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Ten years of CO emissions as seen from Measurements of Pollution in the Troposphere (MOPITT)

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1. Introduction

1 Both as an air pollutant and as a key player in atmospheric chemistry, carbon monoxide (CO) is an active research topic that requires accurate representation of the location and magnitude of CO surface emissions. At northern midlatitudes, CO is mainly emitted by incomplete fossil fuel combustion while biomass burning is its main source in tropics and in the Southern Hemisphere. Given the large spatiotemporal variability of CO emissions, simplifications have to be introduced in order to inventory them for large regions: the inventories are built using national inventory statistics and emission factors per activity type, and they suffer from large uncertainties [Streets et al., 2006; Klimont and Streets, 2007].

1 In this context, attempts have been made to deduce CO emissions from CO atmospheric concentrations by statistical inversion methods. The first inversion studies relied on the concentration measurements from the NOAA/Earth System Research Laboratory (ESRL) surface network [Bergamaschi et al., 2000; Pétron et al., 2002; Kasibhatla et al., 2002]. The operation of the Measurements of Pollution in the Troposphere (MOPITT) instrument on board NASA’s Terra satellite since March 2000 triggered similar studies based on remote sensing data at a global scale [Arellano et al., 2004, 2006; Pétron et al., 2004; Yurganov et al., 2010], at a regional scale [Allen et al., 2004; Headl et al., 2004; Turquety et al., 2008; Tanimoto et al., 2008] or for specific events [Pfister et al., 2005; Turquety et al., 2007]. They were based on the CO retrievals MOPITT Version 3 release and they were restricted to short periods such as 2000–2001 and operated at a subcontinental resolution. The introduction of a variational approach for this kind of study [Stavracou and Müller, 2006; Henze et al., 2007; Chevallier et al., 2009; Kopacz et al., 2009, 2010] allows more flexibility in the inversion: longer periods can be processed at finer resolution with a larger number of interactive chemical species.

4 The retrieval algorithms that generate the CO profiles from the MOPITT radiance observations have also dramatically improved recently, with enhancements of the general formulation of the retrieval and of its individual components, like the radiative transfer model [Deeter et al., 2010]. The latest version (Version 4) of the retrievals has been released in June 2009.

5 This paper takes advantage of these recent developments and presents the first estimation of the global CO emissions from MOPITT Version 4 data for the decade 2000–2009. It extends the preliminary study of Chevallier et al. [2009] that focused on the African continent for years 2000–2006. Our approach is based on a simplified chemical transport model (CTM), LMDz-SACS, embedded
in a variational inversion system. This global CTM explicitly represents the evolution of five interacting chemical tracers in the atmosphere: CO, CH$_4$, HCHO, H$_2$ and methyl chloroform (MCF). OH concentrations link them all together and are constrained by MCF measurements simultaneously. Chemistry of hydrocarbons is also taken into account in this system via a priori computation of the important terms (e.g., secondary CO or HCHO production) using a CTM with more complete chemistry.

The components of the inversion system are described in section 2. Section 3 gives the results of the inversion and explores their main features. The inverted emissions are evaluated by comparison of the modeled concentrations with independent measurements at the surface (NOAA/ESRL, AGAGE, CSIRO, EMPA, SAWs, NIWA and JMA/MRI), from aircrafts (YAK-AEROSIB) and from satellite (Infrared Atmospheric Sounding Instrument, IASI). Section 6 focuses on the advantages and limitations of our method.

2. Methodology

Our inversion method [Chevallier et al., 2009] aims at adjusting the emissions, jointly called $x$, in such a way that they become consistent both with the atmospheric observations $y$ and with some prior state $x_b$, given their respective uncertainties represented by their error covariance matrix $R$ and $B$. The uncertainties are assumed to be Gaussian-distributed. The solution, which will be called “posterior” in the following, is found by the iterative minimization of a Bayesian cost function $J$ defined as

$$J(x) = (x - x_b)^T B^{-1} (x - x_b) + (H(x) - y)^T R^{-1} (H(x) - y).$$

The vector $x$ contains (1) the global CO, CH$_4$, H$_2$ and methylchloroform MCF emissions at an 8 day and at a 3.75° × 2.75° (longitude, latitude) resolution together with their 3D initial conditions, (2) the factors, at the same resolution, to scale at once the surface emissions and the 3-D chemical production (by VOCs) of HCHO, and (3) the scaling factors for the 4-D OH atmospheric concentrations for four latitude bands (90°S–30°S, 30°S–0°, 0°–30°N, 30°N–90°N). The constraints in $y$ are the observations of concentrations of CO and MCF. In this study, CH$_4$, H$_2$ and HCHO are not constrained by dedicated observations. The results of the inversion on these species are very close to the priors. Since CH$_4$, H$_2$ and HCHO are not the species of interest in this study, the results for these species are not shown here.

For technical reasons, the 10 year period considered here is processed in consecutive 13 month chunks with a 1 month overlap from one chunk to the next. About 25 iterations are needed to reduce the norm of the gradient of $J$ by 95% with the M1QN3 limited-memory quasi-Newton minimization algorithm that we use [Gilbert and Lemaréchal, 1989].

2.1. The Chemistry Transport Model

The global 3-D CTM of the atmosphere LMDZ-SACS links the variables to the inferred $x$ and the observed variables $y$. It is derived from the general circulation model LMDZ developed at Laboratoire de Météorologie Dynamique [Hourdin et al., 2006] and of the Simplified Atmospheric Chemistry System (SACS) that has been specifically designed for this inversion system [Pison et al., 2009]. The simplifications of SACS refer to the INCA scheme of Hauglustaine et al. [2004] and Folberth et al. [2006]. As seen in Figure 1, SACS aims to represent the oxidation chain of methane by solving the chemical interaction between a limited set of four species: CO, CH$_4$, HCHO and OH. It is assumed that only the production and the loss of CH$_4$, HCHO, CO and H$_2$ are key for the problematic we
investigate: all other intermediate reactions are considered as instantaneous.

[11] The tangent linear and adjoint codes of LMDz-SACS have been developed by Chevallier et al. [2005] and by Pison et al. [2009].

2.2. Prior Emissions

[12] The prior information $x_b$ provided to the inversion system combines two data sets: version 2 of the Global Fire and Emission database GFED-v2 [Van der Werf et al., 2006] for biomass burning, valid until 2008, and version 3.2 of the Emission Database for Global Atmospheric Research (EDGAR3.2) inventory, valid for 1995, for the anthropogenic emissions (industry, fossil fuel, and industrial biofuel combustion) [Olivier and Berdowski, 2001]. No effort is made here to adapt the 1995 EDGAR3 inventory to the 00’s or the GFED-v2 data to year 2009 in the prior as the inversion can adjust each pixel of the surface, each week. Emissions and chemistry of other species like the biogenic ones or nonmethane volatile organics compounds (NMVOC) are those described by Folberth et al. [2006] for LMDZ-INCA; the anthropogenic NMVOC and NOx emissions are estimated as given by the EDGAR v3.2 database, the biogenic NMVOC emissions were taken from the Global Emissions Inventory Activity (GEIA) database [Guenther et al., 1995].

[13] Considering the large uncertainties that still affect such emission inventories, the error standard deviations assigned to the CO prior emissions in the covariance matrix $B$ are arbitrarily set at 100% of the maximum value of the emission time series during the corresponding year for each grid point. This choice of a relatively large value has been made to relax the constraint on the prior emissions. This was found especially important for highly seasonal emissions such as fires [Chevallier et al., 2009]. Errors are set at 100% of the flux for CH$_4$ and H$_2$, and for the HCHO scaling factors. For MCF emissions, the inventory by Montzka et al. [2000] is rescaled according to an update of the study by Bousquet et al. [2005]. As its emissions are supposed to be well known and in order to constrain OH, errors are set at only 1% of the flux for MCF. The errors assigned to the scale factor of OH are set at 10%, which is consistent with the differences between estimates of the OH concentrations of several studies [Prinn et al., 2001; Krol and Lelieveld, 2003; Bousquet et al., 2005]. For the initial conditions, errors are set at 10% for HCHO and MCF and at only 3% for CH$_4$ and 5% for CO.

[14] Following Chevallier et al. [2007], spatial correlations are defined by an e-folding length of 500 km over land (1000 km over sea), without any correlation between land and ocean grid points. Indeed, the recent study by Carouge et al. [2010] shows that the spatial correlations based on more physical properties did not prove to perform better in estimating surface fluxes by inversion than the e-folding decay approach. We neglect temporal correlations, as the state vector is already aggregated on 8 day basis.

2.3. Methyl Chloroform Observations

[15] CO and OH concentrations influence each other to a large extent. The realism of the OH fields is therefore important for the inverse modeling for CO emissions. In the absence of sufficient direct observations of OH, indirect information is usually exploited, like that from MCF measurements, since MCF has a simple chemistry with OH and well-known sources [Prinn et al., 2001; Krol and Lelieveld, 2003; Bousquet et al., 2005]. For the present study, a set of stations that measured MCF nearly continuously for the 2000–2008 period has been selected from the AGAGE and NOAA/ESRL networks available on the World Data Centre for Greenhouse Gases (WDCGG) site at http://gaw.kishou.go.jp/wdgg/. Note that no MCF data are available at the time of writing for year 2009. In this case, errors of the OH concentrations are set to 0.1%.

2.4. MOPITT V4 Observations

[16] The observational constraint on CO emissions comes from Version 4 of the Level 2 product of the MOPITT retrievals (http://www.acd.ucar.edu/mopitt/). The MOPITT instrument has been operated nearly continuously for cloud free spots since March 2000, except during the period from May 2001 until August 2001 when the instrument experienced a cooler failure. The data before and after this failure are different (four channels were lost) with a change in the bias. These two periods have been termed Phase I (from March 2000 to May 2001) and Phase II (August 2001—…). An anomaly also lead to a 2 month lack of data in summer 2009 (from the end of July to October). To date, the MOPITT record is the longest archive of tropospheric CO observations from space.

2.4.1. Data Processing

[17] We use the 700 hPa–level CO retrievals that have one of the highest sensitivity to the measured radiances in the troposphere [Deeter et al., 2003] together with their associated averaging kernels (AK). The data selection follows the criteria of the MOPITT data quality statement (http://www.acd.ucar.edu/mopitt/products.shtml; access: February 2010): data within 25° from the poles have been left out, as the weight of the a priori CO profile in the MOPITT retrievals increases toward the pole. Nighttime data may be biased and are not exploited [Crawford et al., 2004]. We then averaged the retrievals at the 3.75° × 2.5° resolution of LMDz-SACS at the orbit level, in order to reduce the effect of correlated errors between neighboring observations in the inversion system. As the AK profiles do not vary much within the grid cell, we chose to use the AK profile of the first retrieval when several of them are averaged into a super-retrieval. After this screening, about one million “super-observations” remain in the 13 month inversions.

[18] Error correlations between super-observations can be neglected at the 3.75° × 2.5° resolution used for the inversion [Heald et al., 2004], so that the covariance matrix $R$ of the observation errors is defined as diagonal. Only variances are taken into account. The observation error is the quadratic sum of the measurement error reported in the MOPITT data sets, and of the CTM errors set to 50% of the retrieval values according to Pison et al. [2009].

2.4.2. Differences Between Version 3 and Version 4

[19] Version 4 has been released in June 2009. As shown by Deeter et al. [2010], it benefits from significant advances in radiative transfer modeling, state vector representation, and a priori statistics compared to Version 3 (V3). The retrieval state vector is represented as the logarithm of the volume mixing ratios. In contrast to the V3 products and their seven standard pressure levels, V4 represents the CO
profile on a 10-pressure-level grid: a floating surface plus nine uniformly spaced levels from 900 to 100 hPa for a better representation of the vertical structure of both the CO a priori profiles and the MOPITT weighting functions. The a priori profile varies for each retrieval, which was not the case for V3.

The observed MOPITT V3 (Figure 2a) and MOPITT V4 (Figure 2b) CO concentrations at 700 hPa are displayed in Figure 2 for June 2000. Similar features are seen between the both versions with plumes over Africa and East Asia. However, we can see that the MOPITT V4 concentrations are often smaller than the MOPITT V3 ones. MOPITT V4 has been validated using in situ CO profiles, similarly to previous MOPITT V3 validation studies [Emmons et al., 2004, 2007, 2009]. The retrieval performance has been improved particularly in regions of very high and very low CO concentrations, like over the oceans, where a significant negative bias in clean environments such as the South Pacific Ocean [Emmons et al., 2004] affected the MOPITT V3 retrievals in phase II. As a consequence, the mean bias between our prior modeled concentrations and the MOPITT V4 observations is reduced by about 80% compared to V3. However, the MOPITT retrievals suffer from a time-varying bias in Version 4 [Deeter et al., 2010], as in Version 3 [Yurganov et al., 2008; Emmons et al., 2009; Drummond et al., 2009]. This positive bias drifts by about 0.5 ppbv/yr on average at 700 hPa, in Version 4 [Deeter et al., 2010].

Figure 2. Mean CO concentrations at 700 hPa, in parts per million, for June 2000 with (a) MOPITT Version 3 and (b) MOPITT Version 4.

Figure 3. Mean CO concentrations, in parts per million, for June 2000 in (a) the MOPITT retrievals for the 700 hPa pressure level, (b) the prior concentrations, and (c) the posterior concentrations. The simulations all take the individual MOPITT averaging kernels corresponding to Figure 3a into account. Abscissa and ordinate report the longitudes and latitudes.
should be noted that the positive bias drift is not taken into account in our observation error and may bias the inversion estimate. However, as shown later in section 5.3, the consistency of the MOPITT-based inverted fluxes with the IASI-based ones shows that the impact of the drift remains small.

3. Concentrations

3.1. Spatial Distribution

The month of June 2000 is used as an illustration of the inversion system behavior. The MOPITT retrievals at 700 hPa (Figure 3a), the prior concentrations (Figure 3b) and the posterior CO concentrations (Figure 3c) are displayed in Figure 3. As expected, large discrepancies can be seen between the MOPITT measurements and the prior simulation. In general, the prior underestimates the concentrations. The prior concentrations agree with the MOPITT data rather well in the Southern Hemisphere. In the Northern Hemisphere, an underestimation by about 50% is seen over Canada, South East Asia and over the North Pacific temperate ocean. On the contrary, slight overestimations are observed over the central Africa and South American Temperate regions (60%). Overall, the optimization significantly improves the fit of the simulations to the observations by increasing the emissions.

3.2. Seasonal Cycle

As the scale of interest is regional, prior and posterior monthly averaged CO mixing ratios are compared to the MOPITT measurements, for the 14 continental regions shown in Figure 4, in Figure 5. In the Northern Hemisphere, largest concentrations are seen around March and April whereas the peak of the concentrations occurs in September and October in the southern one. CO posterior concentrations for the 10 year period present consistent seasonal cycles in all regions compared to the corresponding prior, even if the amplitude of these cycles can be different. For example, the prior modeled concentrations are lower than the posterior ones by around 15 ppb over Europe and USA and over the boreal regions in 2008 and 2009. By construction, the optimization brings the model values much closer to the MOPITT measurements. However, as illustrated in Figure 5j with the Indonesian emission peaks in 2002, 2004 and 2006, the model does not exactly fit the observations since the posterior is pulled both toward the observations and toward the prior.

4. Emissions

4.1. Annual and Regional Budget of CO Emissions

This section focuses on the results in terms of inverted emissions from March 2000 to December 2009. The results are summarized per year and per region in Table 1, in Table 2, and in Figure 6.

As expected [Carmichael et al., 2003; Palmer et al., 2003; Pétron et al., 2004; Arellano et al., 2004; Allen et al., 2004; Heald et al., 2004; Wang et al., 2004; Streets et al., 2006], the inversion highlights the importance of South East Asia as a CO source with an average of 280 TgCO/yr, followed by South American Temperate (154 Tg/yr), Northern Africa (143 Tg/yr), Southern Africa (140 Tg/yr), USA (136 Tg/yr) and Europe (130 Tg/yr).

Most of the MOPITT-based inversions published so far have focused on Phase I. Our inverted total CO surface emissions estimates for this period are presented in Table 1. Our posterior budget of 1386 TgCO/yr is 3% higher than the optimized magnitudes obtained by Arellano et al. [2006] with 1342 TgCO/yr (from April 2000 to April 2001). We also compare our estimate with that of Kopacz et al. [2009], who obtained a global budget of 1148 TgCO, which is 16% higher than our estimate. However, our posterior budget of 1276 TgCO/yr (from May 2000 to April 2001) is lower than estimated by Stavrakou and Müller [2006] with 1333 TgCO/yr.

For the months April 2000 to March 2001, our posterior total emissions (1271 TgCO/yr) are also higher than the 1091 TgCO/yr (+16%) of Pétron et al. [2004]. Their
Figure 5. Times series of CO concentrations observed by MOPITT (green) and simulated by the prior (blue) and the posterior (red) concentrations from March 2000 to December 2009. Units are ppm.
optimized value of 207 TgCO/yr for South East Asia is lower than our estimate of 260 TgCO/yr. There are large discrepancies for regions such as Eurasian Boreal, where we found higher emissions (+348% with 93 TgCO), or Australia, where we found lower emissions (−22% with 74 TgCO). For the south Asian region, Pétron et al. [2004] obtain a 121 TgCO annual total, an increase of 21% relative to their prior emissions. In contrast, Heald et al. [2004] estimate the Indian emissions (corresponding to our South Asian region) to be 46 TgCO/yr, that is 50% lower than their prior of 90 TgCO and equivalent to our posterior estimate of 47 TgCO.

Kopacz et al. [2010] recently inverted CO emissions from May 2004 to April 2005, combining CO column measurements from the MOPITT, from the Atmospheric Infrared Sounder (AIRS) and from the SCanning Imaging Absorption SpectroMeter for Atmospheric CartographHY (SCIAMACHY) satellite. Their total surface emissions constitute 1350 TgCO, 9% lower than our estimate of 1474 TgCO for the same period. Kopacz et al. [2010] estimated the annual emissions in Alaska and Canada to be about 24 TgCO, 20% lower than the summer emissions estimate (from June to August) of Pfister et al. [2005] and Turquety et al. [2007] and 70% lower than our posterior value of 41 TgCO. The main difference between our two studies is seen for the USA; our value of 127 TgCO disagrees with their posterior emissions of 46.5 TgCO (~164%) and with the study of Hudman et al. [2008]. Hudman et al. [2008] estimated CO emissions over USA to be about 40 Tg and also mentioned that U.S. road emissions are overestimated in the anthropogenic emission inventory from the U.S. Environmental Protection Agency (EPA). The cause of the large difference is unclear. On the one hand, many uncertainties remain in the inventories (e.g., vehicle cold starts [Parrish, 2006]). On the other hand, the transport of the high CO concentrations from the South American Tropical region over USA, may contribute to a misattribution of emissions to the USA in our inversion.

Table 1. Total CO Emissions in Tg/yr for Years 2000/2001, 2002, 2003, and 2004

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Prior denotes before inversion, and Posterior denotes after inversion. The total estimates include the oceanic CO source amounting to 20 TgCO/yr. All budgets correspond to a 12 month period.

For the south Asian region, Pétron et al. [2004] obtain a 121 TgCO annual total, an increase of 21% relative to their prior emissions. In contrast, Heald et al. [2004] estimate the Indian emissions (corresponding to our South Asian region) to be 46 TgCO/yr, that is 50% lower than their prior of 90 TgCO and equivalent to our posterior estimate of 47 TgCO.

[27] Kopacz et al. [2010] recently inverted CO emissions from May 2004 to April 2005, combining CO column measurements from the MOPITT, from the Atmospheric Infrared Sounder (AIRS) and from the SCanning Imaging Absorption SpectroMeter for Atmospheric CartographHY (SCIAMACHY) satellite. Their total surface emissions constitute 1350 TgCO, 9% lower than our estimate of 1474 TgCO for the same period. Kopacz et al. [2010] estimated the annual emissions in Alaska and Canada to be about 24 TgCO, 20% lower than the summer emissions estimate (from June to August) of Pfister et al. [2005] and Turquety et al. [2007] and 70% lower than our posterior value of 41 TgCO. The main difference between our two studies is seen for the USA; our value of 127 TgCO disagrees with their posterior emissions of 46.5 TgCO (~164%) and with the study of Hudman et al. [2008]. Hudman et al. [2008] estimated CO emissions over USA to be about 40 Tg and also mentioned that U.S. road emissions are overestimated in the anthropogenic emission inventory from the U.S. Environmental Protection Agency (EPA). The cause of the large difference is unclear. On the one hand, many uncertainties remain in the inventories (e.g., vehicle cold starts [Parrish, 2006]). On the other hand, the transport of the high CO concentrations from the South American Tropical region over USA, may contribute to a misattribution of emissions to the USA in our inversion.

[28] One should note that significant differences are observed in any comparison between different global and regional MOPITT-based inverted results, due to the diversity of the inversion systems (such as the space-time resolution of the state vector x, the CTM used, whether or not the OH fields are prescribed in the model, the processing of


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<td>969</td>
<td>1328</td>
<td>969</td>
<td>1318</td>
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</table>
the MOPITT V3 or V4 observations). As an example of the CTM diversity, an intercomparison of 26 CTMs showed a wide range of simulated CO distribution, even though the models all used the same emissions for fuel combustion, industry, and biomass burning [Shindell et al., 2006].

4.2. Error Estimates

[29] In principle, the uncertainty of the posterior emissions can be estimated theoretically using the Monte Carlo approach of Chevallier et al. [2007]. However, a simpler estimation method was preferred here because of the large computational expense of the Monte Carlo approach, in particular when the target period for the CO budget extends to a whole year. As a trade off between computing resources and exhaustivity, we simply estimate the inversions one-sigma uncertainty from the spread of the regional emissions in five sensitivity tests (cases A to E).

[30] 1. In case A, the OH field is replaced by OH-v2 field, different within 5% from the OH reference. It comes from a simulation of the full chemistry model LMDz-INCA, using another realistic emission scenario (the combination of anthropogenic emissions from IIASA, QUANTIFY for ship and GFEDv2 for biomass burning).

[31] 2. In case B, OH prior concentrations errors set at 35%.

[32] 3. In case C, HCHO prior sources increased by 50%.

[33] 4. In case D, CO prior emissions error set at 400%.

[34] 5. In case E, error correlations length set at 1000 km.

[35] The total CO annual emissions for the whole year 2004, found from each test, are given for the 14 regions in Figure 7. These posterior regional results are compared with the reference inversion described in section 2. If the differences between the posterior regional emissions budget as

Figure 6. Total posterior (inverted from MOPITT) CO emissions per year for the 14 regions, in TgCO/yr.

Figure 7. Annual total posterior CO emissions per region obtained by five sensitivity tests for year 2004 compared to the posterior reference. Case A: OH-v2 field; Case B: error on the OH concentrations set to 35%; Case C: the HCHO prior sources increased by 50%; Case D: error on the prior CO emissions set to 400%; Case E: error correlations length set to 1000 km. The prior information combines two data sets: EDGAR 3.2 for anthropogenic emissions and GFEDv2 for biomass burning.
Table 3. Regional and Global Error Estimates, as Estimated From Sensitivity Tests for 2004, in Tg/yr

<table>
<thead>
<tr>
<th>Region</th>
<th>Error Estimates</th>
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<tbody>
<tr>
<td>North Am Boreal</td>
<td>26</td>
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<tr>
<td>USA</td>
<td>26</td>
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<tr>
<td>South Am Trop</td>
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<td>South Am Temp</td>
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<td>Northern Africa</td>
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<tr>
<td>Southern Africa</td>
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<tr>
<td>Western Europe</td>
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<td>Eastern Europe</td>
<td>13</td>
</tr>
<tr>
<td>Eurasian Boreal</td>
<td>22</td>
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<tr>
<td>Middle East</td>
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</tr>
<tr>
<td>South Asia</td>
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<td>South East Asia</td>
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<td>Australia</td>
<td>21</td>
</tr>
<tr>
<td>Globe</td>
<td>166</td>
</tr>
</tbody>
</table>

estimated from this sensitivity study are weak, we assumed that the uncertainty associated with these emissions is small, and conversely.

[36] In case A, we investigate the impact of the prior OH field on the inversion; the inversion experiment is carried out with the OH–v2 field of OH. The impact is small, with differences in yearly emitted masses of a few percent only. In case B, when the OH prior uncertainties were set to 35%, some changes in the estimated regional budgets are observed, in particular over Australia (−20%) and in the Northern Hemisphere (−30% for North American Boreal, −22% for Eurasian Boreal, −20% for Europe and −17% for USA). By relaxing constraints on OH, the MCF observations are able to reduce OH concentrations in the 30°N–90°N latitudinal band, which is counterbalanced with a reduction of the CO emissions in order to keep the fit to the MOPITT retrievals. In case C, the impact of the prior HCHO field is tested by increasing the prior atmospheric source of CO by 50%. The posterior emissions are smaller than in the reference inversion for all regions (−8% for USA, −11% for South American Temperate). When the error on the prior CO emissions is increased to 400% (case D), the posterior yearly emissions are only slightly increased (e.g., for regions South East Asia (+3%) and USA (+2%)). In case E, the impact on the emissions is small (e.g., +2% in USA and −2% in Western Europe).

[37] The difference between the regional inverted emissions in cases D and E relative to the reference inversion is small, indicating that the inversion is largely insensitive to the prior errors statistics. The difference is the largest for case C; the HCHO prior field has a strong influence on the posterior CO emissions, even though its impact is limited to less than 15% in the regions that emit more than 100 TgCO/yr.

[38] As shown in Table 3, the lowest posterior relative uncertainties are found for the regions: Southern Africa, Middle East, South Asia and South East Asia with uncertainties of 6%, 6%, 9% and 10%, respectively. The uncertainty in regions North American Boreal and South American Tropical reaches 40% and 37.5%, respectively.

4.3. Variations Between Prior and Posterior Emissions

[39] Explained by economic developments since 1995 (target year of EDGAR 3.2 inventory), global posterior emissions are higher compared to the prior, by about 13% for Phase I and 37% for Phase II (Tables 1 and 2). All regions contribute to this increase of emissions except three of them (South American Tropical, Indonesia and South Asia). The yearly mean emissions are significantly increased by the inversion over Australia (e.g., by up to 175% (+56 TgCO) in 2002 or 220% (+55 TgCO) in 2004), as also seen by Jones et al. [2009], and over boreal regions.

[40] Figure 8 shows the time series of the prior and posterior monthly CO emissions in each region. The seasonal variation of the posterior source estimates is close to the prior one in most regions, keeping the same interannual variability, even though the amplitude differs. This is the case for regions Indonesia, South East Asia, Middle East, North American Boreal, Eurasian Boreal, South American Temperate, South American Tropical and Northern Africa.

[41] There are, however, some interesting differences in seasonality between prior and posterior cycles. As described by Chevallier et al. [2009] and in agreement with Arellano et al. [2004], Bremer et al. [2004] and Stavroulakis and Müller [2006], one of these differences is seen in Southern Africa: instead of peaking in July–August like the prior, the posterior estimates peak 2 months later, in September–October.

[42] As in findings by Kopacz et al. [2010], our results present a larger-than-expected seasonal variation in CO emissions at northern midlatitudes (for regions South Asia, Middle East and USA). Over USA, the posterior emissions peak late in spring rather than in winter. These maximums present higher amplitude than the prior ones (+7 TgCO in average). This could be explained by the fact that CO emissions from domestic wood burning and from vehicle cold starts are not well treated in the a priori emissions inventories [Miller et al., 2008; Parrish, 2006; Kopacz et al., 2010]. As seen in Figure 8h, the south Asian prior sources are about 7 TgCO per month throughout the years, but the posterior sources range from 2 TgCO/month in summer to 10 TgCO/month in winter. However, it should be reminded that the EDGAR v3.2 data inventory was designed for the year 1995 and has been applied here for years 2000–2009. Since then, Asia has benefitted from an important economic growth and emissions have changed, particularly the contribution of the industry, transport and domestic sectors [Streets et al., 2006; Ohara et al., 2007] and its seasonal cycle may be more similar to that of industrial regions.

4.4. Interannual Variability

[43] The 10 year inversion for CO emissions allows for investigating interannual variability (IAV). A slight decrease in the annual Western European emissions is observed since 2003 whereas the emissions of USA and Eastern Europe seem to be constant. However, global CO emissions show significant interannual variability, in particular in tropical and subtropical regions where the interannual variation is mostly influenced by biomass burning [Szopa et al., 2007], and partly by precipitation rates [Van der Werf et al., 2008].

[44] Boreal regions (North American Boreal and Eurasian Boreal) and Indonesia display a notable IAV. For Indonesia, the annual budgets (Tables 1 and 2) show that the 2006 CO emissions are higher than in other years (96 TgCO/yr, compared with 21 to 49 TgCO/yr for 2005 and 2008–2009). This may be linked to the 2006 El Niño–Southern Oscillation
Figure 8. Times series of prior (blue) and posterior (red) CO emissions from March 2000 to December 2009. IASI posterior is in green, from July 2008 to December 2009. Unit is TgCO/month.
(ENSO) event, which was associated with drought and the greatest rise in wildfire activity in Indonesia since the record-breaking 1997–1998 El Niño [Logan et al., 2008; Chandra et al., 2009].

Global CO emissions range from 1504 TgCO in 2007, to 1318 TgCO in 2009 (Table 2). Indeed, global CO emissions decreased significantly in 2008 and 2009. These low emissions are considered anomalous by Yurganov et al. [2010], who hypothesized that economic recession could have impacted the CO emissions. From Table 2, it can be seen that only two regions are responsible for this diminution: South East Asia and South American Temperate (~103 TgCO and ~70 TgCO, respectively, between 2007 and 2008). For South East Asia, it should be noted that the 2007 biomass burning emissions (particularly the peak in March) were extremely high compared to the other years (e.g., 153, 150, 189, 152 TgCO in GFEDv2 for 2005, 2006, 2007 and 2008), whereas the biomass burning emissions in 2008 could be considered as normal.

Significant IAV in CO emissions was observed in the South American Temperate region since 2000 (137, 170, 184, 138, 198, 128, and 131 TgCO/yr for 2003, 2004, 2005, 2006, 2007, 2008 and 2009). This IAV is well correlated (correlation coefficient of 0.90, using 60 monthly values) with the fire counts of Torres et al. [2010] derived from Aqua-MODIS observations [Giglio et al., 2006], but not with the deforestation rates provided by the Brazilian government remote sensing agency INPE (Instituto Nacional de Pesquisas Espaciais, http://www.obt.inpe.br/prodes/prodes_1988_2009.htm, access: August 2010) (correlation coefficient of 0.44, based on 9 annual values). This apparent inconsistency can be explained by the fact that in this region, biomass burning is the result of the cropland expansion, and consequently to deforestation [Morton et al., 2006], but also to other practices, such as the disposal of agricultural waste.

After low value in 2006 related to unfavorable climate conditions [Gloudemans et al., 2009; Schroeder et al., 2009], there was a peak in CO emissions in 2007, correlated with the largest number of fires detected from space over the last 10 years [Torres et al., 2010]. In 2008, the decrease in South American emissions cannot be explained by climate conditions [Torres et al., 2010], but may be related to socioeconomic factors connected to the economic recession (with unfavorable market conditions for agricultural exports), such as the soybean price collapse in summer 2008.

Although the emissions of the two regions South East Asia and South American Temperate slightly increased from 2008 to 2009 (from 250 to 272 Tg and from 128 to 131 Tg, respectively), global CO emissions still decreased in 2009. Except over Europe, all the Northern Hemisphere regions have lower emissions in 2009 than in 2008. This could be due to lower fossil fuel emissions owing to the economic recession. More extensive work is needed to resolve this issue.

4.5. CO Atmospheric Production and Loss

Although surface emissions are dominant, CO chemical production and loss can play a role in long-term changes in the tropospheric chemical composition and they are optimized in the system. The chemical loss of CO corresponds to CO₂ production according to the reaction with hydroxyl radical OH: CO + OH $\rightarrow$ CO₂ + H. As mentioned in section 2, OH is constrained by the MCF measurements and optimized in the inversion. The yearly averages of the prior and posterior tropospheric OH concentrations are given in Table 4. For the past 10 years, the global OH posterior concentrations value is 8.62 × 10⁻³ mol cm⁻³ on average, 17% lower than the 10.5 × 10⁻³ mol cm⁻³ value of Prinn et al. [2005]. The posterior total masses of OH vary between ~8% (2003) and ~6% (2006) of the prior. These variations are within the expected range of OH interannual variability obtained by other inversion studies [Bousquet et al., 2005; Prinn et al., 2005].

Figure 9 shows five examples of the temporal variations of the chemical production and of the chemical loss of CO over the sea (tropical Pacific Ocean and Indian tropical ocean) and over the continent (Northern Africa, USA, Europe). Prior and posterior CO productions are nearly similar, due to the weak impact of the inversion on HCHO concentrations (not shown). Note that the HCHO prior concentrations used here are smaller than the SCIAMACHY retrievals shown by De Smedt et al. [2008, Figures 13 and 14] and present less variability. The production of CO₂ is large over tropical regions and particularly over tropical oceans which represent about 30% of the global tropospheric CO₂ production. The global CO₂ production from 2000 to 2009 ranges from 1120 TgC/yr to 1072 TgC/yr, with a global average of 1118 TgC/yr, 2% smaller than the Folberth et al. [2005] value of 1137 TgC/yr.

5. Evaluation With Independent Measurements

Our evaluation strategy consists of comparing our posterior-modeled concentrations with several sets of independent measurements at the surface, from aircraft, from other inventories and by satellite.

5.1. Comparison With Surface and Aircraft Measurements

The posterior emissions are evaluated by comparison of the prior and of the posterior modeled CO concentrations with independent and fixed surface measurements from various networks (NOAA/ESRL, AGAGE, CSIRO, EMPA, SAWS, NIWA and JMA/MRI) available on the WDCGG Web site. We have restricted our analysis to 56 sites that present uninterrupted measurements for the 2000–2008 period.
period and, at least, one measure per month. The site locations and results are shown in Figure 10. Stations highlighted in green, for which the ratio of the posterior to the prior bias or root mean square (RMS) is lower than 1, indicate an improvement of the corresponding statistical indicator after optimization. Two different periods are studied here: MOPITT Phase I (March 2000 to May 2001) and Phase II (August 2001 – …).

During Phase I, the absolute biases are reduced by 2% to 154% at 38 stations. The inversion leads to a dramatic improvement relative to the prior simulation for all the high-latitudes sites of the Southern Hemisphere (TDF, CRZ, PSA, HBA, SYO, MAA, MQA, CYA, ARH) and for 76% of the stations in the Northern Hemisphere. This percentage then reaches 84% for Phase II (average ratio for the bias of 0.78), with best results at European stations (PAY, HUN, BSC).

However, in Phase II, the inversion degrades the fit for the NOAA POC boats in the South Pacific Ocean. As shown in Figures 10b and 10d, the ratios of bias and RMS of this station are higher than 1 below the equator. The fit is also degraded at the high-latitude Southern Hemisphere sites, with an average ratio for the bias of 5.22. This feature was already pointed by Arellano et al. [2004] and by Stavrakou and Müller [2006].

We use the airborne campaigns YAK-AEROSIB carried out in April 2006, August 2007 and July 2008 across Siberia by Paris et al. [2008, 2009, 2010] as an additional independent set of measurements for this region not covered

Figure 9. Prior and posterior production and loss of CO from March 2000 to December 2009. The loss of CO corresponds to the production of CO₂. Note that the scale is different between continental regions (USA, Europe, and Northern Africa) and oceanic regions.
by the various networks. The inversion improves the fit at the highest latitudes in Siberia with a reduction by 23% of the absolute bias but degrades it about latitude 50°N (Figures 10b and 10d). As for the fixed station UTA, along the western coast of USA, note that there is little or no MOPITT data in this region (as seen in Figure 2), so that it is only constrained indirectly by data over neighboring regions.

5.2. Comparison With Other Inventories

Large discrepancies exist between different inventories in terms of budgets [Butler et al., 2008] and in terms of seasonal variability, explained by the respective methodologies used in the inventory construction and by the emission factors applied. We compare our posterior western European, American and Asian emissions with several global and regional inventories that correspond to year 2000: (1) the one proposed for the IPCC fifth assessment report AR5 [Lamarque et al., 2010]; (2) the REanalysis of the TROpospheric chemical composition (RETRO) inventory [Pulles et al., 2007], developed by The Netherlands Organisation for Applied Research (TNO) (3) the Cooperative Programme for Monitoring and Evaluation of the Long-range Transmission of Air Pollutants (EMEP) inventory for Europe; (4) the U.S. Environmental Protection Agency (EPA) inventory for USA; and (5) the inventory of Streets et al. [2006] for Asia.

As seen in Figure 11a, these inventories show very similar seasonal cycles for Western Europe and our posterior is in good agreement with IPCC AR5 and EMEP. However, as described in the section 4.3, our posterior cycle shows a strong seasonal variation with large intensities in April for USA and the region South East Asia, which is not seen in the others inventories.

We have seen that south Asian emissions are still misunderstood and several studies contradict each other about total budgets. It is also the case for the seasonal cycle (Figure 11c). Our posterior emissions for this region show...
stronger seasonality whereas the IPCC AR5, EDGAR 3.2 and Streets et al. [2006] inventories do not have any seasonality, the emissions are constant among the year but with different levels: 8 TgCO, 6 TgCO and 5 TgCO, respectively. RETRO presents a strongly different cycle with maximum in winter and minimum in July. For South East Asia, all the inventories are similar, except in spring: our posterior and RETRO peak in April instead of March for the others inventories.

5.3. Comparison With an IASI-Based Inversion

[59] The IASI instrument has been flying as part of the METOP-A platform since October 2006. It provides CO partial columns information [Clerbaux et al., 2009]. Products have been made available by the Laboratoire Atmosphères, Milieux, Observations Spatiales (LATMOS) through the Ether database (http://ether.ipsl.jussieu.fr), for the period starting in June 2008. The retrieval code based on the Optimal Estimation was developed at the Université Libre de Bruxelles [Turquety et al., 2009] and CO data were evaluated against other satellite observations [George et al., 2009].

[60] In a previous study [Fortems-Cheiney et al., 2009], we have obtained similar global CO emissions budgets from IASI and from MOPITT V3, 643 TgCO and 649 TgCO, respectively, from July to November 2008. Consistent with this results, the mass budgets for a 1 year period (from July 2008 to June 2009) are also in good agreement. The global IASI posterior emissions of 1450 TgCO are 10% higher than the MOPITT ones (1304 TgCO). It is mostly due to a 331% (+63 TgCO) and 53% (+80 TgCO) increase of the annual fluxes over regions Indonesia and Northern Africa, respectively, with the IASI inversion, which posterior values of 82 TgCO and 230 TgCO are substantially higher than the MOPITT-based estimate. On the contrary, we notice that the IASI-based emissions in the Southern Hemisphere are lower than the MOPITT ones, in particular in Australia (−43%). For others regions like Western Europe, USA and South East Asia, both the IASI and MOPITT inversions lead to a similar budget (with 70 and 68 TgCO, 110 and 96 TgCO, and 285 and 292 TgCO, respectively).

[61] We compare the impact of the satellite observations on seasonal and regional cycles from July 2008 to December 2009 in Figure 8. The IASI-based posterior emissions also show a strong seasonality in the Northern Hemisphere. Differences are in some cases noticeable like over Western Europe, Middle East and Australia. The IASI posterior emissions also present higher amplitudes over South Asia (Figure 8h) and Indonesia (Figure 8j) than the MOPITT
ones. However, for the others regions, the MOPITT and IASI posteriors present similar cycles (Northern Africa, North American Boreal, South American Tropical, Eastern Europe, South American Temperate, Southern Africa, Eurasian Boreal and South East Asia). This result is all the more encouraging given the known drift of the MOPITT retrievals discussed by Deeter et al. [2010].

6. Conclusions

[62] The inverse modeling approach offers an attractive solution to the problem of estimating the gaseous emissions, that allows one to correct the prior inventories. For the first time, this study aims at estimating the CO emissions for the past 10 years, using a multispecies inversion system and the MOPITT Version 4 retrievals. The “COLSCE-V1” global gridded CO fluxes, at an 8 day and 3.75° × 2.75° resolution, will be available from the CarboScope Web site: http://www.carboScope.eu. Our study indicates that the absolute accuracy of the inverted CO emissions at yearly subcontinental scale should be better than 15% in regions that emit more than 100 TgCO/yr.

[63] Significant modifications of the prior emissions are seen in our study. First, our posterior emissions differ from the prior ones in terms of budget. The global estimate is higher by about 13% compared to the prior for Phase I (March 2000 to May 2001) of the MOPITT acquisition. For Phase II (August 2001–…), this increase reaches about 37%, explained by the fact that the magnitude of the anthropogenic CO emissions have changed since 1995, target year of EDGAR v3.2, which is our prior.

[64] Second, the CO emissions also have some significant interannual variability due to biomass burning emissions, climate, and socioeconomic factors, which requires to be considered for CO modeling. The highest annual budget of the last decade is seen for year 2007 with 1504 TgCO/yr, because of fires over South East Asia and South American Temperate. Then, the CO emissions show a dramatic decrease (1318 TgCO/yr in 2009), that could be connected to the economic recession.

[65] The uncertainty on these emissions ranges from 6% (over regions Southern Africa and Middle East) to 40% over the region North American Boreal. In the case of high latitudes, the inverted emissions improve the fit to the northern sites (i.e., BRW, ALT and CBA with absolute bias reduced by 69%, 69% and 54%, respectively) in spite of their relatively large uncertainties. Indeed, a comparison of our posterior emissions to the independent surface derived CO concentrations shows improved agreement in the Southern Hemisphere for 76% of the stations in the Northern Hemisphere for Phase I. This percentage then reaches 84% for Phase II. However, in Phase II, the inversion fails to improve both the fit to the YAK-AEROSIB measurements about latitude 50°N in Siberia, and the fit to the stations located in the high latitudes of the Southern Hemisphere (with an average ratio for the bias of 2.61) and in Australia (stations CFA, CGO).

[66] Third, as found by Kopacz et al. [2010], some significant seasonal variations of the emissions in the midlatitudes of the Northern Hemisphere are found with both the MOPITT-based and the IASI-based inversions, whereas most anthropogenic inventories do not include any one.

[67] The consistency of the MOPITT-based inverted fluxes with the IASI-based ones shows that the impact of the drift in the MOPITT retrievals does not exclude the possibility of a multidecadal CO monitoring (with the METOP series: METOP-B in 2012, METOP-C in 2016). Nevertheless, an alternative to MCF as a constraint for the OH fields has to be found in the near future, as MCF concentrations are declining rapidly (current values are of only a few parts per trillion) to the extent that MCF soon cannot be used to inform about OH anymore. An alternative to MCF could be the use of 13CO [Krol et al., 2008]. Moreover, together with OH, formaldehyde controls the atmospheric production of CO. Ideally, its concentrations should also be constrained in the inversion. Further, our sensitivity tests highlight HCHO prior fields as the less reliable component in our inversion system. Some improvement is therefore expected by integrating HCHO retrievals from SCIAMACHY or from the Ozone Monitoring Instrument (OMI) [Levelt et al., 2006] in the system.

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