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Top-Down CO Emissions Based On IASI Observations and Hemispheric Constraints on OH Levels

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Abstract Assessments of carbon monoxide emissions through inverse modeling are dependent on the modeled abundance of the hydroxyl radical (OH) which controls both the primary sink of CO and its photochemical source through hydrocarbon oxidation. However, most chemistry transport models (CTMs) fall short of reproducing constraints on hemispherically averaged OH levels derived from methylchlorororm (MCF) observations. Here we construct five different OH fields compatible with MCF-based analyses, and we prescribe those fields in a global CTM to infer CO fluxes based on Infrared Atmospheric Sounding Interferometer (IASI) CO columns. Each OH field leads to a different set of optimized emissions. Comparisons with independent data (surface, ground-based remotely sensed, aircraft) indicate that the inversion adopting the lowest average OH level in the Northern Hemisphere (7.8 × 10^5 molec cm^{-3}, ∼18% lower than the best estimate based on MCF measurements) provides the best overall agreement with all tested observation data sets.

Plain Language Summary Satellite measurements of a pollutant can be used to deduce the emissions of this pollutant to the atmosphere. But often, such estimates have errors due to our uncertain knowledge of the chemical lifetime of pollutants. Here we assess the importance of specifying the correct lifetime for deducing the sources of carbon monoxide. We also show that the measurements can provide new constraints on the lifetime of pollutants. More precisely, we provide constraints on the abundance of OH radicals, often considered to be the main detergent of the atmosphere. Indeed, OH radical is the main oxidant of carbon monoxide, but also methane, which is a potent greenhouse gas.

1. Introduction

The sources of CO include direct emissions and photochemical production due to the oxidation of hydrocarbons. The dominant CO sink being its reaction with the hydroxyl radical (OH), CO has a strong influence on the oxidative capacity of the atmosphere and on the removal of air pollutants. Previous modeling studies used CO measurements in combination with chemistry-transport models (CTMs) to provide top-down constraints on the surface emissions and (in some cases) also the photochemical production of CO (Fortems-Cheiney et al., 2009, 2011; Jiang et al., 2015, 2017; Kopacz et al., 2010; Stavrakou & Müller, 2006). The resulting updated emissions are sensitive not only to the choice of atmospheric data set and inversion setup but also to the modeled OH abundance, primarily because reaction with OH is by far the largest CO sink (also because methane oxidation by OH is among the largest CO sources) (Kopacz et al., 2010; Stavrakou & Müller, 2006). Model-calculated OH fields show large differences among models; for example, the global tropospheric chemical lifetime of methane ranges between 7.1 and 13.9 years (Voulgarakis et al., 2013), whereas the best estimate based on methylchlorororm (MCF) is 11.2 years with an uncertainty range of 9.8–12.5 years (Prather et al., 2012). Furthermore, while models predict higher OH in the Northern than in the Southern Hemisphere, a recent MCF analysis (Patra et al., 2014) indicate an interhemispheric N/S ratio (R_{N/S}) very close to 1 (0.97 ± 0.12), much lower than the Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP) multimodel mean ratio of 1.28 ± 0.10 (Naik et al., 2013). The OH levels being overestimated by most models in the Northern Hemisphere, the total hemispheric top-down CO emissions are likely too high. Here we conduct an inverse modeling assessment of the global CO budget based on Infrared Atmospheric Sounding Interferometer (IASI) retrievals (George et al., 2015) for 2013, taking into account the MCF-derived constraints on global and
Figure 1. Annually averaged CO columns observed by (top left) Infrared Atmospheric Sounding Interferometer (IASI) in 2013, and modeled with IMAGES using (top right) a priori or (bottom panels) optimized emissions. The LN (low OH in Northern Hemisphere) and HN (high OH in NH) runs refer to the OH fields used in the model (Table 1).

2. Methods

2.1. IMAGES

The IMAGES model (Bauwens et al., 2016; Stavrakou et al., 2016) calculates the distribution of 170 chemical compounds at 2° × 2.5° resolution between the surface and the lower stratosphere (44 hPa) using ERA-Interim ECMWF meteorology (Dee et al., 2011). The model uses anthropogenic emissions from HTAPv2 (Janssens-Maenhout et al., 2015), with the nonmethane volatile organic compounds (NMVOC) speciation provided by ACCMIP (Lamarque et al., 2010), fire emissions from GFED4s (van der Werf et al., 2017), and biogenic VOC emissions calculated using MEGAN-MOHYCAN (Guenther et al., 2006; Müller et al., 2008; Stavrakou et al., 2011). Biogenic CO emissions and CO deposition are also taken into account (Müller & Stavrakou, 2005). The chemical degradation mechanism is described in Bauwens et al. (2016) and includes recent isoprene mechanism updates as well as an explicit treatment for 16 other NMVOCs. To minimize the impact of model errors associated with stratospheric chemistry and the boundary condition at the model top, the CO mixing ratios in the seven uppermost layers (44, 55, 67, 80, 96, 113, and 133 hPa) are fixed and set to a reanalysis of Aura Microwave Limb Sounder observations (Santee et al., 2017) by the Belgian Assimilation System for Chemical Observations stratospheric assimilation system (Errera et al. (2017), Figure S1). The flux inversion
Table 1

Table 1: Description of OH Fields Used in the CO Inversions

<table>
<thead>
<tr>
<th>Run label</th>
<th>Description</th>
<th>$\tau_{\text{OH}}$ (years)</th>
<th>$R_{\text{NS}}$</th>
<th>[OH] in NH (10^5 cm$^{-3}$)</th>
<th>[OH] in SH (10^5 cm$^{-3}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>STD</td>
<td>Standard</td>
<td>11.2</td>
<td>0.97</td>
<td>9.2</td>
<td>9.5</td>
</tr>
<tr>
<td>HN</td>
<td>Very high OH in NH</td>
<td>9.8</td>
<td>1.10</td>
<td>11.2</td>
<td>10.1</td>
</tr>
<tr>
<td>LN</td>
<td>Very low OH in NH</td>
<td>12.5</td>
<td>0.85</td>
<td>7.8</td>
<td>9.0</td>
</tr>
<tr>
<td>HS</td>
<td>Very high OH in SH</td>
<td>9.8</td>
<td>0.85</td>
<td>9.8</td>
<td>11.5</td>
</tr>
<tr>
<td>LS</td>
<td>Very low OH in SH</td>
<td>12.5</td>
<td>1.10</td>
<td>8.8</td>
<td>8.0</td>
</tr>
</tbody>
</table>

Note. The parameter $\tau_{\text{OH}}$ denotes the tropospheric chemical lifetime of methane, $R_{\text{NS}}$ is the interhemispheric ratio.

2.2. IASI Observations

The observations are monthly averaged CO total columns from IASI on board Metop-A (Clerbaux et al., 2009; Hurtmans et al., 2012) for 2013 binned onto the CTM resolution (Figures 1, and S2). IASI provides a global coverage twice daily. IASI CO data were previously used in modeling studies (e.g., Fortems-Cheiney et al., 2009; Klonecki et al., 2012) and evaluated against other satellite data sets (George et al., 2009, 2015). The instrument sensitivity is highest in the middle to upper troposphere. Since the vertical information content is coarse, the profiles are not used in the inversion. The model accounts for the IASI averaging kernels and for the temporal sampling in the monthly averages. Since the information content is low at high latitudes during winter, due to the low temperatures and low signal-to-noise ratio (Pommier et al., 2010), we exclude the winter months, defined by the following: October–May (>75° N), November–April (65–75° N), November–March (55–65° N), November–February (48–55° N), and similarly in the Southern Hemisphere (shifted by 6 months). This definition was guided by comparison with Fourier-transform infrared spectroscopy (FTIR) total column data (Figure S3, S4).

The IASI total error (sum of measurement and smoothing error) is typically very low (2–5%) over continents during summer and somewhat higher (5–15%) elsewhere. The number of data per model grid cell and per month is generally very high (200–2,000) and only the systematic part of the retrieval error contributes non-negligibly to the measurement error of the superobservations, namely, the monthly averaged columns at 2° × 2.5°. The large number of measurements per superobservation also reduces greatly the representativity error associated with CO variability within the same grid cell and month (Miyazaki et al., 2012). The dominant error component is the CTM error, which is more difficult to assess. It is here taken to be 35% of the CO column. This choice is supported by the $\chi^2$ diagnostic (i.e., the first term of the cost function divided by the number of observations, cf. supporting information), which should be close to one after optimization (Klonecki et al., 2012). As discussed below its value is of the order of 0.8 after optimization.

2.3. OH Fields Used in CO Inversions

The OH concentrations simulated by IMAGES are replaced in the troposphere by distributions in line with observational constraints based on MCF observations (Patra et al., 2014; Prather et al., 2012). The OH fields rely on the climatological tropospheric distributions of Spivakovsky et al. (2000) (S2000) adjusted to match the MCF constraints. The S2000 distribution achieves $R_{\text{NS}} \approx 1$, with hemispherically averaged OH concentrations $[\text{OH}]_{\text{NH}} = 11.4 \times 10^5 \text{ cm}^{-3}$ and $[\text{OH}]_{\text{SH}} = 11.5 \times 10^5 \text{ cm}^{-3}$ and it implies a global tropospheric chemical methane lifetime of 9.1 years. This field is scaled in each hemisphere to match the hemispherically averaged OH concentrations (Table 1) calculated using

$$[\text{OH}]_{\text{NH}} = X_g \frac{2 R_{\text{NS}}}{R_{\text{NS}} + 1}, \quad [\text{OH}]_{\text{SH}} = X_g \frac{2 R_{\text{NS}}}{R_{\text{NS}} + 1}$$

where $X_g$ is the globally averaged OH concentration consistent with Prather et al. (2012) (best value: 0.932 × 10^6 cm$^{-3}$, uncertainty range = (0.835 – 1.065)×10^6 cm$^{-3}$), and $R_{\text{NS}}$ the N/S concentration ratio from Patra et al. (2014) (0.97, range = 0.85 – 1.1).
3. Results

3.1. Optimized Fluxes

The IASI CO columns are illustrated on Figure 1 along with the IMAGES CO columns with either a priori or optimized emissions. In contrast with previous modeling studies (Fortems-Cheiney et al., 2009; Kopacz et al., 2010; Stavrakou & Müller, 2006), the modeled columns of the LN simulation overestimate the observations over most of the Northern Hemisphere, due to the lower OH concentrations and therefore higher CO lifetime compared to those studies. The optimizations bring the model much closer to IASI data (Figure S5); for example, the \( \chi^2 \) diagnostic (which measures the overall bias between model and observations) is reduced by almost 65% in both hemispheres in the LN optimization. Among the different optimizations, LN achieves the lowest average \( \chi^2 \) (0.70), followed by LS and the standard run STD (both 0.74), whereas the optimizations with higher OH (HN and HS) performed comparatively worse (0.86 and 0.84). The LN inversion is also the simulation which achieves the lowest mean bias in both hemispheres (−0.1% in NH and 0.2% in SH).

Although the major features of the optimized model CO columns are similar in all five source inversions (Figure 1), the corresponding top-down emission estimates show large differences (Figure 2 and Table 2), in particular, for anthropogenic emissions (factor of 1.4 between global emissions in HN and LN). The higher (lower) OH levels in the HN (LN) inversion lead to a shorter (longer) CO lifetime, which is compensated by higher (lower) emissions. The inferred isoprene emissions are also strongly sensitive to the OH levels, and range between 353 (LN) and 454 Tg yr\(^{-1} \) (HN).

The global fluxes range between 2,139 (in LN) and 2,711 Tg CO (in HN, Table 2) and are generally lower compared to previous estimates, for example, 2,748 Tg CO (Stavrakou & Müller, 2006), 2,857 Tg CO (Kopacz et al., 2010), and 2,391 Tg CO (Jiang et al., 2017). The latter study derived an annual anthropogenic flux of 54.3 Tg CO over the United States in 2013, at the upper end of our estimated range, but about half of a previous top-down estimate (97 Tg CO, Jiang et al. (2015)). Low anthropogenic CO emissions over the United States are supported by independent evaluations using aircraft data. Data collected during the aircraft campaign of the International Consortium for Atmospheric Research on Transport and Information in 2004 suggested
Table 2
A Priori and Optimized CO Fluxes (Tg CO yr\(^{-1}\)) in the Five Inversions (Table 1)

<table>
<thead>
<tr>
<th></th>
<th>A priori</th>
<th>STD</th>
<th>HN</th>
<th>LN</th>
<th>HS</th>
<th>LS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anthropogenic</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>North America</td>
<td>66</td>
<td>61</td>
<td>79</td>
<td>50</td>
<td>66</td>
<td>58</td>
</tr>
<tr>
<td>USA</td>
<td>44</td>
<td>39</td>
<td>53</td>
<td>31</td>
<td>42</td>
<td>37</td>
</tr>
<tr>
<td>Europe</td>
<td>30</td>
<td>29</td>
<td>30</td>
<td>27</td>
<td>29</td>
<td>28</td>
</tr>
<tr>
<td>China</td>
<td>166</td>
<td>164</td>
<td>202</td>
<td>134</td>
<td>175</td>
<td>156</td>
</tr>
<tr>
<td>India</td>
<td>63</td>
<td>44</td>
<td>52</td>
<td>37</td>
<td>47</td>
<td>41</td>
</tr>
<tr>
<td>Global</td>
<td>544</td>
<td>511</td>
<td>617</td>
<td>436</td>
<td>564</td>
<td>478</td>
</tr>
<tr>
<td>Open fires</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>North Africa</td>
<td>55</td>
<td>53</td>
<td>57</td>
<td>51</td>
<td>53</td>
<td>53</td>
</tr>
<tr>
<td>South Africa</td>
<td>95</td>
<td>107</td>
<td>113</td>
<td>102</td>
<td>120</td>
<td>96</td>
</tr>
<tr>
<td>Amazonia</td>
<td>16</td>
<td>15</td>
<td>15</td>
<td>14</td>
<td>17</td>
<td>12</td>
</tr>
<tr>
<td>Indonesia</td>
<td>17</td>
<td>10</td>
<td>10</td>
<td>9</td>
<td>10</td>
<td>9</td>
</tr>
<tr>
<td>Global</td>
<td>291</td>
<td>320</td>
<td>352</td>
<td>294</td>
<td>354</td>
<td>293</td>
</tr>
<tr>
<td>Other sources (global)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CH(_4)+OH</td>
<td>718</td>
<td>718</td>
<td>820</td>
<td>644</td>
<td>816</td>
<td>648</td>
</tr>
<tr>
<td>NMVOC</td>
<td>750</td>
<td>733</td>
<td>815</td>
<td>674</td>
<td>813</td>
<td>671</td>
</tr>
<tr>
<td>Ocean/soils</td>
<td>96</td>
<td>98</td>
<td>107</td>
<td>91</td>
<td>108</td>
<td>90</td>
</tr>
<tr>
<td>Total source</td>
<td>2399</td>
<td>2381</td>
<td>2711</td>
<td>2139</td>
<td>2655</td>
<td>2180</td>
</tr>
</tbody>
</table>

that the U.S. EPA (Environmental Protection Agency) overestimates CO emissions by 60% (Hudman et al., 2008), corresponding to an estimate of 26 Tg in 2013 when accounting for the emission trend (www.epa.gov/sites/production/files/2016-12/national_tier1_caps.xlsx). Using a geographically more limited data set, another study deduced a 15% overestimation of U.S. EPA emissions, implying a flux of 42 Tg CO in 2013 (Anderson et al., 2014). The CO top-down estimate of the LN inversion (31 Tg yr\(^{-1}\)) lies between these two campaign-based estimates, whereas the U.S. total of the HN inversion (53 Tg yr\(^{-1}\)) seems largely overestimated.

The annual top-down fluxes over China range between 134 Tg CO (LN) and 202 Tg CO (HN), in good agreement with a previous inversion based on Measurement Of Pollution in the Troposphere (MOPITT) columns (160–177 Tg CO; Jiang et al., 2017). It is difficult to further validate the CO emission inventories, especially in China, due to the sparsity of independent measurements. Estimates from different bottom-up inventories indicate large differences for the same year (2008), 105 Tg CO in EDGARv3.2 (European Commission Joint Research Centre, 2011), 202 Tg in REASv2.1 (Kurokawa et al., 2013), whereas recent evaluations suggest intermediate values (153 Tg in EDGARv4.3.1 and 158 Tg in Saikawa et al., 2017, both for 2008).

The sensitivity of the fire emissionsto OH levels is relatively smaller (Table 2). However, all inversions indicate strongly decreased fluxes over Indonesia during the fire season, with the strongest decrease (factor of 3) occurring in June. This result is supported by an independent top-down inversion of VOC fluxes based on HCHO columns (Bauwens et al., 2016), pointing to a large flux overestimation (factor of 5) in the GFED4s database in this region in June 2013. Similarly, the inferred decrease in Amazonian fire fluxes is corroborated by a similar emission decrease in VOC fluxes in August–September 2013 (ca. 20%, Bauwens et al., 2016).

### 3.2. Comparison to Independent Observations

We evaluate the inversions against ground-based FTIR total CO columns at 16 sites of the Network for the Detection of Atmospheric Composition Change (NDACC) (ftp.cpc.ncep.noaa.gov/ndacc/station, see supporting information). Their estimated systematic errors are typically 2–4%. For simplicity, we focus on the inversions with either the lowest (LN) or the highest (HN) [OH] in the Northern Hemisphere, where most evaluation data are available. In all simulations, the root-mean-square deviation (RMSD) between modeled and observed monthly columns is decreased in both hemispheres in the posterior estimates. The results in the Northern Hemisphere indicate a clear improvement when [OH] is low (RMSD of 6.9% for LN, 8.4% for HN). A large part (about half) of the difference between the RMSD of LN and HN inversions is associated with
Figure 3. Modeled and observed CO columns at Fourier-transform infrared spectroscopy (FTIR) stations. In dark blue: FTIR monthly means with their standard deviations; dashed lines: modeled values using a priori fluxes; solid lines: modeled values using optimized fluxes. HN run in red, LN in green.

The optimizations are further evaluated against surface CO mixing ratios from the NOAA/GMD and GAW networks (Figure S6), for a total of 90 stations providing data in 2013. The uncertainty on those measurements is typically 2–5 ppb (Novelli et al., 2017). The comparison is summarized in Figures 4 and S7. Overall, the best (worst) performance in the Northern Hemisphere is realized by the LN (HN) simulation. The LN simulation provides also the best match with Southern Hemisphere data. The RMSD is strongly decreased in the Southern Hemisphere upon inversion, for example, from 49 to 21% for LN. This is for largely due to the reduction of fire emissions over Indonesia in June, causing a strong reduction of the bias at Bukit Kotobatang in the stations of Toronto and Mauna Loa. At Toronto, located in the direct vicinity of anthropogenic emission sources, the higher CO fluxes of the HN inversion lead to a significant overestimation (+10%, Figure 3). At Mauna Loa, far away from the source areas and where CO is therefore more sensitive to OH levels, HN underestimates the FTIR columns in spite of its higher emissions. The RMSD between the model and FTIR columns is very similar to the RMSD between the model and IASI columns (7.3% for LN, 8.3% for HN in the Northern Hemisphere), suggesting an excellent general consistency between IASI and FTIR CO total columns, at least when high-latitude wintertime IASI data are excluded. In the Southern Hemisphere, both FTIR and IASI data indicate a better agreement (lower RMSD) when OH levels are close to the standard case (runs STD, LN, HN, with RMSD very close to 6.6% against FTIR in all three cases) than when OH is either very high (HS, 8.3%) or very low (LS, 7.7%, Figure 4).
Figure 4. (upper panels) RMSD (in %) between monthly Fourier-transform infrared spectroscopy (FTIR) CO total columns and corresponding model values and (lower panels) between monthly CO mixing ratios at GMD/GAW stations and the optimized model fields in the (left) Northern and (right) Southern Hemispheres.

Sumatra (Figure S7). The RMSD reduction from optimization is smaller in the Northern Hemisphere (from 14.5 to 13.6%). The model generally underestimates the wintertime observations at high northern latitudes, possibly due to the exclusion of high-latitude wintertime IASI data in the inversions. As for the FTIR total columns, the HN simulation overestimates CO over source areas (e.g., over Colorado and Utah) and it underestimates CO at remote oceanic sites (Midway, Mauna Loa and Guam), whereas the LN achieves a comparatively better match at these stations. The general conclusions are therefore consistent with those drawn from total CO column data. Nevertheless, the comparisons for surface mixing ratios show a larger spread (reflected by larger RMSD) than in the case of total columns. This is likely due to the larger model uncertainties for surface mixing ratios than for total columns, which are less affected by errors associated with vertical mixing.

Finally, the modeled CO profiles over North America are evaluated against aircraft measurements from the SEAC4RS (Toon et al., 2016) and DC3 (Barth et al., 2015) campaigns and from the Global Greenhouse Gas Reference Network (GGGRN) (Sweeney et al., 2015) (cf. Figure S8). Despite their very different emissions and OH fields, the three optimizations yield very similar results in the middle and upper troposphere, where IASI sensitivity is highest (4–12 km). Closer to the surface, however, significant differences are found, reaching up to 20% between HN and LN. The higher emissions of the HN inversion lead to near-surface CO overestimation in all campaigns and (for GGGRN) at all seasons, although the discrepancy is very small in spring and fall. The LN achieves the best agreement (lowest RMSD) with observations of both DC3 and SEAC4RS campaigns, as well as of GGGRN data in summer and winter, and it also performs better than the HN inversion against GGGRN data in spring and fall.

4. Discussion and Conclusions

This study demonstrates the strong impact of model uncertainties in OH concentrations on global and regional top-down CO flux estimates. For example, differences of 40% in the top-down global anthropogenic emissions are found when varying the OH levels, and even larger differences are found regionally, for example, over China and the United States. Clearly, all inverse modeling studies addressing CO, CH₄, and other reactive species should report their global mean OH levels and interhemispheric ratios.

Evaluation of our results against IASI and a wide range of independent CO measurements (FTIR total columns, network surface mixing ratios, aircraft in situ data) shows that the inversion adopting the lowest average OH
abundance in the Northern Hemisphere provides the best match with most observational sets, suggesting, in particular, that the interhemispheric (NH/SH) OH ratio might be close to the lower end of the range (0.85–1.1) reported by Patra et al. (2014), whereas the global photochemical methane lifetime might be near the upper end of the range (9.8–12.5 years) given in Prather et al. (2012). This inversion achieves the best agreement with measurements in the Southern Hemisphere, although other source inversions (e.g., STD) achieve very similar results in that hemisphere. While uncertainties exist due to errors in transport, seasonal evolution, and intra-hemispheric distributions of OH, we note that these inverse estimates fit independent aircraft measurements of the Eastern United States best assuming low OH in the Northern Hemisphere.

Acknowledgments

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