

⁹Atmospheric Chemistry Division, National Center for Atmospheric Research, Boulder, Colorado, USA

¹⁰Spectroscopie de l'Atmosphère, Service de Chimie Quantique et Photophysique, Université Libre de Bruxelles (ULB), Brussels, Belgium

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Correspondence to: M. Parrington (mark.parrington@ed.ac.uk)

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Abstract

We analyse the tropospheric ozone distribution over North America and the North Atlantic to boreal biomass burning emissions during the summer of 2010 using the GEOS-Chem 3-D global tropospheric chemical transport model, and observations from in situ and satellite instruments. In comparison to observations from the PICO-NARE observatory in the Azores, ozonesondes across Canada, and the Tropospheric Emission Spectrometer (TES) and Infrared Atmospheric Sounding Instrument (IASI) satellite instruments, the model ozone distribution is shown to be in reasonable agreement with mean biases less than 10 ppbv. We use the adjoint of GEOS-Chem to show the model ozone distribution in the free troposphere over Maritime Canada is largely sensitive to NO_x emissions from biomass burning sources in Central Canada, lightning sources in the central US, and anthropogenic sources in eastern US and south-eastern Canada. We also use the adjoint of GEOS-Chem to evaluate the Fire Locating And Monitoring of Burning Emissions (FLAMBE) inventory through assimilation of CO observations from the Measurements Of Pollution In The Troposphere (MOPITT) satellite instrument. The CO inversion showed that, on average the FLAMBE emissions needed to be reduced to 89 % of their original values, with scaling factors ranging from 12 % to 102 %, to fit the MOPITT observations in the boreal regions. Applying the CO scaling factors to all species emitted from boreal biomass burning sources led to a decrease of the model tropospheric distributions of CO, PAN, and NO_x by as much as -20 ppbv, -50 ppbv, and -20 ppbv respectively. The impact of optimizing the biomass burning emissions was to reduce the model ozone distribution by approximately -3 ppbv (-8 %) and on average improved the agreement of the model ozone distribution compared to the observations throughout the free troposphere reducing the mean model bias from 5.5 to 4.0 ppbv for the PICO-NARE observatory, 3.0 to 0.9 ppbv for ozonesondes, 2.0 to 0.9 ppbv for TES, and 2.8 to 1.4 ppbv for IASI.

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1 Introduction

The importance of ozone as a tropospheric trace constituent in the contexts of climate, air quality, and tropospheric chemistry has become well established in recent years. The main sources of tropospheric ozone are the oxidation of carbon monoxide (CO) and volatile organic compounds (VOCs) in the presence of nitrogen oxides (NO_x), and transport from the stratosphere. Removal of ozone from the troposphere is through dry deposition and photochemical destruction. The photochemical lifetime of ozone in the troposphere ranges from minutes and hours in the planetary boundary layer (PBL) to several days and weeks in the free troposphere where it can be subject to intercontinental transport and affect tropospheric composition far from the source region. The ozone distribution over the North Atlantic is of particular interest in this respect as it is strongly influenced by the transport of precursor emissions, both natural and anthropogenic in origin, from the North American continental PBL and subsequent chemical processes in the outflow (e.g. Fehsenfeld et al., 2006). Quantifying the influence of the different parameters that affect the North Atlantic tropospheric ozone distribution is important as this outflow and its subsequent chemical transformation can have an impact on tropospheric composition and air quality further downwind (e.g. over Europe Derwent et al., 2004). In this paper we evaluate the tropospheric ozone distribution over North America and the North Atlantic during the summer of 2010, and its sensitivity to ozone precursor emissions, particularly from boreal biomass burning sources, using a numerical model of atmospheric chemistry and transport, and in situ and satellite observations.

A number of previous studies have focussed on the distribution of tropospheric ozone, and influence of North American pollution, over the North Atlantic. The first long-term measurements of surface level ozone across the Atlantic Ocean were made from ship-based measurements in the 1970s and 1980s (Winkler, 1988). The North Atlantic Regional Experiment (NARE) (Fehsenfeld et al., 1996) was an airborne and ground-based measurement campaign conducted in the spring and summer of 1993

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to measure the distribution of tropospheric ozone and its precursors over the region. The first intensive measurements of the tropospheric ozone distribution over the North Atlantic were made during NARE with daily ozonesondes launched from Iceland, Newfoundland, Bermuda and the Azores (Oltmans et al., 1996). In 2004, the International Consortium for Atmospheric Research on Transport and Transformation (ICARTT) (Fehsenfeld et al., 2006) sampled North American pollution outflow as it was transported across the North Atlantic with airborne measurements made from the Azores (Lewis et al., 2007) and Europe (Real et al., 2007). In situ measurements of ozone and its precursors have also been made in the lower Atlantic free troposphere from the PICO-NARE observatory on the peak of Mt. Pico in the Azores at 38.47° N, 28.40° W, 2,225 m altitude (Honrath et al., 2004; Helmig et al., 2008). Model studies of the outflow of North American ozone precursor emissions and their impact on the tropospheric ozone distribution over the North Atlantic have been performed. Flatoy et al. (1996) and Kasibhatla et al. (1996) presented 3-D model analyses of the summertime tropospheric ozone distribution over the North Atlantic of the NARE campaign period. More recent analysis used the GEOS-Chem chemistry transport model to evaluate the transport of North American pollution outflow across the Atlantic and its subsequent impact on surface ozone over Europe (Li et al., 2002).

The main focus of these previous studies has been on the transport of ozone precursor emissions, from anthropogenic sources, out from the North American PBL and over the Atlantic. Measurements made during ICARTT did, however, sample plumes of Canadian biomass burning outflow along the eastern seaboard and over the North Atlantic. These measurements showed an unusual mixture of organic compounds within NO_y ($\text{NO} + \text{NO}_2 + \text{PANs} + \text{HNO}_3 + \text{HONO} + \text{NO}_3 + 2\text{N}_2\text{O}_5 + \text{organic nitrates}$) that existed only within biomass burning plumes over the Atlantic (Lewis et al., 2007). Much of this NO_y was held as peroxyacetyl nitrate (PAN) but with an abundance higher than expected and that was highly sensitive to slight changes in altitude and temperature. Measurements of ozone, CO, NO_x , and NO_y at the PICO-NARE observatory in the mid-Atlantic have also been shown to have enhancements that attribute their sources

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to boreal biomass burning (Lapina et al., 2006; Val Martin et al., 2006; Helmig et al., 2008). Years of high boreal fire activity have been shown to increase ozone concentrations in the free troposphere over the Azores by up to 10 ppbv (Lapina et al., 2006) with photochemically aged biomass burning plumes influencing tropospheric oxidant chemistry after 1–2 weeks of transport to the region (Helmig et al., 2008). In situ measurements of biomass burning outflow close to boreal source regions in North America were made during the Arctic Research of the Composition of the Troposphere from Aircraft and Satellites (ARCTAS) campaign in the summer of 2008. Analysis of the fresh plumes observed during ARCTAS showed very little evidence for ozone formation in the aircraft, satellite, and model results (Alvarado et al., 2010). Correlations between ozone and CO were observed in some of the plumes (9 out of 34) with an average ozone enhancement of $+0.5\% \pm 1.9\%$ consistent with a previous analysis of biomass burning outflow over Canada (Mauzerall et al., 1996). Observations of biomass burning plumes by the TES satellite instrument during ARCTAS (Alvarado et al., 2010) and over Siberia in the summer of 2006 (Verma et al., 2009) showed some evidence of ozone production in the smoke plume but also observed ozone depletion and were unable to attribute ozone changes to the fire emissions.

The Quantifying the impact of BOREal forest fires on Tropospheric oxidants over the Atlantic using Aircraft and Satellites (BORTAS) project is the most recent study of the tropospheric composition over the North Atlantic, combining model outputs with in situ and satellite observations of key tracers of boreal biomass burning and tropospheric oxidant chemistry. The initial phase of the BORTAS study took place in the summer of 2010, consisting of ground-based and ozonesonde measurements of biomass burning outflow, and is described in the following section. The second phase of the BORTAS study took place in the summer of 2011 and builds on the first phase with an aircraft measurement campaign including instrumentation to fully evaluate the speciation of NO_y and the chemistry within plumes of boreal biomass burning outflow as that are transported over the North Atlantic.

An overview of the BORTAS project is presented in Sect. 2. Section 3 presents an overview of the GEOS-Chem model of chemistry and transport used in this analysis. Section 4 presents the mean 3-D tropospheric ozone distribution for the first phase of the BORTAS campaign, conducted in the summer of 2010, with output from the GEOS-Chem chemical transport model evaluated against measurements made by ozonesondes and at the PICO-NARE observatory, and satellite observations. The sensitivity of the modelled ozone distribution to the model inputs, focussing on the boreal biomass burning regions, and MOPITT CO inversion, is presented in Sect. 5. We conclude in Sect. 6.

2 The BORTAS-A measurement campaign

The analysis of the summer 2010 tropospheric ozone distribution, presented here, is part of a multi-national project, led by the University of Edinburgh, aimed at Quantifying the impact of BOREal forest fires on Tropospheric oxidants over the Atlantic using Aircraft and Satellites (the BORTAS project, <http://www.geos.ed.ac.uk/research/eochem/bortas/>). The overarching scientific aims of the BORTAS project are to: (1) sample biomass burning outflow from boreal North America over the western boundary of the North Atlantic using ground-based, balloon, and aircraft measurements; (2) describe the observed chemistry within plumes using numerical models of atmospheric chemistry and transport, paying particular attention to the NO_y and organic chemistry; (3) quantify the impact of boreal forest fires on oxidant chemistry over the temperate and subtropical Atlantic Ocean; and (4) detect, validate and quantify the impact of boreal biomass burning on global tropospheric composition using data from space-borne sensors. Central to BORTAS is a measurement campaign with the UK Facility for Airborne Atmospheric Measurements (FAAM) BAe146 research aircraft, to be based out of Halifax, Nova Scotia, Canada, supported by a variety of ground-based measurements at Dalhousie University in Halifax and ozonesondes launched by Environment Canada from a number of sites across eastern Canada. The aircraft measurement campaign

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was originally scheduled to take place in July/August 2010 but was postponed until July/August 2011, due to disruption of air traffic over the Atlantic Ocean by the eruption of the Eyjafjallajökull volcano in Iceland during spring 2010. Although the aircraft measurement campaign was postponed, the support measurements went ahead as planned through July/August 2010 and make up phase A of the project, referred to as BORTAS-A from hereonin. The satellite data central to analysis of the BORTAS-A campaign period are ozone from the Infrared Atmospheric Sounding Instrument (IASI) and Tropospheric Emission Spectrometer (TES), and CO from the Measurements Of Pollution In The Troposphere (MOPITT) instrument. Further satellite observations for BORTAS data analysis are retrieved profiles of ozone, CO, and organic species associated with biomass burning outflow from the Atmospheric Chemistry Experiment Fourier Transform Spectrometer (ACE-FTS) but are beyond the scope of this work and will be analysed in other papers.

The main component of the ground-based campaign is the Dalhousie University Raman Lidar (Duck et al., 2007) which provided continuous, under clear sky conditions, measurements of tropospheric aerosol profiles throughout the campaign period. Additional instrumentation was also available at Dalhousie University (referred to as the Dalhousie Ground Station, or DGS, from hereonin) providing measurements of particulate matter and its speciation, wind profiling, surface ozone and total column of a number of atmospheric tracers. Of greater relevance to the work presented in this paper is a network of seven measurement sites across central and eastern Canada from which daily ozonesondes were launched between 12 July and 4 August 2010 inclusive, the original dates of the BORTAS aircraft campaign. The location of the ozonesonde launch sites, along with the time of the daily launches and the number of sondes launched from each site, are listed in Table 1 and shown in Fig. 2a. Analysis of the other DGS data is beyond the scope of the work presented here and will be reported in other papers (e.g. Tereszchuk et al., 2011).

2.1 Boreal biomass burning activity during BORTAS-A

Boreal biomass burning activity during the BORTAS-A campaign is evaluated using the Fire Locating And Monitoring of Burning Emissions (FLAMBE) inventory (Reid et al., 2009) to estimate the total carbon emissions from boreal biomass burning sources during the summer of 2010. The FLAMBE inventory provides hourly estimates of carbon and aerosol emissions based on fire data from geostationary and polar orbiting satellite platforms. Active fire data from the Geostationary Operational Environmental Satellite (GOES) platforms and Moderate Resolution Imaging Spectroradiometer (MODIS) instruments are combined with a 1 km×1 km land use database. MODIS fire emissions are extrapolated from roughly 6-hourly observations using a diurnal cycle that releases 90% of the total emissions between 09:00 and 19:00 local time (Reid et al., 2004). The FLAMBE inventory is also used in the analysis presented here to provide biomass burning emission estimates for the GEOS-Chem chemical transport model described in the next section.

Figure 1 shows the daily distribution of GOES and MODIS fire hotspots used by the FLAMBE inventory, and the estimated total carbon emissions, in kg, between 1 July and 4 August 2010. Figure 1a shows the daily distribution of fire hotspots, sampled on a horizontal grid with a resolution of 0.5 degree latitude×0.67 degree longitude, across the northern extra-tropics. Only the biomass burning emissions for the boreal region (defined as all latitudes poleward of 50° N, blue box in Fig. 1a) are evaluated here. Subregions of the boreal region are defined for boreal North America (170°–50° W, 50°–75° N), Central Canada (120°–90° W, 52°–62° N), and Eastern Siberia (110°–180° E, 50°–75° N) denoted by gold, red and green boxes in Fig. 1a respectively. Fires in Alaska, southern Central Canada, and eastern Canada were more prominent in the first half of the campaign period, with later fires in British Columbia and northern Saskatchewan in Central Canada. Fires over Eastern Siberia burned throughout the period with widespread activity continuing into the beginning of August.

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Figure 1b shows timeseries of the daily total carbon emissions over the defined time period for each region. Emissions over the whole boreal region increase steadily from values of approximately 10^{13} kg at the beginning of July to a peak value of almost 1.4×10^{14} kg on day 211 (30 July). The Eastern Siberia region is generally the largest contributor to the total boreal carbon emissions apart from a three day period between day numbers 192 and 195 (11 to 14 July) when the North American emissions are larger. The carbon emissions from boreal North America generally remain fairly constant at less than 10^{13} kg with peak values of 2×10^{13} kg on days 194 (13 July) and 205 (24 July). The boreal North American carbon emissions are largely dominated by the Central Canada region until day 208 (27 July) when the North American emissions show an increase up to 2×10^{13} kg while the Central Canada emissions remain fairly constant at approximately 5×10^{12} kg. The total carbon emitted from 1 July to 4 August 2010 is 1.2×10^{15} kg for the boreal region, 2.7×10^{14} kg for North America, 1.8×10^{14} kg for Central Canada, and 6.8×10^{14} kg for Eastern Siberia.

3 The GEOS-Chem global 3-D chemistry transport model

Model analysis of the BORTAS-A campaign period is performed with the GEOS-Chem chemical transport model. GEOS-Chem is a global 3-D model driven by assimilated meteorological observations from the NASA Goddard Earth Observing System version 5 (GEOS-5) from the Global Modeling and Assimilation Office (GMAO). The meteorological fields have a horizontal resolution of 0.5 degree latitude \times 0.67 degree longitude with 72 levels in the vertical, and a temporal resolution of 6 h (3 h for surface fields). In the work presented here, we use v8-02-04 of GEOS-Chem with a horizontal resolution of 2 degree latitude \times 2.5 degree longitude. Biomass burning emissions in the model are from the FLAMBE inventory (Reid et al., 2009) which provides hourly emissions of total carbon at a resolution of less than 5 km \times 5 km. The carbon emissions are aggregated over a generic 1 degree \times 1 degree grid, which is interpolated to the horizontal resolution of the model simulation, and scaled for individual gas and aerosol phase

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biomass burning tracers using emission factors from Andreae and Merlet (2001). The biomass burning emissions are distributed uniformly throughout the PBL using boundary layer heights from GEOS-5. Anthropogenic emissions in the model are provided for North America on a regional basis by: the Environmental Protection Agency (EPA) National Emissions Inventory 1999 (NEI99) updated for the eastern United States (Hudman et al., 2007) for the US; the Big Bend Regional Aerosol and Visibility Observational (BRAVO) study for Mexico (Kuhns et al., 2005); and the National Pollutant Release Inventory (NPRI) Criteria Air Contaminants (CAC) emissions for Canada. We use Asian anthropogenic emissions prepared for the NASA INTEX-B mission in 2006 (Zhang et al., 2009b), and European emissions are from the European Monitoring and Evaluation Programme (EMEP) inventory. Biogenic emissions are from the Model of Emissions of Gases and Aerosols from Nature (MEGAN) version 2.1 (Guenther et al., 2006). The lightning source of NO_x in GEOS-Chem is estimated from lightning flash rates calculated using deep convective cloud top heights (Price and Rind, 1992) provided from GEOS-5, locally redistributed using a seasonally varying climatology of lightning flash counts observed from space by the Optical Transient Detector (OTD) and the Lightning Imaging Sensor (LIS) (Sauvage et al., 2007); the vertical distribution of the source is imposed according to Pickering et al. (1998). Emissions of NO_x from soil sources are estimated using the scheme described by Wang et al. (1998). The stratospheric ozone distribution is represented by a linearized ozone (Linoz) parameterization (McLinden et al., 2000).

4 Model and observed tropospheric ozone distribution over North America and the North Atlantic

Figure 2 shows the mean model tropospheric ozone distribution at 750, 500, and 310 hPa (corresponding to approximate altitudes of 2.2, 5.5, and 8.9 km respectively), across North America and the North Atlantic averaged over the BORTAS-A campaign period. The model ozone distribution at 750 hPa shows an area of relatively high ozone

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concentrations (between 60 and 75 ppbv) localised predominantly over the eastern US and extending across the North American Basin to the 60° W meridian. Eastward of this meridian, a fairly broad filament of moderately high (45–60 ppbv) ozone, compared to background values of approximately 40 ppbv, can be traced across the ocean almost as far as western Europe. Poleward of 50° N, the model shows much lower ozone concentrations with some localised areas of high ozone across Greenland and, more notably, in an area of Canada to the west of Hudson Bay and close to the biomass burning source regions in Central Canada (Fig. 1a). The mid-tropospheric, 500 hPa, ozone distribution similarly shows relatively high ozone values (> 75 ppbv) across the eastern US and North American Basin along with plumes extending across the ocean equatorward of 50° N, with peak values of approximately 80 ppbv between 30° W and western Europe. Moderate model ozone values (60–70 ppbv) are shown across Canada at this pressure level and areas of localised high ozone values, such as those at 750 hPa, are not as apparent. At 310 hPa the model ozone distribution is dominated by high concentrations (>95 ppbv) over the eastern Atlantic ocean and centred on the Azores. Across North America, moderately elevated ozone concentrations (90–95 ppbv compared to background values of 80–90 ppbv) can be seen extending along the US-Canada border, and along the east coast of the US and Canada up to 50° N. Localised areas of ozone at these moderately elevated values are also seen at the higher latitudes to the west and east of Hudson Bay, and to the west of Iceland.

4.1 GEOS-Chem vs. Ozonesonde profiles

Ozonesondes are balloon borne instruments that measure the atmospheric ozone concentration during the balloon ascent through its electro-chemical reaction in an aqueous solution of potassium iodide. The inherent response time of the measurement is about 20–30 s and corresponds to a vertical resolution of 100–150 m for a balloon ascent of 4–5 m s⁻¹. The reported accuracy for ozonesonde measurements in the troposphere is typically less than 10 % (Thompson et al., 2011). A detailed overview of the observing capabilities of ozonesondes, and their role in measurement campaigns

over recent years was recently presented by Thompson et al. (2011).

Throughout the BORTAS-A campaign daily launches of ozonesondes were performed from seven launch sites distributed across Canada between the sources of boreal biomass burning and the Maritime provinces. Figure 2a and Table 1 show the ozonesonde launch sites used during BORTAS-A with their location and the total number of ozonesondes launched from each site during the campaign. Figure 3 shows mean ozonesonde profiles, averaged over the GEOS-Chem vertical levels, and the co-located (in space and time) model ozone profiles, for the individual launch sites and all sites. In general the mean bias between the ozonesonde and model ozone profiles is less than 10 ppbv throughout the lower and free troposphere. In the upper troposphere (altitudes above 8 km) the model generally shows a relatively large (>10 ppbv) negative bias compared to all ozonesonde locations except Walsingham. The largest mean model biases in the lower troposphere tend to occur below 2 km and are generally positive, which in some cases can be in excess of 10 ppbv, and are most prominent at Bratt's Lake, Egbert, Walsingham, and Yarmouth. In the free troposphere at all sites, the mean bias is within ± 5 ppbv. At each launch site, both positive (notably at Walsingham and Sable Island) and negative (Goose Bay) biases are shown in the free troposphere but differences in the under- or over-estimates of the model ozone distribution do not show any apparent geographical dependence. The comparison for Bratt's Lake is unusual in that it shows both positive (below 4 km) and negative (above 5 km) biases. At all launch sites, the mean GEOS-Chem and ozonesonde profiles agree within the $1\text{-}\sigma$ standard deviations, with the model and observations showing comparable variability.

Figure 4 shows the variability in the absolute differences between model ozone profiles co-located to the time and location of each ozonesonde profile (Table 1). In general, the biases are within ± 30 ppbv ($\pm 10\%$) throughout the troposphere at each site. Those sites noted above for showing a relatively higher bias in the PBL, i.e. Bratt's Lake, Egbert, Walsingham, and Yarmouth, all show a persistent positive model bias below 2 km throughout the BORTAS-A campaign period although the biases at Egbert

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are shown to be lower than at the other three sites. The overestimate in the model ozone concentrations in the PBL shows peak values at the surface for all sites apart from Yarmouth, where the peak bias appears to be between 500 and 1500 m, also shown in Fig. 3. Differences in the free troposphere at each launch site show a mixture of positive and negative biases over the course of the campaign period. The mean and median biases, in ppbv, at each site are shown in Table 1. The largest mean positive biases are seen at Walsingham, Egbert, Sable Island, Yarmouth, and Bratt's Lake and correspond to relative biases of 27.2 %, 20.1 %, 10.7 %, 10.1 %, and 16.7 % respectively. These relatively large mean positive biases can be attributed to one or two profiles showing a substantially larger positive model bias throughout the troposphere. Smaller mean biases at the other launch sites suggest that relatively large biases between individual model and ozonesonde profiles are more episodic and localised in nature. At Goose Bay and Montreal, the mean relative biases are 4.2 % and 4.7 % respectively.

Differences between the model ozone distribution and the ozonesonde measurements can potentially be attributed to small-scale vertical structures which can be measured by the ozonesondes and which are not properly resolved by the model. The inability of the model to capture these structures could be related to the relatively coarse vertical resolution of the model levels (approximately 0.7 km in the free troposphere and 1 km in the upper troposphere), or missing emissions/processes in the model. The potential for attributing these differences to the model emissions, transport, and chemistry will be assessed in Sect. 5. A more detailed analysis of the differences between the model and ozonesonde profiles is beyond the scope of this paper and will be addressed in other work.

4.2 GEOS-Chem vs. Pico observatory data

Measurements of ozone and other trace gas (CO, NO, NO₂, NO_y) and aerosol species were made at the PICO-NARE observatory throughout the summer of 2010 in support of BORTAS-A. The PICO-NARE observatory is situated on the peak of Mt. Pico in the

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Azores in the mid-Atlantic, 38.47° N, 28.40° W, at an altitude of 2,225 m (Honrath et al., 2004; Helmig et al., 2008). Ozone was measured using a commercial ultraviolet absorption instrument (Thermo Environmental Instruments, Inc., Franklin Massachusetts, Model 49C) (Owen et al., 2006; Helmig et al., 2008). In the work presented here we use 1 h averaged data. These data have a precision of typically less than 1 ppbv. The data have not been calibrated at the time of writing but are anticipated to have an accuracy of less than ± 1 –2 % similar to the estimated accuracy of ± 3 % previously reported for 30 min averaged ozone data, based on adjustments made to July 2003 raw ozone measurements (Owen et al., 2006).

Figure 5 shows the time series of 1 h averaged ozone concentrations measured at the PICO-NARE observatory between 4 July and 4 August 2010 inclusive, and the time series of model ozone output co-located in time and space to each measured value. The ozone measurements over this time period have a mean value of 42.3 ppbv in the range 20.4 to 81.8 ppbv with a standard deviation of 11.9 ppbv. The model ozone time-series tends to be at the higher end of the measured values and shows less variability reflected in a mean of 49.6 ppbv in the range 37.8 to 71.0 ppbv and a standard deviation of 6.0 ppbv. The mean relative bias between the model and observed ozone is 14 %. The correlation coefficient between the measured and model timeseries is 0.5 indicating that, although the model appears to capture some of the higher measured peak values, for example on days 192, 204, and 214, the relatively coarse horizontal resolution of the model output means that it cannot capture most of the measured variability.

4.3 GEOS-Chem vs. Satellite observations

We extend the evaluation of the model ozone distribution across North America and the North Atlantic with a comparison against tropospheric ozone profiles retrieved from nadir viewing infrared satellite instruments. The model ozone distribution is compared against tropospheric ozone observations from the Tropospheric Emission Spectrometer (TES) and the Infrared Atmospheric Sounding Instrument (IASI). Ozone profiles

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retrieved from satellite measurements are an estimate of the atmospheric state which is generally expressed as follows:

$$\hat{\mathbf{x}} = \mathbf{x}_a + \mathbf{A}(\mathbf{x} - \mathbf{x}_a) + \varepsilon \quad (1)$$

where $\hat{\mathbf{x}}$ is the retrieved ozone profile, \mathbf{x}_a is the a priori profile applied in the retrieval, \mathbf{x} is the true ozone profile, \mathbf{A} is the averaging kernel, and ε is the measurement noise. The averaging kernels provide the sensitivity of the retrieved state to the true state of the atmosphere. The trace of the averaging kernel matrix gives a measure of the number of independent pieces of information available in the measurements, more commonly referred to as the degrees of freedom for signal (DOFS) (Rodgers, 2000). Ozone profile retrievals from the TES and IASI satellite datasets, considered here, are expressed in terms of natural logarithm of volume mixing ratio ($\ln(\text{VMR})$) and partial column VMR. Comparison of the model ozone output against satellite data is presented in terms of $\ln(\text{VMR})$. The IASI averaging kernels are converted from partial column ozone to $\ln(\text{VMR})$ following Appendix A of Zhang et al. (2010). For a $\ln(\text{VMR})$ retrieval, Eq. (1) can be written:

$$\ln \hat{\mathbf{x}} = \ln \mathbf{x}_a + \mathbf{A}'(\ln \mathbf{x} - \ln \mathbf{x}_a) + \varepsilon' \quad (2)$$

Letting x_i denote the VMR for layer i , which is the i -th element of \mathbf{x} , and assume that the difference between x_i and $x_{a,i}$ is relatively small so that

$$\ln x_i - \ln x_{a,i} \approx \frac{x_i - x_{a,i}}{x_{a,i}} \quad (3)$$

with the same relationship holding between \hat{x}_i and $x_{a,i}$. The elements a'_{ij} of the converted averaging kernel \mathbf{A}' are then related to the elements a_{ij} of the averaging kernel \mathbf{A} by:

$$a'_{i,j} = \left(\frac{x_{a,i}}{x_{a,j}} \right) a_{i,j} \quad (4)$$

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Figure 6 shows averaging kernels for ozone profiles retrieved from the TES and IASI data for a cloud-free scene close to Halifax, NS, on 16 July 2010. The IASI averaging kernels have been converted from partial column VMR to $\ln(\text{VMR})$ using Eq. (4). The averaging kernels are colour coded to indicate the relative sensitivities of retrieval levels in different layers of the atmosphere: 0–5 km (red); 5–10 km (green); and 10–16 km (blue).

4.3.1 TES O₃

The TES instrument (Beer et al., 2001) is a high resolution imaging infrared Fourier-transform spectrometer (FTS), launched aboard the NASA EOS Aura satellite in July 2004. The Aura satellite is in a polar Sun-synchronous orbit with a repeat cycle of 16 days. TES utilises a nadir-viewing geometry and an instrument field-of-view at the surface of 8 km×5 km to observe spectral radiances in the range 650–3050 cm⁻¹ at an apodized spectral resolution of 0.1 cm⁻¹. Geophysical parameters are retrieved from the measured radiances based on a Bayesian framework that solves a constrained least squares problem (Bowman et al., 2006) on a forward model grid with 67 vertical levels (TES Science Team, 2009) with a thickness of approximately 0.6–0.7 km in the troposphere. The TES data have been filtered based on the mean and root mean square of the radiance residual and on the cloud top pressure of each profile, following Parrington et al. (2008) and TES Science Team (2009). We use TES global survey level 2 data version 4 (V004), geographical coverage of these data during the summer of 2010 did not extend poleward of 50° N in order to conserve the lifetime of the instrument. TES V004 ozone profile retrievals were validated against ozonesondes during the spring and summer phases of ARCTAS and found to be biased high in the troposphere by up to 15 % (Boxe et al., 2010), comparable to the 3–10 ppbv bias reported for the V002 retrievals at northern mid-latitudes (Nassar et al., 2008). TES ozone profile retrievals have been used in a number of studies evaluating biomass burning outflow in the boreal regions over Siberia (Verma et al., 2009) and North America (Alvarado et al., 2010; Boxe et al., 2010), and outflow from African biomass burning over the Atlantic

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Ocean (Jourdain et al., 2007) and Indonesia (Bowman et al., 2009). Ozone profiles retrieved from the TES data, for cloud-free conditions over the North Atlantic, are typically characterized by the averaging kernels shown in the left-hand plot of Fig. 6. A DOFS value of 1.07 for the troposphere (for a tropopause defined as the 120 ppbv ozone isopleth), compared to a value of 4.12 for the full profile, indicate that there is a single piece of independent information retrieved with the averaging kernels for the 0–5 km atmospheric layer (red lines in Fig. 6) showing the peak sensitivity to be between 3 and 4 km.

Figure 7 shows the comparison of the GEOS-Chem and TES ozone mid-tropospheric (600–400 hPa, approximately 4–6 km) distributions averaged over the BORTAS-A campaign period over North America and the North Atlantic. The model output has been sampled at the time and location of each retrieved TES ozone profile and incorporate the TES observation operator (Jones et al., 2003). A bias of +7.5 %, calculated as the mean of the biases reported in the Table 4 of (Boxe et al., 2010), excluding those from Barrow, is removed from the TES ozone profiles and is assumed to be constant throughout the troposphere. The bottom panel of Figure 7 shows the GEOS-Chem minus TES ozone difference. The ozone output from the model, sampled in the observation space, shows the highest concentrations equatorward of 40° N and centred over the eastern US and North Atlantic Basin. Across the Atlantic Ocean, the model shows moderate (approx. 60 ppbv) ozone concentrations. Relatively high ozone concentrations (>75 ppbv) over the Great Lakes region and eastern Atlantic Ocean are also evident although not as prominent as shown for the 500 hPa pressure level in Fig. 2b. In contrast, the TES ozone does not show the apparent north-south concentration gradient in the model output, with relatively high ozone values shown across the domain. The TES data show the highest ozone concentrations over the North Atlantic Basin and the eastern Atlantic Ocean, with a filament of high ozone extending north-south in between. The biases between the GEOS-Chem and TES ozone distributions are shown to be approximately evenly distributed between positive and negative values across North America and the North Atlantic. The mean and median of the model

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minus TES ozone bias across the domain shown in Fig. 7 are -0.34 and -0.12 ppbv in the range -40.83 to $+34.57$ ppbv.

4.3.2 IASI O₃

The IASI instrument (Clerbaux et al., 2009) is a Fourier transform spectrometer with a 2 cm optical path difference covering a spectral range from $645\text{--}2760\text{ cm}^{-1}$ at an apodized spectral resolution of 0.5 cm^{-1} . It observes the atmosphere in the nadir with an instantaneous field-of-view of four 12 km diameter pixels at the sub-satellite point scanned in a west-east swath of approximately 2200 km, perpendicular to the satellite orbit track. IASI was launched on the European Metop satellite, which is in a sun-synchronous orbit with an equator crossing at 0930 and 2130 local time, in October 2007. Tropospheric ozone profiles are retrieved from the IASI radiance measurements with the FORLI retrieval algorithm (version 20100406) based on using an optimal estimation methodology developed for nadir viewing thermal infrared sounders (Coheur et al., 2005; Boynard et al., 2009) on a retrieval grid with 40 vertical levels and vertical resolution of 1 km in the troposphere. Ozone profiles retrieved from the IASI data are found to be biased low by approximately 8% in the lower troposphere, below 6 km compared to ozonesondes at northern mid-latitudes in 2008 (Dufour et al., 2011). IASI data have been used in studies of biomass burning outflow in the Mediterranean during the summer of 2007 (Coheur et al., 2009). The averaging kernels for an IASI ozone profile retrieval are shown in the right-hand plot of Fig. 6. In contrast to the TES ozone averaging kernels, an ozone profile retrieved from the IASI measurements give lower values of DOFS for both the full and tropospheric profiles (3.33 and 0.55 respectively) with the sensitivity for the 0–5 km layer peaking at approximately 6 km. The differences in sensitivity between the IASI and TES ozone profile retrievals reflects the relatively coarse resolution of the IASI spectra and vertical retrieval grid compared to TES.

Figure 8 shows the comparison of the GEOS-Chem and IASI ozone mid-tropospheric (600–400 hPa, approximately 4–6 km) distributions averaged over the BORTAS-A campaign period over North America and the North Atlantic. Retrieved

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IASI ozone profiles with tropospheric DOFS less than 0.5 have been filtered out of the analysis. The model output has been sampled at the time and location of each valid retrieved IASI ozone profile and smoothed with the IASI averaging kernels. The IASI averaging kernels are converted from a partial column retrieval to $\ln(\text{VMR})$ using Equation 4 as described above. A mean bias profile that linearly increases in altitude from -15% at the lowest retrieval level to -5% at 8 km, and with values of 0% and $+10\%$ at 9 and 10 km respectively, based on comparison to ozonesonde profiles at northern mid-latitudes in 2008 (C. Clerbaux, personal communication, 2011), is added to the IASI ozone profiles. The bottom panel of Fig. 8 shows the GEOS-Chem minus IASI ozone difference. The model ozone output, sampled in the IASI observation space, shows a similar distribution to that shown in Figs. 2 and 7 with the higher ozone concentrations (approximately 60 ppbv) across the US, North Atlantic Basin, and eastern Atlantic Ocean. Poleward of 50°N smoothing of the model ozone output with the IASI averaging kernels leads to lower ozone values over the North Atlantic Ocean than shown at 500 hPa in Fig. 2b, centred around 30°W , due to limited sensitivity of the IASI ozone retrievals to the troposphere. At the higher latitudes over Canada, moderately high ozone concentrations (40–60 ppbv) are shown of similar order of magnitude at 500 hPa in Fig. 2b. The mean ozone distribution observed by IASI shows a similar pattern to that of the GEOS-Chem output sampled in the IASI observation space. The GEOS-Chem minus IASI ozone differences show the model to be higher than the observations over the majority of the geographical region shown. The largest disagreement between the model and IASI are across the central US and central Atlantic Ocean, where the ozone values are at their highest (>50 –60 ppbv), and the closest agreement poleward of 50°N and eastward of 90°W , where the ozone concentrations are relatively low (<30 ppbv). The mean and median of the model minus IASI ozone bias across North America and the North Atlantic are $+1.92$ and $+2.33$ ppbv in the range -23.48 to $+10.36$ ppbv.

5 Adjoint analysis of model inputs

The previous section presented an evaluation of the 3-D model tropospheric ozone distribution over North America and the North Atlantic using in situ and satellite observations. Here we evaluate the sensitivity of the model ozone distribution to the model inputs to isolate and compare their relative influences. The adjoint model approach provides an efficient means for analysing the sensitivity of the model output to each of the model inputs by evaluating the gradient of a cost function J with respect to the input parameters (e.g. emissions estimates, initial conditions). For a sensitivity analysis, the cost function can be simply defined as a set of model predictions g of a particular model state variable, e.g. trace gas profile or integrated column amount, \mathbf{c} :

$$J = \sum_{\mathbf{c} \in \Omega_s} g(\mathbf{c}) \quad (5)$$

where Ω_s is the set of times at which the cost function is evaluated.

When observations of a particular state variable (\mathbf{c}_{obs}) are available, the cost function balances the model and observed state, and the input parameter, e.g. emission inventory, (\mathbf{p}) against a priori knowledge of that parameter (\mathbf{p}_a), and is generally defined as:

$$J = \frac{1}{2} \sum_{\mathbf{c} \in \Omega} [\mathbf{H}(\mathbf{c}) - \mathbf{c}_{\text{obs}}]^T \mathbf{S}_{\text{obs}}^{-1} [\mathbf{H}(\mathbf{c}) - \mathbf{c}_{\text{obs}}] + \frac{1}{2} [\mathbf{p} - \mathbf{p}_a]^T \mathbf{S}_p^{-1} [\mathbf{p} - \mathbf{p}_a] \quad (6)$$

where \mathbf{H} is the observation operator which transforms the model profile into the observation space taking into account the characteristics of the observations (e.g. the averaging kernels for a profile retrieved from satellite data). \mathbf{S}_{obs} and \mathbf{S}_p are the observational and a priori error covariance matrices respectively and will be described below. Ω is the domain (in time and space) over which model predictions and observations are available.

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The optimal solution for the parameter to be estimated minimizes J in Eq. (6) so that the gradient of the cost function with respect to the model state vector is zero:

$$\nabla_{\mathbf{c}} J = \frac{\partial J(\mathbf{c})}{\partial \mathbf{c}} = 0 \quad (7)$$

Solution of the optimization problem requires the gradients of J with respect to the model state vector, and these are efficiently calculated using the adjoint method. The derivation of the adjoint of GEOS-Chem is described in detail by Henze et al. (2007).

The adjoint variable at any given timestep is the sensitivity of the cost function with respect to the model state vector (e.g. $\lambda_c^0 = \nabla_{c_0} J$ gives sensitivity to initial conditions; $\lambda_p^0 = \nabla_p J$ gives sensitivity to other model parameters such as emissions).

We use the GEOS-Chem adjoint for sensitivity calculations, to evaluate the relative contributions of different emissions sources to the modelled ozone distribution over the North Atlantic, and data assimilation, to evaluate the CO emissions estimates used by the forward model. The adjoint of GEOS-Chem was first described by Henze et al. (2007) and the version used here includes model code updates to the most recent version of GEOS-Chem. The model inputs are the same as described for the GEOS-Chem model in Sect. 3. The GEOS-Chem adjoint has previously been applied in model sensitivity studies for global aerosol sources (Henze et al., 2007) and attribution of ozone pollution in the western US (Zhang et al., 2009a), and for inversion of global CO emissions with satellite observations (Kopacz et al., 2010).

5.1 Sensitivity calculations

The sensitivity of the model tropospheric ozone distribution over the BORTAS-A campaign period to the model input parameters is evaluated using the adjoint of GEOS-Chem described above. The cost function for these adjoint sensitivity calculations is defined, in the manner described by Eq. (5), to be the sum of the model ozone distribution in the free troposphere (between the top of the PBL and the tropopause, typically 750–250 hPa at northern mid-latitudes) above the three ozonesonde launch sites in

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Maritime Canada (i.e. Goose Bay, Sable Island, and Yarmouth, see Table 1). The forward and backward simulations are performed from 12 July to 4 August 2010, and the cost function is evaluated at every hour throughout the simulation.

Figure 9 shows the sensitivity of the model ozone distribution over Maritime Canada to the NO_x emissions from biomass burning, lightning, anthropogenic, and combined aircraft, soil and biofuel sources, and CO emissions from biomass burning and anthropogenic sources. The location of the ozonesonde launch sites, where the cost function is initialized are shown by red triangles. The sensitivities shown have been normalised so that relative influence of the different emission sources can be compared against each other. The largest model ozone sensitivity is to anthropogenic NO_x emissions, centred over the northeastern US and southeastern Canada, in Quebec and Nova Scotia. The influence of the anthropogenic emissions extends across North America but the sensitivity is less than half the peak values over the main anthropogenic source regions. Biomass burning is shown to be the next most significant NO_x emission source, with the peak sensitivity localised to the main biomass burning region in Central Canada. The adjoint simulation also shows (although not shown here) some sensitivity of the free tropospheric ozone distribution to biomass burning NO_x emissions over Alaska and Eastern Siberia but is greatly reduced compared to the Canadian source. The model ozone also shows sensitivity to lightning NO_x emissions which, although smaller in magnitude compared to the anthropogenic and biomass burning sensitivity, cover a large area of North America with peak sensitivity shown to be over the central US, to the west of the Great Lakes, and extending into the eastern US and Central Canada. The combined sensitivity to NO_x emissions from other sources (i.e. aircraft, soil, and biofuel) is shown to be much lower than that for the anthropogenic, biomass burning, and lightning sources. For the CO emissions, shown in Fig. 9b, the largest ozone sensitivity is to the biomass burning sources in Central Canada. The peak sensitivity is approximately half that of the ozone sensitivity to the NO_x biomass burning emissions.

Figure 9 highlights some of the key potential sources of uncertainty in the model ozone chemistry and their relative contributions. In the remaining sections of this pa-

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per we evaluate ozone precursor emissions from biomass burning sources in order to isolate its associated uncertainty in the model ozone distribution. Although the anthropogenic sources of NO_x show the largest sensitivities, further quantification of their contribution is beyond the scope of the work presented here and will be evaluated in future work within the context of the BORTAS-B measurement campaign in summer 2011.

5.2 Biomass burning emissions

To evaluate the biomass burning emissions in the model we utilise satellite observations of CO which, due to its considerable longer atmospheric lifetime, of several weeks compared to several hours for NO_x , is a more reliable tracer of pollution outflow on a global scale. Emissions of NO_x , along with other species emitted by biomass burning, are related to the CO emissions through emission factors (e.g. Andreae and Merlet, 2001). Satellite observations of CO are available from a number of different platforms and measurement techniques. We use CO observations made by the Measurements Of Pollution In The Troposphere (MOPITT) instrument as they have been extensively used in inversions with GEOS-Chem in previous studies (e.g. Arellano et al., 2006; Fortems-Cheiney et al., 2009; Jones et al., 2009; Kopacz et al., 2009, 2010; Gonzi et al., 2011; Jiang et al., 2011). The MOPITT instrument is a gas correlation radiometer, launched aboard the NASA EOS Terra satellite in December 1999. MOPITT observes the atmosphere in the nadir with a group of four pixels with an instantaneous field-of-view of $22\text{ km} \times 22\text{ km}$ at the satellite sub-point, which are scanned across the orbit track in a swath approximately 650 km wide. The EOS Terra platform is in a Sun-synchronous orbit with a 10.30 local equator crossing time and a repeat cycle of 16 days, which combined with the MOPITT sampling, provides almost global coverage every 3–4 days. A full instrument description is given by Drummond et al. (2010). In this work we use vertical profiles of CO concentration retrieved using version 4 (V4) of the MOPITT operational processing algorithm (Deeter et al., 2010). In this product, CO profiles are based solely on $4.7\text{ }\mu\text{m}$ band (thermal infrared) radiances and are retrieved

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as quantities of $\log_{10}(\text{VMR})$ on a 10-level vertical pressure grid. Validation of the MOPITT V4 CO retrievals against aircraft flask samples over North America show a bias of approximately 1 % in the lower troposphere and up to -6 % in the mid-troposphere, with a drift in the bias of between approximately 1 and 2 % per year (Deeter et al., 2010).

Figure 10 shows the mean Northern Hemisphere CO emissions from the FLAMBE inventory averaged for the BORTAS-A campaign period. Biomass burning in the boreal regions feature prominently with the largest emissions, in excess of $10^{13} \text{ molec}^{-1} \text{ cm}^{-2} \text{ s}^{-1}$, shown in Central Canada, Eastern Siberia, and western Russia. Table 2 shows the total mass of carbon emitted from biomass burning, in the context of other emission sources of CO, over the BORTAS-A campaign period. The mass of carbon emitted is shown for the globe versus the boreal regions (all latitudes poleward of 50° N), and for the North America (170° to 50° W , 50° to 75° N), Central Canada (120° to 90° W , 52° to 62° N), and Eastern Siberia (110° to 180° E , 50° to 75° N) defined in Fig. 1. For all regions, the biomass burning emissions are the dominant source, accounting for more than 75 % of the total CO emissions. The total mass of nitrogen emitted as NO_x from different sources is shown in Table 3. In the boreal regions, biomass burning sources account for more than 60 % of the total NO_x emissions over North America, and 80 % over Eastern Siberia.

Previous studies using the FLAMBE inventory have shown the CO emissions to be a factor of approximately 2 too high when compared to aircraft observations from the spring (Fisher et al., 2010) and summer (Alvarado et al., 2010) phases of the ARC-TAS campaign in 2008. Optimizing the CO emissions used in the model simulations presented earlier in this work is desirable for two reasons: (1) quantifying the amount of ozone production and transport from the biomass burning source; and (2) reducing the uncertainty in the model ozone distribution associated with the reported over-estimation of the biomass burning source. We optimize the monthly CO emissions estimates for May-August 2010 using an inverse modelling approach with the GEOS-Chem adjoint described above. In the inversion MOPITT CO observations are averaged over

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the 4° latitude \times 5° longitude horizontal resolution of the model. The observation error covariance matrix \mathbf{S}_{obs} is defined as a uniform error of 20 % and is assumed to be diagonal at the $4^\circ \times 5^\circ$ resolution used for the inversion (Kopacz et al., 2010). The a priori error covariance matrix \mathbf{S}_a includes a uniform error of 50 % for combustion sources and 25 % for the global oxidation source following (Kopacz et al., 2010), and are assumed to be uncorrelated. The CO emissions are assumed to be optimized when the inversion has converged to a solution or when the twentieth iteration of the inversion process has been reached, when the change in the cost function is less than 1–2 %.

Figure 11 shows the results of the CO inversion for July 2010. Figure 11a and b show the a priori and a posteriori Northern Hemisphere emissions, in units of $\text{molec}^{-1} \text{cm}^2 \text{s}^{-1}$, for the combined contributions from biomass burning, anthropogenic, biofuel, and monoterpene sources. Figure 11c shows the, unitless, scaling factors derived from the MOPITT CO inversion. In general the CO inversion decreases the total CO emissions across the Northern Hemisphere, reflected by scaling factors of less than 1 in Fig. 11c. The distribution of the scaling factors reflects the location of the a priori CO emissions with the lower values of the scaling factors (i.e. where the emissions are estimated to have the largest over-estimate) coincident to the biomass burning emissions sources, particularly in the boreal regions. The mean scaling factor over the Northern Hemisphere is 0.86 in the range 0.09 to 1.15 and the total CO emitted from biomass burning over the BORTAS-A measurement period is reduced by 47 Tg C (53 %) from the value reported in Table 2, with the largest decrease over sub-tropical Africa. Mean scaling factors for the boreal region and the North America, Central Canada, and Eastern Siberia regions defined for Figure 1 are 0.89 (in the range 0.12–1.02), 0.92 (0.22–1.02), 0.72 (0.22–0.99), and 0.82 (0.20–1.01) respectively reducing the total CO emitted from biomass burning sources in these regions during BORTAS-A by 1.2 (12 %), 66.5 (24 %), 36.5 (43 %), and 9.6 Tg C (4 %) respectively. The mean scaling factor of 0.72 for the Central Canada region is almost double the scaling factor of 37.5 % applied to the amount of biomass consumed to match model CO profiles, using FLAMBE, to aircraft measurements taken during the ARCTAS sum-

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mer campaign in 2008 (Alvarado et al., 2010). The scaling factor applied by Alvarado et al. (2010) is within the range of scaling factors calculated for this region from the MOPITT CO inversion. Figure 11d shows the histogram of the relative bias between the MOPITT and model CO total columns after the first (black dashed line) and twentieth (red solid line) iterations of the CO inversion. The mean MOPITT minus model CO bias after the first iteration is -4.02% with a standard deviation of 14.72% which is reduced to $+2.46\%$ with a standard deviation of 5.90% after the twentieth iteration, indicating an improved fit of the model CO to the observations. The distribution of the CO bias after the twentieth iteration shows a higher frequency of zero bias compared to the distribution after the first iteration.

5.3 Impact on model ozone and precursor distributions

We evaluate the sensitivity of the model ozone distribution to the estimated CO emissions from biomass burning with the full chemistry forward model simulation described in Sect. 3. The scaling factors derived for the CO emissions by the MOPITT inversion performed at $4^\circ \times 5^\circ$ horizontal resolution above are first regridded per unit area to a resolution of $2^\circ \times 2.5^\circ$. The biomass burning CO emissions at each timestep are multiplied by the monthly scaling factors, while the CO emissions from other sources are not modified. In this way the response of the model ozone distribution only to the biomass burning can be evaluated. Figures 10 and 11 show the significant emissions from biomass burning and anthropogenic sources across the Northern Hemisphere are generally quite distinctive from each other, and the anthropogenic emissions more dominant in regions where they coincide.

Figure 12 shows the mean model distributions of ozone, CO, PAN, and NO_x averaged over the BORTAS-A campaign period at approximately 500 hPa from the simulation with the scaled FLAMBE inventory (top row) and the change in the model tracer distributions over the simulation with the original biomass burning emissions (bottom row). The relative differences between the model tracer distributions with the scaled and original biomass burning emissions all show a decrease across North America

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and the North Atlantic, apart from NO_x which shows a decrease across North America and almost zero difference over the Atlantic Ocean eastward of 60° W. All tracers show the largest changes (as much as -3 ppbv (-8%) for ozone, -20 ppbv (-30%) for CO, -50 pptv (-40%) for PAN, and -20 pptv (-40%) for NO_x) localised to the boreal biomass burning sources in Central Canada. For the ozone, CO, and PAN distributions, larger differences are shown to extend poleward from the main source region in northern Saskatchewan and eastward across Hudson Bay to northeast Canada and the North Atlantic. The largest differences (less than half the peak values over the fire regions for all tracers) are generally poleward of 50° N with smaller differences across the rest of the domain. The largest differences in the NO_x distribution are more localised to the source region with smaller differences across Maritime Canada. The differences associated with scaling the non-boreal biomass burning emissions (e.g. in the southern and eastern US) are more prominent in the ozone and CO distributions than for PAN and NO_x . The peak mean PAN and NO_x concentrations are centred over the eastern US and are approximately a factor of 2 for PAN, and 4 for NO_x , higher than their respective distributions over the Canadian biomass burning regions.

To evaluate the impact of scaling the biomass burning emissions on the model ozone distribution, we calculate mean differences between the model ozone profiles, from the original and scaled simulations, and the ozonesonde, TES, and IASI observations. Figure 13 shows mean difference profiles, in ppbv, between the model ozone and (a) ozonesondes, (b) TES, and (c) IASI. In all cases model ozone profiles are sampled at the time and location of each observation with black dashed lines showing the original model simulation and red solid lines showing the simulation with the scaled biomass burning emissions. In Fig. 13a and b the reported biases in the retrieved ozone profiles has been taken into account as described in Sects. 4.3.1 and 4.3.2.

The ozonesonde comparison shows a positive bias throughout the troposphere up to 8 km decreasing from $+9$ ppbv between 0.5–1 km to almost 0 ppbv at 6 km. Above 8 km, the bias becomes negative decreasing to -15 ppbv at 10 km. Scaling the biomass burning emissions leads to a better agreement between the mean model and

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ozonesonde profiles in the lower troposphere, below 5 km, and between 6.5 and 8 km with a mean bias of almost 0 ppbv. At altitudes between 5 and 6.5 km the reduction in the ozone profile, where the original bias was 0 ppbv, leads to a slight negative bias up to −2 ppbv. The mean difference, averaged from 0–10 km, of the bias is reduced from 2.96 ppbv for the original simulation to 0.88 ppbv for the scaled simulation. The TES comparison shows a negative bias between 2 and 6 km, with values down to −2 ppbv, and a positive bias above 6 km which increases to more than 20 ppbv at 10 km. The mean and median of the bias are 1.98 ppbv and 0.04 ppbv. The scaled biomass burning emissions decreases the mean and median of the bias to 0.94 ppbv and −0.73 ppbv respectively. The IASI comparison shows a positive bias throughout the troposphere up to 8–8.5 km, with a peak value of 3.6 ppbv, and a negative bias, of −5 ppbv at 10 km, in the upper troposphere with both a mean and median of the bias of 2.8 ppbv. The scaled simulation reduces the mean and median of the bias to 1.4 ppbv and 1.1 ppbv respectively. The mean bias compared to the PICO-NARE observations, not shown, is also decreased from 5.54 ppbv to 3.95 ppbv averaged over the BORTAS-A campaign period.

6 Conclusions

We have presented an evaluation of the tropospheric ozone distribution over North America and the North Atlantic during the summer of 2010 and its sensitivity to boreal biomass burning emissions. We used the GEOS-Chem chemical transport model to estimate the 3-D tropospheric ozone distribution and its performance relative to tropospheric ozone observed by in situ measurements from ozonesondes across Canada and the PICO-NARE observatory in the mid-Atlantic Ocean, and satellite measurements from TES and IASI. The model ozone distribution is shown to be in reasonable agreement with all of the observations, with mean biases less than 10 ppbv throughout the troposphere.

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We used the adjoint of GEOS-Chem to calculate the sensitivity of ozone in the free troposphere over ozonesonde launch sites in the Maritime provinces of Canada to biomass burning NO_x emissions relative to other sources of NO_x emission in July 2010. Having identified the model ozone sensitivity to biomass burning emissions, the adjoint of GEOS-Chem was further used to optimize the FLAMBE biomass burning inventory used in the GEOS-Chem simulations through an inversion of MOPITT CO profile data using a 4-D variational assimilation framework. The MOPITT CO inversion shows that the FLAMBE inventory overestimates the CO emissions throughout the Northern Hemisphere and the calculated scaling factors show that in the boreal regions, the FLAMBE emissions need to be scaled to as little as 22 % and 20 %, or mean scalings of 72 % and 82 %, of their original values across Canada and Eastern Siberia respectively. Scaling ozone precursor emissions from the FLAMBE inventory with the monthly factors calculated by the CO inversion in a full chemistry forward simulation shows a relatively large impact on the chemical species directly emitted by biomass burning (i.e. CO, PAN, and NO_x), reducing their concentrations at 500 hPa by up to approximately –50 % in the average values calculated over a three week period between 12 July and 4 August 2010.

The impact of the optimized FLAMBE inventory on the model ozone distribution was shown to be smaller relative to the directly emitted tracers, with decreases of up to –3 ppbv or –8 % averaged over the BORTAS-A campaign period. The change in the model ozone, and its mean difference compared to the ozonesonde, TES, and IASI observations, was shown to be a decrease of approximately 2–3 ppbv throughout the troposphere. Despite this relatively small change, the mean bias between the model ozone and ozonesonde profiles was reduced throughout the troposphere between the surface and 8 km from 2.96 ppbv to 0.88 ppbv. A similar improvement in the mean tropospheric bias was shown when compared to the satellite observations when their reported biases are taken into account with a reduction from 1.98 ppbv to 0.94 ppbv for TES and 2.8 ppbv to 1.4 ppbv for IASI.

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The adjoint sensitivity analysis of the model ozone distribution in the free troposphere showed that NO_x emissions from anthropogenic sources and lightning over North America also have a significant influence. A recent study has shown that the anthropogenic NO_x emissions over North America decreased by approximately 6% during 2006–2009 (Lamsal et al., 2011) which is not taken into account in the emissions estimates used by GEOS-Chem. For the lightning emissions, limitations in the model ability to reproduce the distribution of lightning intensities has been shown to underestimate the tropospheric ozone distribution relative to TES observations over North America (Jourdain et al., 2010). Evaluating these uncertainties in the anthropogenic and lightning NO_x emissions estimates were beyond the scope of the work presented here but further research under the BORTAS project will address these emissions sources and quantify their influence on the model ozone distribution.

The second phase of the BORTAS project (BORTAS-B) is an aircraft measurement campaign with the UK FAAM BAe146 research aircraft, which took place from 12 July to 3 August 2011. Instrumentation on the BAe146 provided a comprehensive suite of measurements of species associated with biomass burning in the outflow to the North Atlantic throughout the free troposphere. In particular, a Laser Induced Fluorescence system (Dari-Salisburgo et al., 2009) measured NO_2 and speciated NO_y which will provide valuable constraints for evaluating the model ozone distribution to its precursors and the chemistry within the biomass burning outflow. Analysis of the BORTAS-B campaign data will build on the results of the BORTAS-A campaign presented here, by utilising a more detailed chemical mechanism derived from the Master Chemical Mechanism to interpret tropospheric composition over North America and the North Atlantic observed by ground-based, airborne and satellite measurements.

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Table 1. Location of ozonesonde launch sites used during the BORTAS-A measurement campaign, and the number of ozonesondes launched from each site between 12 July and 4 August 2010. The distribution of the launch sites across Canada is shown in Fig. 2a.

Launch site	Lon./° W	Lat./° N	Launch time/UTC	Number	Mean (median) bias/ppbv
Bratt's Lake, SK	104.7	50.2	14	22	1.1 (4.1)
Egbert, ON	79.8	44.2	17	21	7.8 (9.4)
Walsingham, ON	80.6	42.6	17	17	13.0 (11.9)
Montreal, QC	73.4	45.5	17	26	0.6 (2.8)
Goose Bay, NL	60.1	53.5	11	22	−0.9 (0.2)
Sable Island, NS	60.0	44.0	23	22	4.1 (5.0)
Yarmouth, NS	66.1	43.9	23	23	2.8 (4.8)

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Table 2. Total CO emissions over the BORTAS-A campaign period in units of Tg C. Values in italics have units of 10^{-3} Tg C.

CO emission source	Global	Boreal	<i>North America</i>	<i>Central Canada</i>	<i>Eastern Siberia</i>
Biomass burning	100.40	9.75	<i>274.39</i>	<i>85.29</i>	<i>259.57</i>
Anthropogenic	31.40	0.31	<i>7.02</i>	<i>1.39</i>	<i>1.37</i>
Biofuel	4.97	0.11	<i>0.82</i>	<i>0.18</i>	<i>0.19</i>
Monoterpenes	3.77	0.31	<i>13.88</i>	<i>1.73</i>	<i>6.35</i>
Total	140.54	10.48	<i>296.11</i>	<i>88.60</i>	<i>267.49</i>

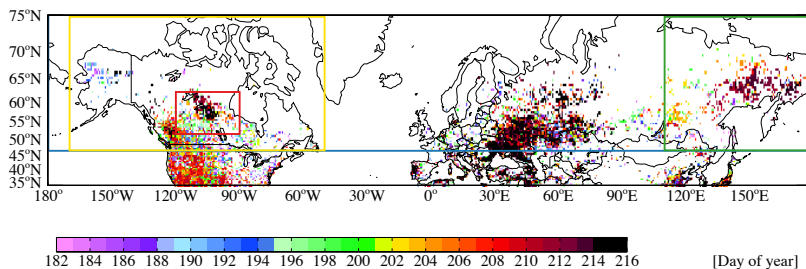
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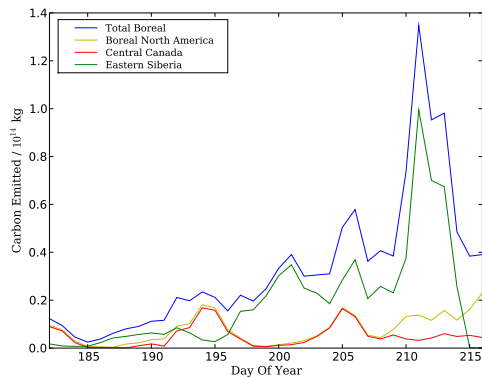
Table 3. Total NO_x emissions over the BORTAS-A campaign period in units of Gg N. Values in italics have units of 10⁻³ Gg N.

NO _x emission source	Global	Boreal	<i>North America</i>	<i>Central Canada</i>	<i>Eastern Siberia</i>
Biomass burning	1552.38	155.04	<i>3778.98</i>	<i>1132.67</i>	<i>4404.59</i>
Anthropogenic	1670.78	25.93	<i>623.30</i>	<i>196.27</i>	<i>86.72</i>
Lightning	765.49	26.28	<i>960.98</i>	<i>174.58</i>	<i>684.97</i>
Soil	574.74	12.39	<i>533.00</i>	<i>33.97</i>	<i>203.62</i>
Fertilizer	81.74	1.53	<i>17.41</i>	<i>0.20</i>	<i>3.22</i>
Aircraft	35.12	1.43	<i>101.09</i>	<i>3.54</i>	<i>9.12</i>
Biofuel	62.43	1.67	<i>11.87</i>	<i>2.60</i>	<i>2.75</i>
Total	4742.67	224.25	<i>6026.64</i>	<i>1543.82</i>	<i>5394.99</i>

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(a)



(b)

Fig. 1. Daily distribution of GOES and MODIS fire hotspots used in the FLAMBE inventory and the estimated total carbon emissions, in kg, from 1 July (day 182) and 4 August (day 216) 2010. Plot (a) shows the distribution of fire hotspots across the northern extra-tropics as a function of day of year. Plot (b) shows timeseries of the daily total carbon emitted for the boreal region (latitudes $>50^{\circ}$ N), boreal North America (170° – 50° W, 50° – 75° N), Central Canada (120° – 90° W, 52° – 62° N), and Eastern Siberia (110° – 180° E, 50° – 75° N). Line colours in plot (b) correspond to the boxes shown in plot (a) in which total emissions for each region are defined.

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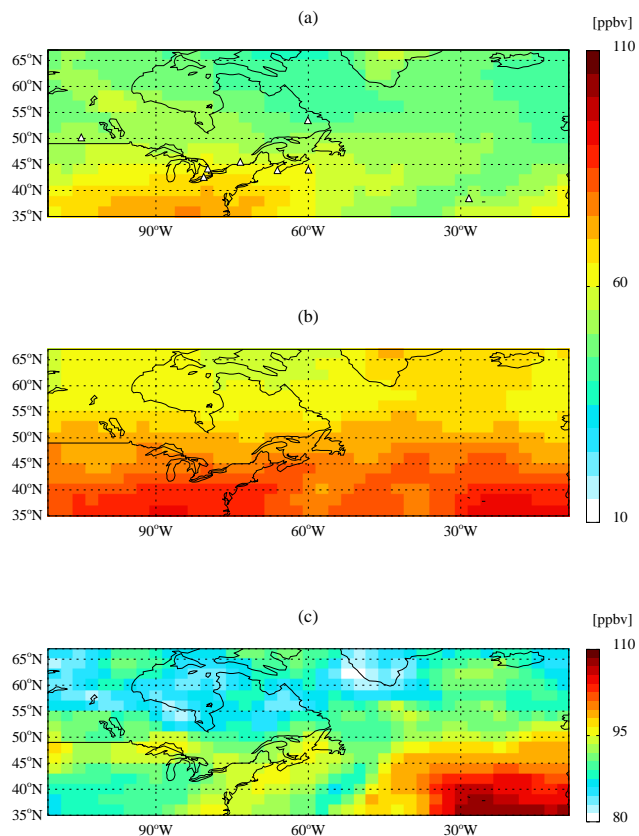


Fig. 2. Mean GEOS-Chem model ozone distribution (ppbv) across North America and the North Atlantic at **(a)** 750 hPa (2.2 km), **(b)** 500 hPa (5.5 km), and **(c)** 310 hPa (8.9 km) from 12 July to 4 August 2010. White triangles in plot **(a)** show the location of ozonesonde launch sites used during the BORTAS-A campaign across Canada, listed in Table 1, and the PICO-NARE observatory in the mid-Atlantic.

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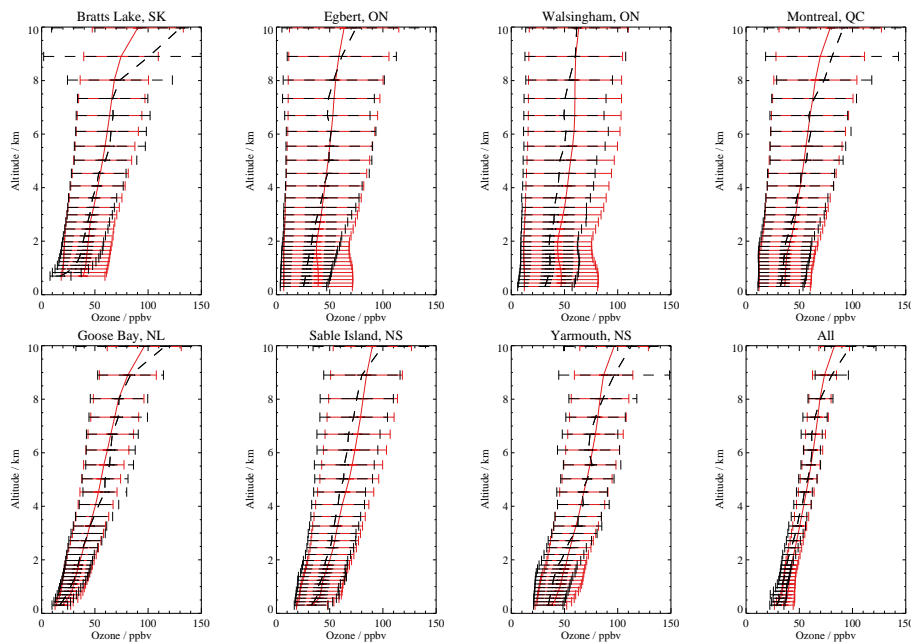


Fig. 3. Mean model and observed tropospheric ozone profiles over each of the ozonesonde launch sites listed in Table 1. Black dashed lines show the mean ozonesonde profile at each site and red solid lines show the mean spatially and temporally co-located GEOS-Chem ozone profile to each ozonesonde. Error bars show the 1- σ standard deviation. The bottom right plot is the mean ozone profile from all 153 ozonesondes launched throughout the campaign.

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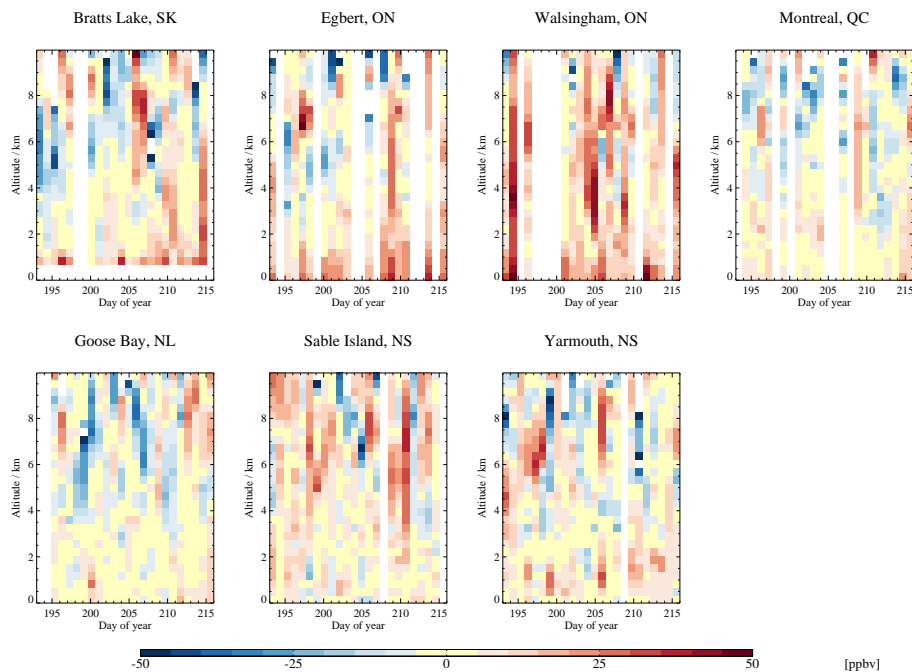


Fig. 4. Time-height cross-sections of the daily ozone differences between GEOS-Chem and ozonesonde profiles over each of the ozonesonde launch sites (Table 1) used during the BORTAS-A measurement campaign (12 July to 4 August 2010). White indicates no data present.

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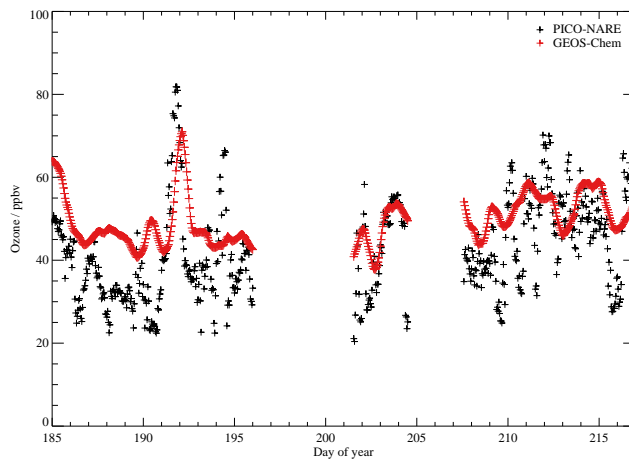


Fig. 5. Ozone time series measured at the PICO-NARE observatory (black symbols), and the co-located model ozone output (red symbols), from 4 July to 4 August 2010 inclusive.

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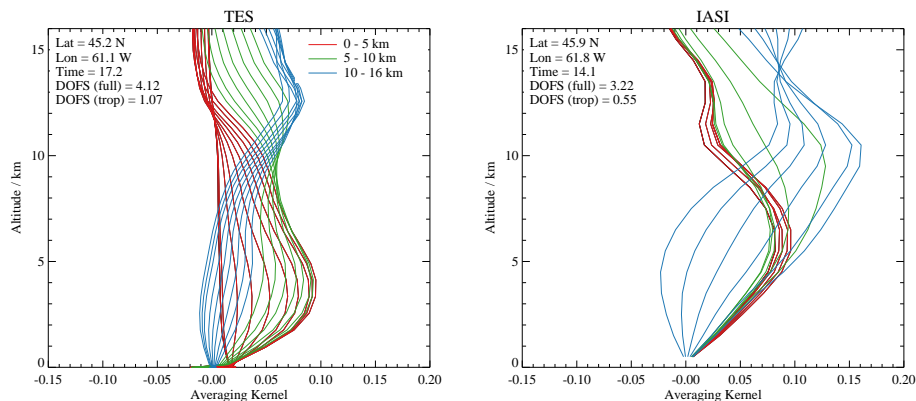


Fig. 6. Averaging kernels for ozone profile retrievals made by TES and IASI co-located to Halifax, NS on 16 July 2010.

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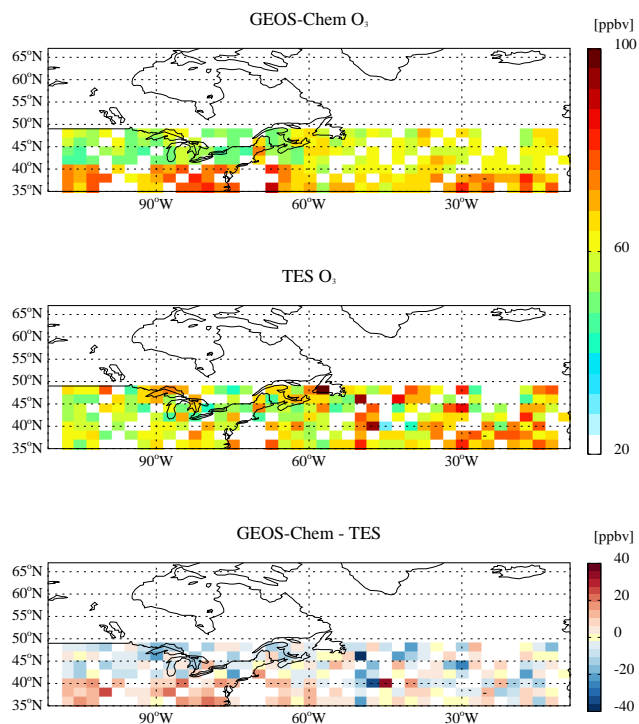


Fig. 7. Comparison of the mean mid-tropospheric (600–400 hPa) ozone distributions modelled by GEOS-Chem and observed by the Tropospheric Emission Spectrometer and the difference (model minus observed). The plots show the mean daytime ozone averaged over 12 July to 4 August 2010.

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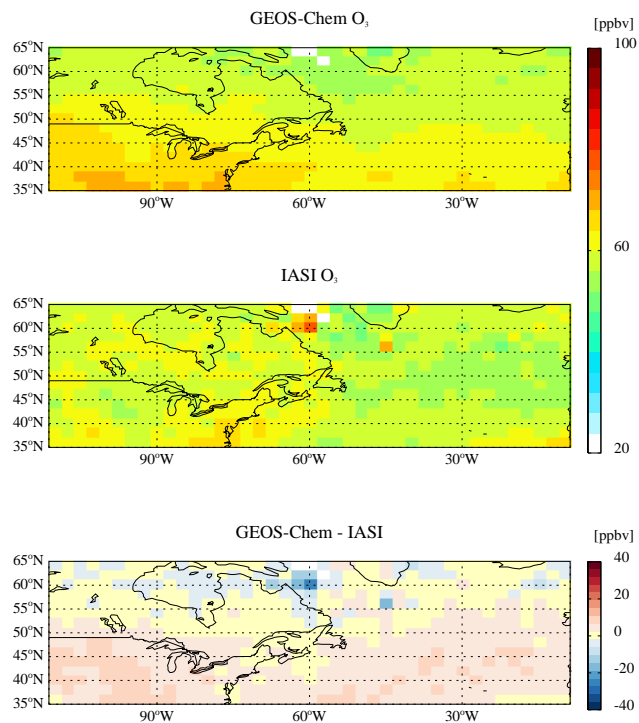


Fig. 8. Same as Fig. 7 for the ozone distribution observed by the Infrared Atmospheric Sounding Instrument.

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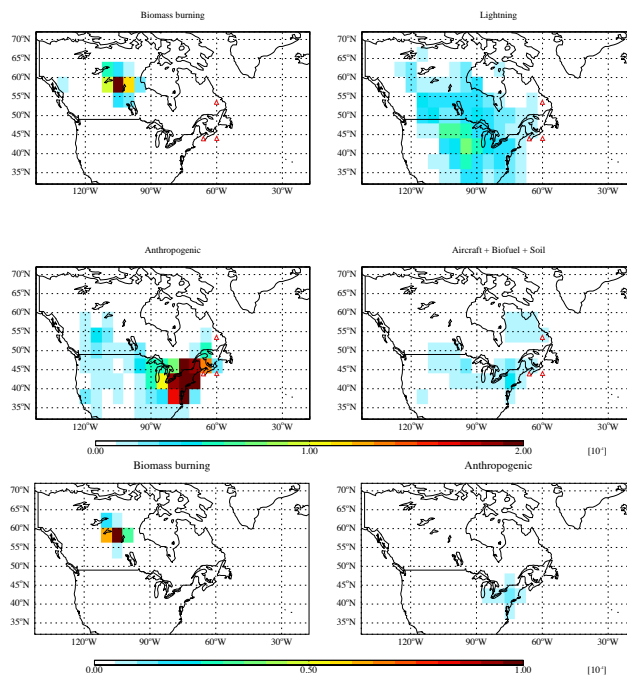


Fig. 9. Normalised cost function sensitivity of model free tropospheric (approximately 750–250 hPa) ozone profiles over ozonesonde launch sites in Maritime Canada to: **(a)** NO_x emissions estimates associated with biomass burning, lightning, anthropogenic, and combined aircraft/biofuel/soil sources; and **(b)** CO emissions estimates associated with biomass burning and anthropogenic sources over the BORTAS-A campaign period (12 July to 4 August 2010). The ozonesonde launch site locations (Goose Bay, Sable Island, and Yarmouth from Table 1) are denoted by red triangles in each plot.

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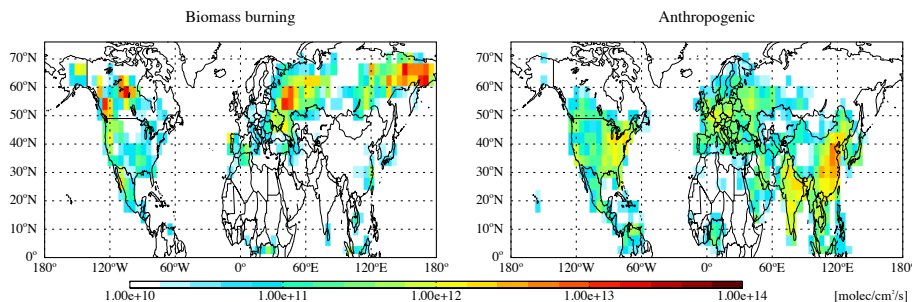


Fig. 10. Mean Northern Hemisphere CO emissions from biomass burning and anthropogenic sources ($\text{molec}^{-1} \text{cm}^{-2} \text{s}^{-1}$) averaged over 12 July to 4 August 2010. Biomass burning emissions are from the FLAMBE inventory (Reid et al., 2009). Anthropogenic emissions in the model are as those described in Sect. 3.

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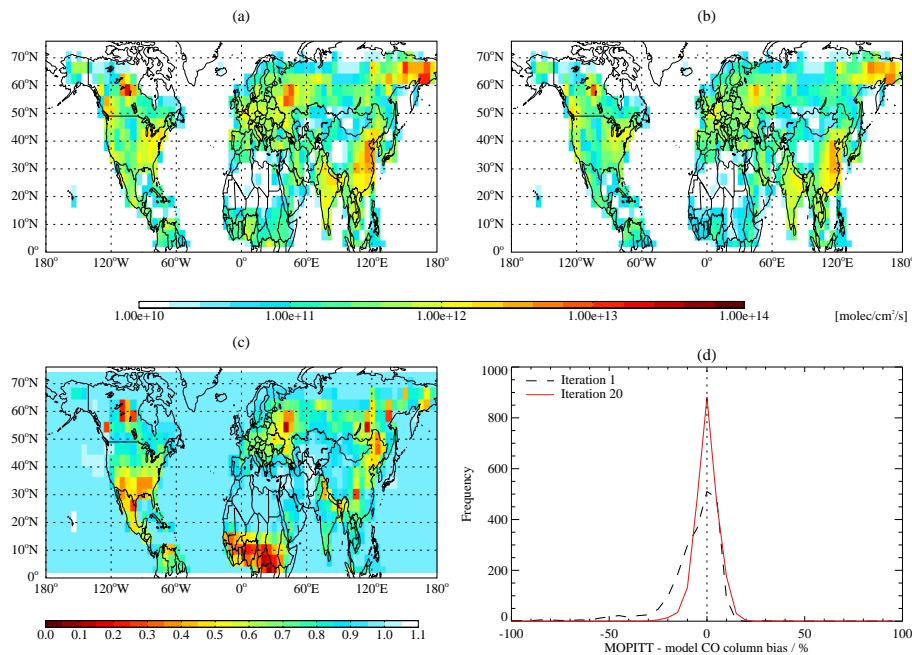


Fig. 11. Mean Northern Hemisphere a priori and a posteriori CO emissions from all sources and scaling factors derived from the assimilation of MOPITT V4 CO observations for July 2010. Plots **(a)** and **(b)** respectively show the a priori and a posteriori CO emissions and plot **(c)** shows the scaling factors. Plot **(d)** shows histograms of the mean MOPITT minus model biases averaged over the northern extra-tropics for the first (black dashed line) and twentieth (red solid line) iterations of the MOPITT inversion.

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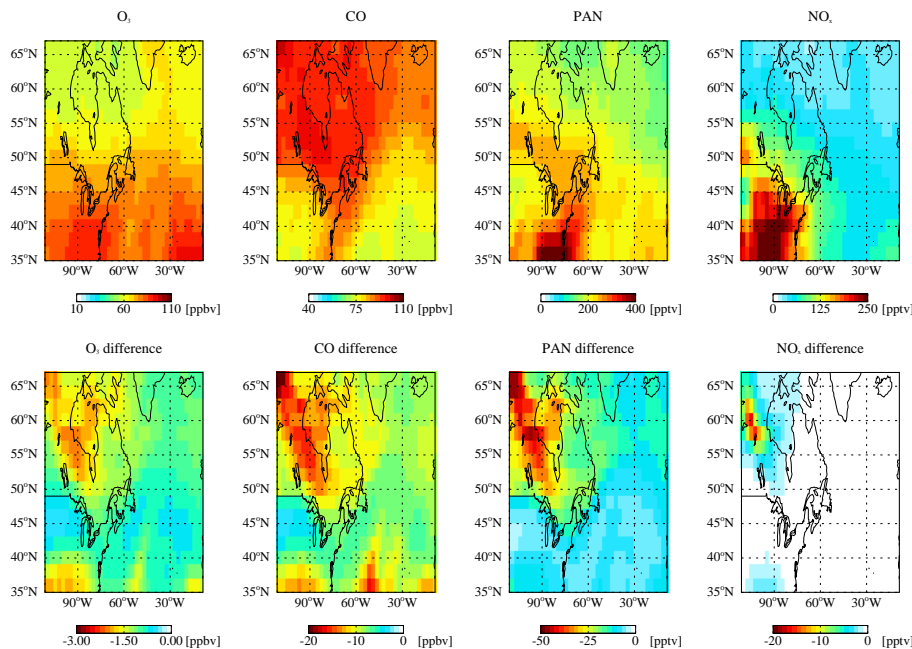


Fig. 12. Model ozone, CO, PAN, and NO_x distributions across North America and the North Atlantic at 500 hPa averaged over the BORTAS-A campaign period, 12 July to 4 August 2010. The top row shows the mean distributions from the model simulation with the scaled FLAMBE inventory derived from the MOPITT CO inversion and the absolute differences compared to the model outputs with the original FLAMBE emissions are shown in the bottom row.

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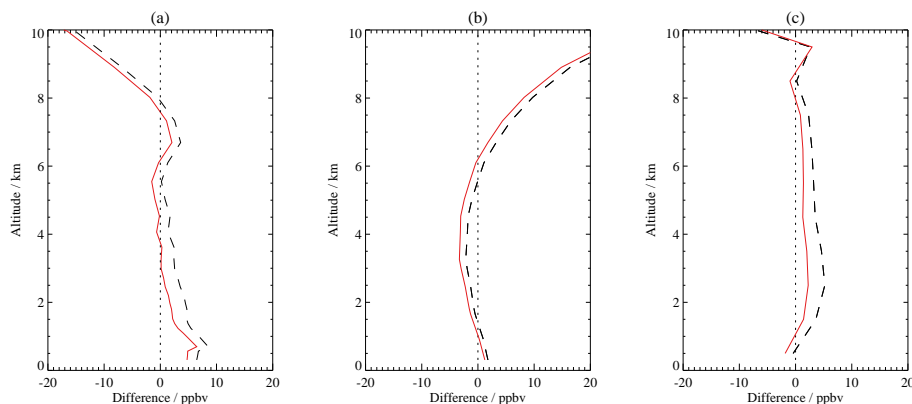


Fig. 13. Mean difference profiles (ppbv) of the model ozone output to **(a)** ozonesonde, **(b)** TES, and **(c)** IASI data averaged over the BORTAS-A campaign period (12 July to 4 August 2010). The model output is sampled at the time and location of the observed profiles for each dataset. Black dashed lines show the mean difference of the model output with the original FLAMBE inventory to the data and red solid lines show the difference of the model output with the scaled FLAMBE emissions derived from the MOPITT CO inversion.

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