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# Historical (1850–2000) gridded anthropogenic and biomass burning emissions of reactive gases and aerosols: methodology and application

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## Abstract

We present and discuss a new dataset of gridded emissions covering the historical period (1850–2000) in decadal increments at a horizontal resolution of  $0.5^\circ$  in latitude and longitude. The primary purpose of this inventory is to provide consistent gridded emissions of reactive gases and aerosols for use in chemistry model simulations needed by climate models for the Climate Model Intercomparison Program #5 (CMIP5) in support of the Intergovernmental Panel on Climate Change (IPCC) Fifth Assessment Report (AR5). Our best estimate for the year 2000 inventory represents a combination of existing regional and global inventories to capture the best information available at this point; 40 regions and 12 sectors are used to combine the various sources. The historical reconstruction of each emitted compound, for each region and sector, is then forced to agree with our 2000 estimate, ensuring continuity between past and 2000 emissions. Simulations from two chemistry-climate models is used to test the ability of the emission dataset described here to capture long-term changes in atmospheric ozone, carbon monoxide and aerosol distributions. The simulated long-term change in the Northern mid-latitudes surface and mid-troposphere ozone is not quite as rapid as observed. However, stations outside this latitude band show much better agreement in both present-day and long-term trend. The model simulations indicate that the concentration of carbon monoxide is underestimated at the Mace Head station; however, the long-term trend over the limited observational period seems to be reasonably well captured. The simulated sulfate and black carbon deposition over Greenland is in very good agreement with the ice-core observations spanning the simulation period. Finally, aerosol optical depth and additional aerosol diagnostics are shown to be in good agreement with previously published estimates and observations.

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## 1 Introduction

In order to perform climate simulations over the historical record, it is necessary to provide climate models with information on the evolution of radiatively active gases and aerosols. Climate models that will contribute to scenario analysis for IPCC AR5 report (Intergovernmental Panel on Climate Change; Fifth Assessment Report) usually start their model calculations in pre-industrial times, specified in this case as being 1850 (Taylor et al., 2008). In order to enable such simulations, knowledge of the evolution of both historic emissions and concentrations of air pollutions is required. Examples of gridded emissions can be found at the GEIA/ACCENT emissions portal (available at: <http://geiacenter.org>). The determination of these emissions require a variety of steps involving the knowledge of the source of emission (e.g. amount of fossil fuel combusted by power plants), an emission factor (e.g. how much of a given chemical species is emitted for a specific mass of a given fuel burned in a specific technological process accounting for the operation of abatement measures) and a procedure for mapping onto a geographical grid (e.g. the location point sources such as power plants). As discussed in Klimont and Streets (2007) the quality of available emission inventories varies, between high quality inventories for point sources in Europe or North America (e.g.  $\text{SO}_2$  from power plants which is based on emissions monitoring) and inventories that are less reliable, especially in developing or industrializing countries due to incompleteness of activity data or lack of test-based emission factors. The resulting uncertainty leads to a range of possible emissions for a given process and base year that varies strongly between regions, sectors, and pollutants (e.g. Streets et al., 2006; Klimont and Streets, 2007). These uncertainties lead to a range of possible emission outcomes for a given source. While this issue applies to all types of emissions, we will focus in this paper on anthropogenic (defined here as originating from energy use in stationary and mobile sources, industrial processes, domestic and agricultural activities) and open biomass burning emissions. And our overall approach to building this new emission dataset is to combine a variety of data sources to maximize the

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information content; this is done through a combination of (1) regional and global inventories in order to define year 2000 emissions (2) long-term global emission datasets to define historical changes in emissions.

In order to perform chemistry simulations with enough resolution to resolve regional structures and their changes, our target is to provide monthly emissions at a horizontal resolution of  $0.5^\circ$  in latitude and longitude every 10 years. All emissions necessary for the simulation of tropospheric ozone and aerosols are provided, including methane ( $\text{CH}_4$ ), carbon monoxide ( $\text{CO}$ ), nitrogen oxides ( $\text{NO}_x$ ), total and speciated non-methane volatile organic compounds (NMVOCs), ammonia ( $\text{NH}_3$ ), organic carbon (OC), black carbon (BC) and sulfur dioxide ( $\text{SO}_2$ ). Compounds relevant for other issues (e.g. CFCs, HCFCs and HFCs, mercury, persistent organic pollutants) have not been included in this inventory as they were not necessary for the stated goal of providing climate models distributions of tropospheric ozone and aerosols for radiative forcing. The emission data are provided at the start of each decade. While interannual changes may be important for the detailed analysis of past “rapid” (i.e. less than a few years) pollution changes, we decided that decadal data are overall better suited to fulfill the needs of AR5 for the following reasons: (1) annual data sets exist only for a limited set of species (Smith et al., 2004) or only for the recent past – for example RETRO, available for the 1960–2000 period (Schultz et al., 2007, 2008) and REAS, which covers the 1980–2003 (Ohara et al., 2007; Smith et al., 2004), (2) emissions become increasingly uncertain going back in time, limiting the usefulness of single year data and (3) future emissions generated by Integrated Assessment Models (IAMs) are usually available with time steps on the order of 10 years (van Vuuren et al., 2010). Finally, while seasonal variations in anthropogenic emissions could also be of importance, (1) no enough information of past emissions available and (2) expected to be of lesser relevance compared to the long-term trends.

The paper is organized as follows: in Sect. 2, we discuss the land-based anthropogenic emissions, defined here as originating from industrial, domestic and agriculture activity sectors. Section 3 described the biomass burning emissions. In Sect. 4,

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we present our reconstruction of ship and aircraft emissions. Application of these emissions in chemistry-climate models and the analysis of the modeled concentrations of ozone carbon monoxide and aerosols are discussed in Sect. 5. Finally, discussion and conclusions are in Sect. 6.

### 5 1.1 Land-based anthropogenic emissions

In the case of land-based anthropogenic emissions (i.e., excluding aircraft and ship emissions), two available datasets on historical emissions, RETRO (1960–2000; Schultz et al., 2007) and EDGAR-HYDE (1890–1990; van Aardenne et al., 2001), provide information on emission changes over the second half of the 20th century for a limited set of compounds (Table 1), or at least. As mentioned in the Introduction, our approach consists of generating first our best estimate for 2000, based on the combination of global and regional datasets. This aggregation is performed using a set of 40 regions (Table 2) and 12 sectors (Table 3). Then, using a combination of RETRO and EDGAR-HYDE, historical trends for each sector in each region are generated. Finally, the historical emissions of reactive gases (ozone precursors only) are computed using the historical trends applied to our 2000 emissions. We discuss those three steps in this section.

While several recent assessments have shown that regional emissions have experienced significant changes between 2000 and present, especially in Southeast Asia (e.g., Richter et al., 2005; Zhang et al., 2009; Klimont et al., 2009), the lack of comparably comprehensive studies for this period in other regions made it impossible to provide detailed calculations that covered the entire globe; and therefore the year 2000 was used as the reference.

### 1.2 Definition of year 2000 emission

25 Anthropogenic emissions of reactive gases (with the exclusion of  $\text{SO}_2$ , see below) in 2000 are defined in terms of a variety of global and regional inventories. In generating

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our emission dataset, published or reviewed regional inventories have been given preference over global inventories in case these were available. This was the case for the EMEP (2006) inventory for Europe, the REAS inventory for Asia and the EPA inventory for North America. In those regions we assume that these inventories more appropriately reflect regional circumstances than the global inventories. Furthermore, the inventories for North America and European countries within the EMEP domain have been extensively evaluated through model and observation studies and inventories reported as Annex I inventories to the United Nations Framework Convention on Climate Change (UNFCCC) are subject to expert review. The EDGARv32 FT2000 dataset (Van Aardenne et al., 2005; Olivier et al., 2005) and preliminary emissions from EDGAR v4.0 for agriculture (EC-JRC/PBL, 2009) are used where regional information is not available. As the various inventories are combined at the level of regional averages, no attempt is made at smoothing potential discontinuities across regional boundaries. Although additional information is available in the literature for several regions (e.g., Zhang et al., 2009; Streets et al., 2003, 2006; Klimont et al., 2009; Cofala et al., 2007; NARSTO, 2006; Garg et al., 2006), we did not attempt to include those as they were less complete than the inventories applied in this study (e.g. only covering 1 country or including limited sectoral information or did not extend to the year 2000) or were already integrated into the datasets we have used (e.g., for East Asia REAS includes results of several specific inventories). While some of these inventories might provide better insight in emissions for one specific sector or compound for individual countries, it goes beyond the scope and available resources of the global inventory presented here to evaluate them on an individual basis and include them in a systematic way into the global inventory. Such approach might provide additional datasets in the future that can be used to improve upon this work.

Emissions of black carbon (BC) and organic carbon (OC) included in the dataset presented here represent an update of Bond et al. (2007) and Junker and Liousse (2008) as harmonization of emission factors was performed for the year 2000 from these papers and the studies they reference. Emissions of sulfur dioxide (Smith et al., 2010;

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S. J. Smith, J. van Aardenne, Z. Klimont, R. Andres, A. Volke and S. Delgado Arias "Historical Sulfur Dioxide Emissions: 1850–2005", 2010) are an update of Smith et al. (2001, 2004), with emissions from the UNFCCC submissions and other regional inventories where available.

For all species, gridding is performed using the EDGAR v4.0 spatial distribution maps specific for each sector at a 0.1° resolution, and aggregated to a 0.5° grid (EC-JRC/PBL, 2009). The 2000 SO<sub>2</sub> map used additional data from EDGARFT for smelting and fuel processing emissions.

### 1.3 Historical reconstruction

As we have two distinct emission datasets for ozone precursors (RETRO and EDGAR-HYDE) with differing trends and there is no a priori reason to dismiss one inventory over the other, we have devised an approach that maximizes the information from both datasets where applicable. Using the RETRO and EDGAR-HYDE historical trends per region and per sector, we generate historical trends for each sector in each region by defining the ratio of the emissions at a specific decade to its value in 2000 (our reference data set). This ratio is a concise representation of the changes in fuel use and emission factor over time and its full history (1850–2000) can be used to scale our 2000 inventory to define emissions in previous decades. The scaling of the anthropogenic emissions for reactive gases (excluding SO<sub>2</sub>) using EDGAR-HYDE and RETRO relies on the assumption that each reconstruction provides a reasonable (albeit sometimes different) representation of the time evolution of emissions; this can clearly only be applied to species available in both emission datasets, i.e. CO, NO<sub>x</sub> and NMVOCs. The main differences between the RETRO and EDGAR-HYDE datasets are in the emission factors variations over time (with RETRO having more technology information) and, to a lesser extent, the completeness of the inventory (e.g. no industrial process emissions in RETRO). Therefore, emissions for decades prior to 2000 can be calculated through a direct scaling (per sector and region) of our 2000 emissions, with a weighting factor defined as a linear combination of the RETRO and EDGAR-HYDE

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scaling factors, increasingly favoring EDGAR-HYDE when going further back in time (as RETRO is only available until 1960). To provide emissions back to 1850, EDGAR-HYDE emissions (which cover 1890 to 1990) are extrapolated to 1850 using global fossil fuel consumption estimates from Andres et al. (1999) and regional scale data for population from the HYDE dataset (Goldewijk, 2005).

In summary, the scaling for each sector and region is computed using the following steps:

1. The 1990–2000 change is computed in RETRO only (since the year 2000 is not included in EDGAR-HYDE).
2. The decadal changes between 1960 and 1990 are a combination of RETRO and EDGAR-HYDE.
3. The decadal changes between 1890 and 1960 are computed from EDGAR-HYDE only (no RETRO estimates prior to 1960).
4. The emissions between 1850 and 1890 are exactly as computed from EDGAR-HYDE and its extrapolation.
5. Smoothing is applied to scaling factors across 1960 and 1990 to limit jumps in the scaling factor.

The advantage of using a scaling approach is that it only requires that the existing emission inventories provide a time history of the specific emission, without having to deal with emission biases between inventories. Clearly, such history is meaningful only within a particular sector and for a specific region as pollution controls vary; the scaling is therefore capturing the change in fuel amount (usually fairly well-constrained) and change in the emission factors. It also relies on the assumption of an unbiased 2000 estimate; however, if such a bias were to be present, the methodology presented here could be applied to an updated set of 2000 estimates.

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The results from this scaling process are illustrated in Fig. 1 (left panel) for the case of CO emissions from the transportation sector in the United States. In this case, the RETRO inventory (red line) indicates a peak in 1970, followed by a rapid decline until 2000; on the other hand, EDGAR-HYDE (green line) does not peak in 1970 but instead is still larger in 1980. Therefore, the combination (black line) leads to a scaling which, going back in time, does not quite reach the RETRO levels in 1970 but instead reaches a smaller maximum in 1980. It represents therefore a combination of the two inventories. In terms of CO emission amount for this specific sector and region (Fig. 1, right panel), the combined use of the 2000 emissions with the scaling factor time evolution (Fig. 1, left panel) provides a new evolution of emissions with time that captures the changes and amplitudes of each original inventory; because of the different 2000 emissions for this sector and region, the combined dataset lies in-between the original sets.

While there can be wide variations for a specific sector or region, the global total amounts of anthropogenic emissions for each compound are actually quite similar to either RETRO or EDGAR-HYDE (Fig. 2), except in the case of agricultural waste burning and NMVOCs. The largest difference (in absolute amounts) is an increase in CO emissions compared to the EDGAR-HYDE estimate. Emissions of nitrogen oxides are quite similar between the two original inventories; in particular, the emissions between 1960 and 1980 are almost identical in RETRO and EDGAR-HYDE. Our combined dataset ends up slightly higher over that time period because our 2000 NO<sub>x</sub> emission estimate is larger than in RETRO. On the other hand, our 2000 NMVOC emissions are smaller than either RETRO or EDGAR-HYDE, again with a peak in 1990, similar to RETRO.

For OC and BC, two historical inventories were available; Bond et al. (2007) and Junker and Liousse (2008). In our study the historical trend of Bond et al. (2007) was used, constrained by our combined 2000 estimate. In addition, agricultural waste burning was computed from the CO estimate (see above) using regional emission factor (OC/CO and BC/CO) based on our 2000 emission estimates. For NH<sub>3</sub> we use the

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reconstruction by Beusen et al. (2008) while for CH<sub>4</sub>, since only one historical inventory (EDGAR-HYDE) exists, the only constraint to the present emission dataset comes from our 2000 estimate. The SO<sub>2</sub> time evolution was consistently calculated from Smith et al. (2010). These are shown in Fig. 3.

5 Soil emissions of nitrogen oxides are clearly affected by the use of fertilizers; it is therefore very difficult to disentangle the natural and anthropogenically-perturbed components to this flux. We have therefore used the EDGAR-HYDE estimate of soil NO<sub>x</sub> emissions prior to 1950 (i.e. before strong growth in man-made fertilizer use; Erisman et al., 2009) as the natural component. The 2000 anthropogenic portion (included in  
10 the agricultural sector, see Table 2) is estimated from EDGAR-v4, while the time evolution is based on Yan et al. (2005).

In all cases, gridding of the emissions for the 1850–2000 period relies on a weighted mean of the distributions obtained using either population (from the HYDE dataset) or the year 2000 gridded emissions. It is applied so that the weighting associated  
15 with the 2000 gridded distributions decreases when going back in time, with emissions after 1980 using the same grid as 2000; this is based on the assumption that, within a region, heavy infrastructure (such as power plants) has a very long (decades) lifespan. Although this approach might lead in few specific areas to shifts in source allocation, e.g., collapse of several economies in Eastern Europe in the 1990s “removed from the map” several industrial sources, we believe this has only limited impact on the  
20 the simulations intended using these historical sets of data.

No vertical emission profile is provided; however, the availability of sectoral emissions (energy, industry, domestic, etc.) in our emission files allows consistent assumptions about stack height to be applied if desired.

25 Speciation of NMVOC emissions is performed using the RETRO inventory. In this case, regional information for the split of the total NMVOC emitted into a set of specific hydrocarbons (Table 4) is available for the year 2000. Because of the lack of additional information, the same ratio (specific hydrocarbon to total NMVOCs at each grid point) is kept constant for the whole historical period.

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## 2 Biomass burning emissions

Only a few inventories provide biomass burning emissions for the past decades (Ito and Penner, 2005; Schultz et al., 2008; Mieville et al., 2010). In this paper, we focus on the following: (1) the RETRO inventory (Schultz et al., 2008) provides emissions from  
5 wildfires for each year during the 1960–2000 period, on a monthly basis; (2) the GICC inventory (Mieville et al., 2010) gives emissions from open biomass burning for the 20th century (1900–2000) on a decadal basis based on Mouillot et al. (2005) (3) the GFEDv2 inventory (van der Werf et al., 2006) covers emissions for the 1997–2006 period.

10 For our study, we have established a best guess estimate of historic biomass burning emissions from a combination of these three datasets: The GICC inventory is used as input data for the construction of the 1900–1950 dataset, the RETRO inventory for the 1960–1990 dataset and the GFEDv2 inventory for the 2000 estimate. The GFEDv2 inventory was favored over the 2000 estimate from RETRO because it is  
15 one of the most state-of-the art global biomass burning dataset currently available that incorporates satellite-based burned area estimates and seasonality.

Given the substantial interannual variability of biomass burning on a global and regional scale (e.g., Duncan et al., 2003; Schultz et al., 2008), it is problematic to use a snapshot dataset from an individual year for the development of a dataset that is  
20 considered to be representative for a decade. We therefore decided to construct historic gridded biomass burning emissions from decadal means (years 0 to 9 of a given decade), except for the 2000 estimate which is calculated from the 1997–2006 average.

In order to enforce consistency of biomass burning emissions over the entire period, carbon emission fluxes from the three datasets are first harmonized, taking the 2000  
25 estimate from GFEDv2; emissions of trace gases and aerosols are then re-calculated from the gridded carbon emission fluxes provided in the three datasets by applying a single set of vegetation-type specific emission factors. The vegetation cover map

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is derived from the MODIS predominant vegetation cover map as provided with the GFEDv2 inventory (van der Werf et al., 2006). It contains a classification of the year 2000 vegetation into the major vegetation classes savanna/herbaceous vegetation, tropical forest and extratropical forest. The emission factors for these classes were  
5 harmonized to those given by Andreae and Merlet (2001, with updates from M. O. Andreae, personal communication, M. O. Andreae, 2008).

Emissions from burning of soil organic matter, notably peat soil, which is ignited by fires in the overlying surface vegetation, may strongly influence emission production in some boreal and tropical regions (Page et al., 2002; Kasischke et al., 2005). Therefore,  
10 peat fires are explicitly taken into account in this new inventory. We assumed that peat fires can contribute up to 45% to the total carbon emissions released per grid cell if the fractional peat cover is 100%. If the fractional peat cover is lower, the relative contribution of carbon emissions from fires in surface vegetation increases accordingly. Note that this is an update from the original RETRO inventory. Information on the  
15 fractional distribution of peat soils is taken from the FAO (2003) WRB Map of World Soil Resources. The assumed maximal contribution of peat fires to the total carbon emission production refers a lower bound estimates provided for boreal and tropical peats (Kasischke et al. 2005; Heil, 2007). Emission factors for peat are taken from Christian et al. (2003) and Iinuma et al. (2007).

A monthly seasonality has been added to the original decadal GICC dataset; it is derived from the GFEDv2 seasonality (1997–2006) (van der Werf et al., 2006). The GFEDv2 seasonality was also used to redistribute in space and time total carbon emissions of the following regions in the attempt to improve the carbon emission patterns of the original RETRO data: Contiguous United States, Central America, South America,  
25 Northern Hemisphere Africa, Southern Hemisphere Africa, India, Continental Southeast Asia and Australia (for region definition; see Schultz et al., 2008). For the RETRO region Siberia and Mongolia, the redistribution was done using combined information from the GFEDv2 seasonality and the monthly Fire Danger Index (FDI) (described in Schultz et al., 2008). The yearly global total biomass burning carbon emissions of the

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original datasets remain unchanged from these corrections (deviations less than 1% from original value), while monthly totals may differ.

Biomass burning emissions are held constant between 1850 and 1900, as no additional information on burned area reconstructions is available (Mouillot et al., 2005).  
5 Furthermore, ice-core and charcoal records (McConnell et al., 2007; Marlon et al., 2008) indicate little variations during this time period. The time evolution of biomass burning emissions for the main compounds of interest is shown in Fig. 4.

In the distributed emission dataset, no information on vertical distribution is provided. As the separate distribution of grassland and forest fires are provided, users can apply  
10 methods similar to Lavoué et al. (2000) (for example) if deemed necessary.

Emissions from fuelwood burning and charcoal production, sometimes also labeled as biomass burning, are accounted for in anthropogenic residential sector emissions (see Sect. 2).

### 3 Ship and aircraft emissions

15 Total ship emissions, including international shipping, domestic shipping and fishing, but excluding military vessels, are taken from a recent assessment by Eyring et al. (2009) to reflect updated information on the fleet and emission factors. In this latter study, estimates of fuel consumption and CO<sub>2</sub> in the year 2000 are based on the International Maritime Organization (IMO) study discussed in Buhaug et al. (2008), while  
20 the best-estimate for non-CO<sub>2</sub> emission totals is derived as a mean of previous studies (Corbett and Köhler, 2003; Eyring et al., 2005; Endresen et al., 2003, 2007). Ship emissions are distributed over the globe using the International Comprehensive Ocean-Atmosphere Data Set (ICOADS; Wang et al., 2007), which provides changing shipping patterns on a monthly basis. NMVOC emissions from crude oil transport (evaporation  
25 during loading, transport, and unloading) from Endresen et al. (2003) were added to the gridded ICOADS ship emission dataset. Consistent with our treatment of other anthropogenic sources, ship emission totals are spread onto the 0.5°×0.5° grid boxes

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without accounting for dispersion, chemical transformation and loss processes on the sub-grid scale. Neglecting the plume processes in global models may lead to an over-estimation of ozone formation (Franke et al., 2008; Charlton-Perez et al., 2009).

To extend the non-CO<sub>2</sub> ship emissions backward in time, the 2000 emission totals from Eyring et al. (2009) are scaled with the historical CO<sub>2</sub> emission time series from Buhaug et al. (2008) back to 1870. Emissions for 1850 and 1860 are estimated by scaling changes in global ship tonnage as collated by Bond et al. (2007). For the historical geographical ship distribution we use ICOADS data back until 1950. Prior to 1950s there was much less activity by ships on Pacific routes, with shipping concentrated on the North and South Atlantic oceans, Mediterranean, and Indian/Indonesian trade routes (personal communication, J. Corbett, 2008). To map emissions before 1950, the Pacific trade routes are therefore constrained to go to zero in 1900, forcing the emissions to be concentrated in the remaining regions. Between 1900 and 1950 a linear interpolation between these patterns is applied to provide decadal gridded ship emissions.

Aircraft emissions of NO<sub>x</sub> and BC are calculated using the FAST model (Lee et al., 2005) for the European Quantify project (<http://www.pa.op.dlr.de/quantify/>). Global scheduled and non-scheduled aircraft movements are taken from the AERO2K database (Eyers et al., 2005) for the year 2002. Fuel consumption is calculated using the industry-standard PIANO aircraft performance model (Simos, 2004) for all the main aircraft types including four categories of turbo-props. Fuel consumption is then assigned to the routes using a great-circle assumption and NO<sub>x</sub> emissions calculated with the Deutsches Zentrum für Luft-und Raumfahrt fuel flow method (Lecht, 1999). BC emissions are calculated using the emission factors developed from Eyers et al. (2005). Monthly distributions were calculated and provided on a 3D grid with an original resolution of 1° × 1° latitude/longitude and vertical discretization of flight levels of 2000 feet, which corresponds to the actual (pressure) levels used by air traffic. The vertical distribution of traffic was parameterized from a statistical analysis of EUROCONTROL air traffic data that provided a relationship between mission distance, aircraft type and

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average cruise altitude (Lee et al., 2005). By convention, aviation “bottom up” inventories underestimate fuel and emissions for a variety of reasons (perfect routing, no stacking, limited data on non-scheduled traffic, military aircraft) when compared with International Energy Agency (IEA) statistics of kerosene sales, so that the three-dimensional inventory is scaled up to the IEA kerosene data to ensure that the global and annual totals are the same.

Historical emissions from aviation are provided on an annual total basis. Data from 1940 to 1995 are taken from Sausen and Schumann (2000) and extended to 2000 using IEA data (the basis of the time series of Sausen and Schumann, 2000). Emissions prior to 1940 are assumed to exponentially decay such that the emissions by 1910 are zero. This is clearly of limited importance since emissions in 1940 and before are believed to be quite small.

#### 4 Application

We have described above (see Figs. 2–4) how changes in emissions at the global scale have been very significant between 1850 and 2000. It is however important to remember that these changes have very different regional characteristics (Fig. 5 for the specific case of NO<sub>x</sub> emissions) and only global three-dimensional chemistry-climate models can fully capture the implication. Therefore, in this section, we discuss the application of the emissions described above to the simulation of tropospheric composition changes between 1850 and present by two chemistry-climate models: CAM-Chem and G-PUCCINI. Of those two models only CAM-chem includes an interactive representation of aerosols (i.e. G-PUCCINI reads in previously generated aerosol distributions). Analysis of the modeled results against available observations provides an initial understanding of successes and limitations of the emissions described in this paper.

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## 4.1 Model description and simulation setup

### 4.1.1 CAM-Chem

We use the Community Atmosphere Model version 3.5 (Gent et al., 2009) modified to include interactive chemistry (i.e. with feedback to the radiation calculation in the atmosphere) to calculate distributions of gases and aerosols. The model configuration used in this study includes a horizontal resolution of 1.9° (latitude) by 2.5° (longitude) and 26 hybrid levels, from the surface to ≈40 km with a timestep of 30 min; the transient simulation was performed continuously between 1850 and 2009. In order to simulate the evolution of the atmospheric composition over the recent past, the chemical mechanism used in this study is formulated to provide an accurate representation of both tropospheric and stratospheric chemistry (Lamarque et al., 2008). Specifically, to successfully simulate the chemistry above 100 hPa, we include a representation of stratospheric chemistry (including polar ozone loss associated with stratospheric clouds) from version 3 of MOZART (MOZART-3; Kinnison et al., 2007). The tropospheric chemistry mechanism has a limited representation of non-methane hydrocarbon chemistry in addition to standard methane chemistry, extended from Houweling et al. (1998) with the inclusion of isoprene and terpene oxidation and updated to JPL-2006. This model has a representation of aerosols based on the work by Tie et al. (2001, 2005), i.e. sulfate aerosol is formed by the oxidation of SO<sub>2</sub> in the gas phase (by reaction with the hydroxyl radical) and in the aqueous phase (by reaction with ozone and hydrogen peroxide). Furthermore, the model includes a representation of ammonium nitrate that is dependent on the amount of sulfate present in the air mass following the parameterization of gas/aerosol partitioning by Metzger et al. (2002). Because only the bulk mass is calculated, a lognormal distribution is assumed for all aerosols using different mean radius and geometric standard deviation (Liao et al., 2003). The conversion of carbonaceous aerosols (organic and black) from hydrophobic to hydrophilic is assumed to occur within a fixed 1.6 days. Natural aerosols (desert dust and sea salt)

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are implemented following Mahowald et al. (2006a and b), and the sources of these aerosols are derived based on the model calculated wind speed and surface conditions.

At the lower boundary, the time-varying (monthly values) zonal-averaged distributions of CO<sub>2</sub>, CH<sub>4</sub>, H<sub>2</sub> and all the halocarbons (CFC-11, CFC-12, CFC-113, HCFC-22, H-1211, H-1301, CCl<sub>4</sub>, CH<sub>3</sub>CCl<sub>3</sub>, CH<sub>3</sub>Cl and CH<sub>3</sub>Br) are specified following the datasets used in Garcia et al. (2007). In addition, the monthly-mean time-varying sea-surface temperatures (SSTs) and sea-ice distributions are taken from a 20th century CCSM-3 simulation (Meehl et al., 2008); as this simulation only extended from 1870 to present, the sea-surface temperature and ice extent between 1850 and 1870 are assumed to be the same as 1870.

### 4.1.2 G-PUCCINI

Simulations are performed with the Goddard Institute for Space Studies (GISS) model for Physical Understanding of Composition-Climate INteractions and Impacts (G-PUCCINI) (Shindell et al., 2006b). Its behavior in the GISS AR4 version of the climate model has been documented and extensively compared with observations (e.g. Dentener et al., 2006a; Shindell et al., 2006a, b; Stevenson et al., 2006). Tropospheric chemistry includes basic NO<sub>x</sub>-HO<sub>x</sub>-O<sub>x</sub>-CO-CH<sub>4</sub> chemistry as well as peroxyacetyl nitrates and the hydrocarbons isoprene, alkyl nitrates, aldehydes, alkenes, and paraffins. The lumped hydrocarbon family scheme was derived from the Carbon Bond Mechanism-4 (CBM-4) and from the more extensive Regional Atmospheric Chemistry Model (RACM), following Houweling et al. (1998). To represent stratospheric chemistry, the model includes chlorine- and bromine-containing compounds, and CFC and N<sub>2</sub>O source gases. The chemistry used here is quite similar to that documented previously, with a few additions: acetone has been added to the hydrocarbons included in the model following (Houweling et al., 1998), polar stratospheric cloud formation is now dependent upon the abundance of nitric acid, water vapor and temperature (Hanson and Mauersberger, 1988), and a reaction pathway for HO<sub>2</sub>+NO to yield HNO<sub>3</sub> has been

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added (Butkovskaya et al., 2007). Chemical calculations are performed seamlessly throughout the troposphere and stratosphere. The full scheme includes 156 chemical reactions among 50 species with a time step of 20 min. Photolysis rates are calculated using the Fast-J2 scheme (Bian and Prather, 2002), whereas other chemical reaction rate coefficients are from JPL-2000 (Sander et al., 2000).

The chemistry model is fully embedded in the GISS modelE climate model (Schmidt et al., 2006). For the simulations described here, we have used the development version of the model near its “frozen” state for AR5 simulations. This version of the model has an equilibrium climate sensitivity of 3.7 °C for a doubling of CO<sub>2</sub>. The model was run at 2° latitude by 2.5° longitude Cartesian horizontal resolution, with increased effective resolution for tracers by carrying higher order moments at each grid box. This configuration had 40 vertical hybrid sigma layers from the surface to 0.01 hPa (≈80 km). Tracer transport uses a non-diffusive quadratic upstream scheme (Prather, 1986). Time-slice simulations were performed every 20 years during the 1850–1930 time period, and every 10 years from 1930–2000. Values were then interpolated to give decadal means. Simulations were carried out for 8 years, with the average of the last five used for analysis. The GCM was driven by observed decadal mean sea-surface temperatures and sea-ice distribution (Rayner et al., 2003) and prescribed abundances of long-lived greenhouse gases.

## 4.2 Evaluation of model results

In this section, we focus our evaluation to long-term trends in surface and mid-troposphere ozone (both models), surface concentration of carbon monoxide (both models) and aerosol optical depth and aerosol deposition (CAM-chem only); indeed, the main purpose for the emission dataset described above is to be used for studies of long-term changes in tropospheric composition of relevance to climate radiative forcing. Note that both model simulations use constant 2000 emissions after year 2000 (up to 2009). Emissions beyond year 2000 will be discussed elsewhere (van Vuuren et al., 2010).

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### 4.2.1 Surface ozone

Observations of surface ozone over the last decades indicate a significant rise in the Northern Hemisphere (Parrish et al., 2004; Oltmans et al., 2006; Derwent et al., 2007). We focus here on stations with at least 20 years of observations (Table 5), providing timeseries for comparison with model results. Model ozone fields are interpolated to the location of the observations, including altitude. Because the observations over the United States Pacific coast (see Table 5) are filtered to only provide background conditions (Parrish et al., 2004), the analysis of the model results is performed using surface ozone from approximately 200 km west of the actual station location. In addition, in the case of Mace Head, background (Derwent et al., 2007) and unfiltered surface ozone (from <http://tarantula.nilu.no/projects/coc/emepdata.html>) observations are available; in this case, we use the unfiltered data, as they are more representative of the modeled field.

Over the European sites (Fig. 6, top), both models (with small inter-model difference) provide a good representation of present-day observed surface ozone, except at Arkona where G-PUCCINI tends to overpredict the observed concentration by approximately 8 ppbv.

Going back in time, until 1990, the model results and the observations agree quite well with each other, indicating that the recent trends in emissions are probably well captured for the this region. Prior to 1990, the observations tend to decrease substantially faster (approximately twice as fast) than the models simulate. In particular, observations in Arkona during the mid-1980s show very low values (less than 10 ppbv). Without any indication of problem with the observations, it is clear that the models are not able to represent such distribution. A possible explanation is that highly localized emission patterns not captured by our emission dataset could have led to those very low ozone concentrations. The overall agreement between the two models in the simulated long-term tendencies (Fig. 6) indicate that the smaller-than-observed modeled trend prior to 1990 is most likely due to a limitation of our emission datasets.

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Over the US Pacific Coast, the models are again similar to one another, but neither matches the rapid increase in surface ozone seen in observations in recent years (Parrish et al., 2004). This is perhaps not surprising as emissions changes were only determined until 2000.

5 However, additional long-term records of surface ozone (Barrow, Alaska; Mauna Loa, Hawaii; Cape Matatula, American Samoa; South Pole, Antarctica, Fig. 7) show a different picture, in which changes in ozone in the 1980s are not increasing as rapidly, if at all. In particular, the record at Samoa actually indicates a long-term decrease in ozone, contrary to the findings of Lelievre et al. (2004). In all those places, both  
10 models perform quite well in their capture of the long-term trends (note that, for visualization purposes, a constant bias of 6 ppbv was added to the observed record at Barrow to match the simulated levels; this is likely due to the specific environment at Barrow, at the edge of the Arctic Ocean, which is difficult to capture with a coarse-grid global model). At Samoa, climate trends may have played a substantial role in the apparent decrease between ~1990 and ~2000 in the observations, as the models have rather different trends despite the same emissions data. The use of observed SSTs in the G-PUCCINI simulations may have allowed it to capture local climate changes that could have contributed to the recent ozone decline seen in that remote Western Pacific location. At the South Pole, there is indication of the impact of stratospheric ozone depletion, bringing minimal levels during the mid-1990s, followed by a slight recovery and leveling-off since 2000 (Chipperfield et al., 2007). CAM-chem is able to capture this trend, while G-PUCCINI tends to underestimate ozone in 2000, apparently due to an overestimate of downward transport of air having experienced Antarctic stratospheric ozone depletion (unlike the surface, stratospheric ozone did not recover to the values  
25 seen in the 1970s and early 1980s by 2000; Chipperfield et al., 2007).

#### 4.2.2 Surface carbon monoxide

Comparison (Fig. 8) of modeled and observed (averaged 1990s conditions) surface carbon monoxide at Mace Head (a useful comparison since this station also provides

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surface ozone, Fig. 6) indicates a negative bias (approximately 20 ppbv) and a reduced seasonal cycle, with a larger negative bias during winter. These features are present in both simulations and are found at most of the Northern Hemisphere stations (not shown); agreement during summer conditions seems to always be slightly  
5 better than in the winter. Biases in the Southern Hemisphere are much smaller (especially for CAM-chem, not shown). The overall Northern Hemisphere negative bias in both model simulations indicates that the overall emissions of carbon monoxide (and possibly NMVOCs) are underestimated in this bottom-up inventory; this underestimation is likely in the anthropogenic emissions as this bias is present year-round.  
10 However, owing to the long lifetime of carbon monoxide during winter (up to a few months; Edwards et al., 2005), it is also possible that biomass burning emissions in the latter part of the year over Russia are not well enough characterized to provide the wintertime maximum (Edwards et al., 2005). Underestimation of CO emissions could also be caused by the underestimation of fuelwood combustion (activity data problem) or an overestimation of combustion technologies (too high combustion efficiency). However, further analysis (beyond the scope of this paper) is required to fully understand the reason for this low bias. The long-term change (between 1990 and present, Fig. 9) in carbon monoxide at Mace Head (using unfiltered observations, see <http://tarantula.nilu.no/projects/coc/emepdata.html>) shows that the models capture the recent change relatively well; it is clear however that this analysis suffers from the lack of long-term (>30 years) records. Interestingly, the simulated change in surface CO at Mace Head between 1960 and 1990 is quite different between the two models, contrary to very similar ozone change over the same period.

25 The lifetimes of CO and CH<sub>4</sub> can be used as global measure of the OH content of the atmosphere. For the 2000 conditions, the methane chemical lifetime (i.e. not including the small deposition flux as the simulations were performed with specified bottom layer methane concentration) is 8.9 years for CAM-chem and 8.6 years for G-PUCCINI, in excellent agreement with the IPCC AR-4 estimates of 8.7±1.3 years (Denman and Brasseur, 2007). For the same period, the CO lifetime is 1.7 months for CAM-chem,

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in good agreement with Horowitz et al. (2002). There is therefore no indication that a significant bias in OH in these models could explain the low bias in CO.

### 4.2.3 Mid- and upper-tropospheric ozone

A compilation of mid-tropospheric ozone observations from a variety of platforms (Cooper et al., 2010) indicates that Western United North America (25°–55° N, 130°–90° W, 3–8 km) has experienced a recent (1995–2008) increase in ozone concentration (Fig. 10), most likely associated with Asian emissions. The model results indicate again that they are performing very well in estimating the present-day (2000) ozone concentration. Evidently, they are not able to reproduce inter-annual variability but our 5- to 10-yr averages are very close to the observed values. Similarly to Fig. 6, the two models exhibit a very consistent long-term evolution. The post-2000 observations are showing a rapid increase that the models, owing to their use of constant 2000 emissions, are not capturing. Furthermore, the (limited) 1984 dataset provides a much lower ozone concentration than the models indicate, similar to the surface ozone analysis above.

### 4.2.4 Aerosol optical depth, burden and lifetime

A useful measure of the radiative impact of aerosols can be evaluated through the calculation of the aerosol optical depth (Schulz et al., 2006). We display in Fig. 11 the CAM-chem simulated annual average total aerosol optical depth (AOD) at 550 nm for 1850 and 2000. The occurrence of widespread pollution over the Northern Hemisphere is clearly identifiable, while the natural contributions of dust, sea-salt and background sulfate (from dimethylsulfide oxidation and sulfur dioxide non-eruptive volcanic emissions, as both are kept constant during the simulations) are unchanged. In terms of the global average, the 2000 AOD simulated value is 0.12, which represents an increase of 0.033 over the 1850 conditions (0.087). This anthropogenic increase is very much in agreement with the average AEROCOM results (Schulz et al., 2006). We also compare our annual average aerosol optical depths to AERONET sun photometer

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site data at 173 sites globally (Holben, et al., 1998). Sun photometry data represents some of the highest quality data for assessing total aerosol optical depth. We include all stations where monthly averages at 500 nm were available for all 12 months. The model able to capture much of the variability, but underestimates the aerosols optical depth at high observed values (Fig. 12). The correlation coefficient between modeled values and observations is 0.67. In terms of dust and sea-salt aerosols, comparison with surface observations (using iron deposition as a proxy, see Fig. 13) indicates a reasonable representation of present-day conditions.

An additional important evaluation for aerosol is their global burden and lifetime. Results for the 2000 conditions are summarized in Table 6. Compared to the AEROCOM results (Schulz et al., 2006), the lifetime of carbonaceous aerosols is approximately 2 days shorter (from approximately 7.5 days to 5.5 days), leading to a smaller burden. On the other hand, sulfate lifetime is almost exactly the same, as is the anthropogenic contribution (i.e. the difference between 2000 and 1850 burdens).

### 4.2.5 Aerosol ice-core deposition

Ice core measurement of aerosol and gas content can provide information on long-term changes in deposition and concentration. In particular, Greenland ice cores have been recently used to study the importance of black carbon in the Arctic (McConnell et al., 2007). The model results (wet and dry deposition of sulfate) are interpolated to the model grid point nearest to the D4 ice core site (71.4° N, 44° W) with the closest model topography altitude to D4 (approx. 100 km north of the actual D4 location); indeed, precipitation patterns (and therefore deposition) exhibit a strongly decreasing latitudinal gradient across the Greenland ice sheet. There is a remarkable agreement (Fig. 14, top) between the observations and the simulated deposition. In terms of sulfate, the maximum deposition rate (40 mg/m<sup>2</sup>/year in the observations) occurs in 1980, when the global emissions (but especially over the United States and Russia) peaked (Fig. 3). There is also indication of a local maximum sulfate deposition at the beginning of the 20th century in both the observations and the model field.

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Similarly (Fig. 14, bottom), black carbon (hydrophilic only) deposition at D4 has peaked in the early part of the 20th century. We find that, using the same sampling procedure as for sulfate, the model captures that feature quite well (albeit not as strongly as the observations suggest), along with the overall changes over the simulated period. This is again indicative of adequate regional emission changes in North America as Greenland deposition is most strongly influenced by emissions in that region (Shindell et al., 2008), in this case related to changes (in both anthropogenic emissions increasing, see Fig. 3) and biomass burning (decreasing, see Fig. 4).

## 5 Discussion and conclusions

We have presented in this paper a new set of historical anthropogenic (defined here as originating from industrial, domestic and agriculture activity sectors) and biomass burning emissions of reactive gases and aerosols covering 1850–2000. This dataset is the result of a large (and first of its kind) community effort to bring together expertise from various sectors and existing inventories; as such, it builds upon and complements previous inventories. Our dataset represents a combination of existing regional and global inventories, and the combination of information was performed on a regional (40 regions) and sectoral (13 sectors) representation. Some detailed inventories were not included as they had a limited scope (e.g. national inventories for single or limited number of compounds or sectors). Furthermore, the use of a variety of inventories precludes full consistency between carbon dioxide, reactive gases and aerosol emissions for anthropogenic, biomass burning, land-use and natural emissions. It is unclear how important this lack of full consistency is, but it will be important to focus on this issue in future similar emission datasets. Finally, no emission uncertainty is provided; based on the recent studies by Bond et al. (2004, 2007) and Smith et al. (2010; S. J. Smith, J. van Aardenne, Z. Klimont, R. Andres, A. Volke and S. Delgado Arias, “Historical Sulfur Dioxide Emissions: 1850–2005”, 2010), it can be expected to be large (a factor of 2 or larger) for some compounds and

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regions. All data are publicly available at <http://www.iiasa.ac.at/web-apps/tnt/RcpDb> and <ftp://ftp-ipcc.fz-juelich.de/pub/emissions/>.

The primary purpose of this inventory is to provide emissions for chemistry-climate simulations (with the Climate Model Intercomparison Program #5 in support of the IPCC AR5 as the overall focus) for the study of long-term changes in atmospheric composition. In particular, the emissions for year 2000 serve as an anchor point for historical emissions (as discussed in this paper) and future emissions (as discussed in van Vuuren et al., 2010). This ensures continuity in emission datasets throughout the IPCC period of interest (1850–2100). Because of its focus on long-term changes, this dataset provides emissions every 10 years and does not attempt to reproduce interannual variability, which can be significant, particularly for biomass burning emissions.

Using two chemistry-climate models, we have performed 1850–2009 simulations (transient or time-slice experiments) in order to provide a first-order evaluation of the emissions. The focus of this evaluation is on long-term changes of tropospheric species relevant to climate forcing. In particular, we find that the model simulations for the 1990–2000 conditions represent quite well the observed surface and mid-troposphere ozone distributions. There is however indication that the modeled long-term increase since the early 1980s is not as strong and rapid as recent publications indicate (Parrish et al., 2004; Cooper et al., 2010). On the other hand, comparison with other long-term ozone records (Barrow, Mauna Loa, Samoa and South Pole) shows good agreement for the available period 1970–2000; there is therefore clearly a need for understanding ozone changes at the regional scale. We found that carbon monoxide is biased low in both models; the reason for this bias (present in many sites over the Northern Hemisphere but not so much in the Southern Hemisphere) is not clear at this point.

Ice-core deposition of sulfate and black carbon over Greenland is well simulated (albeit only the CAM-chem model has simulated aerosols) in both amplitude and long-term trend. In particular, the black carbon maximum at the turn of the 20th century is a combination of increasing anthropogenic and decreasing biomass burning emissions. In addition, global measures of aerosol content are inline with the AEROCOM

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estimates for present-day burden and lifetime, especially for sulfate. Finally, aerosol optical depth comparison with AERONET observations indicates a reasonably good simulation of present-day conditions.

The observations of long-term changes in atmospheric composition clearly indicate large regional variations. As discussed in our paper, modeling these changes is a difficult challenge that combines the role of changing emissions and changing climate; but it is the key to our understanding of future air quality (Dentener et al., 2006b).

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**Table 1.** List of multi-decadal inventories used in this study.

Species	EDGAR-HYDE	RETRO	Smith et al.	Bond et al.	Junker and Liousse
	EDGAR	1960–2000	1850–2000	1850–2000	1860–1997
CH <sub>4</sub>	X				
CO	X	X			
NO <sub>x</sub>	X	X			
NMVOC	X	X			
NH <sub>3</sub>	X				
SO <sub>2</sub>			X		
OC				X	X
BC				X	X

5000

**Table 2.** List of regions.

Region number	Name of Region
1	Canada
2	USA
3	Mexico
4	Rest of Central America
5	Brazil
6	Venezuela
7	Argentina
8	Rest of South America
9	Northern Africa
10	Western Africa
11	Eastern Africa
12	Rest of Southern Africa
13	South Africa
14	France
15	Germany
16	Italy
17	UK
18	Rest of Western Europe
19	Rest of Central Europe
20	Baltic States (Estonia, Latvia, Lithuania)
21	Turkey
22	Ukraine
23	Asia-Stan
24	Russia
25	Middle East
26	India
27	Rest of South Asia
28	South Korea (Republic of Korea)
29	North Korea (Democratic People's Republic of Korea)
30	China
31	Taiwan
32	Thailand
33	Rest of Southeastern Asia
34	Indonesia
35	Japan
36	Australia
37	New Zealand
38	Rest of Oceania
39	Greenland
40	Antarctica

5001

**Table 3.** List of sectors.

Sector number	Sector name
1	Energy production and distribution
2	Industry (combustion and non-combustion)
3	Land transport
4	Maritime transport
5	Aviation
6	Residential and commercial
7	Solvents
8	Agriculture
9	Agricultural waste burning on fields
10	Waste
11	Open vegetation fires in forests
12	Open vegetation fires in savanna and grasslands
13	Natural emissions

5002

**Table 4.** NMVOC speciation.

name	molecular weight	comments
alcohols	46.2	assumed C <sub>2</sub> H <sub>5</sub> OH
ethane	30.0	
propane	44.0	
butanes	57.8	
pentanes	72.0	
hexanes and higher	106.8	
ethene	28.0	
propene	42.0	
ethyne	26.0	
other alkenes and alkynes	67.0	
benzene	78.0	
toluene	92.0	
xylene	106.0	
trimethyl benzenes	120.0	
other aromatics	126.8	
esters	104.7	surrogate species = CH <sub>3</sub> C(O)O(CH <sub>2</sub> ) <sub>n</sub> CH <sub>3</sub>
ethers	81.5	surrogate species = CH <sub>3</sub> CH <sub>2</sub> O(CH <sub>2</sub> ) <sub>n</sub> CH <sub>3</sub>
chlorinated HC	138.8	
methanal (CH <sub>2</sub> O)	30.0	
other alkanals	68.8	
ketones	75.3	
acids	59.1	
other VOC	68.9	use median C ratio of other compounds

5003

**Table 5.** List of stations.

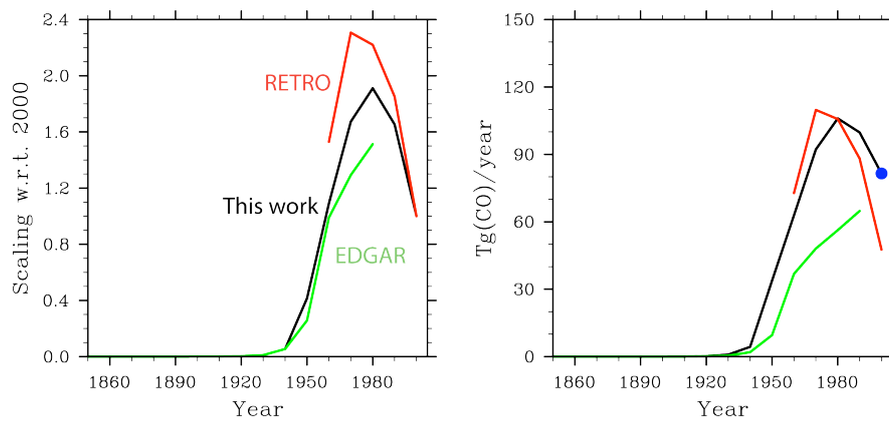
Station name	Latitude	Longitude	Altitude (m)	Period
Arkona/Zingst	54.4 N	13.26 E	42	Sep 1956–Jun 2004
Barrow	71.32 N	156.6 W	11	Mar 1973–Dec 2005
Cape Matatula	14.24 S	170.57 W	42	Sep 1975–Dec 2004
Hohenpreissenberg	47.89 N	11.02 E	985	Jan 1995–Dec 2007
Mace Head	53.33 N	9.9 W	25	Nov 1987–Sep 2006
Maunao Loa	16.54 N	155.58 W	3397	Sep 1973–Dec 2004
South Pole	89.90 S	24.8 W	2810	Jan 1975–Dec 2004
US Pacific Coast	N/A	N/A	N/A	Mar 1988–Feb 2007
Zugspitze	47.42 N	10.98 E	2960	Jan 1995–Dec 2002

5004

**Table 6.** Global burden and lifetime for anthropogenically-perturbed aerosols in 1850 and 2000 for the CAM-chem simulation.

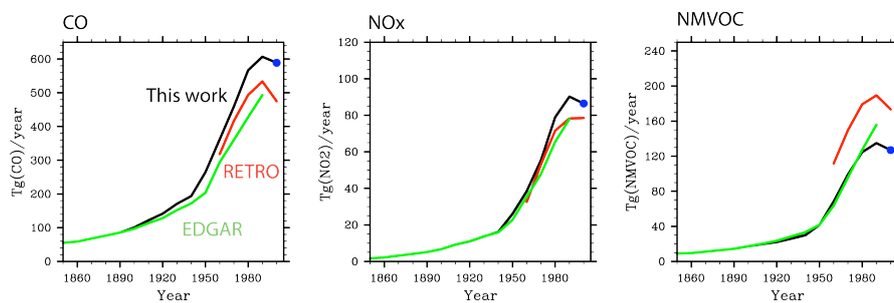
	1850	2000
<b>Sulfate</b>		
Burden ( $\text{mg}(\text{SO}_4)/\text{m}^2$ )	1.55	3.65
Lifetime (days)	3.4	3.6
<b>Black carbon</b>		
Burden ( $\text{mg}(\text{C})/\text{m}^2$ )	0.09	0.24
Lifetime (days)	5.6	5.8
<b>Organic carbon</b>		
Burden ( $\text{mg}(\text{C})/\text{m}^2$ )	0.64	1.04
Lifetime (days)	5.2	5.4

5005



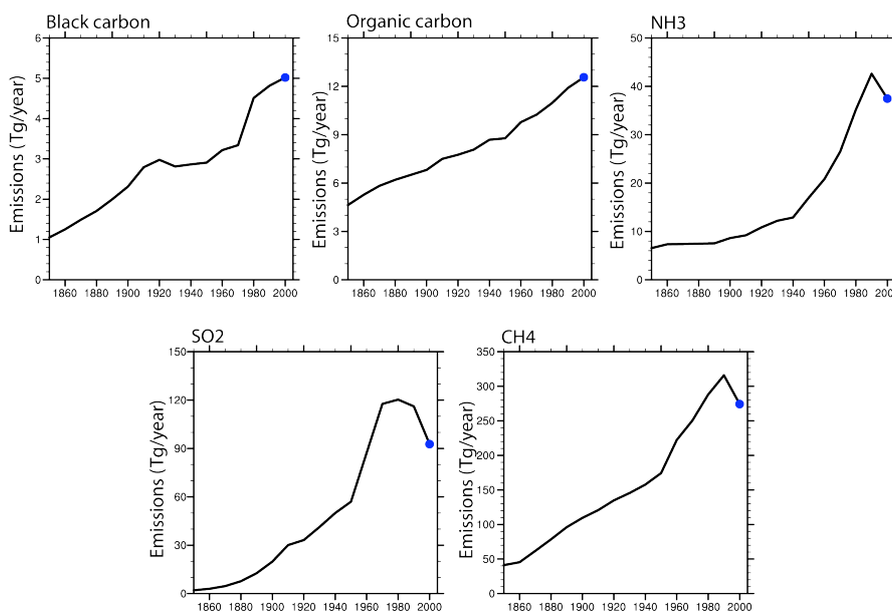
**Fig. 1.** Time evolution of the United States CO transportation sector emissions. Left panel is scaling with respect to 2000 (dimensionless). Right panel shows actual emissions ( $\text{Tg}(\text{CO})/\text{year}$ ).

5006



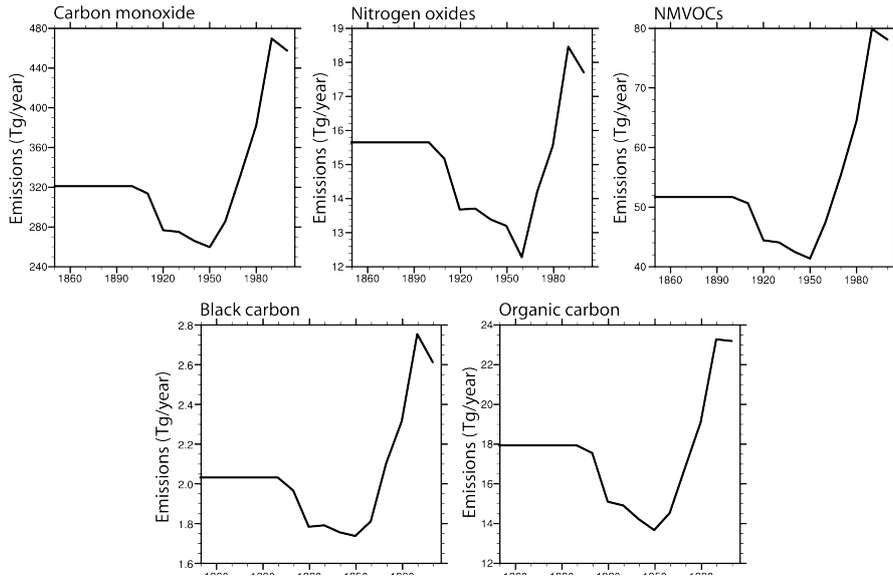
**Fig. 2.** Time evolution of the total (sum of all sectors but agricultural waste burning) land anthropogenic emissions for CO (Tg(CO)/year), NO<sub>x</sub> (Tg(NO<sub>2</sub>)/year) and total NMVOC (Tg(NMVOC)/year).

5007



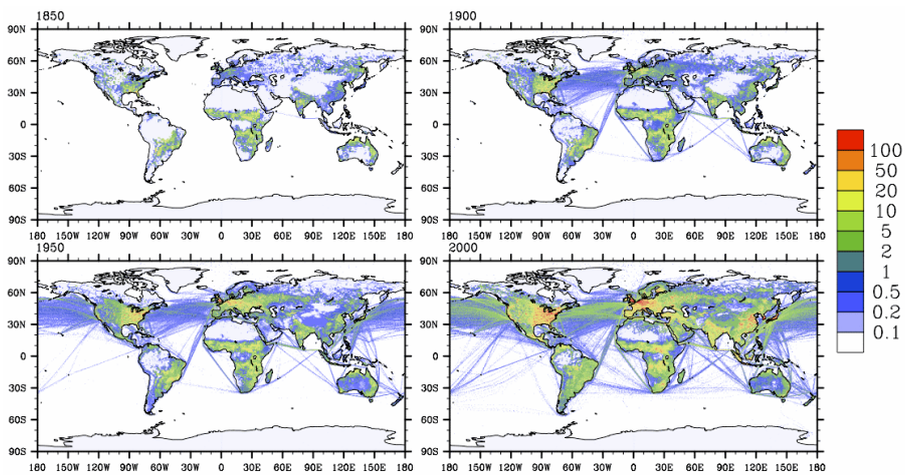
**Fig. 3.** Time evolution of the total (sum of all sectors) land anthropogenic emissions for black carbon (Tg(C)/year), organic carbon (Tg(C)/year), ammonia (Tg(NH<sub>3</sub>)/year), sulfur dioxide (Tg(SO<sub>2</sub>)/year), and methane (Tg(CH<sub>4</sub>)/year).

5008



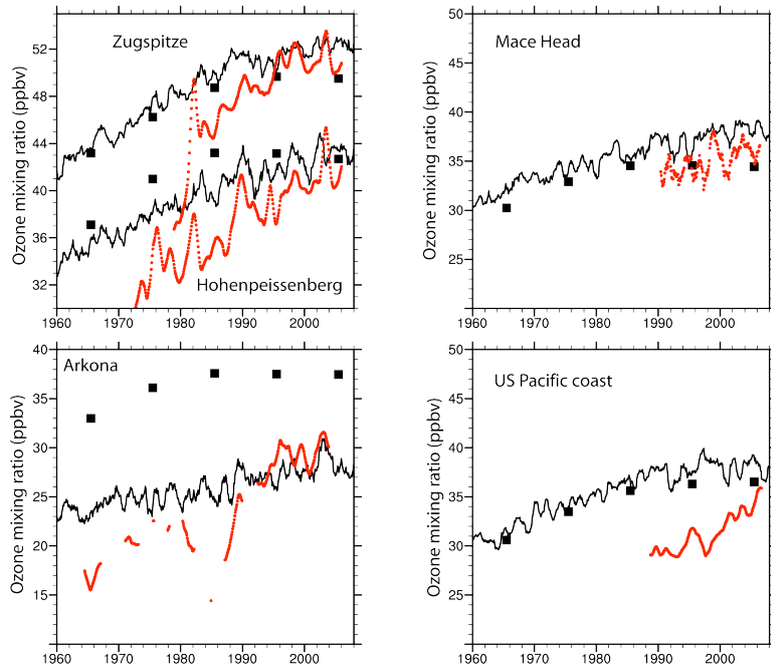
**Fig. 4.** Time evolution of the total open biomass burning (forest and grassland) emissions for carbon monoxide (Tg(CO)/year), NO<sub>x</sub> (Tg(NO<sub>2</sub>)/year), NMVOC (Tg(NMVOC)/year), black carbon (Tg(C)/year) and organic carbon (Tg(C)/year).

5009



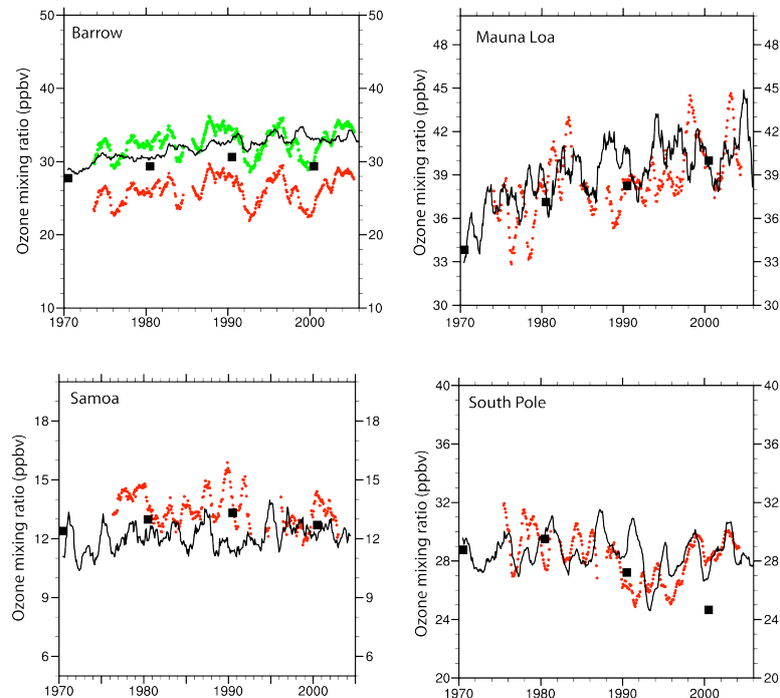
**Fig. 5.** Total annual emissions (anthropogenic, shipping and biomass burning) of NO<sub>x</sub> (Tg(N)/year) for 1850 (top left), 1900 (top right), 1950 (bottom left) and 2000 (bottom right).

5010



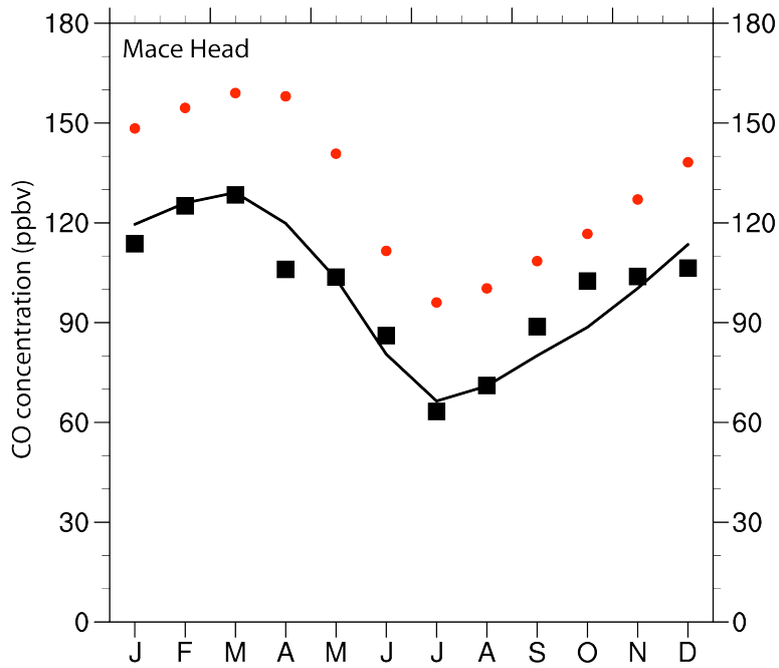
**Fig. 6.** Time evolution (1960–2009) of surface ozone (12-month running mean) at a variety of sites. Observations are in red, CAM-chem results are in black (solid line) and G-PUCCINI results are in black (solid squares). Note the emissions beyond 2000 are kept at their 2000 level.

5011



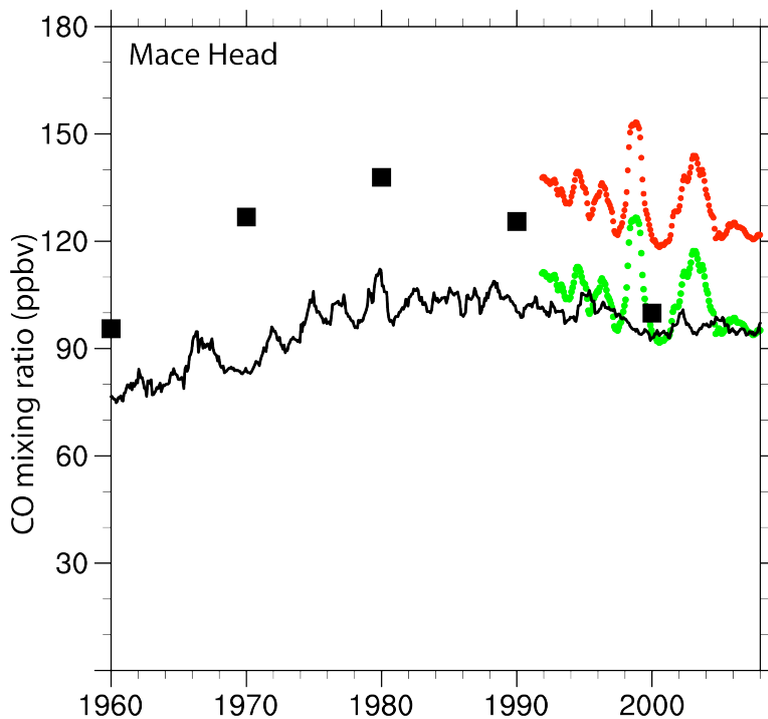
**Fig. 7.** Time evolution (1970–2009) of surface ozone (12-month running mean) at a variety of sites. Observations are in red, CAM-chem results are in black (solid line) and G-PUCCINI results are in black (solid squares). Note the emissions beyond 2000 are kept at their 2000 level. A constant value of 6 ppbv was added to the Barrow observations to take into account model deficiencies; this is shown as green dots.

5012



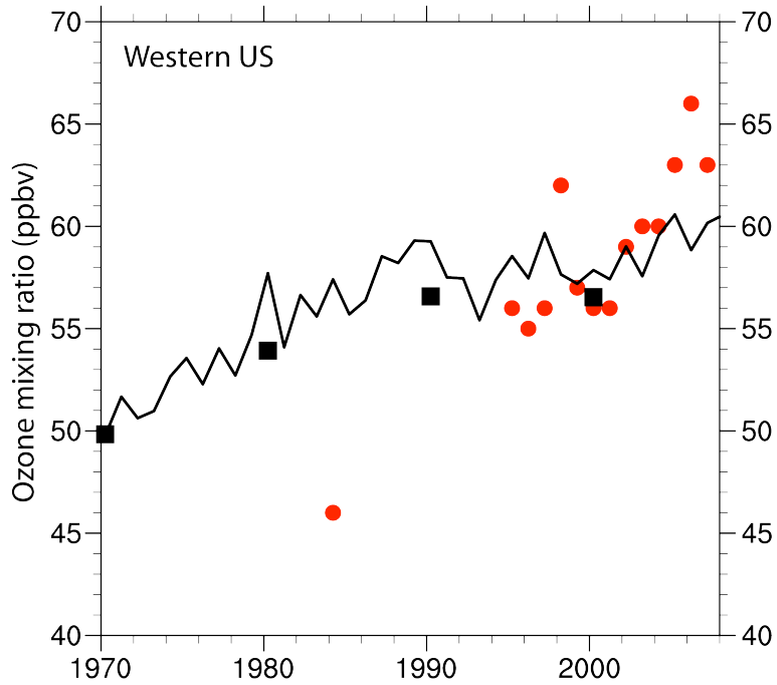
**Fig. 8.** Seasonal cycle of carbon monoxide (ppbv) at Mace Head for the 1990s. Observations (averaged 1991–1999) are in red, CAM-chem results are in black (solid line) and G-PUCCINI results are in black (solid squares).

5013



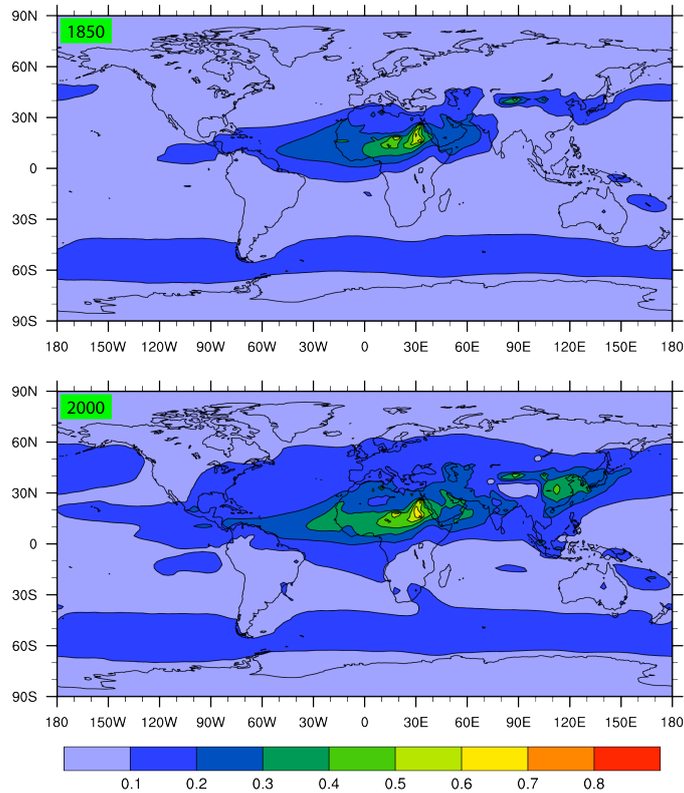
**Fig. 9.** Time evolution (1960–2009) of surface CO (12-month running mean) at Mace Head. Observations are in red, CAM-chem results are in black (solid line) and GPUCCINI results are in black (solid squares). Note the emissions beyond 2000 are kept at their 2000 level. A constant value of 25 ppbv was subtracted from the observations to take into account model deficiencies; this is shown as green dots.

5014



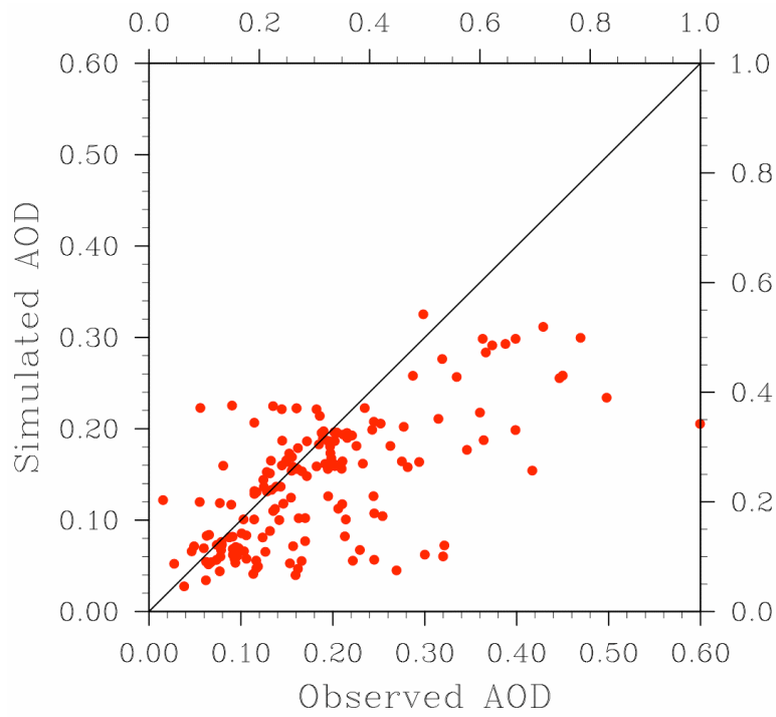
**Fig. 10.** Time evolution (1970–2009) of mid-troposphere ozone (springtime mean) from Cooper et al. (2010). Observations are in red, CAM-chem results are in black (solid line) and G-PUCCINI results are in black (solid squares). Note the emissions beyond 2000 are kept at their 2000 level.

5015



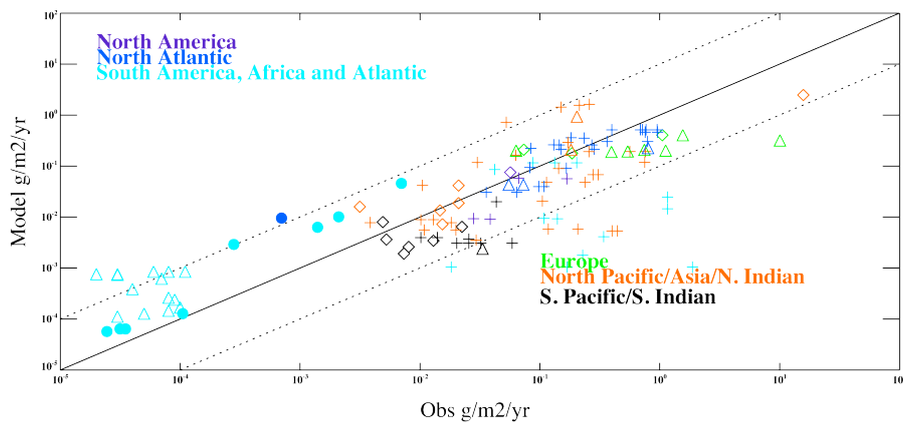
**Fig. 11.** Total (natural and anthropogenic) CAM-chem simulated aerosol optical depth at 550 nm (decadal average) for 1850 and 2000.

5016



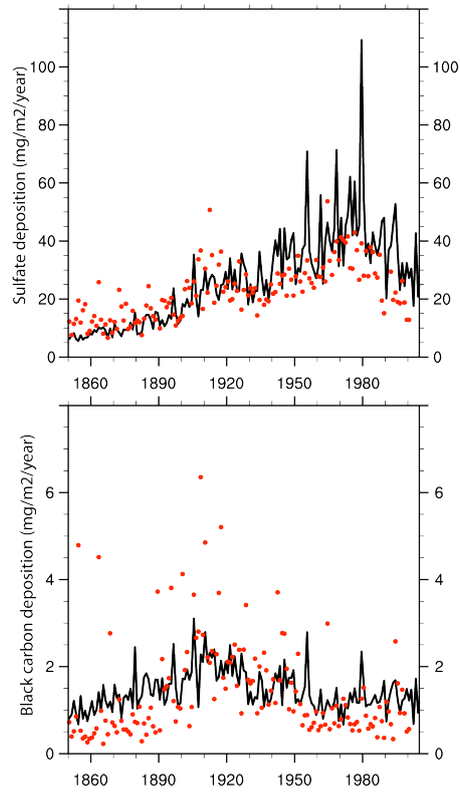
**Fig. 12.** Comparison between observed and modeled (present-day) annually aerosol optical depth at 500 nm. The observed values are based on annually averaged AERONET optical depths (Holben et al., 1998).

5017



**Fig. 13.** Comparison between observed and modeled (present-day) iron concentration at a variety of sites ( $\text{g}/\text{m}^2/\text{year}$ ). The compiled observations are available from Mahowald, et al. (2009).

5018



**Fig. 14.** Deposition (annual average) over Greenland (D4 site) of sulfate and black carbon. Ice-core observations are in red and CAM-chem results are in black.