-discuss.net/10/14513/2010/ acpd-10-14513-2010-supplement.pdf). It should be noted that the different EC concentrations measured either directly or derived from aerosol light absorption measurements (often referred to as black carbon or "BC") agree reasonably well at both T0 and T1. Paredes-Miranda et al. (2009) showed that the different BC measurements also agreed well at T0. Since at T0 only low-time resolution measurements of EC are available (12 or 24 h), a combination of measurements was used to produce a high-resolution EC surrogate, using the thermal-optical Sunset labs analyzer data from Stone et al. (2008), and the aethalometer measurements of BC by Marley et al. (2009). At T1 the thermal-optical measurements of EC were used, which have 1-h time resolution (de Gouw et al., 2009). For some filters, the estimated fC_{M}^{OC} exceeds 1. This is however physically possible as discussed above due to the nuclear bomb excess.

In their samples, Marley et al. (2009) did not correct for the enrichment of ¹⁴C due to nuclear bomb radiocarbon, which results in the reported fC_M being larger than fC_{NF} by approximately 1.16 for wood burning emissions and 1.055 for biogenic aerosols (Szidat et al., 2009b). To estimate the non-fossil carbon fraction (fC_{NF}) contained in OC we correct for this effect by using an average value of 1.1 under the assumption that modern carbon comes in equal proportions from sources with similar ages as wood and recently photosynthesized biogenic material (Table 3).

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2.3 Estimates of fC_{NF} in PM₁₀ OC at the T0 site

The second dataset (called "AIK") consists of four PM₁₀ filters collected at T0 (Aiken et al., 2009b) during (1) 21 March 09:04 a.m.-22 March 09:05 a.m., (2) 22 March 09:20 a.m.-23 March 09:20 a.m., (3) 26 March 09:40 a.m.-27 March 09:40 a.m., (4) ₅ 29 March 11:04 a.m. –30 March 11:05 a.m. The filters were analyzed directly for modern carbon content in EC and OC. Details of the analyses are described by Aiken et al. (2009b). Although only four 24-h filters are available, they were collected and analyzed completely independently, and thus are extremely valuable to compare to the Marley et al. (2009) data reported at the same site. Furthermore, the direct separation for ¹⁴C analysis into OC and EC is of special importance for source apportionment, as both carbonaceous particle fractions originate from different sources. fC_{NE} was estimated from the $fC_{\rm M}$ measurements using the same convention as above.

2.4 TC and OA component analysis

TC filter measurements were performed simultaneously as part of the ¹⁴C measurements (Table 3). The MAR reported values vary from 10-25 µgC/m³ at T0 and from 3-20 µgC/m³ at T1 with a few exceptionally elevated values at T0 on 11 March (30 μgC/m³), and at T1 on 9 and 13 March (30 and 96 μgC/m³). TC peak values (>30 µgC/m³) correspond most likely to local biomass burning events, which however have not been recorded by any other instrument operating at T0 and T1 during MILA-GRO.

AMS data were acquired at the T0 and T1 supersites. Experimental details, intercomparisons to other instruments, and the methodology and results of the component analysis using positive matrix factorization (PMF, Ulbrich et al., 2009) are presented in Aiken et al. (2008, 2009ab), Paredes-Miranda et al. (2009), Huffman et al. (2009) and de Gouw et al. (2009). The main components determined by PMF are hydrocarbonlike OA (HOA), a surrogate for primary combustion OA; biomass-burning OA (BBOA), thought to be dominated by primary biomass burning OA; and oxygenated OA (OOA),

3 Modeling approach

3.1 Model description

For this study, the mesoscale chemical transport model CHIMERE was run from 11 to 31 March 2006 over the Mexico City region at both regional (35×35 km²) and urban (5×5 km²) grid scales using the same configuration and the same forcing (i.e. meteorology, emissions, boundary and initial conditions) as described by Hodzic et al. (2009, 2010). Here we provide a brief summary of key sources and processes that influence the amount of modeled carbonaceous aerosols.

The CHIMERE model simulates the emissions of primary EC and OA, the chemistry and gas/aerosol partitioning of secondary organic species and their gaseous precursors, in addition to their transport, boundary layer mixing, and dry and wet deposition processes. Similar to Hodzic et al. (2009, 2010), primary gaseous and aerosol species arise from (i) anthropogenic sources as reported by the 2002 official Mexico City Metropolitan Area (MCMA) emission inventory (CAM, 2004) and the National Emissions Inventory (NEI) of 1999 emissions outside of the city (http: //mexiconei.blogspot.com/), (ii) biomass burning emissions as estimated from satellite data by Wiedinmyer et al. (2006), (iii) biogenic VOC emissions calculated online using the MEGAN model (Guenther et al., 2006), and (iv) PBAP from fungal spores. The PBAP emissions had not been included in our previous studies and were calculated following the emission algorithm of Heald and Spracklen (2009). The formation of secondary organic species from traditional anthropogenic and biogenic precursors (i.e. volatile organic compounds, VOCs) is modeled as by Pun et al. (2006). As described by Hodzic et al. (2010), the model also includes the gas-phase chemistry and partitioning of semi-volatile and intermediate-volatility primary organic vapors (SVOC and IVOC, or together as S/IVOC) based on two alternative parameterizations: the first ACPD

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proposal of Robinson et al. (2007), and the updated parameterization from Grieshop et al. (2009). The carbon (OC) and oxygen fractions for each S/IVOC lumped species are explicitly modeled according to the specifications of the Robinson and Grieshop parameterizations. In addition to carbonaceous species, CHIMERE accounts for windblown dust, secondary inorganic species (sulfate, nitrate and ammonium), particulate water, and other primary anthropogenic particles. The size distribution of all species is represented using a sectional approach with 8 size bins (40 nm to 10 microns in physical diameter) with internal mixing within each bin. Further information on the model formulation can be found on http://www.lmd.polytechnique.fr/chimere/.

3.2 Non-fossil OC estimation from the model results

The model simulates OA mass concentrations, which in addition to OC also include oxygen, hydrogen and nitrogen mass. Therefore, the OC concentration at each point in space and time arising from source i (OC_i) is estimated from the predicted OA concentration for that source (OA_i) as:

$$OC_i(\mu g C m^{-3}) = \frac{OA_i(\mu m^{-3})}{(OM/OC)_i}$$

where (OM/OC), represents the source-specific ratio of total OA to OC concentrations. The term OM/OC is retained here for consistency with most of the previous literature, even though the term OA/OC would be more consistent with the terminology of this paper. The value of OM/OC for ambient particles depends on the source and the degree of oxidation of the aerosol. Recent studies have reported values of 1.6 and 2.1 for mixed OA at urban and rural sites, respectively (Turpin and Lim, 2001; Aiken et al., 2008). In this study, we use source-specific OM/OC ratios from Aiken et al. (2009a) measured in Mexico City for the surrogates of anthropogenic combustion POA (1.38), biomass burning POA (1.55), and total SOA (1.95). For biogenic SOA formed in a smog chamber Shilling et al. (2009) reported an OM/OC of 1.6 at the OA levels in Mexico City (~20 µg m⁻³, Aiken et al., 2009a). Here we use a value of 1.7 for biogenic SOA to conservatively account for additional aging that occurs in the atmosphere and is not fully captured in smog chambers (Aiken et al., 2008; Ng et al., 2010). The OC content in the SOA formed from S/IVOC was directly determined from modeled species as both carbon and oxygen fractions are explicitly modeled.

The model allows separate tracking of POA and SOA concentrations resulting from various sources including urban emissions, biomass burning, biogenic or PBAP emissions. Once the OC concentrations arising from all modeled OA sources are calculated at a given point in space and time, the fraction of non-fossil carbon $fC_{\rm NF}^{\rm OC}$ is derived according to the following equation:

$$fC_{NF}^{OC} = \frac{SOC_{BSOA} + POC_{BB} + SOC_{BB} + POC_{PBAP} + 0.2 \times (POC_{urb} + SOC_{urb})}{\sum_{i} OC_{i}}$$

where SOC_{BSOA} is the OC in biogenic SOA, and the last term estimates the impact of non-fossil carbon emissions from urban sources, due to the impact of sources such as food cooking, tire and brake wear, resuspended road dust, trash burning, biofuel use, cigarette smoke, etc. (Hildemann et al., 1994; Christian et al., 2010). It is assumed that ~20% of the carbon from urban sources is non-fossil, as an average of the values determined by Hildemann et al. (1994) for the Los Angeles basin. A rough estimate of this ratio in Mexico City based on the MCMA emission inventory and the source-specific non-fossil contributions determined by Hildemann et al. (1994) suggests that about 18% of the total urban OC emissions should be considered as non-fossil, which may be a lower limit since the emission inventory does not properly account for sources such as trash burning and biofuel use (Christian et al., 2010). This parameter is very uncertain and further research should be aimed at its quantification. The sensitivity of our results to this parameter is examined in Sect. 4.4.

The OC associated with biomass burning, biogenic SOA and PBAP emissions is considered fully as non-fossil. However, it should be noted that biogenic and biomass burning emissions do not always result from natural activities. For example agricultural crops emit biogenic VOCs, and burning of agricultural residues, prescribed or arson-related wildfires are anthropogenic sources of modern carbon. In addition it has been

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suggested that biogenic SOA formation may be more efficient when biogenic VOCs mix with anthropogenic pollution due to perhaps higher oxidants, POA, NO_x, or acidity (de Gouw et al., 2005; Weber et al., 2007; Carlton et al., 2008; de Gouw and Jimenez, 2009). Thus one should not equate "non-fossil OC" with "OC from natural emissions" or "uncontrollable OC".

Carbonates can also be a source of fossil or non-fossil carbon. It is necessary to distinguish between dust-related carbonates that are fossil, vs. marine carbonates that are non-fossil. In Mexico City at T0 there were about $0.46\,\mu gC$ m⁻³ from carbonate (Querol et al., 2008), which is 3% of the total carbon in OC+EC. The non-fossil contribution from marine origin is likely negligible in this region. In addition, it is reasonable to consider that OC analysis should be minimally affected by carbonates as some of the carbonates are not dissolved by the HNO₃ digestion used in this analysis (hydrofluoric acid digestion is needed for full dissolution) and the CO₂ release from most carbonates needs higher temperatures than those used in this analysis. Overall the error introduced by carbonates in the analyses of the AIK filters is estimated to be less than 1%. Thus in the model, the contribution of carbonates is not considered.

3.3 Model application during MILAGRO

Results of three model simulations are discussed here. The main focus of this paper is on the results from the "ROB" and "GRI" simulations. These simulations include SOA formation from traditional VOC precursors, and also explicitly treat SOA formation from aging of primary organic S/IVOC following the approaches by Robinson et al. (2007) and Grieshop et al. (2009), as discussed by Hodzic et al. (2010). These two simulations are also compared to the reference ("REF") simulation from Hodzic et al. (2009) where SOA is only formed from VOCs using a traditional two-product model, and in which POA is assumed to be inert and non-volatile.

For comparison with measurements, the simulated parameters are spatially and temporally interpolated at the location of the measurement sites. The first day of the simulation corresponds to the "spin-up" time and the model results are analyzed from 12

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to 31 March. Measured and predicted concentrations of all species are reported under ambient conditions of pressure and temperature; mass concentrations should be multiplied by about 1.42 for conversion to STP conditions (1 atm, 273 K).

The CHIMERE model was recently evaluated against meteorological and gas-phase 5 observations of the MIRAGE field study (Hodzic et al., 2009), and also used to examine OA formation pathways in the Mexico City region by anthropogenic and biogenic traditional precursors (Hodzic et al., 2009), as well as primary semi-volatile organic vapors (Hodzic et al., 2010).

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Model-measurement comparison for fC_{NF}^{TC}

Figure 2 compares the predicted and observed fractions of non-fossil carbon contained in carbonaceous aerosols for the REF, ROB, and GRI simulations. The observed values from both datasets range from 0.24 to 0.67 within the city and from 0.50 to 0.86 downwind (Table 3), suggesting that fossil sources are major contributors to TC within Mexico City, whereas biogenic and biomass burning contributions represent a larger fraction 20 km downwind of the city center.

The model $fC_{\rm NF}^{\rm TC}$ results are quite consistent (with some scatter) with the AIK data. As fC_{NE}^{TC} is about 0.15 larger for the MAR data compared to the AIK data, the model shows a similar discrepancy vs. the MAR dataset, with the model having lower fC_{NE}^{TC} by ~0.15 at both sites, again consistent across the REF, ROB and GRI simulations. The variability of fC_{NE}^{TC} in the model at T0 is much lower than observed for both the AIK and MAR data. No increase in fC_{NF}^{TC} between urban (T0) and suburban (T1) locations is seen in the REF simulation vs. 0.15 in the MAR measurements, while the ROB and GRI simulations capture the difference between the urban and suburban sites better. Model results also show that the choice of the SOA formation mechanism significantly

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impacts the modeled fC_{NF}^{TC} values. As discussed in Hodzic et al. (2009, see Fig. 11) the traditional approach can predict biogenic SOA fairly well when compared to specific tracers, which is consistent with studies at other locations. However SOA formation associated with urban emissions is severely underpredicted by the REF simulation leading to larger relative fraction of non-fossil TC within the city. The GRI simulation produces 0.05–0.1 lower fC_{NF}^{TC} (more fossil aerosol) than ROB due to differences in the precursor and mechanism details. As the results of ROB and GRI simulations are very similar (within 10%) in terms of fC_{NF}^{TC} and the ROB simulation predicts OA concentrations slightly better during MILAGRO (Hodzic et al., 2010), in the rest of the paper we will only focus on the analysis of the ROB simulation.

Figure 2 also shows a surprisingly large difference between the two sets of measurements with a substantially lower non-fossil OC fraction for AIK, as already discussed by Aiken et al. (2009b). Although sampling times were not coincident for the two sets of data, the average fC_{NF}^{TC} value for AIK filters (0.34) is much lower than the one for MAR (0.54). The main known difference between the two sets of filters is their size cut, with a PM₁₀ cut for the AIK data and a PM₁ cut for the MAR filters. However this factor is unlikely to explain the difference as the amount of OC in coarse particles is very small (Querol et al., 2008; Aiken et al., 2009b). The coarse mode is dominated by crustal particles, and likely contains some vegetative detritus (Stone et al., 2008) or PBAP (see below) that would contribute to measured non-fossil carbon. Paved road dust and break wear also contribute to the supermicron mode but have a large fraction of modern OC (Hildemann et al., 1994). The contribution of carbonates can be neglected for Mexico City as already discussed in Sect. 3.2. In the following some possible reasons for discrepancies found between the two sets of observations, and between the model and MAR observations will be discussed.

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4.3 Assessment of carbonaceous aerosol predictions

Another possible reason for model disagreement with measured $fC_{\rm NF}^{\rm TC}$ could be the errors in the EC or OC simulations in this region. The measured fC_{NE}^{TC} presented in

The presence of PBAP in supermicron particles could be a reason for differences in fC_{NIE}^{TC} for PM₁ vs. PM₁₀, although in the opposite direction as observed. For the comparisons with the AIK data presented in Fig. 2 and also below, the modeled fC_{NF} values 5 were calculated for PM₁₀ aerosols, and PBAP were included. PBAP have been found to contribute to the global budget of OA (e.g. Mahowald et al., 2008). Recently, Heald and Spracklen (2009) estimated the contribution of fungal spores as 23% of total primary OA emissions, or 7% of the fine-mode source. In this study PBAP emissions were calculated as a function of leaf area index (LAI, m² of leaf surface per m² of ground surface) and water vapor mixing ratios (H₂O_{vap} dimensionless) following the emission algorithm from Heald and Spracklen (2009) (C. Heald, personal communication, 2009): E_{PBAP} (g/m²/s)= α ·LAI·H₂O_{vap} where α equals 5.18×10⁻⁸ and 1.55×10⁻⁷ g/m²/s for PM_{2.5} and PM₁₀ respectively. Figure 3 shows the spatial distribution of the predicted average PBAP in the vicinity of Mexico City from 15-30 March 2006. Within the city basin the average values are close to 0.2 µg/m³. These concentrations are similar to the average concentrations predicted by Heald and Spracklen (2009) using GEOS-Chem. However, greater variability is found during the day, with minimum values in range 0.05–0.1 μg/m³ in the early afternoon, and high values in range 0.3–0.4 μg/m³ at night. It should be noted that this average PBAP concentration of 0.2 µg/m³ is about 10–15% of the estimated biogenic SOA (~1.5 μg/m³, Hodzic et al., 2009) and therefore is not expected to dominate the non-fossil OC fraction. The presence of PBAP cannot explain, but also does not greatly increase, the differences in fC_{NE}^{TC} between the two sets of filters.

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Figure 2 accounts for both EC and OC, so the model ability to simulate both carbonaceous aerosol components within Mexico City must be evaluated. This is first examined using real-time measurements of BC and OA. Because non-fossil carbon samples are available for 12 h day- and nighttime intervals (MAR data), here we compare the corresponding 12h-averaged predictions. This comparison is intended to complement the evaluation of predicted POA and SOA time series for March 2006 presented in our previous work (Hodzic et al., 2010), focusing here on the specific time periods that are relevant to ¹⁴C measurements, and including the comparison of EC which was not presented before.

4.3.1 Comparison with EC measurements

Observed and predicted concentrations of elemental carbon (EC) are shown in Fig. 4. The comparison of EC diurnal profiles at T0 indicates that the modeled values are a factor of 2 lower during the morning peak, and that this gap is somewhat reduced in the afternoon and eliminated overnight. The comparison performed for MAR sampling periods confirms the model underprediction for many filters, especially at T0. Similar model behavior is encountered at the T1 site. Our results are consistent with those of Fast et al. (2009) and this error is most likely due to a too low emission inventory of EC as discussed in Fast et al. (2009). The better prediction overnight does not contradict that conclusion, as it most likely results from cancellation of errors as the too low EC emissions are balanced by the too low mixing layer in the model (Fast et al., 2009; Hodzic et al., 2009). If the model underprediction of EC was rectified (e.g. by increasing EC emissions by a factor of 2), the simulated fC_{NIF}^{TC} would decrease on average by about 0.02 as fC_{NF}^{EC} is only 0.04–0.13 (Table 3).

Comparison with OC measurements

The model skill in predicting OA was examined with the high time-resolution data and model results by Hodzic et al. (2009, 2010), and is re-examined using OC and specif-

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ically for the filter period averages in Fig. 5. At the urban T0 site, the observed AMS daytime (nighttime) OC concentrations vary from 8 to 20 μgC/m³ (3–17 μgC/m³) with rather low values found during the more ventilated periods of e.g. 19-20 March and higher levels observed during more stagnant days e.g. 16, 21, 22 or 30 March (Fast 5 et al., 2009; Hodzic et al., 2009). On most days, SOC contributes between 40 and 60% of the total OC (not shown). Both model runs capture relatively well the observed temporal relative variations, with the exception of the high daytime value observed on the 18th or the nighttime one from the 20th March. These peak values coincide with intense advection of biomass burning plumes from nearby wildfires (Aiken et al., 2009a, b) that are generally underpredicted in the model as discussed by Hodzic et al. (2010). The predicted OC from the ROB simulation is in a reasonable agreement with observed values i.e. within 10-30% most of the time. The REF model simulation using the traditional SOA approach predicts substantially lower OC levels because of a severe underprediction of SOA from anthropogenic and BB sources (not shown here, see Hodzic et al., 2009, 2010).

At the suburban T1 site, OC modeling appears to be more challenging, as already discussed by Hodzic et al. (2010). The underprediction by the REF simulation is more severe ($\sim \times 2$), and there is more scatter between the model and the observations. A reasonable agreement is obtained for the ROB simulation especially during the day, with the exception of the 15-16 March when the dispersion of pollutants in the model is too low (Hodzic et al., 2010). Some underpredictions are observed in some cases of wildfire impacts (e.g. the night of 18 March and 23 March daytime).

The important result of these comparisons is that the ROB simulation provides a fairly reasonable mass closure for OC in particular within the city, which is a requirement that needs to be met before examining the model ability to predict the relative proportions of fossil and non-fossil OC. One should also keep in mind that for a few high BB events that are not captured by the model (e.g. daytime filter on the 21st at T0), the model is likely to underpredict the fC_{NF}^{OC} for those particular filters.

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Fossil and non-fossil contributions to OC

Now that we have established the reasonable model skill in predicting OC levels, we focus on determining whether or not the predicted non-fossil OC fraction is reasonable. as shown in Fig. 6. Good agreement with measurements is observed for the AIK data. The observed fC_{NF}^{OC} varies from 0.37 to 0.55 within the city, and show less day-to-day variability than the MAR data. The differences among the four AIK filters are thought to be mainly due to variability in BBOA impacts (Aiken et al., 2009b). Higher $fC_{\text{NIE}}^{\text{OC}}$ values (0.47 and 0.55) for the first two samples correspond to higher fire impact period (as shown by higher BBOC values Hodzic et al., 2010, Figs. 2-3), and feature 0.10-0.15 higher fC_{NF}^{OC} values than filters from 26 and 29 March (fC_{NF}^{OC} of 0.37 and 0.40) taken during lower fire influence. However, it should be noted that the fC_{NIF}^{TC} and fC_{NIF}^{EC} for these filters show a clear fire signature only for the first day (21 March), with higher values by 0.19 and 0.08 respectively than during the rest of the time which is dominated by fossil sources (fC_{NF}^{TC} <0.33 and fC_{NF}^{EC} <0.05).

PM₁ OC at T0 as estimated from the MAR dataset has fC_{NE}^{OC} in the range 0.49–1.08, whereas the values are even higher (0.57-1.07) at the suburban location T1. The values of $fC_{NF}^{OC} > 1.0$ for some samples may be due to measurement noise, and suggest that the contribution of noise to scatter in the MAR data is likely to be at least 0.10. Alternatively, perhaps the assumptions for converting fC_M into fC_{NF} are too conservative (too low assumed average $fC_M/fC_{NF}=1.10$) for the whole dataset or for some samples with high BB impact. At T0, nighttime values are slightly higher (fC_{NE}^{OC} in range 0.59– 1.08) than during daytime (AM: 0.49-0.67) suggesting that background biogenic SOA, BBOA from smoldering fires, and other non-fossil sources are more important contributions at night. This is consistent with the results of Hodzic et al. (2009) that have already shown a higher relative contribution of biogenic SOA during nighttime when urban SOA and POA emissions from urban activities are at lower. BBOA peak values were also observed during the nighttime and early morning hours due to the advection

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of smoke from smoldering fires in the shallow nighttime boundary layer (Aiken et al., 2009a, b; Hodzic et al., 2010). Also, the highest $fC_{\rm NF}^{\rm OC}$ values, exceeding 1.0, were collected overnight on March 21 during high BB impact events. Somewhat lower difference between day and nighttime values is observed at T1, e.g. 0.23 at T0 and 0.08 at T1 on average, due to a more limited urban influence. Similar to the $fC_{\rm NF}^{\rm TC}$ comparison, the simulated $fC_{\rm NF}^{\rm OC}$ are lower for the ROB simulation compared to the MAR dataset, as shown in Fig. 6. This gap is particularly pronounced at T1. The simulated $fC_{\rm NF}^{\rm OC}$ range from 0.25 to 0.55 at T0, and up to 0.65 at T1 and are thus on average 0.20–0.30 lower than the MAR dataset.

The origin of the high fC_{NF}^{OC} values reported for MAR filters does not seem to find explanation in our modeling results. Unlike for the filter measurements, the predicted fC_{NF}^{OC} values for PM₁ and PM₁₀ aerosols are different by less than 0.02 (Fig. 6a). Thus the model difference between the predicted submicron and supermicron values cannot explain the large gap found between the two sets of filters. If the AIK measurements were correct that would suggest that MAR measurements are likely biased towards high fC_{NF}^{OC} values.

The $fC_{\rm NF}^{\rm OC}$ measurements were also combined with the measured or estimated OC concentrations (see Table 3) to estimate the absolute concentrations of fossil and nonfossil OC. The predicted fossil OC concentrations at both T0 and T1 sites are slightly lower than the AIK data and higher than the MAR data at T0, while data and model are more consistent at T1. Non-fossil OC is somewhat underpredicted when compared to the AIK data and shows a large underprediction at T1 with respect to MAR data. For T0 the lower fossil OC of the MAR data (vs. the model) is compensated by higher non-fossil OC than the model. At T1, where only the MAR dataset is available the nonfossil OC is higher by $5-8\,\mu{\rm gC/m}^3$ compared to the observations on most of the days. Given the reasonable agreement of the model with the observed AMS values shown on Fig. 5b, this comparison suggests inconsistencies between the PM₁ OC filter data and the AMS measurements at T1.

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Filters collected between 23 and 24 March during the low BB period illustrate well these inconsistencies. Filter data seems to indicate an underprediction of the background non-fossil concentrations of about 4 µgC/m³, i.e. while fossil OC concentrations seem to match the observed values. As the BB influence is very limited during this period within the city, a possible interpretation would be that biogenic emissions are substantially underpredicted. However this is not consistent with results reported by Hodzic et al. (2009) that showed a good general model agreement with measurementbased estimates of biogenic SOA. On some days the fossil OC derived from the MAR filters is close to zero or zero, which seems unrealistically low for this suburban site located next to a highway, and is also contradictory with the presence of fossil-dominated EC. One should also keep in mind that day-to-day comparison of filters is a difficult task because the local meteorology plays an important role on the dispersion of pollutants within the boundary layer.

4.4.1 Sensitivity analysis

For the model results discussed above, we assumed that fC_{NF}^{OC} from urban emissions is 0.20 as an average of the values determined by Hildemann et al. (1994) for the Los Angeles basin. This fraction is uncertain and could vary highly from city to city. In particular it may be biased low for Mexico City where sources such as trash burning and biofuel use are not included in the emissions inventory. Therefore, here we test the model sensitivity to this assumption by considering lower and upper limits for fC_{NE}^{OC} of urban emissions of 0 and 0.40, as shown in Fig. 7. The lower limit of 0 is thought to be unrealistic but helps provide information on the modeled trend, while a value ~0.30 may be more realistic after accounting for the effects of biofuel use and trash burning (as discussed above), and 0.40 may be considered an upper limit. The comparison indicates a strong sensitivity to this parameter, with a near doubling of the modeled fC_{NE}^{OC} values when increasing the contribution of urban emissions from 0 to 0.40. This is not surprising given the fact that urban sources are the major contributor to predicted

OC in and near Mexico City (Hodzic et al., 2010). The predicted fC_{NF}^{OC} values for the AIK filters increase from 0.25–0.30 to 0.55–0.60 for the extreme sensitivity cases, and are either 0.2 below the observed fC_{NF}^{OC} , or 0.2 above it. The scatter in the comparison with the MAR data does not improve for most of the points although a factor of 2 higher fC_{NF}^{OC} is predicted when considering urban fC_{NF}^{OC} of 0.40 instead of 0.

4.5 Spatial distribution of non-fossil OC in the Mexico City area

Model results also provide valuable information on the relative importance of non-fossil and fossil sources at the regional scale. Figure 8 presents the average spatial distribution of the fC_{NF}^{OC} in the vicinity of Mexico City as predicted by ROB simulation during the second half of March 2006. Based on the biomass burning intensity two distinct periods have been selected, i.e. a high BB (17-22 March) and a low BB (24-29 March) period. For both periods, fC_{NF}^{OC} displays a strong spatial gradient between urban and remote areas. Lower fC_{NF}^{OC} contribution of 0.30–0.50 is predicted over the Mexico City valley and its dominant outflow region to the north/northeast. These values are due to the importance of urban POA and SOA in this region, which has fC_{NF}^{OC} = 0.20 in the model. Higher fC_{NF}^{OC} (0.65–0.85) are predicted over the mountains surrounding the city, and over the western part of the plateau, which are less influenced by urban and industrial activities. Substantial differences in fC_{NF}^{OC} amplitude can also be seen between high and low BB periods. During the low BB period, $fC_{\rm NF}^{\rm OC}$ ranges from 0.35–0.40 in the valley, while this contribution increases up to 0.55-0.65 over the mountains and exceeds 0.70 west of Mexico City. During high BB periods, the contribution of nonfossil carbon is enhanced by about 0.10-0.20 within the city and the surrounding hills. The fire plumes show a strong signature in fC_{NE}^{OC} , an their effects can be seen over the mountains.

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4.6 Source apportionment of OA with ¹⁴C/AMS data

The combined ¹⁴C (AIK) and AMS aerosol measurements allow for an estimation of the sources of OA (Szidat et al., 2004), beyond the evaluation of the total non-fossil and fossil fractions of OC (Lanz et al., 2008). We assume that fossil-/non-fossil fractions are the same for the AIK filters and for AMS data. HOA is assumed to be 0.80 fossil and 0.20 non-fossil, in accordance with the assumptions above. The same assumption is applied to the SOA from urban precursors.

Figure 9a shows the model predictions for the daily average of the whole period. while Fig. 9b shows the model results over the AIK filter periods, and Fig. 9c shows the results from the AIK data. The AMS results indicate that biomass burning organic aerosol (BBOA) represents about 5 and 18% of OA during low and high BB respectively. According to this method, the estimated fraction of non-fossil OOA (OOA_{NE}) accounts for ~35% of the OA and it is larger than the fossil fraction (8-12%). Primary organic species (HOA) from fossil fuel burning represents about 30% while the contribution of the non-fossil HOA is less than 8%. Other unidentified organic sources represent a minor fraction (<8%).

Model predictions for the 14-30 March period show a somewhat different OA apportionment. The main difference is the larger proportion of fossil secondary organics i.e. 25-30%, with a 15% higher model prediction, whereas the non-fossil fraction is about 5-10% lower in the model. The amount of primary organic aerosols from fossil and non-fossil sources is close to the observations. The non-fossil SOA fraction is composed of about 20% urban, 50% biogenic and 30% biomass burning sub-fractions in average, with some variability between low and high biomass burning periods (see Fig. 9b). The biomass burning SOA is mostly produced from the photochemistry of S/IVOC precursors emitted by fires that occurs in the vicinity of Mexico City (Hodzic et al., 2010). The model also predicts a factor of two lower proportion of primary biomass burning aerosols with the predicted fraction varying from 2 to 8%, which is consistent with our earlier study (Hodzic et al., 2010).

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The CHIMERE model has been applied to the Mexico City metropolitan area during March 2006 and compared with aerosol ¹⁴C and AMS measurements to investigate the origin of elevated levels of non-fossil (NF) carbon aerosols observed in this urban area. The followings findings emerged from the study:

- i) The model results suggest that the relatively high fraction of non-fossil carbon found in Mexico City seems to arise from the combination of biogenic SOA sources, biomass burning POA and SOA, as well as non-fossil urban POA and SOA. The study shows, in agreement with previous assessments, that the differences between the two ^{14}C datasets could not be explained by the different size cuts between the two sets of filters, implying larger than expected uncertainties in field measurements of ^{14}C . Much smaller difference (<2%) between PM $_1$ and PM $_{10}f\text{C}_{\text{NF}}^{\text{OC}}$ values is suggested from model results. The contribution of primary biological particles (PBAP) is relatively small both in terms of surface concentrations (~0.2 $\mu\text{g/m}^3$) and non-fossil carbon. A 10–20% increase in $f\text{C}_{\text{NF}}^{\text{OC}}$ was observed for both sets of filters as well as modeled values during enhanced wildfire activity in comparison to periods when fires were suppressed by rain, quantifying the contribution of that source to the measured OC levels.
- ii) Modeling results using the most complete SOA model (ROB simulation) for OC and $fC_{\rm NF}^{\rm OC}$ show reasonable agreement with the AIK dataset but are lower than the MAR dataset. None of the simulations could explain the elevated values of $fC_{\rm NF}$ reported by the MAR dataset, especially at T1. If AIK measurements were the most accurate ones that would imply that our modeling of the organic aerosol mixture is reasonable especially during low biomass-burning periods. If on the contrary, the MAR data were more accurate that would indicate that the model predictions are too low for reasons that we have not been able to explain in this work. To make further progress it is urgent to understand and resolve the mea-

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iii) Similar fC_{NF}^{TC} predictions are found for the GRI run, with values generally 0.05-0.10 lower than for ROB simulations. The fraction of non-fossil carbon in urban emissions has a large impact on the simulations, and this parameter needs to be further constrained by additional source and ambient measurements. The model results also indicate a strong spatial pattern on fC_{NE}^{OC} , with lower values in the urban area and larger values in regional air, which is also observed in the measurements.

surement discrepancy.

iv) The combined analysis of radiocarbon and AMS data allowed assessing the model skill (ROB simulation) in reproducing the source-specific OA composition. It was found that the contribution of fossil sources to secondary organic aerosols was overpredicted by 10-20%, while that of non-fossil sources was reproduced within 5-10%. The comparison also confirms that the biomass burning contribution to POA is a factor of 2 lower than observed as already seen by Hodzic et al. (2010).

Our results highlight the benefits of a complementary use of aerosol ¹⁴C measurements and AMS data for air quality modeling studies, as well as the critical need for higher temporal resolution of ¹⁴C measurements i.e. ≤3-h instead of 12 h or 24 h filters and ¹⁴C measurement intercomparisons, in order to better quantify the ¹⁴C measurement noise, diurnal cycle, possible effects of plumes and "hot" sources, etc. in future studies.

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Table 1. ¹⁴C apportionment studies showing the contribution of non-fossil carbon to TC.

Location	Site	Date	Size	fC _{NF} (%)	TC (μgm ⁻³)	Reference	
		Sites	located in the US	SA			
	URB	Jan-Dec 82		29	9		
Los Angeles, CA	SUB	Jan-Dec 82	PM _{2.1}	38	10	Hildemann et al., 1994	
Denver, CO	SUB	Aug 96 Dec 96-Jan 97		42 26	6.1 9.2		
Brighton, CO	RUR	Dec 96-Jan 97	$PM_{2.5}$	26	5.9	Currie et al., 1994 Klinedinst and Currie, 1999	
Albuquerque, NM	URB	Dec 85		76	35	Klinedinst and Curre, 198	
Houston, TX	SUB	Aug 02		48	4.1	Lemire et al., 2002	
Nashville, TN	SUB	Jun-Jul 99	PM _{2.5}	64	5.0	Lewis et al., 2002	
Tampa, FL	SUB	May 02	1 1412.5	70	3.1	Lewis and Stiles, 2006	
N. Birmingham, AL	URB	Sep 03-Jan 04	PM _{2.5}	30	13	·	
Chattanooga, TN	URB	Mar-Apr 03	DM. 57 5.8 Taillere		Tanner et al., 2004		
Centreville, VA	RUR	Sep 03-Jan 04	PM _{2.5}	72	6.6	Ke et al., 2007	
Look Rock, TN	RUR	Summer 01-02	PM ₁₀ /PM _{2.5}	74	5.5	Zheng et al., 2006	
	URB	Summer 00-05		56	2.3		
Phoenix, AZ	UKD	Winter 01-06	PM _{2.5}	49	7.3	Bench et al., 2007	
	RUR	Summer 00-05	F IVI _{2.5}	95	0.8	Schichtel et al., 2008	
Grand Canyon, AZ		Winter 01-06		100	0.2	, = + + +	
Libby, MT	VAL	Nov 03-Feb 04	$PM_{2.5}$	82	20	Ward et al., 2006	
		Sites	s located in Europ	е			
Stockholm, S	URB	Nov 05	TOD	50	2.8	7	
Aspvreten, S	RUR	Nov 05	TSP	80	1.7	Zencak et al., 2006	
Gothenburg, S	URB-BCG	Feb-Mar 05	PM ₁₀	56	3.0	Szidat et al., 2009b	
Gothenburg, S	URB-BCG	Jun-Jul 06	PM _{2.5}	53	2.7	52idat et al., 2009b	
Aveiro, P	RUR RUR	Summer 02-03		63	4.0		
Aveilo, F		Winter 02-04		81	14.0		
K-Puszta, H		Summer 02-03		82	5.0		
K-i uszta, i i		Winter 02-04	PM _{2.5}	69	11.0	Gelencser et al., 2007	
Sonnblick, A	BCG	May-Jun 03	PIVI _{2.5}	78	1.6	Gelencser et al., 2007	
Soffiblick, A	BCG	Oct-Dec 02		65	0.21		
Dun do Domo ED	BCG	Jun 03		87	4.92		
Puy de Dome, FR	ВСС	Jan 02+Jan+Dec 03		62	0.86		
Zurich, CH	URB-BCG	Aug-Sep 02	PM ₁₀	54	5.9	Szidat et al., 2004,2006	
		Feb 03		59	19	32ldat et al., 2004,2000	
Sedel, CH	RUR	Jan-Feb 06	PM ₁₀	63	26	Szidat et al., 2007	
Roveredo, CH	RUR	Jan 04	PM ₁₀	83	16	Perron et al., 2010	
rtoveredo, err	NON	Nov-Dec 04	PM ₁	85	15		
		(Other locations				
	URB	Jun+Aug 02		30	5.5		
Tokyo, J	URB	Apr+Oct 02	PM ₁₃	38	4.8	Yamamoto et al., 2007	
	URB	Dec 02+Feb 03	1.0	40	6.3		
Dallian C		Jun 01		48	27.8	V	
Beijing, C	URB	Jan-Feb 01	PM _{2.5}	33	50.9	Yang et al., 2005	
	URB	March 06	DM	35	16.7	Marley et al., 2009	
Mexico City, MX	SUB	March 06	PM ₁	63	20.5		
**	URB	March 06	PM ₁₀	33	18.5	Aiken et al., 2009b	

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Table 2. Terminology used for the various fractions and sources of organic compounds.

Organic compounds					
VOC	Volatile organic compounds				
SVOC	Semi-volatile organic compounds; the subset of VOC which according to their vapor pressure can partition between the gas and the aerosol phases (Robinson et al., 2007).				
IVOC	Intermediate-volatility organic compounds, those with volatility in between that of SVOC and the speciated VOC (Robinson et al., 2007)				
S/IVOC	SVOC+IVOC				
OA POA	Organic aerosol; includes both primary and secondary fractions Primary organic aerosol, here treated as non-volatile (REF run) and semi-volatile species (ROB and GRI runs)				
SOA EC	Secondary organic aerosol Elemental carbon				
OC	Organic carbon contained in the organic aerosol				
SOC	Secondary organic carbon contained in the SOA				
TC	Total carbon; EC+OC				
PBAP	Primary biological particles				
AMS	Aerodyne aerosol mass spectrometer				
PMF	Positive matrix factorization, a mathematical technique used to extract components from AMS spectra (Ulbrich et al., 2009, and references therein)				
HOA	Hydrocarbon-like organic aerosols, an OA component derived from PMF analysis of AMS spectra. It is generally used as a surrogate for urban combustion POA (Jimenez et al., 2009, and references therein)				
OOA	Oxygenated organic aerosols, an OA component derived from PMF analysis of AMS spectra and characterized by its high oxygen content. It is generally used as a surrogate for SOA (Jimenez et al., 2009, and references therein)				
BBOA	Biomass burning organic aerosols, an OA component derived from PMF analysis of AMS spectra. It is generally used as a surrogate for urban combustion POA (Jimenez et al., 2009, and references therein)				

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Table 3. Measured concentrations of TC and EC/BC aerosols, and corresponding non-fossil fractions at T0 and T1 during MILAGRO.

Date	Period ^a	TC μgC/m ³	EC/BC ^b μgC/m ^d	fC _{NF} ^{EC}	$fC_{NF}^{TC^{c}}$	fC _{NF}				
		at T0 (fr	om Marley et al., 2	(800						
3/11	AM	30.1	5.79		0.55	0.67				
3/11	PM	22.5	5.99		0.60	0.80				
3/13	PM	15.5	4.25		0.59	0.79				
3/17	PM	18.4	2.10		0.55	0.61				
3/21	PM	10.0	4.36	$0.05^{(3)}$	0.63	1.08				
3/23	AM	21.4	3.11		0.46	0.52				
3/23	PM	9.7	2.45		0.67	0.88				
3/27	AM	13.6	3.59		0.37	0.49				
3/27	PM	8.8	1.85		0.48	0.59				
at T0 (from Aiken et al., 2008b)										
3/21	Daily	15.8	2.29	0.13	0.49	0.55				
3/22	Daily	24.3	7.86	0.04	0.33	0.47				
3/26	Daily	9.6	2.26	0.05	0.32	0.40				
3/29	Daily	24.4	9.28	0.04	0.24	0.37				
			om Marley et al., 2	(800						
3/11	AM	17.1	1.74		0.67	0.74				
3/11	PM	19.3	1.33		0.63	0.68				
3/13	AM	96.9	1.89		0.68	0.69				
3/13	PM	12.6	1.31		0.78	0.86				
3/15	AM	16.4	1.60		0.59	0.65				
3/15	PM	15.1	0.77		0.64	0.67				
3/17	AM	14.4	1.46		0.66	0.72				
3/17	PM	15.1	1.43		0.68	0.74				
3/19	PM	10.1	0.83	0.05	0.78	0.84				
3/21	AM	14.7	1.59	0.00	0.70	0.78				
3/21	PM	1.9	0.60		0.75	1.07				
3/23	AM	15.3	2.24		0.72	0.83				
3/23	PM	8.9	1.34		0.86	0.99				
3/25	AM	11.0	2.05		0.84	1.02				
3/25	PM	3.6	0.85		0.63	0.81				
3/27	AM	10.1	1.53		0.50	0.57				
3/27	PM	2.5	0.80		0.60	0.86				

a Sampling time period corresponds to 06:00-18:00 LT for AM filters, to 18:00-06:00 for PM filters, and to 09:00 to 09:00 LT for daily-averaged filters.

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b EC/BC data used at T0 for the MAR are from Aethalometer measurements for the time period before 17 March, and from EC real-time measurement afterwards. For AIK EC are from filter measurements. At T1 all data are from direct real-time measurements of EC.

^c fC_{NF} are corrected for the bomb effect, assuming that current values are 11% overpredicted.

^d For MAR fraction of modern carbon in BC is assumed to be 5%.

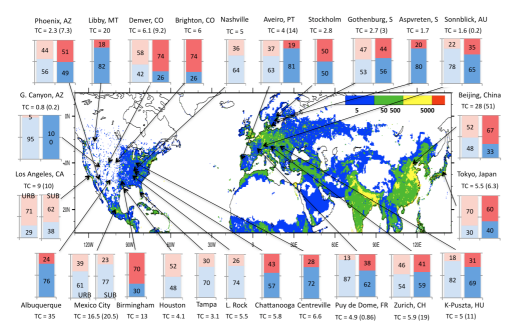


Fig. 1. Population density (km $^{-2}$) and the relative non-fossil (blue bars) and fossil (red bars) carbon reported in recent studies for the Northern Hemisphere. Summer (lighter colors) and winter (darker colors) are distinguished. When available, total carbon concentrations (μ gC m $^{-3}$) are also indicated for the summer and winter (in parenthesis) samples.

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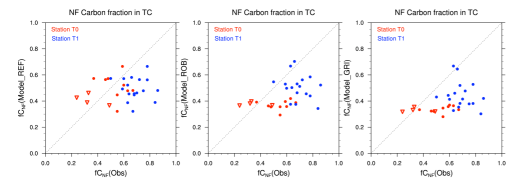


Fig. 2. Comparisons of predicted and observed non-fossil carbon fractions (fC_{NE}) determined for Total Carbon filters. The submicron (MAR) filters are represented by dots, while PM₁₀ (AIK) filters are indicated by the triangles. Modeled values are based on REF (left), ROB (middle) and GRI (right) simulations. Both MAR and AIK data have been corrected for effects of historical nuclear bomb testing (see text for details).

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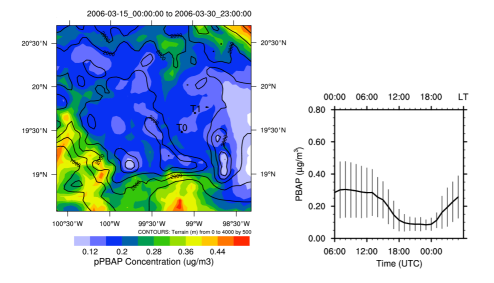


Fig. 3. Spatial distribution **(a)** and average diurnal profile at T0 **(b)** of primary biological particles (PBAP, μ g/m³) as simulated by the model over the Mexico City area between 15 and 31 March 2006.

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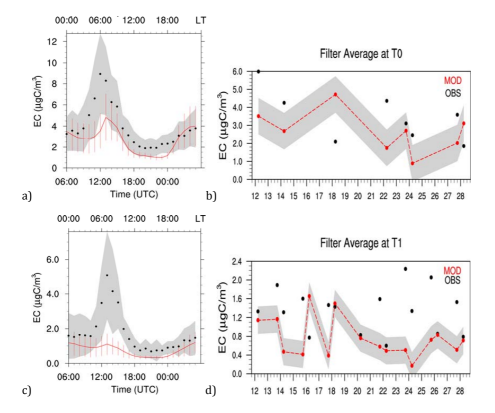


Fig. 4. Comparison of EC diurnal profiles (left panels: **a, c**) and averaged EC concentrations (right panels: **b, d**) for MAR filters as predicted by the CHIMERE model (red line) and as measured (Aethalometer-Marley, OCEC Doran, see Fig. S1, see http://www.atmos-chem-phys-discuss.net/10/14513/2010/acpd-10-14513-2010-supplement.pdf) (black dots) at T0 (top panel: a, b) and T1 (bottom panel: c, d). Sampling time periods are given in Table 3. In plot (b), model values are given only for available submicron filters to allow a more focused comparison with radiocarbon data.

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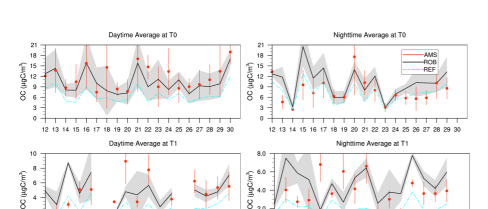


Fig. 5. Comparison of day- and nighttime averaged concentrations of submicron organic carbon particles as predicted by the CHIMERE model for the ROB (black line) and REF (blue line) simulations, and as measured by the AMS instrument (red dots) at T0 and T1 sites. The variability (±1 sigma) around the average value is also indicated for both modeled (grey area) and observed (vertical bars) values. Day- and nighttime averaging intervals correspond to MAR sampling periods given in Table 3.

14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30

0.0

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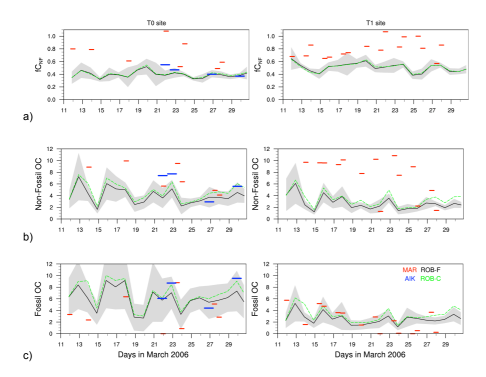


Fig. 6. (a) Fractions of non-fossil organic carbon (fC_{NF}) and associated (b) non-fossil and (c) fossil OC concentrations (µqC/m³) as predicted by CHIMERE and derived from MAR (red) and AIK (blue) filters measurements at T0 and T1. Filter-sampling time frame is given in Table 3. Model outputs from the ROB simulation are daily-averaged from 09:00 to 09:00 LT (the consecutive day) for submicron (black line ROB-F) and PM₁₀ (green line ROB-C) size distributions.

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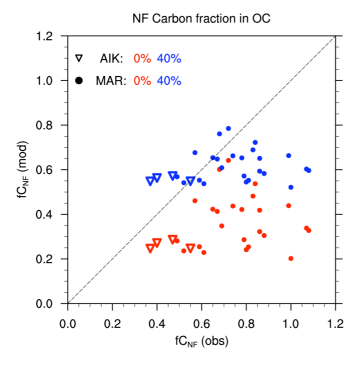


Fig. 7. Predicted (ROB) and observed $fC_{\rm NF}$ for MAR (dots) and AIK (triangles) for all filters available during MIRAGE 2006 at T0 and T1. Model sensitivity to assumed fraction of modern carbon in urban anthropogenic emissions is tested: no modern carbon is assumed (lower limit = red color), and 40% modern carbon content in urban anthropogenic emissions is considered (upper limit = blue color).

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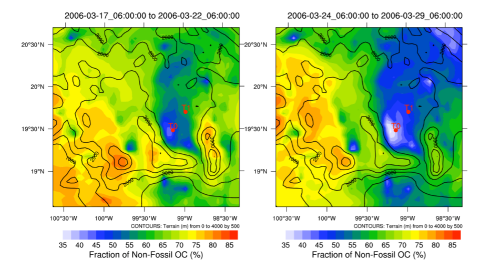


Fig. 8. Model predicted fraction of non-fossil carbon based on the ROB simulation. Fraction of non-fossil carbon was averaged over two distinct periods i.e. high biomass burning period (from 17–22 March 00:00 LT, left panel) and low biomass burning period (from 24–29 March 00:00 LT, right panel).

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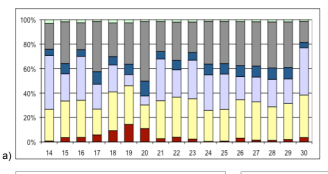
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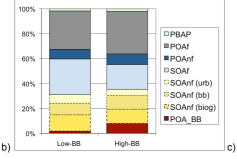
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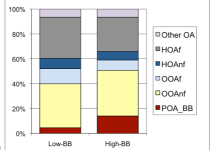


Fig. 9. Source specific contributions to organic PM₁₀ aerosols at T0 as predicted by the ROB simulation during (a) the second half of March 2006, (b) the high and low biomass burning periods, and (c) as derived from four AIK filters.