We thank the reviewers for their thoughtful and detailed comments. Below we respond to the individual comments.

Reviewer #1

The authors make an interesting contribution to the quantification of CO surface emissions and of their trend over the past 15 years. I recommend its publication provided the following issues are addressed. Most of them are minor, but a couple of them deserve much more attention.

Thank you for your comments. Modifications have been made to improve this manuscript.

Q1: *l.* 80: The authors anticipate on their results, which is not really appropriate in an introduction (it breaks the logic flow).

Changed.

Q2: *l.* 97: measurement and model systematic errors can be damped but not suppressed.

Changed.

Q3: *l.* 98: "systematic biases" -> "systematic errors".

Changed.

Q4: *l.* 143: the previous example of the SCIAMACHY bias is time-dependent. The authors should explain why they think that the MOPITT bias does not vary much with time (mostly with the season).

The limited measurements provided by the HIPPO aircraft will result in uncertainties in the correction factors, which is more significant in the seasonal average than annual average. On the other hand, we are focusing on the interannual variation of CO emissions. The seasonal variation of CO emissions is not very important in this work. Consequently, we decided to use the annual mean correction factor. More description has been added.

Q5: *l.* 185: the authors seem to neglect the error statistics provided by the retrieval product. We can understand that they prefer raising them at 20% to be conservative, given likely systematic errors, but ignoring the vertical correlations is really surprising. This point is important because it bears most of the credibility of the following profile/lower profile inversion results vs. column inversion results. In addition, the ad-hoc uncorrelated observation budget used here is not internally consistent: when summing the profile level (error) covariances, one does not get the column (error) variance. This inconsistency basically suppresses the possibility to compare the two types of results meaningfully. Last, model errors are very likely correlated in the vertical and even uncertain large or medium vertical correlations (let us say 0.5 for instance) for this term of the observation error budget are better than the null correlations assumed here. A very good question! We have compared the discrepancies associated with two types of error covariance matrix in the preparation stage of this work: 1) diagonal matrix (this work); 2) full error covariance matrix including vertical correlation, based on MOPITT error covariance. Our results show that the difference in the scaling factors is small, perhaps due to the large amount of satellite measurements in our global scale inversion. Because we are focusing on the mitigation of effects of systematic errors, we used the diagonal matrix to keep consistency with our previous studies. However, as the reviewer indicated, a better description for the error covariance matrix is important. We will improve our methodology in our future study.

Q6: *l.* 186: the authors seem to combine combustion and VOC sources of CO together but later in Section 4.2 they show result by source type. They should explain how they split the information on the source type with simple column or profile retrievals of CO. In particular, I cannot see how VOC sources and their trends can be separated from the rest.

As the reviewer indicated, we cannot completely separate the a posteriori emission estimates from different sources. However, the various spatial and temporal distribution of emissions sources (e.g. anthropogenic vs. biomass burning) provides valuable information to distinguish the contribution from each category. In order to further isolate the influences of biomass burning, the months dominated by biomass burning (biomass burning CO > 50% of total CO emission in an individual grid) are excluded in the trend analysis for anthropogenic and VOC sources (Figure 5). More description has been added.

Q7: *l.* 215-216: This sentence (": : : indicated that regional inversions have more advantages than global inversions : : : better controlled") is unnecessarily polemical and may actually be wrong depending on how we understand "better controlled". There are pros and cons and the statement cannot leave the impression that the case has been closed.

Thank you for your suggestion! The statement has been changed.

Q8: *l.* 219: "model" -> "models".

Changed.

Q9: *l.* 228: the authors need to be clear that they do not use the same land data in the first and in the second step. Otherwise they would correlate boundary condition errors and observation errors in the second step and possibly induce weird side effects on their results (because those correlations are not accounted for).

I am sorry for the confusion. In the two-step approach:

Step 1: We directly modify CO concentrations using sequential Kalman filter assimilation. Both MOPITT data over land and ocean are used.

Step 2: We constrain CO emissions over land with MOPITT data over land only. The boundary condition is from step 1.

The objective of Step 1 is to provide the best global CO fields, based on MOPITT. We need to assimilate MOPITT data over land in the first step to keep the consistency between boundary conditions and emissions.

Q10: *l.* 254: Montzka et al. (2011) is recalled, but these authors wrote "Despite the much lower atmospheric CH3CCl3 mixing ratios in recent years ('13 ppt in 2007), they remained precisely measured through 2007. Precision for the analysis of CH3CCl3 (0.5 to 0.75% as repeatability) has remained comparable to the nearly constant (on a relative basis) standard deviation of paired flask means collected within a month at remote stations of 0.7 1.1% through 2007. Data after the end of 2007 are not included in this report owing to instrumental problems that developed in 2008." The present authors should give the same level of detail and clarify the fact that the instrumental problem does not affect their results.

The website (NOAA) shows: "NOAA flask data obtained by the GCMS for some compounds analyzed during the 2008.5-2009.5 period are subject to some small biases owing to instrumental issues during that period. Data obtained for CH3CCl3 during that time period, for example, should not be used for deriving hydroxyl radical concentrations"

According to Figure 4, we believe the influence of the instrumental problems (2008.5-2009.5) on our analysis (2001-2015) is small.

Q11: l. 276: "demonstrate" is too strong.

Changed.

Q12: 1. 296: there is also an initial increase in the measurements that should be commented.

The initial increase at 2001-2002 could be caused by uncertainties in the data. We are trying to avoid to make a conclusion about trend based on short (2 years) period data. A sentence has been added for this issue.

Q13: *l.* 313: this is only true for the profile results.

As shown in Table 1, an increase of Chinese emissions from 2001 to 2004 is shown by all three analyses.

Q14: *l.* 335: large PBL height errors happen everywhere over the globe. Why should they just affect India and SE Asia?

Thank you for pointing out this issue. We have removed "PBL height" in the discussion.

Q15: *l.* 374: these 2014 and 2015 studies are not "more recent" than Field et al.(2016). Actually, the authors could discuss the "more recent" study of Yin et al. (2016) that seems to well overlap with their approach.

The discussion has been changed. We didn't cite Yin's work here, because we hope to demonstrate the consistency between our inversion results with studies using different approach (not an inverse modelling).

Q16: l. 376: extra comma.

Changed.

Q17: *l.* 396: the above-mentioned issue in the observation error statistics is also a likely explanation.

The lower tropospheric profile data includes the lowest three levels (1000hPa, 900hPa and 800 hPa). The influence of correlation of these three levels should be small.

Q18: *l.* 464: to be fair and consistent with the second part of the sentence, the authors should also speak of an update about this question, since it has been (imperfectly) addressed before.

Changed.

Q19: References should be ordered.

Changed.

Reviewer #2

The work investigates the possible cause of the observed trend of a reduction of Carbon Mononixde (CO) emissions over the last 15 years over the northern hemisphere and parts of China. This trend is somewhat mitigated by an increased trend of CO emissions over India. The authors use global MOPITT remote sensing data of CO in the thermal infrared region to constrain model forecasts of CO concentrations and surface emissions. The model being used is the adjoint of the off-line global chemistry transport model GEOS-Chem.

The authors make 4 big assumptions: 1) Unknown model biases can be handled with by providing independent boundary conditions of CO concentrations over oceans each month from a Kalman Filter inversion run, 2) Local continental scale emissions can be estimated then by a 4dvar method constrained by MOPITT observations over land (and constrained by the boundary conditions of CO concentrations over the oceans), 3) The inversion system works best by removing a latitudional bias in MOPITT retrievals as derived from the HIAPER Pole to Pole Observations campaign (HIPPO), 4) The hydroxyl radical (OH) variability cannot explain the decrease in CO emissions if we put trust in the method of using MCF (methyl chloroform) measurements as a proxy for estimating atmospheric OH concentration change.

Thank you for your comments. Modifications have been made to improve this manuscript.

Q1: *Chapter: 2.1 MOPITT: Did you do any data thinning on the MOPITT data and how did you screen the MOPITT data?*

We employed the same data quality control as our previous studies. Detailed description has been added in Section 2.1.

Q2: line 176-178: You need to describe the 4dvar adjoint method in more detail. What are typical numbers of N and it is not clear from the equation (line 178) or Figure 3 how you defined the length of the assimilation window in your 4dvar system. In GEOS-Chem met fields are typically updated every 6 hours – does this also correspond to your assimilation window (e.g. 6 hour window)? Or is your assimilation window a full month and observations are sampled every hour?

Thank you for your suggestion! More description has been added.

In order to match model output, the high resolution MOPITT measurements are averaged temporally (one-hour resolution) and spatially ($4^{\circ}x5^{\circ}$ resolution) to produce grid mean observations. The length of assimilation window is one month. The number (*N*) of grid mean observations in one month is around 10000.

Q3: line 186-189: Cite: D.B.Jones, et al: Potential of observations from the Tropospheric Emission Spectrometer to constrain continental sources of carbon monoxide, doi:10.1029/2003JD003702, J. Geopys. Res, 2003

It is not clear to me why the authors cannot follow the method of constructing the observation error covariance matrix as outlined in the above paper (Dylan et al 2003). Of course TES and MOPITT are different products but as far as I remember MOPITT will also let you construct a retrieval error matrix as part of their released data products (they come with the averaging kernels). It is true that there is some vertical correlation in the averaging kernels but cannot account for the information loss of a uniform or flat construed observation error.

Jones et al. (2003) used the NMC method to assess the model transport errors. This approach uses pairs of model forecasts, of different length, but which are valid for the same time, to characterize the model errors. We do not have such forecasts available during this analysis period.

We have compared the discrepancies associated with two types of error covariance matrix in the preparation stage of this work: 1) diagonal matrix (this work); 2) full error covariance matrix including vertical correlation, based on MOPITT error covariance. Our results show that the difference in the scaling factors is small, perhaps due to the large amount of satellite measurements in our global scale inversion. Because we are focusing on the mitigation of effects of systematic errors, we used the diagonal matrix to keep consistency with our previous studies. However, as the reviewer indicated, a better description for the error covariance matrix is important. We will improve our methodology in our future study.

Q4: 190-196: Reword and emphasise that posterior emissions estimates (e.g. Table 1) do not have uncertainty reduction error bars because of the way the adjoint method works and ask Daven Henze if there is a reference for that.

Thank you for your suggestion! The discussion has been modified.

Q5: line 194-196: "As shown by Heald et al (2004), different assumptions about the inversion configuration can produce differences in the source estimates that are significantly larger than the a posteriori errors." Is this statement related to the bias correction in the next paragraph (line 197-209)? Why is this important here?

We hope to demonstrate that the actual a posteriori uncertainty (including systematic errors) is much larger than the a posteriori uncertainty calculated based on Gaussian assumption (random errors).

Q6: line 197-198: "Removing the bias in initial conditions is essential for inverse analysis, and can be performed with various data assimilation techniques." Have you got a reference for this? I have heard people claiming (I am not one of them) that in a good inversion system there is no bias correction needed. Have you tested your system without bias correction?

We have tested the effects of initial condition in our previous study. As shown in Figure 4a of Jiang et al. (2013), there are large discrepancies between MOPITT and original model simulation due to the accumulation of model errors prior to the assimilation window. We cannot use the biased initial condition for the inverse analysis.

"a good inversion system there is no bias correction needed" is valid for the ideal condition. However, there are always systematic biases, and we cannot ignore them. For example, Figure 1 shows noticeable discrepancies between MOPITT and HIPPO. We have to mitigate these discrepancies using latitude dependent correction factors, although we know the best approach is an update of retrieval algorithm.

Q7: line 218-220: "They demonstrated that the systematic bias associated with North American CO emissions due to OH distribution can be reduced by up to 50% with optimised boundary conditions. Similar optimisation on the boundary condition can also be employed in global model, for example, Pfister et al. (2005) constrained biomass burning CO emissions from boreal North America with optimised CO fields outside the impacted region."

How does this relate to your work? Your are using pre-calculated OH fields from a full chemistry run. Are you making the point here that the influence of the badly understood

OH bias can be reduced by optimised CO 3D boundary conditions (e.g. from your Kalman Filter at the beginning of each month)? Please clarify.

We hope to demonstrate that the influences of systematic errors can be mitigated by the optimization on the boundary condition. We have changed the statement to make it more concise.

As the reviewer indicated, the optimization on the boundary conditions (e.g. around North America) can really mitigate the influences of OH bias on a posteriori estimation of North American CO emissions. Although the OH distribution over North America continent is still biased in a reginal inversion, the adverse effects of biased OH distribution on the CO inflow from outside of North America can be significantly reduced.

Q8: Figure 3: This needs clarification in the Figure caption or text. If I am right to assume that your Kalman filter runs from 1_{st} of March until 31nd December first and is completely independet of the 4dvar inversion in the assimilation window? And there is no feedback of the 4dvar inversion results to the boundary conditions of the following months?

Thank you for your suggestion! The Figure caption has been changed.

Q9: 4.1 Long-term variation of global tropospheric OH. Krol et al. found a somewhat different result of OH trends based on MCF measurements and model studies. Admittedly for a different study period (1978-1998). You could (or should) cite that paper: M. Krol et al., 1998: Global OH trend inferred from methylchloroform measurements, 103, p.10,697–10,711, 1998, J. Geopys. Res.

The citation has been added.

Q10: 4.2 Long-term variation of global CO emissions. It would be a good idea if you split the section into different smaller subsections: 4.2.1 Emission;s US 4.2.2 Emissions EU; 4.2.3 Emissions India + South East Asia; 4.2.4 Biomas Burning Emissions etc.

Thank you for your suggestion! Two subsections "Regional analysis for anthropogenic emissions" and "Regional analysis for biomass burning emissions" have been added.

Q11: line 425-427: "In a recent study, Schnell et al. 92015) evaluate surface O3 concentrations simulated by multi-models for North America and Europe. They found most models can provide good simulations for the patterns of O3 but cannot reproduce the magnitude." I do not think citing an ozone study supports your argument in terms of CO.

This citation has been removed.

Q12: *line 466-468. Reformulate the part including 'MCF'. I do not think you have used MCF to 'evaluate changes in the sources and sinks of atmospheric CO ... '.*

The statement has been modified.

Q13: Table 1. Add a fifth column of global total posterior emissions to the 3 individual sub tables: 'MOPITT Columns (Tg/year)', 'MOPITT Profile (Tg/year)' and 'MOPITT Lower Profile (Tg/year)'.

Add a sixth column to the 3 individual sub tables for posterior CH4 and VOC production. Also append 4 single columns for the global prior emissions in each year. e.g. Year, US, EU, China, India, CH4, VOC, US, EU, China, India, CH4, VOC, US, EU, China, India CH4, VOC, PRIOR ANTHRO, PRIOR CH4, PRIOR VOC, PRIOR, TOTAL, And comment on these global budgets in the main text.

Thank you for your suggestion! Three columns of global total anthropogenic emissions have been added in Table 1. A new table (Table 2) was added to show the annual variation of biomass burning emissions.

We didn't provide values for CO sources from VOC and CH₄ oxidization because our results for these two sources are inconclusive. The values for a priori emissions are also excluded because the tables are already complex.

Q14: Figure 11. I am not convinced your method of singling out the meteorological effects works as intented. Firstly, what exactly is being defined as meteorological conditions? I think the accumulation of surface CO, especially over the tropical regions and to a lesser extened the slight loss of CO at higher latitudes is an artifact and CO builds up, unrealistically, in GEOS-Chem tagged tracer mode. I am not asking you to conduct more model calculations. However, it would have beend interesting to see if a full global 4x5 GEOS-Chem CO chemistry run gives a similar answer than Figure 11a and 11b.

A very good question! Our forward model simulation, based on various versions of the meteorological fields (i.e. GEOS-4, GEOS-5 and GEOS-FP), is not an ideal tool for the analysis of influences of meteorological fields. We have modified the text to emphasize on this point.

1 2	A fifteen year record of CO emissions constrained by MOPITT CO observations
3	
4 5 6	Zhe Jiang ^{1,2} , John R. Worden ¹ , Helen Worden ² , Merritt Deeter ² , Dylan B. A. Jones ³ , Avelino F. Arellano ⁴ , Daven K. Henze ⁵
7 8 9 10 11 12 13 14 15	 ¹Jet Propulsion Laboratory, California Institute of Technology, Pasadena, CA, USA ²National Center for Atmospheric Research, Boulder, CO, USA ³Department of Physics, University of Toronto, Toronto, ON, Canada ⁴Department of Hydrology and Atmospheric Sciences, University of Arizona, Tucson, AZ, USA ⁵Department of Mechanical Engineering, University of Colorado, Boulder, CO, USA
16	
17	
18	
19	
20	
21	
22	
23	
24	
25	
26	
27	
28	

29 Abstract

30 Long-term measurements from satellites and surface stations have demonstrated a 31 decreasing trend of tropospheric carbon monoxide (CO) in the Northern Hemisphere over the past decade. Likely explanations for this decrease include changes in anthropogenic, fires, and/or 32 33 biogenic emissions or changes in the primary chemical sink hydroxyl radical (OH). Using 34 remotely sensed CO measurements from the Measurement of Pollution in the Troposphere 35 (MOPITT) satellite instrument, in-situ methyl chloroform (MCF) measurements from World Data 36 Centre for Greenhouse Gases (WDCGG), and the adjoint of the GEOS-Chem model, we estimate 37 the change in global CO emissions from 2001-2015. We show that the loss rate of MCF varies by 38 0.2% in the past 15 years, indicating that changes in global OH distributions do not explain the 39 recent decrease in CO. Our two-step inversion approach for estimating CO emissions is intended 40 to mitigate the effect of bias errors in the MOPITT data as well as model errors in transport and 41 chemistry, which are the primary uncertainties when quantifying CO emissions using these 42 remotely sensed data. Our results confirm that the decreasing trend of tropospheric CO in the 43 Northern Hemisphere is due to decreasing CO emissions from anthropogenic and biomass burning 44 sources. In particular, we find decreasing CO emissions from the United States and China in the 45 past 15 years, unchanged anthropogenic CO emissions from Europe since 2008, We find decreasing trends of biomass burning CO emissions from boreal North America, boreal Asia and 46 47 South America, but little change over Africa. In contrast to prior results we find positive trend in 48 CO emissions is likely for India and southeast Asia.

Deleted:, and likely a positive trend from India and southeast Asia, in contrast to recently reported results.

Deleted: The inconsistency between our analysis with recent study suggests more efforts are needed for robust conclusion about the variation of anthropogenic Deleted: Southeast

50 1. Introduction

49

51 Tropospheric CO is a product of incomplete combustion and a byproduct of the oxidation

58 of hydrocarbons. It plays a key role in atmospheric chemistry because it is the main sink for OH, 59 and an important precursor for tropospheric ozone (O_3) . Recent studies demonstrated significant 60 change in tropospheric CO abundance in the past decade. Using Atmospheric Infrared Sounder 61 (AIRS) CO measurements, Warner et al. (2013) indicated that Northern Hemispheric CO mixing 62 ratio decreased by 1.28 ppb/year in the period of 2003-2012. Worden et al. (2013) demonstrated Northern Hemispheric CO column measurements from MOPITT show a decrease of ~0.92%/year 63 64 in the period of 2000-2011. Using observations from Mt. Bachelor Observatory, Gratz et al. (2015) 65 also show a negative trend of CO concentration by 1.9%/year in the period of 2004-2013. 66 However, the reason for the large variation of tropospheric CO abundance is still unclear; for 67 example, Strode et al. (2016) found decreases in modeled CO abundance over North America and 68 Europe, but increases over China, based on bottom-up emissions.

69 There is currently much effort focused on accurately quantifying emissions of CO. For 70 fossil fuels and biofuels, energy consumption statistics and emission factors are usually used to 71 construct the emission inventories (e.g. Streets et al. 2006; Ohara et al. 2007; Zhang et al. 2009; 72 Zhao et al. 2012). Biomass burning emissions are commonly calculated as the product of burned 73 area, fuel loads, combustion completeness and emission factors (e.g. van der Werf et al. 2006, 74 2010; van Leeuwen and van der Werf 2011). Because of the large uncertainties in the emission 75 inventories, space-based remotely sensed measurements and surface/aircraft in-situ observations have been assimilated to provide "top-down" constraints on CO emissions (e.g., Arellano et al., 76 77 2006; Chevallier et al. 2009; Jones et al., 2009; Kopacz et al., 2010; Jiang et al., 2011; Fortems-Cheiney et al. 2011; Hooghiemstra et al. 2012; Miyazaki et al. 2015). In a recent study, Yin et al. 78 (2015) constrained global CO emissions for the period 2002-2011 to investigate the possible 79 80 reasons for the decreasing CO abundance in the Northern Hemisphere. Using MOPITT column

Deleted: The budget of tropospheric CO is determined by its sources and sinks.

83	data (version 6J) over the whole globe, Yin et al. (2015) indicate that the negative trend in the		
84	Northern Hemisphere is driven by decreasing anthropogenic emissions from North America,		
85	Europe and China,	D	Deleted: , similar to our result.
86	The major sink of tropospheric CO is OH. Because of its high variability and short lifetime		
87	(about one second), it is difficult to assess the spatial and temporal variation of global OH through		
88	direct measurements (Spivakovsky et al. 2000; Lelieveld et al. 2004). Alternatively, Montzka et		
89	al. (2011) demonstrated small interannual variability of global OH for the period 1997-2007 by		
90	using the loss rate of MCF as a proxy. The measurements of MCF are assimilated in recent CO		
91	inversion studies to provide updated OH (e.g. Fortems-Cheiney et al. 2011, 2012; Yin et al. 2015),		
92	but the estimates are adversely affected by the sparse distribution of measurements.		
93	The objective of this work is to investigate the dominant reasons for the decreasing CO		
94	trend in the Northern Hemisphere, and to provide updated CO emission estimates for model studies.		
95	Using methods and results from our prior work, our approach for estimating emissions is intended	D	Deleted: Our
96	to reduce the effects of model errors of transport and chemistry, as well as bias errors in the data,		
97	on our conclusions about CO emissions; these are the primary uncertainties that affect CO		
98	emissions estimates. For example, bias errors as a function of latitude in MOPITT data can have a		
99	substantial impact on emissions estimates (Deeter et al., 2014). Model errors of transport and		
100	chemistry will have variable and substantial effects on CO emissions in different parts of the globe		
101	due to seasonal and latitudinal variations in convection, advection, and boundary layer height		
102	(Jiang et al., 2013, 2015a, 2015b).		
103	In order to <u>reduce</u> the influences from these measurement and model transport systematic		Deleted: suppress
104	errors, we performed a two-step inversion by combining sequential Kalman Filter (Jiang et al.	D	Deleted: systematic
105	2013 2015a 2015b) with four-dimensional variational (4D-Var) assimilation (Henze et al. 2007)		

in this work, using the GEOS-Chem model. Instead of optimizing the CO concentrations and 111 112 emissions simultaneously (e.g. Fortems-Cheiney et al. 2011, 2012; Yin et al. 2015), our first step, 113 the sequential Kalman Filter, modifies the atmospheric CO concentration directly to provide low 114 bias initial (monthly) and boundary (hourly) conditions, whereas the second step (4D-Var) 115 constrains CO emissions assuming perfect initial and boundary conditions. We also apply bias 116 corrections to MOPITT and compare the surface CO concentrations obtained by constraining the 117 model with either MOPITT profile, total column, or lower troposphere data to test which data type 118 provides the most accurate comparison with independent surface in-situ measurements.

This paper is organized as follows: in Section 2 we describe the MOPITT instruments and the GEOS-Chem model used in this work. In Section 3 we outline the inverse method. We then investigate the long-term variations of global tropospheric OH and CO emissions in Section 4, and we discuss the changes in tropospheric CO, and the contributions from emissions and meteorological conditions. Our conclusions follow in Section 5.

124 2. Observations and Model

125 **2.1. MOPITT**

126 The MOPITT instrument was launched on December 18, 1999 on the NASA/Terra 127 spacecraft. The satellite is in a sun-synchronous polar orbit of 705 km and crosses the equator at 128 10:30 local time. The instrument makes measurements in a 612 km cross-track scan with a 129 footprint of 22 km x 22 km, and provides global coverage every three days. The MOPITT data 130 used here were obtained from the joint (J) retrieval (V6J) of CO from TIR (4.7µm) and NIR (2.3µm) 131 radiances using an optimal estimation approach (Worden et al., 2010; Deeter et al., 2011). The 132 retrieved volume mixing ratios (VMR) are reported as layer averages of 10 pressure levels (surface, 900, 800, 700, 600, 500, 400, 300, 200 and 100 hPa). The relationship between the retrieved CO 133

134 profile and the true atmospheric state can be described as:

135
$$\hat{z} = z_a + A(z - z_a) + G\epsilon \tag{1}$$

where z_a is the MOPITT a priori CO profile, z is the true atmospheric state, Ge describes the 136 retrieval error, and $A = \partial \hat{z} / \partial z$ is the MOPITT averaging kernel matrix, which gives the sensitivity 137 138 of the retrieval to the actual CO in the atmosphere. The MOPITT V6 data have been evaluated by 139 Deeter et al. (2014) using aircraft measurements from HIAPER Pole-to-Pole Observations (HIPPO) 140 and the National Oceanic and Atmospheric Administration (NOAA). For the TIR/NIR multi-141 spectral retrievals, they found negative bias drift (-1.27%/year) at lower troposphere (800 hPa), 142 and positive bias drift (1.64%/year) at upper troposphere (200 hPa). The bias drift in the total 143 column is negligible (0.003%/year). Following our previous studies (Jiang et al. 2013; 2015a; 2015b), we reject MOPITT data with CO column amounts less than 5x10¹⁷ molec/cm² and with 144 145 low cloud observations. The threshold of 5×10^{17} molec/cm² was selected to prevent unrealistically 146 low CO columns from adversely impacting the inversion analyses. Since the NIR radiances 147 measure reflected solar radiation, only daytime data are considered here. 148 Figure 1 shows the comparison between MOPITT CO retrievals and HIPPO aircraft

149 measurements. The aircraft measurements are smoothed with MOPITT averaging kernels. The 150 comparison demonstrates a negative bias of MOPITT CO retrievals in the tropics and a positive 151 bias at the middle latitudes in the lower troposphere. Opposite bias is observed in the upper 152 troposphere. Similar latitude dependent biases in remote sensing retrievals have been revealed for 153 methane (CH₄) observations from Scanning Imaging Absorption Spectrometer for Atmospheric 154 Chartography (SCIAMACHY, Bergamaschi et al. 2007, 2009; Meirink et al. 2008), Greenhouse 155 Gases Observing Satellite (GOSAT, Turner et al. 2015), and CO observation from MOPITT (version 4, Hooghiemstra et al. 2012). Similar to previous studies, we reduce the adverse effect of 156

157	the latitude dependent bias by applying latitude dependent correction factors to MOPITT CO
158	retrievals, based on the black solid line in Figure 1, which represents a 4-order polynomial curve
159	fitting (in a least-squares sense) for all data points. It should be noted that the possible seasonal
160	variations of MOPITT retrieval biases are not included in our analysis because we are focusing on
161	the interannual variation of CO emissions.

162 2.2. GEOS-Chem

163 The GEOS-Chem global chemical transport model (CTM) [www.geos-chem.org] is driven 164 by assimilated meteorological fields from the NASA Goddard Earth Observing System (GEOS-5) at the Global Modeling and data Assimilation Office. For the simulations in this work, various 165 166 versions of GEOS meteorological fields are used, including GEOS-4 (2000-2003), GEOS-5 (2004-2012) and GEOS-FP (2013-2015). We use version v35j of the GEOS-Chem adjoint, which is based 167 168 on v8-02-01 of the forward GEOS-Chem model, with relevant updates through v9-02-01. Our analysis is conducted at a horizontal resolution of 4°x5° with 47 vertical levels and employs the 169 170 CO-only simulation in GEOS-Chem, which uses archived monthly OH fields from the full 171 chemistry simulation. The OH fields used in this work are from GEOS-Chem version v5-07-08, 172 with a global annual mean OH concentration of 0.99x10⁶ molec/cm³ (Evans et al. 2005). The 173 potential long-term variation of global tropospheric OH is evaluated in section 4.

The global anthropogenic emission inventory is from EDGAR 3.2FT2000 (Olivier et al., 2001), but are replaced by the following regional emission inventories: the US Environmental Protection Agency National Emission Inventory (NEI) for 2008 in North America, the Criteria Air Contaminants (CAC) inventory for Canada, the Big Bend Regional Aerosol and Visibility Observational (BRAVO) Study Emissions Inventory for Mexico (Kuhns et al. 2003), the Cooperative Program for Monitoring and Evaluation of the Long-range Transmission of Air

180 Pollutants in Europe (EMEP) inventory for Europe in 2000 (Vestreng et al. 2002) and the INTEX-181 B Asia emissions inventory for 2006 (Zhang et al. 2009). Biomass burning emissions are based on 182 the Global Fire Emission Database (GFED3, van der Werf et al. 2010). The a priori biomass 183 burning emissions in Sep-Nov 2006 were applied to Sep-Nov 2015 over Indonesia. Additional CO 184 sources come from oxidation of methane and biogenic volatile organic compounds (VOCs) as 185 described in previous studies (Kopacz et al. 2010; Jiang et al. 2013). The biogenic emissions are simulated using the Model of Emissions of Gases and Aerosols from Nature, version 2.0 186 (MEGANv2.0, Guenther et al. 2006). The distribution of the annual mean CO emissions for 2001-187 2015 is shown in Figure 2. The annual global sources are 892 Tg CO from fossil fuel, biofuel and 188 189 biomass burning, 623 Tg CO from the oxidation of biogenic VOCs, and 876 Tg CO from the 190 oxidation of CH₄.

191 **3. Inversion Approach**

We use the 4D-var data assimilation system in GEOS-Chem (Henze et al. 2007) to constrain the CO sources. In this approach, we minimize the cost function defined as:

194	$J(x) = \sum_{i=1}^{N} (F_i(x) - z_i)^T S_{\Sigma}^{-1} (F_i(x) - z_i) + (x - x_a)^T S_a^{-1} (x - x_a) $ [Formatted: Right]
195	where x is the state vector of CO emissions, N is the number of MOPITT observations that are
196	distributed in time over the assimilation period, z_i is a given MOPITT measurement, and $F(x)$ is
197	the forward model. The temporal resolution of forward model output $(F(x))$ is one hour, and
198	consequently, the high resolution MOPITT measurements are averaged temporally (one-hour
199	resolution) and spatially (4°x5° resolution) to produce grid mean observations. The number (N) of
200	grid mean observations in our assimilation window (one month) is around 10000.
201	The error estimates are assumed to be Gaussian, and are given by S_{Σ} , the observational Formatted: Indent: First line: 0.5"
202	error covariance matrix, and S_a , the a priori error covariance matrix, respectively. The Gaussian

203	assumption excludes important systematic errors, such as biases in OH distribution, long-range	
204	transport and satellite retrievals in the cost function. Due to lack of meaningful information about	
205	the systematic errors, we assume a uniform observation error of 20% without spatial correlation.	
206	The combustion CO sources (fossil fuel, biofuel and biomass burning) and the oxidation source	
207	from biogenic VOCs are combined together, assuming a 50% uniform a priori error. We optimize	
208	the source of CO from the oxidation of CH_4 separately as an aggregated global source, assuming	
209	an a priori uncertainty of 25%.	
210	Without consideration of systematic errors, the a posteriori error covariance matrix is the	 Deleted: Because
211	inverse of the Hessian matrix, which is not stored in the 4D-var optimization scheme, Bousserez	 Deleted: does not store the full Hessian matrix, we do not
212	et al. (2015) presented an approach to construct the a posteriori error covariance matrix, using the	 Deleted: , which is
213	approximation of Hessian matrix. As opposed to earlier studies using surface measurements, the	Deleted: inverse
214	high spatial density of measurements from satellite instruments can effectively suppress the	Deleted: the
215	contribution from random errors in the cost function, leaving systematic errors as the critical factor	
216	in the uncertainty. As shown by Heald et al. (2004), different assumptions about the inversion	
217	configuration (systematic errors) can produce differences in the source estimates that are	
218	significantly larger than the a posteriori errors, calculated based on random errors. Consequently,	 Deleted:
219	estimates of a posteriori uncertainties are not provided in this work (e.g. Table 1 and Table 2).	
220	Removing the bias in initial conditions is essential for inverse analysis (Jiang et al. 2013),	 Deleted: ,
221	and can be performed with various data assimilation techniques. Model simulations driven by	
222	optimized emissions can provide good initial conditions (e.g. Gonzi et al. 2011; Bruhwiler et al.	
223	2014; Deng et al. 2014; Houweling et al. 2014). Alternatively, tracer concentrations can be	
224	modified directly to avoid the effect from long-range transport error (e.g. Kopacz et al. 2009; Jiang	
225	et al. 2013, 2015a). There are also efforts to optimize emissions and concentrations simultaneously	

(e.g. Fortems-Cheiney et al. 2011, 2012; Bergamaschi et al. 2013; Yin et al. 2015), however, the
contributions from emissions and concentrations to model bias may be hard to be distinguished.
Figure 3 shows the methodology of our assimilation system. Following our previous studies (Jiang
et al. 2013, 2015a, 2015b), we produce initial conditions at the beginning of each monthly
assimilation window by assimilating MOPITT data using a sequential Kalman filter. For the results
presented here, the Kalman filter assimilation was carried out from March 1, 2000 to December
31, 2015.

240 Systematic errors have critical influences on inverse analysis. Jiang et al. (2013) found that 241 the modeled CO concentration from a 10-day forecast simulation have large discrepancy with 242 assimilated CO fields, because of bias in model convective transport. Jiang et al. (2015a) 243 demonstrated that free tropospheric CO is more susceptible to the influences of OH bias than lower 244 tropospheric CO due to the process of long-range transport. Previous studies suggest the influences 245 of systematic errors can be mitigated by enhancing the contributions from local emissions to the 246 discrepancy between model and data, while keeping the influence from long-range transport as 247 low as possible due to sources of uncertainties that are difficult to quantify. For example, Pifster 248 et al. (2005) constrained biomass burning CO emissions from boreal North America with 249 optimized CO fields outside the impacted region; Jiang et al. (2015b) indicated that the results of 250 regional inversions are more reliable when the boundary conditions are optimized. 251 In this work, we designed a two-step inversion to reduce the effects of these systematic 252 errors. As shown in Figure 3, we define the ocean scene (red grids) as boundary conditions. In the 253 first step of our inverse analysis, sequential Kalman filter assimilation, we directly modify CO

254 concentrations without any change to emissions in order to provide an optimized CO fields as

255 consistent as possible with MOPITT. In the second step, the optimized CO fields are used to

Deleted: On the other hand, Jiang et al. (2015b) indicated that regional inversions have more advantages than global inversions because the boundary conditions can be better controlled. They demonstrated that the systematic bias associated with North American CO emissions due to OH distribution can be reduced by up to 50% with optimized boundary conditions. Similar optimization on the boundary condition can also be employed in global model, for example, Pifster et al. (2005) constrained biomass burning CO emissions from boreal North America with optimized CO fields outside the impacted region.

Deleted: (e.g. emission uncertainty in the upstream continent, uncertainties in the chemical sink and convective transport in the transport pathway), that are difficult to quantify.

rewrite CO concentrations over the ocean every hour, while 4D-var inversion is employed to constrain CO emissions, without any change on CO distribution over ocean. Only MOPITT data over land (white grids) were assimilated to constrain CO emissions in the second step. With the fixed/optimized boundary conditions, the global inversion system has been converted to a combination of several regional inversions. Consequently, the emission and transport errors from one continent (e.g. North America) will not affect the emission estimation of another continent (e.g. Europe).

279 4. Results and Discussion

280 4.1. Long-term variation of global tropospheric OH

281 The distribution of tropospheric OH has significant influence on the inverse analysis of CO 282 emissions (Jiang et al. 2011). Various approaches have been employed to improve the OH 283 distribution in previous studies. Jiang et al. (2013) assimilated MOPITT CO retrievals in full chemistry model simulation to provide updated OH fields. Miyazaki et al. (2015) demonstrated 284 285 that assimilation of Tropospheric Emission Spectrometer (TES) O₃, Ozone Monitoring Instrument 286 (OMI) NO2, and MOPITT CO can provide a better description of tropospheric OH. There are also recent efforts that have assimilated surface in-situ MCF measurements (Fortems-Cheiney et al. 287 288 2011, 2012; Yin et al. 2015). However, because of the uncertainties in model chemistry schemes, 289 potential bias drifts in satellite remotely sensed observation, and sparse distribution of surface in-290 situ measurements, OH abundances provided by these approaches may not be ideal for the 291 estimation of long-term CO variation. 292 Emissions of MCF are regulated by the Montreal Protocol agreement. The loss rate of MCF

has become a good tool to evaluate the variation of tropospheric OH (e.g. <u>Krol et al. 1998;</u>
Bousquet et al. 2005; Prinn et al. 2005; Montzka et al. 2011). Using the same approach as Montzka

295 et al. (2011), we assess the variation of tropospheric OH in the period of 2001-2015. Figure 4a 296 shows the locations of WDCGG sites with MCF measurements, and Figure 4b shows the global 297 mean MCF concentration in the past 15 years. Similar as Montzka et al. (2011), our result shows 298 a exponential decrease of MCF concentration. The loss rate of MCF, derived from 12-month apart 299 of monthly means [e.g., ln(MCF_{Jan2007}/MCF_{Jan2006})] varies by 0.2% in the past 15 years (Figure 300 4c). The interannual variation is more likely due to the sparsity and discontinuity of measurements. 301 The small variation of loss rate of MCF demonstrates that the long-term variation of global 302 mean OH distributions is negligible in the past 15 years. Consequently, the decreasing trend of 303 tropospheric CO in North Hemisphere is driven by decreasing CO sources, rather than sinks. For 304 this reason, the default monthly OH fields of GEOS-Chem model (Evans et al. 2005), without 305 interannual variability, are used in this work to constrain the long-term variation of CO emissions. 306 Because the abundances of tropospheric OH have large regional discrepancies (e.g. Jiang et al. 307 2015a), it is possible that the actual OH is more variable at regions lacking MCF measurements 308 (e.g. India and southeast Asia). Futhermore, the magnitude and seasonality of the default monthly 309 OH fields could also have uncertainty. Consequently, the magnitude of CO emissions in our 310 analysis may still be affected by biases in OH, although the two-step assimilation system is 311 designed to suppress their influence.

312 4.2. Long-term variation of global CO emissions

In this work, we performed monthly inversions for the period of 2001-2015, using MOPITT column, profile and lower tropospheric profile (lowest three retrieval levels) data to investigate the influences associated with vertical sensitivity of satellite instrument and model transport error. Figure 5 shows the CO emission trends for 2001-2015 constrained by these different datasets. Because of the combination of various emission categories (i.e. anthropogenic,

biomass burning and VOC oxidation) in our methodology, we cannot completely separate the a posteriori emission estimates from different sources. However, the various spatial and temporal distribution of emissions sources (e.g. anthropogenic vs. biomass burning) provides valuable information to distinguish the contribution from each category. In order to further isolate the influences of biomass burning, the months dominated by biomass burning (biomass burning CO > 50% of total CO emission in an individual grid) are excluded in the trend analysis for anthropogenic and VOC sources (Figure 5).

325 For anthropogenic sources, all three analysis show significant emission reduction from 326 North America, Europe and China. The emission estimates constrained with MOPITT column and 327 profile data suggest increasing CO emissions from India and Southeast Asia. Conversely, the 328 emission estimate constrained with MOPITT lower tropospheric profile data shows a decreasing 329 trend in this region, and this decreasing trend is also obtained by Yin et al. (2015). As shown in 330 Jiang et al., (2013), errors in model convection in this region have a large effect on CO emissions 331 estimates, and information about the vertical profile of CO has a stronger influence on the results. 332 For biomass burning sources, we found a negative trend over boreal North America, boreal 333 Asia and South America, and a positive trend over Indonesia that is primarily due to the strong 334 impacts of El Nino in 2006 and 2015 on biomass burning in this region (e.g. Field et al., 2016). 335 Our results for biogenic VOCs are inconclusive; the emission estimates constrained with MOPITT 336 column and profile data show moderate positive trends in the tropics, and slight negative trends in 337 mid-latitude regions, whereas the emission estimate constrained with MOPITT lower tropospheric 338 profile data shows a negative trend globally.

339 4.2.1. Regional analysis for anthropogenic emissions

Figure 6a shows the regional variation of anthropogenic emissions from the United States

Deleted: demonstrate

342 (US). The emission estimates constrained with MOPITT column and profile data match very well 343 with the a priori emissions, whereas the emission estimate constrained with MOPITT lower 344 tropospheric profile data is much higher. All three analyses demonstrate a significant emission 345 reduction over our study period. As shown in Table 1, the total anthropogenic CO emission 346 (constrained with MOPITT profile data) from US is 56.8 Tg in 2015, which is 35% lower than that 347 in 2001 (87.7 Tg). Figure 7a shows the monthly mean CO concentrations from WDCGG stations 348 in US, which demonstrates a similar decreasing trend as our analysis. The initial increase at 2001-349 2002 could be caused by uncertainties in the data. The decreasing trend is consistent with the US 350 Environmental Protection Agency (EPA) Emissions Trends Data (https://www.epa.gov/air-351 emissions-inventories/air-pollutant-emissions-trends-data), and other observation records for 352 western US (Gratz et al. 2015), southeast US (Hidy et al. 2014) and North Atlantic (Kumar et al. 353 2013).

354 Figure 6b shows the regional variation of anthropogenic emissions from Europe. All three 355 analyses show an underestimation of a priori emissions, suggesting the CO emissions in the EMEP 356 inventory are too low. Our results show that anthropogenic emissions decrease during the period 357 of 2001-2007, but are almost unchanged in the following years, which is consistent with the 358 observations from WDCGG stations (Figure 7b). Recent studies (Hilboll et al. 2013; Schneider et 359 al. 2015) showed that NO₂ over Europe from SCIAMACHY is decreasing in the period of 2002-2008, and almost unchanged in the period of 2008-2011. Henschel et al. (2015) indicated that the 360 361 unchanged NO₂ over Europe could be caused by European emissions that are failing to achieve 362 the expected reduction standards. Because anthropogenic CO and NO₂ share some of the same combustion sources, it is possible that the unchanged CO emission in our analysis is also due to a 363 364 failure of emission controls.

365 Figure 6c shows the regional variation of anthropogenic emissions from east China. We 366 found Chinese anthropogenic emissions are increasing in the period of 2001-2004. Accompanied 367 with the global economy recession, the total anthropogenic CO emission (constrained with 368 MOPITT profile data) from east China decreases to 175.4 Tg in 2008, which is 15% lower than 369 that in 2004 (205.6 Tg). Our analysis shows a temporary increase of Chinese emissions in 2009 370 (185.9 Tg), followed by continuous decrease. The total Chinese anthropogenic CO emission is 159.0 Tg in 2015, which is 7% lower than that in 2001 (170.4 Tg). Using surface in-situ 371 372 measurements at Hateruma Island, Tohjima et al. (2014) constrained CO emissions from China 373 for the period 1999-2010. They found Chinese CO emission increases from 1999-2004, and 374 decreases since 2005. Using a "bottom-up" approach, recent studies (Zhao et al. 2012; Xia et al. 375 2016) indicated that the growth trend of Chinese CO emissions has been changed since 2005 376 because of improvements in energy efficiency and emission control regulations (e.g. Liu et al. 377 2015). Figure 7c shows the observation records from 2 stations in the East China outflow region, 378 which demonstrate similar variations.

379 Figures 6d-6e show the regional variation of anthropogenic emissions from India and 380 Southeast Asia. The emission estimates constrained with MOPITT column and profile data 381 demonstrate significant positive trend in our study period, whereas the emission estimate 382 constrained with MOPITT lower tropospheric profile data shows a decreasing trend. Schneider et al. (2015) showed that NO₂ over south Asia from SCIAMACHY is increasing in the period of 383 384 2003-2011. Using OMI NO₂ measurements, recent studies (e.g., Duncan et al. 2016) demonstrated that NO₂ over India has a positive trend during 2005-2015. Observations from Cape Rama (CRI) 385 station (Figure 7d) demonstrate that CO concentration in 2010-2013 is significantly higher than 386 387 that in 2001-2002. For these reasons, we have more confidence in our results that indicate

388 increasing anthropogenic CO emissions from India and Southeast Asia in the past 15 years. The 389 trend based on the MOPITT lower-tropospheric data is incorrect because of model error in 390 convection in this dynamically varying region, and the negative bias drift in MOPITT lower 391 tropospheric retrievals (Deeter et al., 2014). The total anthropogenic CO emission (constrained 392 with MOPITT profile data) from India and Southeast Asia is 130.4 Tg in 2015, which is 34% 393 higher than that in 2001 (97.5 Tg). It should be noted that the inconsistency between our analysis 394 with Yin et al. (2015) suggests more studies are needed for robust conclusion about the variation 395 of anthropogenic CO emissions for this region.

396 Although our inverse analysis (constrained with MOPITT profile data) suggests similar 397 anthropogenic CO emissions from East China in 2008 and 2014, Figure 7c demonstrates that mean 398 CO concentrations over the outflow region of East China are 6 ppb higher in 2014 compared to 399 2008. Our previous study (Jiang et al. 2015c) indicated that anthropogenic emissions from India 400 and southeast Asia have an important influence on pollutant concentrations in the east China 401 outflow region. It is possible that the increase of CO concentration observed by WDCGG stations 402 in this region is caused by the significant increase of anthropogenic CO emission from India and 403 southeast Asia. In the most recent 5 years (2011-2015), our results (constrained with MOPITT 404 profile data) suggested a 20.5 Tg emission reduction from East China, and a 10.1 Tg emission increase from India and Southeast Asia. Assuming a fixed emission growth rate, projected 405 406 anthropogenic CO emissions from India and Southeast Asia will overtake Chinese emissions in 407 2020, resulting in serious socioeconomic issues on both local and global scales.

408 4.2.2. Regional analysis for biomass burning emissions

Figure 8 and Table 2 show the regional variation of biomass burning emissions. There are

Deleted: shows

16

significant decreasing trends in three regions (i.e. boreal North America, boreal Asia, and South

Deleted: and boundary layer height

414	Alaska and western Canada) in 2004 (Figure 8a), which have been reported by previous studies
415	(e.g. Pfister et al. 2005; Turquety et al. 2007), and also from boreal Asia during 2001-2003 (Figure
416	8b) due to significant fire activity in Siberia (e.g., Yurganov et al., 2005, Stroppiana et al., 2010).
417	For South America (Figure 8c), we found higher biomass burning emissions in the periods of 2004-
418	2007 and 2010, consistent with fire activity reported in previous studies (e.g. Hooghiemstra et al.
419	2012; Bloom et al. 2015).
420	Figure 8d shows the regional variation of biomass burning emissions from Africa. The fire
421	activities in Africa demonstrates obvious seasonality: peak in boreal winter for Northern
422	Hemispheric Africa, and in austral winter for Southern Hemispheric Africa. Similar to previous
423	studies (e.g. Chevallier et al. 2009; Tosca et al. 2015), there is no obvious emission trend in Africa
424	in the past 15 years. This is also consistent with the burned area trends described by Andela et al.
425	(2014) which show opposite directions for Northern Africa (decreasing) versus Southern Africa Deleted:
426	(increasing) and would have cancelling effects in the trend for the continent as a whole.
427	Our results exhibit two strong biomass burning events in Indonesia, 2006 and 2015,
428	individually (Figure 8e). Previous studies (e.g. Logan et al. 2008; Zhang et al. 2011; Worden et al.
429	2013b, 2013c, Field et al., 2016) demonstrate the direct relationship between strong Indonesian
430	fires and El Niño. Recent studies (Huang et al. 2014: Inness et al. 2015) confirm low biomass
431	ines and Er Millo. Second studies (mang et al. 2014) inness et al. 2015) comminities belefed. Mole recent
	burning activities in Indonesia in the period of 2007-2012. CO emissions from the Indonesian fires
432	burning activities in Indonesia in the period of 2007-2012. CO emissions from the Indonesian fires associated with the 2015 El Niño were 92 Tg, (for October, 2015, as constrained with MOPITT Deleted: ,
432 433	burning activities in Indonesia in the period of 2007-2012. CO emissions from the Indonesian fires associated with the 2015 El Niño were 92 Tg, (for October, 2015, as constrained with MOPITT Deleted: , profile data), and were about three times higher than the October 2006 El Nino driven fire
432 433 434	burning activities in Indonesia in the period of 2007-2012. CO emissions from the Indonesian fires associated with the 2015 El Niño were 92 Tg, (for October, 2015, as constrained with MOPITT Deleted: , profile data), and were about three times higher than the October 2006 El Nino driven fire emissions (32 Tg). Not including the 2015 El Niño driven fires, our analysis indicates a negative Deleted:
432 433 434 435	burning activities in Indonesia in the period of 2007-2012. CO emissions from the Indonesian fires associated with the 2015 El Niño were 92 Tg, (for October, 2015, as constrained with MOPITT profile data), and were about three times higher than the October 2006 El Nino driven fire emissions (32 Tg). Not including the 2015 El Niño driven fires, our analysis indicates a negative trend of global biomass burning emissions in the past 15 years, as shown in Figure11f.

America). Our results show high biomass burning emissions from boreal North America (mainly

440 4.3. Changes in tropospheric CO during 2001-2015

441 In this section, we evaluate our inversion results using independent long-term surface insitu measurements from WDCGG stations. Figure 9a shows the annual trend of surface CO 442 443 concentration for 2001 - 2015 from WDCGG sites, and from model simulations driven with a 444 priori emissions. Most WDCGG sites exhibit negative trends in the past 15 years, confirming the 445 decreasing trend of global tropospheric CO, which is consistent with satellite observations (e.g. 446 Warner et al. 2013; Worden et al. 2013). There are also stations with positive trends, for example, 447 Tae-ahn Peninsula (TAP, Korea), Ascension Island (ASC, equtorial Atlantic Ocean), Cape Rama 448 (CRI, India), Bukit Koto Tabang (BKT, Indonesia) and Cape Grim (CGO, Australia). Globally, 449 the a priori model simulation is in reasonable agreement with WDCGG measurements: both show 450 negative trends in middle/high latitude, and positive trends in some tropical regions. However, 451 there are noticable discrepancies, for example, the surface observation from Yonagunijima (YON, 452 east China sea) shows a negative trend in our study period, suggesting decreasing trend from 453 Chinese CO emission, whereas the a priori simulation demonstrates significant positive trend. 454 Figure 9b-9d show the model simulations driven with a posteriori emissions. The a 455 posteriori emissions constrained with MOPITT lower tropospheric profile data (Figure 9d) results

in unrealistic large CO reduction, which could be caused by the negative bias drift of MOPITT retrievals at lower troposphere (Deeter et al. 2014) and the influence from possible variability in model convective transport. The a posteriori emissions constrained with MOPITT column and profile data have similar comparisons. For example, both of them suggest a negative trend over east China, consistent with observations from YON, and positive trend over northeast Asia, consistent with observations from TAP.

462

2 In order to better compare the discrepancy between model simulation and surface

463	observations, Figure 9e-9g show the improvement due to a posteriori emissions, derived by
464	$abs(Trend_{aposteriori} - Trend_{WDCGG}) \text{ - } abs(Trend_{apriori} \text{ - } Trend_{WDCGG}). Blue (red) means the a posteriori of the second s$
465	emissions improves (degrades) the agreement with WDCGG measurements compared to the
466	simulated surface CO using a priori emissions, while white indicates no change from the prior. As
467	shown in Figure 9f, the CO emissions constrained with MOPITT profile data improved the model
468	simulation for most WDCGG sites in the Northern Hemisphere. The a posteriori emissions
469	constrained with MOPITT column data are somewhat worse, particularly over Europe, while CO
470	emissions constrained with MOPITT profile data over Europe give improved comparisons to
471	WDCGG surface CO measurements. Worden et al. (2010) demonstrated that the degrees of
472	freedom for signal (DFS) of MOPITT multi-spectral profile retrievals (TIR+NIR) is about 1.5-2.0
473	over land, which is reduced to about 1 DFS when converted to a total column. This reduction in
474	vertical information in MOPITT column data can affect the the reliability of inverse analysis
475	results (Jiang et al., 2015a). It should be noticed that the vertical correlation in model simulation
476	is not considered in our assimilation, which could be another possible reason for this discrepancy.
477	Figure 10a-10d show the long-term mean value of surface CO concentration for 2001 -
478	2015 from WDCGG sites, and model simulations driven with a priori and a posteriori emissions.
479	All simulations provide similar results for long-term mean value. Figure 10e-10g show the
480	improvement due to a posteriori emissions, derived by $abs(CO_{aposteriori} - CO_{WDCGG})$ - $abs(CO_{apriori})$
481	- $\mathrm{CO}_{\mathrm{WDCGG}}$). Figure 10f demonstrates that CO emissions constrained with MOPITT profile data
482	improved the model simulation in about half of the sites in the Northern Hemisphere, whereas the
483	a posteriori emissions constrained with MOPITT column data are somewhat worse (Figure 10e).
484	Evaluating modeled tracer concentrations using surface in-situ measurements is more challenging
485	than evaluating long-term trends. Important sources of uncertainty include the representation error
1	

Deleted: In a recent study, Schnell et al. (2015) evaluate surface O_3 concentrations simulated by multi-models for North America and Europe. They found most models can provide good simulations for the patterns of O_3 but cannot reproduce the magnitude.

491 (e.g. Chang et al. 2015; Kharol et al. 2015) and vertical mixing of boundary layer (e.g. Castellanos

492 et al. 2011; Cuchiara et al. 2014).

512

493 Because our a posteriori simulation, particularly using emissions constrained with 494 MOPITT profile data, results in significant improvement in the long-term trend, and moderate 495 improvement in the mean value, we believe these a posteriori estimates provide a better description 496 for the long-term variation of global CO emissions. A remaining question is to explore how 497 changes in meterological conditions affect the long-term variation. By fixing CO emissions to 498 2001 levels, Figure 11a-11b show the long-term trend of modeled surface and column CO during 499 2001-2015, due only to changes in meterological conditions. At the surface level (Figure 11a), we 500 found changes in meterology result in a moderate positive trend in the Northern Hemisphere, 501 particularly, over northeast Asia, consistent with observation records from the TAP station; and 502 significant positive trend in tropics, consistent with observation record from ASC station. On the 503 other hand, the influence of meterological conditions on column CO (Figure 11b) is much weaker. 504 The discrepancy between surface and column CO suggests the possible contribution from variable 505 convective transport. It should be noted that our analysis for the contributions from meterological 506 conditions could be affected by the discrepancies among various versions of the meterological 507 fields (i.e. GEOS-4, GEOS-5 and GEOS-FP), and the lack of consistency in model physics of 508 GEOS-5 (e.g. the transition from GEOS 5.1.0 to GEOS 5.2.0 in late 2008). 509 Figure 11c-11h show the variation of global tropospheric CO due to changes in emissions. 510 Yin et al. (2015) indicated that the negative trend of tropospheric CO in the Northern Hemisphere 511 is driven by decreasing anthropogenic emissions from North America, Europe and China. Along

513 burning emissions from boreal North America and boreal Asia (Figure 11e, 11f) to be an important

with reductions in anthropogenic emissions (Figure 11c, 11d), we found the decrease of biomass

Deleted: , which could be associated with changes in the frequency of deep convection (Tan et al. 2015) or the change from El Niño to La Niña in

Deleted: study period (Andela et al. 2014). In order to assess the influence of

Deleted:) on the trend analysis, we reploted (not shown) Figure 11a-11b for the period 2004-2012 with

Deleted: meterological fields, and obtained similar significant positive trend in tropics, which suggests limited influence

Deleted: meterological field version differences on the trend analysis.

526 factor for this negative trend. In constrast to the emission reduction from North America, Europe 527 and China, we found increasing anthropogenic emissions from India and southeast Asia, which 528 result in a pronounced positive trend of tropospheric CO, while Yin et al. (2015) obtain a negative 529 trend for this region. This discrepancy requires further study and we will need to test the relative 530 importance of the primary differences in our methods, i.e., models and inversion approaches, 531 climatological OH (this study) vs. assimilated surface measurements of CH₄ and MCF to update 532 OH (Yin et al.) and the use of MOPITT profile vs. column CO retrievals (Yin et al., assimilate 533 only column CO).

534 **5. Summary**

535 The objective of this work is to investigate the dominant reasons for the observed variation 536 of global tropospheric CO over the past 15 years. We provide an update for this critical question 537 and also an updated CO emission estimates for model studies. In particular, we use surface 538 measurements of MCF to evaluate changes in the sinks of atmospheric CO, and constrain the 539 sources using MOPITT CO measurements to explain the observed decrease in CO concentrations. 540 Our two-step approach for estimating global CO emissions mitigates the effects of model errors 541 from transport and chemistry, as well as measurement bias error. 542 Using the same approach as Montzka et al. (2011), we assess the variation of tropospheric OH (the primary CO sink) in the period of 2001-2015 using MCF measurements from WDCGG 543 544 stations. Our result demonstrates negligible variation of global tropospheric OH in the past 15 years, and consequently we suggest that the global sink of CO due to chemical loss through OH 545 546 has not likely changed during this time period. We therefore expect the decreasing trend of tropospheric CO in North hemisphere (e.g. Warner et al. 2013; Worden et al. 2013; Gratz et al. 547 548 2015) to be driven by decreasing CO sources. Total anthropogenic CO emissions from the US

 Deleted: , and to
 Deleted: a combination of MOPITT CO measurements and
 Deleted: sources and
 Deleted: with the goal of explaining

were 56.8 Tg in 2015, which are 35% lower than emissions in 2001 (87.7 Tg). Total anthropogenic 553 554 CO emissions from East China were 159.0 Tg in 2015, which are 7% lower than 2001 emissions 555 (170.4 Tg) and 23% lower than 2004 emissions (205.6 Tg). This pronounced decrease of emissions 556 from US and China is an indication of progress for fuel efficiency and emission control regulations. 557 Conversely, our results demonstrate that anthropogenic emissions from Europe decreased from 558 2001 to 2007 but are almost unchanged during 2008-2015. We also found a significant increase of 559 anthropogenic emissions for India and Southeast Asia. The total anthropogenic CO emission from 560 India and southeast Asia is 130.4 Tg in 2015, which is 34% higher than that in 2001 (97.5 Tg). Assuming the same emission growth rate as 2011-2015, we expect that anthropogenic CO 561 562 emissions from India and Southeast Asia will be larger than Chinese emissions by 2020.

563 In a recent study, Yin et al. (2015) indicated that the decreasing tropospheric CO in the 564 Northern Hemisphere is caused by the decrease of anthropogenic emissions from North America, 565 Europe and China. We find that a decrease of biomass burning emissions from boreal North 566 America and boreal Asia is also an important contributor for the negative trend. Globally, our 567 analysis indicates a negative trend of biomass burning emissions in the past 15 years, except in 568 Indonesia due to the strong biomass burning event in 2015 associated with El Niño. Our results 569 demonstrate a significant decrease of biomass burning emissions from South America, which 570 could be associated with the reduction of deforestation in Brazil (Reddington et al. 2015), and the 571 predominant change from El Nino to La Nina in our study period (Andela et al. 2014). For Africa, 572 there is no obvious CO emission trend in the past 15 years, consistent with previous results 573 (Chevallier et al. 2009; Tosca et al. 2015; Andela et al., 2014). Our results are inconclusive in 574 characterizing the CO sources from oxidation of biogenic VOCs. More efforts are needed in the 575 future to better understand the mechanism for tropical CO emissions.

576	Our analysis highlights the importance of space-based instruments for monitoring changes
577	in global pollutant emissions. Our results demonstrate successful emission controls in US and
578	China over the past 15 years, and suggest that emission controls in Europe may need re-evaluation.
579	We also recommend more efforts in the future to better understand the regional and global effects
580	of increasing pollutant emissions from India and Southeast Asia.

582 Acknowledgments.

583 We thank the World Data Centre for Greenhouse Gases (WDCGG) for providing their CO 584 and MCF data. The National Center for Atmospheric Research (NCAR) is sponsored by the 585 National Science Foundation. The NCAR MOPITT project is supported by the National 586 Aeronautics and Space Administration (NASA) Earth Observing System (EOS) Program. The MOPITT team also acknowledges support from the Canadian Space Agency (CSA), the Natural 587 588 Sciences and Engineering Research Council (NSERC) and Environment Canada, along with the 589 contributions of COMDEV (the prime contractor) and ABB BOMEM. MOPITT data sets used in 590 this study are publicly available at http://reverb.echo.nasa.gov and at 591 https://eosweb.larc.nasa.gov/datapool.

592

593 Data availability

- 594 The MOPITT data is available at ftp://l5eil01.larc.nasa.gov/MOPITT/MOP02J.006. The MCF and
- 595 CO measurements from WDCGG is available at http://ds.data.jma.go.jp/gmd/wdcgg/.

596

597 **References**

598 Andela, N., and van der Werf, G.: Recent trends in African fires driven by cropland expansion and

Moved (insertion) [1]

599	El Niño to	La Niña	transition,	Nature	Climate	Change 4.	791-795.	doi:10.1038/nclimate2313.

600 <u>2014.</u>

605

601	Arellano, A., Kasibhatla, P., Giglio, L., Werf, G., Randerson, J. and Collatz, G.: Time-dependent
602	inversion estimates of global biomass-burning CO emissions using Measurement of Pollution ir
603	the Troposphere (MOPITT) measurements, J Geophys Res Atmospheres 1984 2012, 111(D9)
604	doi:10.1029/2005JD006613, 2006.

- 606 Bergamaschi, P., Frankenberg, C., Meirink, J., Krol, M., Dentener, F., Wagner, T., Platt, U.,
- 607 Kaplan, J., Körner, S., Heimann, M., Dlugokencky, E. and Goede, A.: Satellite chartography of 608 atmospheric methane from SCIAMACHY on board ENVISAT: 2. Evaluation based on inverse 609 J model simulations, Geophys Res Atmospheres 1984 2012, 112(D2), 610 doi:10.1029/2006JD007268, 2007.
- 611 Bergamaschi, P., Frankenberg, C., Meirink, J., Krol, M., Villani, M., Houweling, S., Dentener, F.,
- 612 Dlugokencky, E., Miller, J., Gatti, L., Engel, A. and Levin, I.: Inverse modeling of global and
- regional CH4 emissions using SCIAMACHY satellite retrievals, J Geophys Res Atmospheres
 1984 2012, 114(D22), doi:10.1029/2009JD012287, 2009.
- 615 Bergamaschi, P., Houweling, S., Segers, A., Krol, M., Frankenberg, C., Scheepmaker, R.,
- 616 Dlugokencky, E., Wofsy, S., Kort, E., Sweeney, C., Schuck, T., Brenninkmeijer, C., Chen, H.,
- 617 Beck, V. and Gerbig, C.: Atmospheric CH4 in the first decade of the 21st century: Inverse
- 618 modeling analysis using SCIAMACHY satellite retrievals and NOAA surface measurements, J
- 619 Geophys Res Atmospheres, 118(13), 7350–7369, doi:10.1002/jgrd.50480, 2013.
- 620 Bloom, A., Worden, J., Jiang, Z., Worden, H., Kurosu, T., Frankenberg, C. and Schimel, D.:
- 621 Remote-sensing constraints on South America fire traits by Bayesian fusion of atmospheric and

Moved up [1]: Andela, N., and van der Werf, G.: Recent trends in African fires driven by cropland expansion and El Niño to La Niña transition, Nature Climate Change 4, 791–795, doi:10.1038/nclimate2313, 2014.

- 626 surface data, Geophys Res Lett, 42(4), 1268–1274, doi:10.1002/2014GL062584, 2015.
- 627 Bousquet, P., Hauglustaine, D. A., Peylin, P., Carouge, C., and Ciais, P.: Two decades of OH
- variability as inferred by an inversion of atmospheric transport and chemistry of methyl
 chloroform, Atmos. Chem. Phys., 5, 2635-2656, doi:10.5194/acp-5-2635-2005, 2005.
- Bousserez, N., Henze, D., Perkins, A., Bowman, K., Lee, M., Liu, J., Deng, F. and Jones, D.:
- 631 Improved analysis-error covariance matrix for high-dimensional variational inversions:
- application to source estimation using a 3D atmospheric transport model, Q J Roy Meteor Soc,
 141(690), 1906–1921, doi:10.1002/qj.2495, 2015.
- Bruhwiler, L., Dlugokencky, E., Masarie, K., Ishizawa, M., Andrews, A., Miller, J., Sweeney, C.,
- Tans, P., and Worthy, D.: CarbonTracker-CH₄: an assimilation system for estimating emissions
- of atmospheric methane, Atmos. Chem. Phys., 14, 8269-8293, doi:10.5194/acp-14-8269-2014,
 2014.
- 638 Castellanos, P., Marufu, L., Doddridge, B., Taubman, B., Schwab, J., Hains, J., Ehrman, S. and
- 639 Dickerson, R.: Ozone, oxides of nitrogen, and carbon monoxide during pollution events over the
- eastern United States: An evaluation of emissions and vertical mixing, J Geophys Res
 Atmospheres 1984 2012, 116(D16), doi:10.1029/2010JD014540, 2011.
- 642 Chang, K.-L., Guillas, S., and Fioletov, V. E.: Spatial mapping of ground-based observations of
- 643 total ozone, Atmos. Meas. Tech., 8, 4487-4505, doi:10.5194/amt-8-4487-2015, 2015.
- 644 Chevallier, F., Fortems, A., Bousquet, P., Pison, I., Szopa, S., Devaux, M. and Hauglustaine, D.:
- 645 African CO emissions between years 2000 and 2006 as estimated from MOPITT observations,
- 646 Biogeosciences, 6(1), 103–111, doi:10.5194/bg-6-103-2009, 2009.
- 647 Cuchiara, G. C., Li, X., Carvalho, J. and Rappenglück, B.: Intercomparison of planetary boundary
- 648 layer parameterization and its impacts on surface ozone concentration in the WRF/Chem model

- 649 for a case study in Houston/Texas, Atmos Environ, 96, 175–185,
 650 doi:10.1016/j.atmosenv.2014.07.013, 2014.
- Deeter, M., Worden, H., Gille, J., Edwards, D., Mao, D. and Drummond, J.: MOPITT multispectral
 CO retrievals: Origins and effects of geophysical radiance errors, J Geophys Res Atmospheres
 1984 2012, 116(D15), doi:10.1029/2011JD015703, 2011.
- 654 Deeter, M. N., Martínez-Alonso, S., Edwards, D. P., Emmons, L. K., Gille, J. C., Worden, H. M.,
- 655 Sweeney, C., Pittman, J. V., Daube, B. C., and Wofsy, S. C.: The MOPITT Version 6 product:
- algorithm enhancements and validation, Atmos. Meas. Tech., 7, 3623-3632, doi:10.5194/amt-73623-2014, 2014.
- 658 Deng, F., Jones, D. B. A., Henze, D. K., Bousserez, N., Bowman, K. W., Fisher, J. B., Nassar, R.,
- 659 O'Dell, C., Wunch, D., Wennberg, P. O., Kort, E. A., Wofsy, S. C., Blumenstock, T., Deutscher,
- 660 N. M., Griffith, D. W. T., Hase, F., Heikkinen, P., Sherlock, V., Strong, K., Sussmann, R., and
- 661 Warneke, T.: Inferring regional sources and sinks of atmospheric CO₂ from GOSAT XCO₂ data,
- 662 Atmos. Chem. Phys., 14, 3703-3727, doi:10.5194/acp-14-3703-2014, 2014.
- Duncan, B., Lamsal, L., Thompson, A., Yoshida, Y., Lu, Z., Streets, D., Hurwitz, M. and
 Pickering, K.: A space-based, high-resolution view of notable changes in urban NOx pollution
- around the world (2005–2014), J Geophys Res Atmospheres, 121(2), 976–996,
 doi:10.1002/2015JD024121, 2016.
- 667 Evans, M. J., and Jacob, D. J.: Impact of new laboratory studies of N₂O₅ hydrolysis on global
- model budgets of tropospheric nitrogen oxides, ozone, and OH, Geophys. Res. Lett., 32, L09813,
- 669 doi:10.1029/2005GL022469, 2005.
- 670 Field, R. et al., 2015 Indonesian fire activity and smoke pollution show persistent non-linear
- 671 sensitivity to El Niño-induced drought, PNAS, 2016, 9204–9209, doi: 10.1073/pnas.1524888113

- 672 Fortems-Cheiney, A., Chevallier, F., Pison, I., Bousquet, P., Szopa, S., Deeter, M. and Clerbaux,
- 673 C.: Ten years of CO emissions as seen from Measurements of Pollution in the Troposphere
- 674 (MOPITT), J Geophys Res Atmospheres 1984 2012, 116(D5), doi:10.1029/2010JD014416,
 675 2011.
- 676 Fortems-Cheiney, A., Chevallier, F., Pison, I., Bousquet, P., Saunois, M., Szopa, S., Cressot, C.,
- 677 Kurosu, T. P., Chance, K., and Fried, A.: The formaldehyde budget as seen by a global-scale
- 678 multi-constraint and multi-species inversion system, Atmos. Chem. Phys., 12, 6699-6721,
- 679 doi:10.5194/acp-12-6699-2012, 2012.
- 680 Gonzi, S., Feng, L. and Palmer, P.: Seasonal cycle of emissions of CO inferred from MOPITT
- 681 profiles of CO: Sensitivity to pyroconvection and profile retrieval assumptions, Geophys Res
- 682 Lett, 38(8), n/a–n/a, doi:10.1029/2011GL046789, 2011.
- 683 Gratz, L. E., Jaffe, D. A. and Hee, J. R.: Causes of increasing ozone and decreasing carbon
- monoxide in springtime at the Mt. Bachelor Observatory from 2004 to 2013, Atmos Environ,
 109, 323–330, doi:10.1016/j.atmosenv.2014.05.076, 2015.
- 686 Guenther, A., Karl, T., Harley, P., Wiedinmyer, C., Palmer, P.I., and Geron, C.: Estimates of global
- 687 terrestrial isoprene emissions using MEGAN (Model of Emissions of Gases and Aerosols from
- 688 Nature), Atmos. Chem. Phys., 6, 3181-3210, doi:10.5194/acp-6-3181-2006, 2006.
- 689 Heald, C., Jacob, D., Jones, D., Palmer, P., Logan, J., Streets, D., Sachse, G., Gille, J., Hoffman,
- 690 R. and Nehrkorn, T.: Comparative inverse analysis of satellite (MOPITT) and aircraft (TRACE-
- P) observations to estimate Asian sources of carbon monoxide, J Geophys Res Atmospheres 1984
- 692 2012, 109, D23306, doi:10.1029/2004JD005185, 2004.
- 693
 Henschel, S., Tertre, A., Atkinson, R., Ouerol, X., Pandolfi, M., Zeka, A., Haluza, D., Analitis, A.,
 Moved (insertion) [2]

 694
 Katsouyanni, K., Bouland, C., Pascal, M., Medina, S. and Goodman, P.: Trends of nitrogen

ť	695	oxides in ambient air in nine Eu	opean cities between 1999 and 2010	Atmos Environ, 117,	234-
н			*		

696 <u>241, doi:10.1016/j.atmosenv.2015.07.013, 2015.</u>

699

- 697 Henze, D. K., Hakami, A., and Seinfeld, J. H.: Development of the adjoint of GEOS-Chem, Atmos.
- 698 Chem. Phys., 7, 2413-2433, doi:10.5194/acp-7-2413-2007, 2007.

Hidy, G., Blanchard, C., Baumann, K., Edgerton, E., Tanenbaum, S., Shaw, S., Knipping, E.,

- 701 Tombach, I., Jansen, J. and Walters, J.: Chemical climatology of the southeastern United States,
- 702 1999–2013, Atmos Chem Phys, 14(21), 11893–11914, doi:10.5194/acp-14-11893-2014, 2014.
- Hilboll, A., Richter, A., and Burrows, J. P.: Long-term changes of tropospheric NO₂ over [Moved (insertion) [3]
 megacities derived from multiple satellite instruments, Atmos. Chem. Phys., 13, 4145-4169,
 doi:10.5194/acp-13-4145-2013, 2013.
- 706 Hooghiemstra, P., Krol, M., Leeuwen, T., Werf, G., Novelli, P., Deeter, M., Aben, I. and
- 707 Röckmann, T.: Interannual variability of carbon monoxide emission estimates over South
- America from 2006 to 2010, J Geophys Res Atmospheres 1984 2012, 117(D15), n/a–n/a,
 doi:10.1029/2012JD017758, 2012.
- 710 Houweling, S., Krol, M., Bergamaschi, P., Frankenberg, C., Dlugokencky, E. J., Morino, I.,
- 711 Notholt, J., Sherlock, V., Wunch, D., Beck, V., Gerbig, C., Chen, H., Kort, E. A., Röckmann,
- 712 T., and Aben, I.: A multi-year methane inversion using SCIAMACHY, accounting for
- 713 systematic errors using TCCON measurements, Atmos. Chem. Phys., 14, 3991-4012,
- 714 doi:10.5194/acp-14-3991-2014, 2014.
- 715 Huang, L., Fu, R., and Jiang, J. H.: Impacts of fire emissions and transport pathways on the
- 716 interannual variation of CO in the tropical upper troposphere, Atmos. Chem. Phys., 14, 4087-
- 717 <u>4099, doi:10.5194/acp-14-4087-2014, 2014.</u>

 $\label{eq:product} \begin{array}{l} P.: \mbox{ Long-term changes of tropospheric NO}_2 \mbox{ over megacities derived from multiple satellite instruments, Atmos. Chem. Phys., 13, 4145-4169, doi:10.5194/acp-13-4145-2013, 2013. \end{array}$

Moved down [3]: Hilboll, A., Richter, A., and Burrows, J.

Moved (insertion) [4]

- 722 Inness, A., Benedetti, A., Flemming, J., Huijnen, V., Kaiser, J. W., Parrington, M., and Remy, S.:
- 723 The ENSO signal in atmospheric composition fields: emission-driven versus dynamically
- 724 induced changes, Atmos. Chem. Phys., 15, 9083-9097, doi:10.5194/acp-15-9083-2015, 2015.
- Jiang, Z., Jones, D., Kopacz, M., Liu, J., Henze, D. and Heald, C.: Quantifying the impact of model
- rrors on top-down estimates of carbon monoxide emissions using satellite observations, J
- 727 Geophys Res Atmospheres 1984 2012, 116(D15), doi:10.1029/2010JD015282, 2011.
- Jiang, Z., Jones, D., Worden, H., Deeter, M., Henze, D., Worden, J., Bowman, K., Brenninkmeijer,
- 729 C. and Schuck, T.: Impact of model errors in convective transport on CO source estimates
- inferred from MOPITT CO retrievals, J Geophys Res Atmospheres, 118(4), 2073–2083,
 doi:10.1002/jgrd.50216, 2013.
- 732 Jiang, Z., Jones, D., Worden, H. and Henze, D.: Sensitivity of top-down CO source estimates to
- the modeled vertical structure in atmospheric CO, Atmos Chem Phys, 15(3), 1521–1537,
- 734 doi:10.5194/acp-15-1521-2015, 2015a.
- 735 Jiang, Z., Jones, D., Worden, J., Worden, H., Henze, D. and Wang, Y.: Regional data assimilation
- of multi-spectral MOPITT observations of CO over North America, Atmos Chem Phys, 15(12),
 6801–6814, doi:10.5194/acp-15-6801-2015, 2015b.
- 738 Jiang, Z., Worden, J. R., Jones, D. B. A., Lin, J.-T., Verstraeten, W. W., and Henze, D. K.:
- 739 Constraints on Asian ozone using Aura TES, OMI and Terra MOPITT, Atmos. Chem. Phys., 15,
- 740 99-112, doi:10.5194/acp-15-99-2015, 2015c.
- 741 Jones, D., Bowman, K., Logan, J., Heald, C., Liu, J., Luo, M., Worden, J. and Drummond, J.: The
- zonal structure of tropical O3 and CO as observed by the Tropospheric Emission Spectrometer
- in November 2004 Part 1: Inverse modeling of CO emissions, Atmos Chem Phys, 9(11), 3547–

744 3562, doi:10.5194/acp-9-3547-2009, 2009.

745	٨		Moved up [2]: Henschel, S., Tertre, A., Atkinson, R., Querol, X., Pandolfi, M., Zeka, A., Haluza, D., Analitis, A., Katsouyanni, K., Bouland, C., Pascal, M., Medina, S. and Goodman, P.: Trends of nitrogen oxides in ambient air in nine European cities between 1999 and 2010, Atmos Environ 117, 234–241
746	۸		
/4/			doi:10.1016/j.atmosenv.2015.07.013, 2015.
748	۸		Moved up [4]: Huang, L., Fu, R., and Jiang, J. H.: Impacts of fire emissions and transport pathways on the interannual
749	Kharol, S. K., Martin, R. V., Philip, S., Boys, B., Lamsal, L. N., Jerrett, M., Brauer, M., Crouse,		variation of CO in the tropical upper troposphere, Atmos. Chem. Phys., 14, 4087-4099, doi:10.5194/acp-14-4087- 2014, 2014.
750	D. L., McLinden, C. and Burnett, R. T.: Assessment of the magnitude and recent trends in		Inness, A., Benedetti, A., Flemming, J., Huijnen, V., Kaiser, J. W., Parrington, M., and Remy, S.: The ENSO signal in atmospheric composition fields: emission-driven versus
751	satellite-derived ground-level nitrogen dioxide over North America, Atmos Environ, 118, 236-		dynamically induced changes, Atmos. Chem. Phys., 15, 9083-9097, doi:10.5194/acp-15-9083-2015, 2015.
752	245, doi:10.1016/j.atmosenv.2015.08.011, 2015.	/	Moved down [5]: Kopacz, M., Jacob, D.,
753	Kopacz, M., Jacob, D., Henze, D., Heald, C., Streets, D. and Zhang, Q.: Comparison of adjoint		Moved down [0]: Henze, D., Heald, C., Streets, D. and Zhang, Q.: Comparison of adjoint and analytical Bayesian inversion methods for constraining Asian sources of carbon monoxide using satellite (MOPITT) measurements of CO
754	and analytical Bayesian inversion methods for constraining Asian sources of carbon monoxide		columns, J Geophys Res Atmospheres 1984 2012, 114(D4), doi:10.1029/2007JD009264, 2009
755	using satellite (MOPITT) measurements of CO columns, J Geophys Res Atmospheres 1984		Moved (insertion) [6]
756	<u>2012, 114(D4), doi:10.1029/2007JD009264, 2009.</u>		
757	Kopacz, M., Jacob, D., Fisher, J., Logan, J., Zhang, L., Megretskaia, I., Yantosca, R., Singh, K.,		Moved (insertion) [5]
758	Henze, D., Burrows, J., Buchwitz, M., Khlystova, I., McMillan, W., Gille, J., Edwards, D.,		
759	Eldering, A., Thouret, V. and Nedelec, P.: Global estimates of CO sources with high resolution		
760	by adjoint inversion of multiple satellite datasets (MOPITT, AIRS, SCIAMACHY, TES), Atmos		
761	Chem Phys, 10(3), 855-876, doi:10.5194/acp-10-855-2010, 2010.		
762	Krol, M., vanLeeuwen, P.J., and Lelieveld, J.: Global OH trend inferred from methylchloroform		Moved (insertion) [7]
763	measurements, J. Geophys. Res., 103(D9), 10697-10711, doi:10.1029/98JD00459, 1998.		
764	Kuhns, H., Green, M. and Etyemezian, V.: Big Bend Regional Aerosol and Visibility		
765	Observational (BRAVO) Study Emissions Inventory, Report prepared for BRAVO Steering		

- 766 Committee, Desert Research Institute, Las Vegas, Nevada, 2003.
- 767 Kumar, A., Wu, S., Weise, M. F., Honrath, R., Owen, R. C., Helmig, D., Kramer, L., Val Martin,

- 792 M., and Li, Q.: Free-troposphere ozone and carbon monoxide over the North Atlantic for 2001-
- 793 2011, Atmos. Chem. Phys., 13, 12537-12547, doi:10.5194/acp-13-12537-2013, 2013.
- Lelieveld, J., Dentener, F., Peters, W. and Krol, M.: On the role of hydroxyl radicals in the selfcleansing capacity of the troposphere, Atmos Chem Phys, 4(9/10), 2337–2344, doi:10.5194/acp-
- 796 4-2337-2004, 2004.
- 797 Liu, F., Zhang, Q., Tong, D., Zheng, B., Li, M., Huo, H., and He, K. B.: High-resolution inventory
- of technologies, activities, and emissions of coal-fired power plants in China from 1990 to 2010,
 Atmos. Chem. Phys., 15, 13299-13317, doi:10.5194/acp-15-13299-2015, 2015.
- Logan, J., Megretskaia, I., Nassar, R., Murray, L., Zhang, L., Bowman, K., Worden, H. and Luo,
 M.: Effects of the 2006 El Niño on tropospheric composition as revealed by data from the
 Tropospheric Emission Spectrometer (TES), Geophys Res Lett, 35(3),
 doi:10.1029/2007GL031698, 2008.
- 804 Meirink, J., Bergamaschi, P., Frankenberg, C., Amelio, M. d', Dlugokencky, E., Gatti, L.,
- 805 Houweling, S., Miller, J., Röckmann, T., Villani, M. and Krol, M.: Four-dimensional variational
- data assimilation for inverse modeling of atmospheric methane emissions: Analysis of
 SCIAMACHY observations, J Geophys Res Atmospheres 1984 2012, 113(D17),
 doi:10.1029/2007JD009740, 2008.
- 809 Miyazaki, K., Eskes, H. and Sudo, K.: A tropospheric chemistry reanalysis for the years 2005-
- 810 2012 based on an assimilation of OMI, MLS, TES, and MOPITT satellite data, Atmos Chem
- 811 Phys, 15(14), 8315–8348, doi:10.5194/acp-15-8315-2015, 2015.
- Montzka, S. A., Krol, M., Dlugokencky, E., Hall, B., Jöckel, P., Lelieveld, J.: Small Interannual
 Variability of Global Atmospheric Hydroxyl, Science, 331(6013), 67–69,
 10.1126/science.1197640, 2011.
 - 31

- 815 Ohara, T., Akimoto, H., Kurokawa, J., Horii, N., Yamaji, K., Yan, X. and Hayasaka, T.: An Asian
- emission inventory of anthropogenic emission sources for the period 1980–2020, Atmos Chem
- 817 Phys, 7(16), 4419–4444, doi:10.5194/acp-7-4419-2007, 2007.
- 818 Pfister, G., Hess, P., Emmons, L., Lamarque, J. -F., Wiedinmyer, C., Edwards, D., Pétron, G.,
- 819 Gille, J. and Sachse, G.: Quantifying CO emissions from the 2004 Alaskan wildfires using
- 820 MOPITT CO data, Geophys Res Lett, 32(11), doi:10.1029/2005GL022995, 2005.
- 821 Prinn, R., Huang, J., Weiss, R., Cunnold, D., Fraser, P., Simmonds, P., McCulloch, A., Harth, C.,
- 822 Reimann, S., Salameh, P., O'Doherty, S., Wang, R., Porter, L., Miller, B. and Krummel, P.:
- 823 Evidence for variability of atmospheric hydroxyl radicals over the past quarter century, Geophys
- 824 Res Lett, 32(7), n/a–n/a, doi:10.1029/2004GL022228, 2005.
- 825 Reddington, C., Butt, E., Ridley, D., Artaxo, P., Morgan, W., Coe, H. and Spracklen, D.: Air
- quality and human health improvements from reductions in deforestation-related fire in Brazil,
- 827 Nat Geosci, 8(10), 768–771, doi:10.1038/ngeo2535, 2015.
- Schneider, P., Lahoz, W. A., and van der A, R.: Recent satellite-based trends of tropospheric
- nitrogen dioxide over large urban agglomerations worldwide, Atmos. Chem. Phys., 15, 12051220, doi:10.5194/acp-15-1205-2015, 2015.
- 831 Spivakovsky, C. M., Logan, J. A., Montzka, S. A., Balkanski, Y. J., Foreman-Fowler, M., Jones,
- 832 D. B. A., Horowitz, L. W., Fusco, A. C., Brenninkmeijer, C. A. M., Prather, M. J., Wofsy, S. C.
- 833 and McElroy, M. B.: Three-dimensional climatological distribution of tropospheric OH Update
- and evaluation, J. Geophys. Res., 105(D7), 8931–8980, doi:10.1029/1999JD901006, 2000.
- 835 Streets, D., Zhang, Q., Wang, L., He, K., Hao, J., Wu, Y., Tang, Y. and Carmichael, G.: Revisiting
- 836 China's CO emissions after the Transport and Chemical Evolution over the Pacific (TRACE-P)
- 837 mission: Synthesis of inventories, atmospheric modeling, and observations, J Geophys Res

Deleted: Schnell, J. L., Prather, M. J., Josse, B., Naik, V., Horowitz, L. W., Cameron-Smith, P., Bergmann, D., Zeng, G., Plummer, D. A., Sudo, K., Nagashima, T., Shindell, D. T., Faluvegi, G., and Strode, S. A.: Use of North American and European air quality networks to evaluate global chemistry-climate modeling of surface ozone, Atmos. Chem. Phys., 15, 10581-10596, doi:10.5194/acp-15-10581-2015, 2015.



- 846 Atmospheres 1984 2012, 111(D14), doi:10.1029/2006JD007118, 2006.
- 847 Strode, S., Worden, H., Damon, M., Douglass, A., Duncan, B., Emmons, L., Lamarque, J.-F.,
- 848 Manyin, M., Oman, L., Rodriguez, J., Strahan, S. and Tilmes, S.: Interpreting space-based trends
- in carbon monoxide with multiple models, Atmos Chem Phys, 16(11), 7285–7294,
 doi:10.5194/acp-16-7285-2016, 2016.
- 851 Stroppiana, D., Brivio, P. A., Grégoire, J.-M., Liousse, C., Guillaume, B., Granier, C., Mieville,
- 852 A., Chin, M., and Pétron, G.: Comparison of global inventories of CO emissions from biomass
- burning derived from remotely sensed data, Atmos. Chem. Phys., 10, 12173-12189,
- doi:10.5194/acp-10-12173-2010, 2010.
- Tohjima, Y., Kubo, M., Minejima, C., Mukai, H., Tanimoto, H., Ganshin, A., Maksyutov, S.,
- 856 Katsumata, K., Machida, T., and Kita, K.: Temporal changes in the emissions of CH₄ and CO
- 857 from China estimated from CH₄ / CO₂ and CO / CO₂ correlations observed at Hateruma Island,
- Atmos. Chem. Phys., 14, 1663-1677, doi:10.5194/acp-14-1663-2014, 2014.
- 859 Tosca, M., Diner, D., Garay, M. and Kalashnikova, O.: Human-caused fires limit convection in
- tropical Africa: First temporal observations and attribution, Geophys Res Lett, 42(15), 6492–
 6501, doi:10.1002/2015GL065063, 2015.
- 862 Turquety, S., Logan, J., Jacob, D., Hudman, R., Leung, F., Heald, C., Yantosca, R., Wu, S.,
- 863 Emmons, L., Edwards, D. and Sachse, G.: Inventory of boreal fire emissions for North America
- 864 in 2004: Importance of peat burning and pyroconvective injection, J Geophys Res Atmospheres
- 865 1984 2012, 112(D12), doi:10.1029/2006JD007281, 2007.
- 866 Turner, A. J., Jacob, D. J., Wecht, K. J., Maasakkers, J. D., Lundgren, E., Andrews, A. E., Biraud,
- 867 S. C., Boesch, H., Bowman, K. W., Deutscher, N. M., Dubey, M. K., Griffith, D. W. T., Hase,
- 868 F., Kuze, A., Notholt, J., Ohyama, H., Parker, R., Payne, V. H., Sussmann, R., Sweeney, C.,

Deleted: Tan,

33

Moved up [7]: J.

Deleted: Jakob, C., Rossow, W. and Tselioudis, G.: Increases in tropical rainfall driven by changes in frequency of organized deep convection, Nature, 519(7544), 451–454, doi:10.1038/nature14339, 2015.

- 875 Velazco, V. A., Warneke, T., Wennberg, P. O., and Wunch, D.: Estimating global and North
- 876 American methane emissions with high spatial resolution using GOSAT satellite data, Atmos.
- 877 Chem. Phys., 15, 7049-7069, doi:10.5194/acp-15-7049-2015, 2015.
- van der Werf, G. R., Randerson, J. T., Giglio, L., Collatz, G. J., Kasibhatla, P. S., and Arellano Jr.,
- A. F.: Interannual variability in global biomass burning emissions from 1997 to 2004, Atmos.
- 880 Chem. Phys., 6, 3423-3441, doi:10.5194/acp-6-3423-2006, 2006.
- 881 van der Werf, G. R., Randerson, J. T., Giglio, L., Collatz, G. J., Mu, M., Kasibhatla, P. S.,
- 882 Morton, D. C., DeFries, R. S., Jin, Y., and van Leeuwen, T. T.: Global fire emissions and the
- contribution of deforestation, savanna, forest, agricultural, and peat fires (1997–2009), Atmos.
- 884 Chem. Phys., 10, 11707–11735, doi:10.5194/acp-10-11707-2010, 2010.
- 885 van Leeuwen, T. T. and van der Werf, G. R.: Spatial and temporal variability in the ratio of trace
- gases emitted from biomass burning, Atmos. Chem. Phys., 11, 3611-3629, doi:10.5194/acp-113611-2011, 2011.
- 888 Vestreng, V. and Klein, H.: Emission data reported to UNECE/EMEP. Quality assurance and trend
- analysis and Presentation of WebDab, Norwegian Meteorological Institute, Oslo, Norway, MSCW Status Report, 2002.
- 891 Warner, J., Carminati, F., Wei, Z., Lahoz, W., and Attié, J.-L.: Tropospheric carbon monoxide
- variability from AIRS under clear and cloudy conditions, Atmos. Chem. Phys., 13, 12469-12479,
- 893 doi:10.5194/acp-13-12469-2013, 2013.
- 894 Worden, H., Deeter, M., Edwards, D., Gille, J., Drummond, J. and Nédélec, P.: Observations of
- 895 near-surface carbon monoxide from space using MOPITT multispectral retrievals, J Geophys
- 896 Res Atmospheres 1984 2012, 115(D18), doi:10.1029/2010JD014242, 2010.
- 897 Worden, H. M., Deeter, M. N., Frankenberg, C., George, M., Nichitiu, F., Worden, J., Aben, I.,

- 898 Bowman, K. W., Clerbaux, C., Coheur, P. F., de Laat, A. T. J., Detweiler, R., Drummond, J. R.,
- 899 Edwards, D. P., Gille, J. C., Hurtmans, D., Luo, M., Martínez-Alonso, S., Massie, S., Pfister, G.,
- and Warner, J. X.: Decadal record of satellite carbon monoxide observations, Atmos. Chem.
 Phys., 13, 837-850, doi:10.5194/acp-13-837-2013, 2013.
- 902 Worden, J., Wecht, K., Frankenberg, C., Alvarado, M., Bowman, K., Kort, E., Kulawik, S., Lee,
- 903 M., Payne, V., and Worden, H.: CH₄ and CO distributions over tropical fires during October
- 904 2006 as observed by the Aura TES satellite instrument and modeled by GEOS-Chem, Atmos.
- 905 Chem. Phys., 13, 3679-3692, doi:10.5194/acp-13-3679-2013, 2013b.
- 906 Worden, J., Jiang, Z., Jones, D., Alvarado, M., Bowman, K., Frankenberg, C., Kort, E., Kulawik,
- 907 S., Lee, M., Liu, J., Payne, V., Wecht, K. and Worden, H.: El Niño, the 2006 Indonesian peat
- fires, and the distribution of atmospheric methane, Geophys Res Lett, 40(18), 4938–4943,
 doi:10.1002/grl.50937, 2013c.
- 910 Xia, Y., Zhao, Y. and Nielsen, C.: Benefits of China's efforts in gaseous pollutant control indicated
- by the bottom-up emissions and satellite observations 2000–2014, Atmos Environ, 136, 43–53,
 doi:10.1016/j.atmosenv.2016.04.013, 2016.
- 913 Yin, Y., Chevallier, F., Ciais, P., Broquet, G., Fortems-Cheiney, A., Pison, I. and Saunois, M.:
- Decadal trends in global CO emissions as seen by MOPITT, Atmos Chem Phys, 15(23), 13433–
 13451, doi:10.5194/acp-15-13433-2015, 2015.
- 916 Yurganov, L. N., Duchatelet, P., Dzhola, A. V., Edwards, D. P., Hase, F., Kramer, I., Mahieu, E.,
- 917 Mellqvist, J., Notholt, J., Novelli, P. C., Rockmann, A., Scheel, H. E., Schneider, M., Schulz,
- 918 A., Strandberg, A., Sussmann, R., Tanimoto, H., Velazco, V., Drummond, J. R., and Gille, J. C.:
- 919 Increased Northern Hemispheric carbon monoxide burden in the troposphere in 2002 and 2003
- detected from the ground and from space, Atmos. Chem. Phys., 5, 563-573, doi:10.5194/acp-5-

- 921 563-2005, 2005.
- 922 Zhang, Q., Streets, D., Carmichael, G., He, K., Huo, H., Kannari, A., Klimont, Z., Park, I., Reddy,
- 923 S., Fu, J., Chen, D., Duan, L., Lei, Y., Wang, L. and Yao, Z.: Asian emissions in 2006 for the
- 924 NASA INTEX-B mission, Atmos Chem Phys, 9(14), 5131–5153, doi:10.5194/acp-9-5131-2009,
- 925 2009.
- 926 Zhang, L., Li, Q. B., Jin, J., Liu, H., Livesey, N., Jiang, J. H., Mao, Y., Chen, D., Luo, M., and
- 927 Chen, Y.: Impacts of 2006 Indonesian fires and dynamics on tropical upper tropospheric carbon
- 928 monoxide and ozone, Atmos. Chem. Phys., 11, 10929-10946, doi:10.5194/acp-11-10929-2011,
- 929 2011.
- 930 Zhao, Y., Nielsen, C., McElroy, M., Zhang, L. and Zhang, J.: CO emissions in China: Uncertainties
- and implications of improved energy efficiency and emission control, Atmos Environ, 49, 103–
- 932 113, doi:10.1016/j.atmosenv.2011.12.015, 2012.
- 933

934 Tables and Figures

Table 1. Annual total anthropogenic CO emission in different regions, from 2001 to 2015,
 constrained with MOPITT column, profile and lower tropospheric data. The region definition is
 shown in Figure 2e.

Table 2. Annual total biomass burning CO emission in different regions, from 2001 to 2015, constrained with MOPITT column, profile and lower tropospheric data. The region definition is shown in Figure 2f.
 942

Figure 1. Difference between MOPITT CO retrievals and HIPPO aircraft measurements. The
aircraft measurements are smoothed with MOPITT averaging kernels. The black solid line shows
the 4-order polynomial curve fitting, which is used to correct MOPITT data in this work.

Figure 2. (a-d) Mean a priori CO emissions from combustion sources and the oxidation of biogenic
VOCs and CH₄ from 2001 to 2015. The unit is 10¹² molec/cm²/sec. (e-f) Region definitions for (e)
anthropogenic and (f) biomass burning sources.

Figure 3. Schematic diagram for methodology of the assimilation system. Sequential Kalman
 Filter was run from March 1 2000 to December 31 2015 to produce the optimized initial conditions

- (monthly) and boundary conditions (hourly). Monthly 4-DVAR inversions were performed with
 the optimized initial conditions. Only MOPITT data over land (white grids) were assimilated in
- the 4-DVAR inversions, while the CO abundances over ocean (red grids) were defined as
 boundaries and rewritten using the optimized hourly CO fields from Kalman Filter. <u>The Kalman</u>
 filter run is completely independent of the 4-DVAR inversions. There is no feedback of the 4DVAR inversion results to the boundary conditions.

963

- Figure 4. (a) Locations of WDCGG sites with MCF measurements. (b) Global mean MCF
 concentration. (c) Exponential loss rate of MCF, derived from 12-month apart of monthly means
 [e.g., ln(MCF_{Jan2007}/MCF_{Jan2006})]. The black solid line shows the 12-month mean value.
- Figure 5. CO emission trends for 2001 2015, constrained with MOPITT column, profile and
 lower tropospheric profile data. The months dominated by biomass burning emissions are excluded
 from the trend calculation for anthropogenic and biogenic VOC emissions.
- Figure 6. 12-month mean value of anthropogenic CO emissions (with unit Tg/month) for 2001 –
 2015: a priori emission (green) and a posteriori emissions constrained with MOPITT column data
 (black), MOPITT profile data (blue) and MOPITT lower tropospheric profile data (red). The green
 dash line shows the monthly a priori anthropogenic CO emissions. The region definition is shown
 in Figure 2e.
- Figure 7. Monthly mean CO concentrations (green) and 12-month mean value (black) from
 WDCGG stations for 2001 2015. (a) 15-station average in United States (b) 20-station average
 in Europe (c) 2-station (YON and JMA) average in east China outflow (4) Cape Rama (CRI) in
 India.
- Figure 8. Monthly biomass burning CO emissions (with unit Tg/month) for 2001 2015: a priori
 emission (green) and a posteriori emissions constrained with MOPITT column data (black),
 MOPITT profile data (blue) and MOPITT lower tropospheric profile data (red). The region
 definition is shown in Figure 2f.
- Figure 9. Panels (a-d): long-term trend (annual) of surface CO concentration for 2001 2015 from
 WDCGG sites, and model simulations driven with a priori and a posteriori emissions. Panels (e g): effect of a posteriori emissions, derived by abs(Trend_{aposteriori} Trend_{WDCGG}) abs(Trend_{apriori} –
 Trend_{WDCGG}); blue (red) means the a posteriori emissions improves (degrades) the agreement with
 WDCGG measurements compared to the a priori emissions, while white indicates no change from
 the priori. Only stations with more than 10 year observations (the time range between the first and
 last observations) during 2001-2015 are included.
- Figure 10. Panels (a-d): long-term mean value of surface CO concentration for 2001 2015 from
 WDCGG sites, and model simulations driven with a priori and a posteriori emissions. Panels (eg): effect of a posteriori emissions, derived by abs(CO_{aposteriori} CO_{WDCGG}) abs(CO_{apriori} CO_{WDCGG}); blue (red) means the a posteriori emissions improves (degrades) the agreement with
 WDCGG measurements compared to the a priori emissions, while white indicates no change from
 the priori. Only stations with more than 10 year observations (the time range between the first and
 last observations) during 2001-2015 are included.

- **Figure 11.** Long-term trend (annual) of modeled surface and column CO for 2001 2015 with (a-b) all emission sources are fixed at 2001 level. (c-d) variable anthropogenic emissions; (e-f) variable biomass burning emissions; (g-h) variable biogenic VOCs emissions; The variable emissions are constrained with MOPITT profile data.

Page 10: [1] Deleted	Zhe Jiang	1/17/17 10:43:00 AM				
On the other hand, Jia	ang et al. (2015b) indicated that	regional inversions have more				
advantages than global inversions because the boundary conditions can be better controlled. They						
demonstrated that the systematic bias associated with North American CO emissions due to OH						
distribution can be reduced	by up to 50% with optimized	boundary conditions. Similar				

optimization on the boundary condition can also be employed in global model, for example, Pifster et al. (2005) constrained biomass burning CO emissions from boreal North America with optimized CO fields outside the impacted region.

In order to reduce the effects of systematic errors, we designed a two-step inversion to enhance