Increased Northern Hemispheric carbon monoxide burden in the troposphere in 2002 and 2003 detected from the ground and from space


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Abstract. Carbon monoxide total column amounts in the atmosphere have been measured in the High Northern Hemisphere (30°–90° N, HNH) between January 2002 and December 2003 using infrared spectrometers of high and moderate resolution and the Sun as a light source. They were compared to ground-level CO mixing ratios and to total column amounts measured from space by the Terra/MOPITT instrument. All these data reveal increased CO abundances in 2002–2003 in comparison to the unperturbed 2000–2001 period. Maximum anomalies were observed in September 2002 and August 2003. Using a simple two-box model, the corresponding annual CO emission anomalies (referenced to 2000–2001 period) have been found equal to 95 Tg in 2002 and 130 Tg in 2003, thus close to those for 1996 and 1998. A good correlation with hot spots detected by a satellite radiometer allows one to assume strong boreal forest fires, occurred mainly in Russia, as a source of the increased CO burdens.

1 Introduction

The importance of biomass burning for atmospheric composition, including the concentrations of greenhouse gases on global and hemispheric scales, is widely recognized (Andreae and Merlet, 2001; Galanter et al., 2000 and references therein). Flannigan et al. (1998) proposed a possible positive feedback between the emissions of greenhouse gases and wild forest fires. According to Holloway et al. (2000), during a normal year (i.e., without catastrophic wildfires) biomass burning contributes 748 Tg CO/year globally. Andreae and Merlet (2001) estimated a contribution from extratropical forest fires to be 68 Tg CO per year. The IPCC report of 2001 (Ehhalt et al., 2001) considered all of the available information and adopted for global conditions 700 Tg CO per year as emitted by the biomass burning and 650 Tg CO per year due to combustion of fossil and domestic fuels. Petron et al. (2004) inverted the MOPITT (Measurements Of Pollution In The Troposphere) satellite measurements between April 2000 and March 2001 (i.e., during a period of relatively low fire activity in the Northern Hemisphere (NH), see

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Table 1. CO monitoring locations.

<table>
<thead>
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<th>Name</th>
<th>Agency</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Altitude, m</th>
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Notes: 1) Asterisks indicate mountain stations, latitude and longitude are in degrees, negative longitude corresponds to West from Greenwich. 2) Monitoring agencies: CMDL = Climate monitoring and Diagnostics Laboratory, Boulder, Colorado, USA CSIRO = Commonwealth Science and Industry Research Organization, Canberra, Australia JMA = Japan Meteorological Agency, Tokyo, Japan NIES = National Institute of Environmental Studies, Tsukuba, Japan IMK-IFU = IMK-IFU, Forschungszentrum Karlsruhe, Garmisch-Partenkirchen, Germany IMK-ASF = IMK-ASF, Forschungszentrum Karlsruhe, Karlsruhe, Germany AWI = Alfred-Wegener-Institute, Potsdam, Germany UB = University of Bremen, Germany CUT = Chalmers University of Technology, Göteborg, Sweden IAP = Institute of Atmospheric Physics, Moscow, Russia IAG = Institute of Astrophysics and Geophysics, Liège, Belgium

results of this paper). They derived 408 Tg CO per year from the vegetation fires and 683 Tg CO per year from fossil fuel and biofuel use combined. It is well known that the CO emission from the mid and high latitude NH boreal forest fires experiences significant interannual variations (Wotawa et al., 2001). Kasischke et al. (2005) bottom-up calculations of CO amounts emitted by the biomass burning in the boreal areas in 2002 and 2003 were higher than those for the previous two years (i.e., 60–150 Tg per year versus 30–90 Tg per year in 2000–2001). These values may be compared to the industrial/transportation emission in the HNH estimated by Kasischke et al. (2005) for the period between 1998 and 2003 as high as 291 Tg CO per year.

In a recent paper (Yurganov et al., 2004) the CO tropospheric burden anomaly in the High Northern Hemisphere (HNH) between 1996 and 2001 was determined from total column spectrometric measurements combined with surface in situ data. Using a simple box model, the CO emission anomaly in 1998 in comparison to the period 1996–2001, without 1998, was estimated at 38 Tg/month in August and 96±29 Tg for the entire year of 1998. This result is in agreement with the emission anomaly calculated for 1998 by Kasischke et al. (2005): 45–113 Tg, but higher than most other bottom-up estimates referenced by Yurganov et al. (2004). The upper limit of the error due to possible changes in CO sinks was estimated at ±20%.
Jaffe et al. (2004) found an enhancement in June-July-August 2003 CO background surface concentrations of 23–37 ppb at 10 sites in Alaska, Canada and Pacific Northwest and explained this by a transport from Siberia. Jaffe et al. (2004) calculated the CO emission from Siberian fires in 2003 to be 68 Tg CO using a bottom-up approach.

3-dimensional chemical transport models (CTM) are now widely used for forward and inverse modeling of CO sources (e.g., Arellano et al., 2004, Petron et al., 2004 and references therein). The CTM models allow studying specific sources and specific areas on the Globe. However, the models are very complicated and simple box models applied to large reservoirs can be used as well. The CO burden (total mass) in the HNH is regulated by the sources inside the semi-hemisphere and removal processes. The most important of the latter are i) the transport into the subtropics, where CO concentrations are usually lower (Novelli et al., 2003) and, ii) a reaction with OH that is responsible for 90% of the CO chemical sink (Holloway et al., 2000). The CO lifetime in the HNH due to these two sinks, (1.5–3 months and 1.4–12 months, respectively, depending on the season (Yurganov et al., 2004)) makes mixing inside the reservoir very effective in spite of an extremely inhomogeneous distribution of wild fires. This box-model study does not aim to retrieve the rates of regional CO sources, which would require a more sophisticated CTM and higher temporal resolution. On the contrary, a simple box model and a synthesis of all available data for the entire HNH allows one to retrieve the integrated imbalance of the burden of CO resulting from anomalies of the integrated source on a monthly basis.

This paper presents updated results of surface-based total column and in-situ measurements of CO burdens supplemented by new satellite data (Edwards et al., 2004). The CO burdens in summer-autumn of 2002 and 2003 were found to be higher than in 2000 and 2001, and the corresponding CO emission anomalies have been updated using a methodology similar to that used by Yurganov et al. (2004).

2 Observational stations and methods

In this work we present for the first time CO total column amounts derived from infrared solar absorption observations performed between January 2002 and December 2003 with Fourier transform spectrometers (FTIR) operated at stations located in the European Arctic, Scandinavia, Western Russia, the European Alps, and the subtropical Atlantic (Table 1 and Fig. 1). The high resolution spectra (better than 0.005 cm$^{-1}$) were fitted with calculated spectra, and CO vertical profiles were retrieved. Total column amounts derived from the profiles have been used in this paper. Spectra recorded by a Russian home-made grating spectrometer with a resolution of 0.2 cm$^{-1}$ just allowed determination of total column amounts of CO (Yurganov et al., 2002). Rinsland et al. (1998) estimated the random and systematic error of a single total column measurement of CO as ±2–3% and ±5%, respectively. Retrieval techniques are based on the same absolute spectroscopic constants of absorption lines and do not need a calibration; different fitting algorithms applied to the same spectra agree within 1% (Hase et al., 2004). Details of retrieval procedures can be found in the papers by Hase et al. (2004), Yurganov et al. (2002), and Yurganov et al. (2004). Results of comparisons between grating and FTIR spectrometers installed at Egbert, Ontario, Canada and at Lamont, Oklahoma, USA, were described by Yurganov et al. (2002); the maximum systematic disagreement between the instruments with different spectral resolution amounted to 5%. Yurganov et al. (2002) also compared the grating spectrometer data in Zvenigorod, Russia with CO mixing ratios, measured using an aircraft between 0 and 6 km of altitude in the same area on 7 June 1995, 17 August 1995, and 25 March 1996. In spite of differences in locations up to 160 km, differences in the sampling time up to several hours, and a diurnal CO cycle of 10–15% in magnitude, the results show a reasonable agreement between two different modes of measurements (differences between spectroscopic data and in situ data were from 0 to 8%). It should be noted also than an analysis of anomalies
Fig. 2. Monthly mean total column amounts (triangles; in molecules/cm$^2$) of carbon monoxide above various sites. The repeating every year solid blue lines (here and in Fig. 3) correspond to the monthly means, averaged over the period between March, 2000, and February, 2002.

(not the absolute values) employed in this paper allows one to minimize the influence of systematic errors introduced by retrieval techniques or differences due to geographic locations. These data were used for a calculation of the anomalies of CO burden (total mass) in the reservoir 30$^\circ$N–90$^\circ$N between 0 and 10 km altitude.

Air samples collected in the surface atmospheric layer by the NOAA/CMDL Cooperative Air Sampling Network and by other programs (Table 1 and Fig. 1) were analyzed for local CO concentrations. The CMDL analysis of the air samples were made by gas chromatography/HgO reduction detection using instruments from Trace Analytical, Inc. They were referenced to the so-called CMDL Reference Scale, which was based on two sets of primary standards produced at CMDL during the late 1980s and early 1990s (Novelli et al., 2003). At the two Japanese stations, CO mixing ratios were measured continuously using non-dispersive gas correlation IR instruments (Tanimoto et al., 2000, WMO, 2004) and calibrated using standards of the Chemical Evaluation and Research Institute, Japan, or Nippon Sanso company. In both cases the calibration was cross-checked against the National Center of Atmospheric Research standard (Tanimoto et al., 2000). The uncertainty of measurements is estimated as ±(1–3) ppb (Novelli et al., 2003) and ±7% (Tanimoto et al., 2000).

The satellite-borne MOPITT instrument is a thermal IR nadir-viewing gas correlation radiometer described in detail by Drummond (1992) and Deeter et al. (2003). MOPITT uses a cross-track scan, which allows for almost complete coverage of the Earth’s surface in about 3 days, with individual pixels of 22 km×22 km horizontal resolution. Accuracy of the measured CO total column amount is ±10% but the sensitivity of the instrument significantly drops in the boundary layer (Deeter et al., 2004). The measured total column, therefore, depends on an a priori profile. In polluted boundary layer conditions the standard background a priori profile used instead of a more realistic one leads to an underestimation of the total column (cf., Edwards et al., 2004).

3 Results and discussion

Monthly mean CO total column (TC) amounts above 7 ground-based sites are plotted in Fig. 2. Ny-Alesund (Spitsbergen), Kiruna (Sweden), Harestua (Norway) and Zvenigorod (Russia) are low altitude stations and the measured total column amounts represent concentrations integrated over the entire atmosphere, thus including the boundary layer (BL), the free troposphere (FT), and the stratosphere. Total columns measured at the Alpine mountain sites Zugspitze,
Fig. 3. Examples of monthly mean mixing ratios (in parts per billion in mole fraction) at the surface layer sampling sites. Jungfraujoch, and at the high altitude site Izaña (Canary Isles) are CO amounts integrated between the altitude of the station and the top of the atmosphere (note a change in the vertical scale).

It is known that 90% of the CO column amount resides below 12 km (Zhao et al., 2002). We assume here that the fire-induced variations of CO column amounts with durations of a few months occur mainly in the troposphere: the time constant for the vertical exchange between the troposphere and stratosphere is more than one year (Brasseur et al., 1999).

Monthly mean values in Fig. 2 (triangles) may be compared to values averaged over the 24-month-long reference period between March 2000 and February 2002 (solid blue lines). The particular choice of the beginning of the reference period is determined by the start of the MOPITT measurements in March, 2000. 24 following months were characterized by low CO levels, comparing to previous and following time. A common reference seasonal cycle for three stations (Harestua, Kiruna, and Ny-Alesund) was calculated using the data for all three locations. This was done because of some gaps in the measurements at Harestua and Ny-Alesund during the reference period. These raw data indicate higher than normal CO total columns during summer-autumn periods of 2002 and 2003 at all sites. A similar effect (even more clearly due to averaging over the entire semi-hemisphere) is found in the MOPITT data also presented in Fig. 2.

Record high CO column amounts were measured at Zvenigorod in summer-autumn 2002 with a maximum in September. According to the International Forest Fire News (IFFN, 2003), there was smoke pollution from peat and forest fires in the region around Moscow between July and September, 2002. This caused a dramatic reduction in visibility, to less than 100 m in the city, and also had detrimental impacts on the health of the Muscovite population. The fires reached a peak on 6 September. The observed CO spikes (up to 7.3 E18 molecules/cm² on 10 September 2002) reflected intrusions of this polluted air (there were no visible fires around the station itself). The horizontal extent of the highly polluted area remains uncertain. In this paper the days with these spikes were omitted. There were nine days in July–September, 2002 with CO columns that exceeded the summer time 2002 lowest daily value (2.02 E18 molecules/cm²) by more than 4 standard deviations, i.e., above 2.9 E18 molecules/cm². Standard deviation of daily mean values, 0.22 E18 molecules/cm², was calculated for the reference period of 2000–2001. Specifically, we omitted 1 out of 12 measurement days in July, 1 out of 6 days in August, and 7 out of 10 days in September.

Increased CO mixing ratios were found in the boundary layer over many remote sites of HNH in 2002 and 2003 as well. Figure 3 demonstrates examples for the Arctic, two Atlantic sites and an East Asian location. However, the increase of CO mixing ratio in Ryori, Japan was less pronounced than for other sites.

The further analysis will be carried on for anomalies to exclude seasonal variations of CO and to highlight the differences between the 2002–2003 and 2000–2001 periods. The relative anomalies in CO abundance (monthly means divided by the values measured during the reference period) are plotted in Fig. 4 (the top panel is for total column contents, the bottom panel is for mixing ratios in the BL). The anomalies in CO total columns for both data sets have two maxima, in September–October, 2002, and in August, 2003. The CO anomalies at the southernmost station Izaña (28° N) are
Fig. 4. Anomalies of CO total column amounts (A) and surface mixing ratios (B). The measured monthly means were divided by the averages over the period between March 2000 and February 2002. MOPITT data are averaged over the HNH, mean BL is the average over the data of 12 low-level stations. Extreme daily values exceeding 2.9 E18 molecules cm$^{-2}$ in Zvenigorod in July–September, 2002, were omitted (see text).

lower than those at other sites. The lower CO anomalies support the assumption that the northern mid-latitude wildfires play a decisive role in the anomaly in hemispheric CO burdens (cf., a conclusion of Yurganov et al. (2004) for 1998).

The anomalies averaged over in situ BL stations in Fig. 4b (maximum 35% in 2002 and 36% in 2003) were higher than corresponding TC anomalies in Fig. 4a (19% and 25%, respectively). A similar effect was observed in August, 1998: 71% for the BL and 48% for the total column relative to 2000–2001 (Yurganov et al., 2004). The larger surface anomalies are expected if the source of additional CO is on the surface and the vertical mixing is not instantaneous. Moreover, a photochemical removal of anomalous CO occurs during its uplifting.

Fig. 5. Monthly sums of hot spot numbers for selected geographical areas (the locations of the areas are shown in Fig. 1). The numbers of hot spots averaged over the period between March, 2000, and February, 2002 used as a reference are designated by a dashed line.

The total column anomaly measured by MOPITT and integrated over the HNH, has a similar temporal shape (Fig. 4a). Edwards et al. (2004) analyzed the MOPITT data in conjunction with simultaneously measured aerosol optical depths (AOD) from the Moderate-resolution Imaging Spectroradiometer (MODIS) and fire counts, determined by special algorithm based on several channels of MODIS. In September 2002, enhanced CO, AOD, and fire counts were observed concurrently over the European part of Russia. In 2003, forest fires detected by MODIS started burning in early spring and were most intense in Siberia near the Baikal Lake in May. Later on the fires moved northwards through Siberia during the summer.

A convenient tool for fire detection is the number of hot spots on the Earth’s surface, detected at night time by the Along Track Scanning Radiometer (ATSR) or recently launched Advanced Along Track Scanning Radiometer (AATSR) (the “Algorithm 1” was described by Arino and Plummer, 2001). The total number of hot spots was counted separately for three geographical areas (Fig. 5) with boundaries shown in Fig. 1. Although Siberia experienced a larger variation of hot spots, the relative anomalies of CO total column measured by MOPITT and averaged over the same areas do not show significant differences (Fig. 6). It seems that the atmospheric transport effectively mixes this anomalous CO in the semi-hemisphere horizontally.

The anomalies of the CO burden (in Tg) in the reservoir $30^\circ$N–$90^\circ$N, 0–10 km were calculated using three data sets (Fig. 7, top panel and left scale). Firstly, the anomalies in CO total column amounts at four low altitude stations, Harestua, Kiruna, Zvenigorod, and Ny Alesund were averaged and multiplied by the surface area of the reservoir (red open circles in Fig. 7). Secondly (blue rectangles in Fig. 7), the average anomalies in CO mixing ratios in the BL were multiplied by the number of air molecules in the lowest 1.5 km layer.
(0.355 E25 molecules cm$^{-2}$, according to the 1976 US Standard Atmosphere) and by the surface area of the reservoir. These CO burden anomalies were added to the FT anomalies derived from the measurements at the Alpine Jungfraujoch and Zugspitze stations. The data from the Izaña mountain station were not considered here, because the station is located outside the chosen reservoir. And thirdly, the CO burden anomalies for the same reservoir were calculated from the global MOPITT data (full green triangles). All three data sets have similar patterns and are very close in their absolute values. A good general agreement of the data, obtained by different instruments, in different locations is noteworthy. This can be explained by a significant averaging of the data and also by a good horizontal mixing. The only period of disagreement is January–April, 2003, when the total column data resulted in approximately 10 Tg higher values of the CO burden than other data sets. Local effects (mostly due to transport between the semi-hemispheres) may be a cause of this disagreement.

A box model (Yurganov et al., 2004) was applied to estimate the monthly mean anomalies in the CO source in the HNH taking into account transport to the Low Northern Hemisphere (LNH, 0°–30° N, 0–10 km of altitude) and chemical removal.

\[
P_{HNH}' = \frac{dM_{HNH}'}{dt} + L_{trans}' + L_{chem}' \quad (1)
\]

\[
L_{trans}' = \frac{(M_{HNH}' - M_{LHNH}')}{TAU_{trans}} \quad (2)
\]

\[
L_{chem}' = \frac{M_{HNH}'}{TAU_{chem}} \quad (3)
\]

\[
TAU_{chem} = \frac{1}{k[OH]} \quad (4)
\]

\[
k = 1.5*E - 13(1 + 0.6*p) \text{ cm}^3 \text{ mol}^{-1} \text{ s}^{-1} \quad (DeMore et al., 1997) \quad (5)
\]

where \(P_{HNH}'\) is the CO emission rate (in Tg/month), \(M_{HNH}'\), and \(M_{LHNH}'\) are CO tropospheric burdens in the HNH and LNH reservoirs (in Tg); \(dM_{HNH}/dt\) is the change in the tropospheric CO burden during the month, \(L_{trans}'\) and \(L_{chem}'\) are loss terms due to the transport between the semi-hemispheres and OH-consumption, respectively; \(p\) is air pressure in atm; the prime (’) designates the anomaly, i.e., the deviation from the average over the reference period between March 2000 and February 2002.

The interannual variations of the sinks (in other words, \(TAU_{chem}'\) and \(TAU_{trans}'\)) were neglected. \(TAU_{trans}\) was calculated using the GEOS-CHEM global 3-D CTM (Bey et al., 2001) with assimilated 1998 meteorology (Yurganov et al., 2004). \(TAU_{chem}\) was calculated using OH fields as a function of season, latitude, and altitude (Spivakovsky et al., 2000).

Burden anomalies for LNH \((M_{LHNH}')\) in 1996–1999 were calculated from the surface measurements by Novelli et al. (2003), in 2002–2003 they were assumed from the MOPITT measurements. They were taken as the average of the two data sets mentioned above in 2000 and 2001.

Estimates of the CO emission anomalies in the HNH, derived from the model (Eqs. 1–5) using the three data sets described before are plotted on the middle panel of Fig. 7 (the reference period is indicated by the shaded area). Results
Table 2. Annual CO emission anomalies in Tg CO/year for the HNH retrieved from the low-altitude FTIR ($P'(MOPITT)$), and in situ BL, combined with mountain sites ($P'(BL+FT)$) using a box model. The averages of two (1996–1999) or three (2000–2003) estimates are also given. The accuracy is estimated from standard deviation for eight runs of the box model for perturbed input parameters described by Yurganov et al. (2004). For Kasischke et al. (2005) both emissions and anomalies in reference to 2000–2001 are presented.

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<th>Year</th>
<th>$P'(FTIR)$</th>
<th>$P'(MOPITT)$</th>
<th>$P'(BL+FT)$</th>
<th>Average anomaly</th>
<th>Accuracy, Tg/yr</th>
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</tbody>
</table>

The bottom panel of Fig. 7 illustrates the interannual variations of total numbers of hot spots detected by the ATSR after July 1996. The monthly anomalies of hot spots were calculated similarly to CO anomalies. The years 2002 and 2003 were characterized by positive anomalies of hot spots in August, 2002 and May–July, 2003. Positive anomalies of hot spots were observed also in 1996 and 1998. During all of these years higher numbers of fire counts were found mostly in Siberia (see Fig. 5). Note that the high numbers of counts observed in 2000 was distributed almost equally over the three geographical areas (Fig. 5), and the CO level was normal during that year. This apparently implies that the Siberian forest fires influence the hemispheric CO levels disproportionately more than the wild fires in other continental areas.

Plots of the correlation between CO emission anomalies and the fire counts anomalies are displayed in Fig. 8. The slopes of the plots indicate that on a monthly basis 144 fire counts detected correspond to 1 Tg CO emitted. The HNH CO emission anomalies correlate better to the fire counts in Siberia ($R^2=0.51$), than to the fire counts in the entire hemisphere ($R^2=0.36$). This is in agreement with Kasischke et al. (2005), who pointed out that the average emission per unit area for fires in the boreal region were higher in Eastern Russia than in North America, mainly because of higher aboveground biomass levels and deeper organic soil layers.

The average anomalies derived from all three data sets were converted into absolute emission rates assuming a “normal” emission seasonal cycle. Biomass burning emission rates in 2000 (50.7 Tg CO/year) calculated for the MOZART-2 CTM (M. Schultz and C. Granier, personal communication, 2003; Schultz, 2002; Olivier et al., 1996) were used to define a “normal” seasonal cycle (Fig. 9). In 1996 and 1998 maximum emissions were observed in August. In 2002 emissions had a maximum in July–August. In 2003 the emissions presented in Figs. 2, 4a, and 7 are given also in a tabular form in the Electronic Supplement. All three data sets clearly reveal abnormally high CO emission rates in 2002 and 2003. The yearly emission anomalies are presented in Table 2. Since 1996 there have been four years with high CO emission of similar magnitude (1996, 1998, 2002, and 2003) and four years with low emission (1997, 1999, 2000, and 2001).

Kasischke et al. (2005) obtained bottom-up estimates of CO emissions from boreal fires. Attempts were undertaken to quantify the depth of burning layer more accurately than previously. To take into account types of fires different burn severity scenarios were assumed. CO emissions for the high severity case (the estimates for two other scenarios were lower), are presented in Table 2 together with their anomalies referenced to 2000–2001. Our top-down estimates usually are higher then the bottom-up one. The difference between the studies is less for 2002 (the inversion of the MOPITT data agrees with the bottom-up value), but amounts to 100% in 2003. Yurganov et al. (2004) estimated that up to 11% of the additional CO burden measured in the atmosphere may be a result of a photochemical conversion from the pyrogenic non-methane hydrocarbons. This secondary CO source was not considered by Kasischke et al. Some contribution in resolving the discrepancy may also be obtained from taking into account a depletion of [OH] caused by higher [CO].

Daniel and Solomon (1998) estimated that a 1% increase of [CO] led to a 0.34% decrease of [OH]. Yurganov et al. (2004) estimated that halving [OH] leads to 22% lower values of $P'$ in August, 1998. Therefore, the observed 30% increase of [CO] in summer times of 2002 and 2003 would result in a 10% drop of [OH]. This consequently leads to less that 10% lower values of $P'$ retrieved using our model and assumptions for 2002 and 2003. Both effects diminish the difference, but do not resolve the disagreement completely.
started to grow in May–June and reached a peak in July, in accordance with the fire pattern in Siberia (Fig. 5); these three months were characterized by very high numbers of fire counts. However, the August emission was still higher than normal in contrast to a low number of hot spots. There may be two explanations for this effect. Firstly, as previously mentioned, up to 11% of the additional CO in the HNH may be a result of the photochemical conversion from hydrocarbons. A maximum input of this secondary CO source may be expected to occur with some delay after the emission of hydrocarbons, depending on their chemical life-times. Secondly, ATSR can easily detect the flaming fires but the smoldering fires may be under its detection limit.

4 Conclusions

Spectroscopic measurements of carbon monoxide total column amounts from ground based stations in the Arctic and Europe reveal an increased CO abundance in summer and autumn of 2002 and 2003 in comparison with the previous two years. Similar increases were observed in 1996 and 1998 (Yurganov et al., 2004). Increased CO concentrations near the surface were also observed by the CMDL network. Moreover, space-based MOPITT measurements with a full HNH coverage reveal a similar pattern in 2002 and 2003. An inversion of emission source anomalies from monthly burden anomalies (assuming a stable sink) has been done using a simple box model. The annual emission anomalies calculated in 2002 and 2003 are in the range 70–160 Tg CO/year which are similar in magnitude to those in 1996 and 1998. The estimate for 2002 is close to the most recent bottom-up calculations by Kasischke et al. (2005), but our top-down estimate for 2003 is a factor of two higher than that by Kasischke et al. (1995). The anomalies in the CO emission rates correlate with anomalies of fire counts, detected by a satellite-based radiometer. It is most likely that strong wildfires are responsible for the interannual variations of the hemispheric CO burden.

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