Comments on revision of "Forecasting Carbon Monoxide on a Global Scale for the ATom-1 Aircraft

Mission: Insights from Airborne and Satellite Observations and Modeling" by Strode et al.

The authors have sufficiently addressed my comments. However, I would like the authors to consider again the relation between the recent negative trend in CO columns and the lower than average CO values in 2016. It is good to discuss the recent CO trend. But, technically, you cannot make the link, even with a "maybe", between the trend and the value in an individual year. Moreover, in Worden et al. (2013) the trend was calculated over 2000-2012. I suggest the authors to

remove the statement "This negative trend may be contributing to the low values in 2016, ..." for clarity.

We thank the referee for the thoughtful comment. We have removed the statement that "This negative trend may be contributing to the low values in 2016, although there is also substantial IAV in CO."

# Forecasting Carbon Monoxide on a Global Scale for the ATom-1 Aircraft Mission: Insights from Airborne and

- 3 Satellite Observations and Modeling
- 4
  - 5 Sarah A. Strode<sup>1,2</sup>, Junhua Liu<sup>1,2</sup>, Leslie Lait<sup>2,3</sup>, Róisín Commane<sup>4</sup>, Bruce Daube<sup>4</sup>, Steven Wofsy<sup>4</sup>, Austin
  - 6 Conaty<sup>2,5</sup>, Paul Newman<sup>2</sup>, Michael Prather<sup>6</sup>
  - 7
  - 8 <sup>1</sup>Universities Space Research Association, Columbia, MD, USA
  - 9 <sup>2</sup>NASA GSFC, Greenbelt, MD, USA
  - 10 <sup>3</sup>Morgan State University, Baltimore, MD, USA
  - 11 <sup>4</sup>Harvard University, Cambridge, MA, USA
  - 12 <sup>5</sup>SSAI, Greenbelt, MD, USA
  - 13 <sup>6</sup>University of California, Irvine, CA, USA
  - 14
  - 15 Correspondence to: Sarah A. Strode (sarah.a.strode@nasa.gov)

# 16

Abstract The first phase of the Atmospheric Tomography Mission (ATom-1) took place in July-August of 17 18 2016 and included flights above the remote Pacific and Atlantic oceans. Sampling of atmospheric 19 constituents during these flights is designed to provide new insights into the chemical reactivity and processes 20 of the remote atmosphere and how these processes are affected by anthropogenic emissions. Model 21 simulations provide a valuable tool for interpreting these measurements and understanding the origin of the 22 observed trace gases and aerosols, so it is important to quantify model performance. GEOS-5 forecasts and 23 analyses show considerable skill in predicting and simulating the CO distribution and the timing of CO enhancements observed during the ATom-1 aircraft mission. We use GEOS-5's tagged tracers for CO to 24 25 assess the contribution of different emission sources to the regions sampled by ATom-1 to elucidate the 26 dominant anthropogenic influences on different parts of the remote atmosphere. We find a dominant contribution from non-biomass burning sources along the ATom transects except over the tropical Atlantic, 27 where African biomass burning makes a large contribution to the CO concentration. One of the goals of 28 29 ATom is to provide a chemical climatology over the oceans, so it is important to consider whether August 30 2016 was representative of typical boreal summer conditions. Using satellite observations of 700 hPa and 31 column CO from the Measurement of Pollution in the Troposphere (MOPITT) instrument, 215 hPa CO from 32 the Microwave Limb Sounder (MLS), and aerosol optical thickness from the Moderate Resolution Imaging

33 Spectroradiometer (MODIS), we find that CO concentrations and aerosol optical thickness in Aug. 2016

34 were within the observed range of the satellite observations, but below the decadal median for many of the

35 regions sampled. This suggests that the ATom-1 measurements may represent relatively clean but not

36 exceptional conditions for lower tropospheric CO.

#### 37 1 Introduction

38 The first phase of the NASA Atmospheric Tomography Mission (ATom-1) 39 (https://espo.nasa.gov/atom) took place in July-August 2016. The aircraft completed a circuit beginning in 40 Palmdale, California and traversing the remote Pacific and Atlantic oceans, providing an unprecedented 41 picture of the chemical environment at a wide range of latitudes over the remote oceans. Major goals of the 42 Atom mission include identifying chemical processes that control the concentrations of short-lived climate 43 forcers, quantifying how anthropogenic emissions affect chemical reactivity globally, and identifying ways 44 to improve the modeling of these processes, 45 Chemical forecasts from the GEOS-5 model provided insight into the chemical environments and sources of pollution for the diverse regions sampled during the ATom-1 campaign. GEOS-5 forecasts help 46 47 determine the source regions and emission types that contribute to the trace gas and aerosol concentrations 48 measured during ATom, which is directly relevant to the goal of quantifying how anthropogenic emissions 49 affect global chemical reactivity. GEOS-5 supports numerous aircraft missions, and validation of the model 50 forecasts is important for developing confidence in and understanding the limitations of chemistry forecasting 51 for aircraft missions. The ATom dataset, which uses unbiased sampling rather than chasing plumes, provides a unique opportunity to validate the overall performance of the GEOS-5 model on a global scale. 52 53 One of the goals of ATom is to provide an observation-based climatology of important atmospheric 54 constituents and their reactivity in the remote atmosphere. Consequently, it is important to examine whether 55 the ATom observations are temporally and spatially representative of the broader remote atmosphere. Prather 56 et al. (2017) examined the ability of observations from a single path to represent the variability of a broader 57 geographic region, but noted that year-to-year and El Nino/Southern Oscillation (ENSO) variability could 58 also be important. Year to year variability in meteorology and emissions both contribute to interannual 59 variability in trace gases and aerosols, so it is important to consider the temporal representativeness of a 60 single season sampled by ATom. For example, ENSO is a major driver of variability in ozone distributions 61 (Ziemke and Chandra, 2003), and large biomass burning events during El Nino years increase concentrations 62 of trace gases including CO and CO2 (Langenfelds et al., 2002). Biomass burning plays a particularly strong 63 role in driving the interannual variability of CO (e.g. Novelli et al., 2003; Kasischke et al., 2005; Duncan and 64 Logan, 2008; Strode and Pawson, 2013; Voulgarakis et al., 2015). The impacts of large biomass burning 65 events during El Nino events are visible in satellite observations of CO (e.g. Edwards et al., 2004; Edwards 66 et al., 2006; Logan et al., 2008, Liu et al., 2013). Pfister et al. (2010) used a chemistry transport model (CTM) 67 as well as satellite data to examine the CO sources and transport over the Pacific during the INTEX-B mission

Deleted: ,

Λ	Formatted: Font: Not Italic
λ	Deleted: ,
( )	Formatted: Font: Not Italic
//	Formatted: Font: Not Italic
//(	Formatted: Font: Not Italic
//(	Formatted: Font: Not Italic
//(	Formatted: Font: Not Italic
7,(	Formatted: Font: Not Italic
//	Formatted: Font: Not Italic
Ά	Deleted: ;
Â	Formatted: Font: Not Italic
. (	Formatted: Font: Not Italic
1	Formatted: Font: Not Italic
(	Formatted: Font: Not Italic
~(	Formatted: Font: Not Italic

compared to previous years. They found biomass burning to be the largest contributor to interannual
variability, despite its lower emissions compared to fossil fuel sources. Here we show how the time and place
of ATom-1 measurements fit into a global, multi-year climatology of CO. In particular, we assess the extent
to which measurements from the ATom-1 period represent the CO and aerosol distributions over the last
decade and a half.
In this study, we place the August 2016 ATom observations in the context of interannual variability and
assess the contributions of different emission sources to the various regions sampled during the campaign.

We focus on CO, a tracer of incomplete combustion whose lifetime of 1-2 months allows long-range transport to the remote oceans. Section 2 describes the model and observations used in this analysis. Section 3 compares the GEOS-5 CO to observations. Section 4 discusses the global distribution of CO<sub>e</sub> and presents.

81 the relative CO source contributions to the regions sampled by ATom. Section 5 presents an analysis of the 82 interannual variability in CO and aerosol optical thickness seen in satellite observations to assess how well

August 2016 observations represent climatological August conditions. Section 6 summarizes our
 conclusions

#### 85 2 Observations and Model

## 86 2.1 ATom Observations

87 ATom-1 flew transects through the Pacific, Southern, Atlantic and Arctic oceans with the NASA DC8 88 aircraft in August 2016. Each of the 11 flights included sampling from the boundary layer to the top of the 89 aircraft range (39 kft). We use the ATom-1 data (July-August 2016) (Atom Science Team, 2017; Wofsy et 90 al., 2018), for comparison with the model forecasts and analyses. 91 We take ATom-1 CO observations from the Harvard QCLS instrument (Santoni et al., 2014), which 92 has a history of successful measurements during the HIAPER Pole-to-Pole Observations (HIPPO) campaign. 93 Briefly, the instrument uses a pulsed quantum cascade laser at 2160 cm<sup>-1</sup> to measure absorption of CO through an astigmatic multi-pass sample cell (with 76 m path length), with detection using a liquid nitrogen cooled 94 HgCdTe detector. A separate laser and detector are used to measure methane and nitrous oxide in the same 95 cell. Inflight calibrations were conducted with gases traceable to the NOAA WMO (X2014) scale, with 96 97 calibration of tanks before ATom1 and after ATom2 (February 2017) showing no significant change in the 98 CO concentration in the gas standards. The inlet for the instrument was specially designed for the DC-8 99 aircraft. The QCLS observations have an accuracy and precision of 3.5 ppb and 0.15 ppb, respectively. The 100 QCLS observations used in this analysis are being archived at the ORNL DAAC 101 (https://doi.org/10.3334/ORNLDAAC/1604). 102

Deleted:

Deleted: [	
Formatted: Font: Not Italic	
Deleted: ]	
Formatted: Font: Not Italic	

#### 106 2.2 Satellite Observations

107 We use satellite observations that cover more than a decade to examine the interannual variability 108 of CO and aerosols. We focus on satellite observations because they provide broad coverage over the oceans, 109 where surface data is sparse. The Measurement of Pollution in the Troposphere (MOPITT) instrument, which 110 flies on the Terra satellite, provides CO observations beginning in 2000 (Edwards et al., 2004). We use the 111 version 6 thermal infrared (TIR) level 3 product (Deeter et al., 2014). The MOPITT TIR averaging kernels 112 show high sensitivity to CO between 700 and 500 hPa (Emmons et al., 2007). We use the CO column and 113 700 hPa CO retrievals. 114 The Microwave Limb Sounder (MLS) (Waters et al., 2006), which flies on the Aura satellite, 115 provides useful observations of CO down to 215 hPa (Livesey et al., 2008) beginning in 2004. We use the 116 Version 4.2 level 2 data for the 215 hPa level with the recommended quality, status, precision, and 117 convergence criteria. Although MLS data overlap with ATom only at the highest flight levels, MLS and 118 MOPITT provide complementary views of CO in the upper troposphere/lower stratosphere (UTLS) and 119 lower troposphere, respectively. The MOPITT averaging kernels include some sensitivity to the 200 hPa

120 level, implying a small overlap between the MOPITT and MLS observations.

121 The Moderate Resolution Imaging Spectroradiometer (MODIS) instrument on the Aqua satellite

122 provides column aerosol optical thickness (AOT) data beginning in 2002. We use MODIS data in this

analysis because it provides a relatively long data record. We use the Collection-6 level 2 (MYD04\_L2)

124 (Levy et al., 2015) 550 nm AOT data over oceans aggregated into 0.5 degree grid boxes, and then take

125 monthly means with the daily data weighted according to the QA.

# 126 2.3 Model Description

127 We use chemical forecasts and analyses from the GEOS-5 Forward Processing (FP) system to 128 quantify the contribution of different emission sources to the observed CO distribution and to identify the origin of observed plumes. A global model is necessary for this analysis since CO is transported globally. 129 The FP stream from the Global Modeling and Assimilation Office (GMAO) generates GEOS-5 forecast 130 131 products as well as assimilation products using the most current system approved for near-real-time 132 production. We use the FP system in our study because it is the system used to generate forecasts that are 133 used during ATom and other aircraft missions, and is thus relevant to future mission and flight planning. The 134 GEOS-5 model (Molod et al., 2015) is a global general circulation model (GCM) with 72 vertical levels 135 reaching from the surface to 1 Pa. The assimilation system is described in (Rienecker et al., 2008, 2011), 136 and includes assimilation of ozone measurements from the Ozone Monitoring Instrument (OMI) and MLS, 137 and aerosol optical depth as well as meteorological variables. The forward processing system produces 138 output on 72 model levels or 42 pressure levels with 5/16 by 1/4 degree horizontal resolution. Our study uses 139 the pressure level output. 140 The GEOS-5 FP system (Lucchesi, 2017) simulates the transport of CO as well as tagged CO tracers

141 from specific regions and sources, which helps track the transport of pollution outflow. Tagged tracers are

Formatted: Font: Not Italic	
Formatted: Font: Not Italic	
Formatted: Font: Not Italic	

Formatted: Font: Not Italic

Deleted: both
Deleted: it
Deleted: lower troposphere and

#### Deleted: [

Formatted: Font: Not Italic

(	Formatted: Font: Not Italic
-(	Formatted: Font: Not Italic
~(	Deleted: ; Rienecker et al.

Formatted: Font: Not Italic

148 available for biomass burning (BB) globally as well as biomass burning from Eurasia, North America, Africa, 149 and Central and South America; and for non-BB sources globally and from Europe, Asia, and North America. 150 Non-BB sources include fossil fuels, biofuels, CO from oxidation of biogenic VOCs, and CO from methane 151 oxidation, as described in Ott et al. (2010). Supplemental Figure S1 shows the regions included in each 152 tagged tracer. Bian et al. (2013) used observations of dichloromethane and acetonitrile from the ARCTAS 153 mission to validate the anthropogenic and biomass burning CO tracers, respectively. 154 Daily-varying biomass burning emissions come from the Quick Fire Emission Dataset (QFED) 155 version 2 (Darmenov and da Silva, 2015), which is based on fire radiative power from the MODIS instrument. 156 Thus the BB emissions include day-to-day and interannual variability, but the non-BB sources and the OH fields use monthly means and lack daily-scale variability and interannual variability. Table 1 presents the 157 158 August emission inputs for the major regions considered. CO emissions are then scaled up by 20% for fossil 159 fuels and 11% for biomass burning to account for CO production from co-emitted VOCs, since VOC's are 160 not explicitly carried in the GEOS-fp chemical mechanism. This approach was developed by Duncan et al. 161 (2007) to account for the CO source from non-methane hydrocarbon oxidation. 162 CO from methane oxidation is included in the non-BB tagged tracers for the regions in which 163 oxidation occurred. For example, if methane is oxidized over North America, the resulting CO is included 164 in the North American non-BB tracer. The monthly mean methane fields come from a GMI Chemistry and 165 Transport Model (CTM) simulation, which uses prescribed zonal mean surface concentrations. CO is lost by reaction with OH using fixed monthly OH fields archived from the GMI CTM. Supplemental Figure S2 166 shows the methane and OH fields included in the FP system. 167 168 3 **GEOS-5** Chemical Forecasting for ATom

169 During the ATom mission, the GEOS-5 model was engaged to provide chemical forecasts for each 170 flight that include the major chemical species and, for CO, tagged tracers for different sources. The chemical 171 forecasts were used together with meteorological forecasts for day-to-day flight planning, although flight 172 tracks were intentionally not altered to chase specific chemical features to avoid a highly biased sampling of 173 pollution. The chemical forecasts provide the ATom team with a preview of the chemical environments that 174 the flight is expected to sample, including the location of pollution, biomass burning, or dust plumes; regions 175 of substantial but well-mixed anthropogenic pollution; and cleaner regions. The forecasts also provide a 176 broader spatial context for the observations, since the 3-dimensional model output shows the spatial extent 177 of features that intersect the flight track. 178 We examine the performance of the GEOS-5 forecasts by comparing the simulated CO to the QCLS 179 observations. The forecasts provided during the mission used forecast wind fields, with the forecast lead 180 time varying depending on the timing of the flight. For consistency, the results shown here use the CO 181 simulated with the assimilated wind fields, but we note that similar features were seen for the CO simulated 182 with the forecast winds, as further discussed in section 3.1.2. For the model results, we do not apply temporal

183 interpolation between the model output frequency (6 hours). Instead, we sample the model forecasts at the

Formatted: Font: Not Italic	
Formatted: Font: Not Italic	

 Deleted:	ſ
Deleteu.	L

Formatted: Font: Not Italic

Deleted: ]

Formatted: Font: Not Italic

Deleted: i

Deleted: ar

time closest to the mid-point of each flight segment. To compare with observations, the 3D model forecast was interpolated to the longitude, latitude and pressure given in the 10-second merges of the ATom measurements.

### 191 3.1 Analysis of CO along the Meridional Flight Tracks

We compare CO from GEOS-5 to the QCLS CO observations for specific flights, using the 10second merge files. The GEOS-5 CO is taken from the 3D field at the time closest to the mid-point of the flight and interpolated in space to the flight track. The ATom flight tracks are shown in Supplemental Figure S3. We focus on two sections of the ATom-1 circuit: the North to South flights through the Pacific, and the South to North flights through the Atlantic, although we briefly discuss the other flights as well. These two transects allow us to examine the transition from northern hemispheric to tropical to southern hemispheric influence.

#### 199 3.1.1 Pacific legs

Figure 1 shows CO from the three Pacific flights spanning Anchorage, Alaska to Christchurch, New 200 201 Zealand. The top panels show the GEOS-5 curtain of CO along the flight track, with the QCLS observations 202 overplotted in circles. The observations show higher values of CO in the first half of the Anchorage-Kona 203 flight compared to the other portions of the Pacific, and this feature is reproduced in GEOS-5 as well. GEOS-204 5 agrees well with the observed mean value for CO on this flight (Table 2). Tagged tracers (Fig. 1 bottom 205 panels) show that non-BB sources, especially from Asia, are the dominant contributor to CO levels throughout the Pacific, and the decrease in Asian non-BB CO explains the observed decrease in CO as the 206 207 flights move south.

208 The observations show plumes of enhanced CO scattered throughout all three Pacific flights, 209 although they are most intense in the north Pacific, as seen in the Anchorage-Kona flight. GEOS-5 typically 210 reproduces the timing of these plumes, but the magnitude is usually underestimated, particularly for the 211 strongest plumes. This leads to an underestimate of the observed standard deviation of the CO on the 212 Palmdale-Anchorage (Fig. S4) and Anchorage-Kona flights (Table 2). In addition to biases in emissions, 213 observations often show fine-scale structures too small for the model to resolve (Hsu et al., 2004), and 214 underestimating the concentrations in strong plumes is a common problem for global models (e.g. Heald et 215 al., 2003). Either biases in emissions or insufficient vertical or horizontal model resolution may thus be 216 responsible for the underestimate. The tagged tracer for biomass burning shows a small increase at the time 217 of the underestimated plumes near hour 22 of the Anchorage-Kona flight (Fig. 1d,g), suggesting that those 218 underestimates are due to the insufficient magnitude of the simulated biomass burning plumes. An exception 219 is in the tropical Pacific (Kona-Pago Pago flight), in which GEOS-5 predicted some enhancements, driven 220 by fossil fuels, not seen in the observations. Tagged tracers indicate that Asian non-BB CO drove many of 221 the observed enhancements, while others were due to biomass burning. 222

In the south Pacific (Pago Pago to Christchurch segment), the flight sampled the stratosphere three times, with CO levels decreasing to approximately 30 ppb, as shown in Fig. 1c. As expected from Formatted: Font: Not Italic

224 stratospheric chemistry and seen in previous observations, both ATom-1 and GEOS-5 show a strong decrease

225 in CO as the flight rises above the tropopause, with GEOS-5 underestimating the observed gradient. Both

226 the model and measurements show tropospheric CO less than 90 ppbv along the flight route with slightly

227 elevated CO above 600 hPa around T22:00 and T24:00. For this flight and the subsequent flight to Punta

228 Arenas, all observations are in the Southern Hemisphere and the mean values for both ATom-1 and GEOS-

5 agree within the range 54-57 ppb (Table 2, Fig S5).

# 230 3.1.2 Atlantic Legs

231 The ATom flights traversed the Atlantic from South to North, beginning in Punta Arenas, Chile and 232 ending in Kangerlussuaq, Greenland. Figure 2 shows the Atlantic flights from Punta Arenas to Ascension Island to the Azores to Kangerlussuaq. GEOS-5 has an excellent simulation of background CO values seen 233 234 on these flights, with the mean values falling within 2 ppb of the observations (Table 2) while the mean observed values for each flight shift from 69 to 101 to 88 ppb. The observations show plumes of high CO 235 236 intersecting the flight track on all three flights. GEOS-5 also shows plumes of enhanced CO at these 237 locations, but the magnitude is often underestimated (Fig. 2d-f), especially for the Azores-Kangerlussuaq 238 flight. Supplemental Figure S6 shows the CO results for the Azores-Kangerlussuag flight using forecast 239 wind fields and illustrates the temporal evolution of CO plumes along the flight track. Comparison of Fig. 240 S6 with Fig. 2f shows that the impact of using analysis versus forecast wind fields is small for this flight 241 since the forecasts already capture the timing of the plumes. 242 Non-BB sources dominate the background CO levels on all three flights. However, biomass burning 243 plays a dominant role in the plumes of high CO (Fig. 2g-i). South American biomass burning leads to CO enhancements between T14 and T16 of the Punta Arenas to Ascension flight. In the later portion of that 244 flight, biomass burning from Africa leads to strong CO plumes. Strong plumes of African biomass burning 245 are also seen at the beginning of the Ascension to Azores flight. GEOS-5 shows a strong plume around 800 246 247 hPa for the first hour of the flight, which agrees well with observations (Fig. 2b,d). The observations show 248 additional strong plumes in the next hour between 600 and 700 hPa. These plumes are present but underestimated in GEOS-5, possibly due to errors in the magnitude of the emissions or the placement or 249 250 extent of the plumes. The placement and strength of simulated plumes is sensitive to the injection height of 251 the biomass burning, which is a source of uncertainty. In addition, plumes in models tend to dissipate more 252 quickly than in observations due to the numerical effects of limited model resolution (Eastham and Jacob, 253 2017). 254 The non-BB contribution to CO in the Atlantic reflects a mixture of global sources. Asian sources

255make a notable contribution to the non-BB CO variability in the tropics (first half of the Ascension to Azores256flight), but as expected N. American sources become more dominant in the second half. In the northern257(later) portion of the Azores to Kangerlussuaq flight (Fig. 2), GEOS-5 attributes the observed plumes to

Eurasian biomass burning, but underestimates their magnitude. This flight also crosses the tropopause, and

259 both ATom-1 and GEOS-5 show a corresponding dip in CO concentrations. GEOS-5 predicts a plume of

Deleted:

Formatted: Font: Not Italic

Deleted:

262 enhanced CO due to N. American emissions around 11Z of the Azores to Kangerlussuaq flight that is not

seen in the observations (Fig. 2f,i). A similar error is made in the Kangerlussuaq to Minneapolis flight (Fig.

264 S5). This could be due to either an error in the assumed N. American sources, or to misplacement of the

265 plume by the model. A large overestimate of CO at the end of the Minneapolis to Palmdale flight also points

266 to a potential error in North American emissions from either fossil fuels or biomass burning.

## 267 3.2 Model Evaluation Summary

268 We summarize the comparison between the CO simulated by the GEOS-5 analyses and the QCLS observations in Figure 3. The majority of points lie near the one-to-one line, indicating good overall 269 270 agreement between the GEOS-5 and observed CO distributions. The higher concentrations in the tropical 271 Atlantic compared to the tropical Pacific are evident in both the observations and model. Fig. 3 also reveals 272 occasional model overestimates of CO on flights over North America (green triangles), as well as 273 underestimates of high CO plumes over the North Pacific and Tropical Atlantic. An underestimate of 274 Eurasian biomass burning contributes to the model underestimates in the North Pacific and North Atlantic, 275 and has implications for ozone production in aged BB plumes. Globally, the correlation of simulated and 276 observed CO with 5-minute binning is r=0.69. Correlations for the Pacific, Atlantic, and North America are 277 0.72, 0.80, and 0.80, respectively, while the correlation for the southern ocean is 0.053. The poor correlation 278 for the southern ocean reflects the very low variability of CO in this region. The model performs far better 279 at capturing the larger gradients present in the other regions. In general, the good agreement between model 280 outputs and observations testify the model forecasting skill and suggest the suitability of using GEOS-5

# 281 forecast products to guide the design and execution of aircraft campaigns.

#### 282 4 Source Contributions to the Global CO Distribution

# 283 4.1 Global CO Distribution

Figure 4 compares CO from GEOS-5 to the QCLS CO observations for the ATom-1 circuit including the 11 total flight segments. The GEOS-5 CO is taken from the analysis closest to the mid-point of the flight time and interpolated to the flight track following the longitude, latitude and pressure given in the observations. We average both model CO and ATom measurements into one point per 360-seconds for easier visualization.

Both model simulations and measurements show polluted air with higher CO mixing ratios in the northern hemisphere than that in the southern hemisphere in August 2016. Over the northern hemispheric polar region, the observations indicate highly polluted air with CO maxima occurring over Alaska and northwest Canada, features also seen in the GEOS-5 simulation. Over the Atlantic section, CO maxima with slightly lower values occur around the same latitude over west Greenland as shown both in observations and model simulation. CO over the northern most locations along the ATom-1 circuit see some low values both in model and observations, particularly north of 30°N and south of 40°S, due to the measurements occurring **Deleted:** [Liu et al., in prep] **Formatted:** Font: Not Italic

297 in the stratosphere or occurring in the upper troposphere with stratospheric influence. Both model and 298 observations indicate that the air is relatively clean over the Pacific south of 30N with CO less than 70 ppb 299 and the CO minimum around 60S over the southeast Pacific. Over the Atlantic section, both model and observations show low CO concentration south of 30S, but show a strong CO maximum over the tropical 300 301 Atlantic (5S-5N) with CO greater than 120 ppb. This high CO is mainly driven by southern hemisphere BB. 302 CO is slightly lower between 30N and 60N compared to that over tropical Atlantic and the Greenland. The similarity between GEOS-5 and ATom-1 variability in neighboring points is due in part to the vertical 303 304 profiling which places horizontally extensive biomass burning layers in both model and presumably the 305 atmosphere at the same point along the track.

#### 306 4.2 CO Source Contributions

We calculate the contribution of different CO sources to the total simulated CO using the GEOS-5 tagged CO tracers sampled along the ATom flight tracks. This analysis provides a picture of the dominant sources affecting the constituent concentrations observed during ATom-1 for different regions of the atmosphere. The tagging of CO sources includes both biomass burning (BB) and non-biomass burning (non-BB) from four continental areas, with all other sources put into the "other" bin. Other BB sources are small, but other non-BB sources are quite large as they include all natural sources as well as atmospheric photochemical sources such as methane oxidation.

314 Figure 5 shows the contribution of each tagged tracer over the Pacific Ocean from 120°E to 110°W, 315 averaged over 5 degree latitude bins. Non-biomass burning sources dominate at all latitudes, due in part to 316 the inclusion of CO from methane oxidation in addition to fossil fuel sources in these tracers. The oxidation 317 of methane over the remote oceans contributes to the large magnitude of "other non-BB" sources over the southern latitudes of the Pacific. Asian non-BB sources make the largest contribution to middle and upper 318 tropospheric CO (Fig. 5a) at the mid-latitudes of the North Pacific, with smaller contributions from N. 319 320 American and European non-BB sources. The largest biomass burning contribution comes from Africa in 321 the Southern Hemisphere and Tropics, switching to Eurasia in the northern latitudes.

322 Figure 5b shows the relative contributions in the lower troposphere, including the marine boundary 323 layer and defined here as pressures greater than 850 hPa. Missing bars indicate latitudes where no ATom-1 324 measurements were made in the lower troposphere. Asian non-BB CO makes a smaller contribution in the 325 lower troposphere than in the middle and upper troposphere. A strong CO maximum around 30°N is more 326 pronounced in the lower troposphere than above. This bin is not representative of the remote Pacific as it 327 includes Palmdale, California, with large contributions from local North American BB and non-BB sources. 328 The Atlantic flights (0°-60°W) show a large contribution from other non-BB sources in the Southern 329 Hemisphere with increasing contributions from Asian, N. American, and European non-BB CO as the flight moves northward (Fig. 6), similar to the picture over the Pacific. However, the Atlantic receives a larger 330 331 contribution from biomass burning, particularly from Africa, over the Tropics. The contribution from African BB is strong throughout the troposphere, but is particularly pronounced in the lower troposphere, where it exceeds 100 ppb in the bins centered at 10°S and 5°N.

334 We also examine the tagged tracer contributions for each flight, including all altitudes sampled by the flight (Fig. 7, Supp. Table S1). Flights occurring in the tropics and southern hemisphere (Flt. 1, 4-8) 335 336 receive 44-75% of the total CO from other non-BB sources. Other non-biomass burning sources include all 337 non-biomass burning sources located outside North America, Europe, and Asia. The contribution from 338 methane oxidation in addition to southern hemisphere emissions explains this large contribution. Flight 8 339 has a somewhat lower percent contribution from other non-BB sources than the other southern hemisphere 340 and tropical flights due to the higher percent contribution from African biomass burning. In contrast, the Northern Hemisphere flights have a larger contribution from northern hemisphere source regions. Asian 341 342 non-BB explains over a third of the total CO for the northern Pacific flights (Flt. 2-3), while Asian and N. 343 American non-BB sources make comparable contributions to the North Atlantic and N. American flights (Flt. 344 9-11).

Since Figs. 5 and 6 reveal differences in source contributions between the lower troposphere and the middle and upper troposphere, we also examine the source contributions to each flight for the lower troposphere (Pressure > 850 hPa) only (Supp. Fig. S7). Asian sources make a larger percent contribution to the Pacific flights (Flt. 0-4) when all flight altitudes are considered rather than the lower troposphere alone. Regional sources such as African biomass burning for flights 6 and 7 and N. American sources for Flights 9 and 10 make a larger percent contribution in the lower troposphere.

#### 351 5 August 2016 in the Context of Interannual Variability (IAV)

352 One of the major goals for the ATom campaign is to produce a climatology based on un-biased, 353 representative samples (Prather et al., 2017). It is therefore important to consider whether August 2016 is a 354 "typical" boreal summer/austral winter month. Prather et al. 2018, found differences of 8-10% in the 355 chemical reactivity of model simulated air parcels when considering other years compared to 2016. We focus 356 here on the temporal representativeness of the ATom-1 campaign. August of 2016 was ENSO neutral, with 357 a multivariate ENSO index (MEI) (Wolter and Timlin, 1993); (https://www.esrl.noaa.gov/psd/enso/mei) of 358 0.175 for July/August. However, it was preceded by strong El Nino conditions in 2015 and early 2016 359 (Blunden and Arndt, 2016). We therefore consider whether the CO concentrations in August 2016 are typical 360 or anomalous. 361 Multi-year satellite records provide a valuable tool for determining how CO concentrations in the

regions of the ATom-1 flights compare to previous years. We focus our analysis of CO interannual variability on several regions traversed by the ATom flights. Figure 8 shows these regions in black squares overplotted on the MOPITT CO column for August 2016. We also examine the IAV in BB sources from nearby regions, outlined in red on Fig. 8. Figure 9 shows box-and-whisker plots of the mean, minimum, 25<sup>th</sup>, 50<sup>th</sup>, and 75<sup>th</sup> percentiles, and maximum in monthly mean August CO for each region over the 2000-2016 period for MOPITT (CO column and CO at 700 hPa) and 2004-2016 for MLS (CO at 215 hPa). The corresponding

(	Formatted: Font: Not Italic
(	Deleted: [
$\sim$	Deleted: ]
Y	Formatted: Font: Not Italic
·····	<b>Deleted:</b> Spatial representativeness is investigated in <i>Liu et al.</i> [in prep].
$ \rightarrow $	Formatted: Font: Not Italic
(	Formatted: Font: Not Italic

#### Deleted:

373 time series are shown in Fig. 10. The variability in CO BB emissions from the Global Fire Emissions 374 Database version 4 (GFEDv4) (van der Werf et al., 2017) for 2000-2016 is also shown for BB regions that 375 may affect the ATom flights. The BB emissions are averaged over June through August to account for the 376 persistence of CO in the atmosphere. 377 Among the regions mapped here, the tropical Atlantic shows the highest average CO values, as well 378 as the highest 2016 CO values, in both MOPITT and MLS observations (Fig. 9). This is consistent with large 379 biomass burning emissions from southern hemisphere (SH) Africa transported into the tropical Atlantic. While SH Africa has the largest magnitude of biomass burning, its relative variability (variability relative to 380 the mean) is smaller than for the other regions (Fig. 9b). Similarly, the IAV in the MOPITT CO column and 381 700 hPa level over tropical Atlantic is smaller than that of the North Atlantic and Alaska regions. Although 382 383 the variability of CO over tropical Atlantic is relatively small, the MOPPIT CO column shows a statistically

384 significant anti-correlation between the MOPITT CO column over the tropical Atlantic and the MEI (r--

385 0.52). This relationship is not significant for the MOPITT 700 hPa level.

The time series of August MOPITT CO columns for both the North Atlantic and Alaska, regions 386 387 that show high variability, show a small but significant temporal correlation with summertime Siberian biomass burning (r=0.52 for the North Atlantic and r=0.59 for Alaska). Slightly lower values are seen for 388 389 the 700 hPa MOPITT level. The time series of August MOPITT 700 hPa CO shows an increase in 2003 for 390 the North Atlantic and in 2002 and 2003 for Alaska (Fig. 10b,c). Previous studies attribute peaks in these 391 years to the presence of large forest fires in western Russia and Siberia, respectively (Edwards et al., 2004; 392 Yashiro et al., 2009; van der Werf et al., 2006). MOPITT CO values were below average in 2016 for both 393 the North Atlantic and Alaska even though Siberian biomass burning was above average in 2016 (Fig. 9a, b).

Since ENSO is known to drive large biomass burning variability in Indonesia (van der Werf et al.,
 2006), we consider whether it may influence CO concentrations over the New Zealand region. Although the
 MOPITT CO column over the Indonesia region does correlate with the MEI (r=0.64), there is no significant

397 correlation between June-Aug biomass burning in Indonesia and MOPITT CO over New Zealand. However,

398 August is not the peak season for Indonesian biomass burning (Duncan et al., 2003b). The large Indonesian

399 fires that occurred during the strong 1997/1998 El Nino peaked during September to November (Duncan et

400 al., 2003a) and active fire detections for the 2015 Indonesian fires peaked in September and October (Field

401 et al., 2016). Thus we might expect Indonesian biomass burning variability to have a greater influence on

402 CO variability during the September-October season, which was sampled in ATom-3.

How does 2016 compare to previous years? The MOPITT CO column shows tropical Atlantic CO
was near the 75<sup>th</sup> percentile, while the 700 hPa MOPITT level shows it close to the median. This difference
arises because the MOPITT column also includes information from the upper troposphere, and the MOPITT
200 hPa level (not shown) suggests CO levels for 2016 were near the 75<sup>th</sup> percentile. In contrast, MLS shows
that 2016 CO in the upper troposphere was much lower than average, near the 25<sup>th</sup> percentile. The MOPITT
v6 TIR product has a small positive bias drift in the upper troposphere of 0.78 % yr-1 for the 200 hPa level

Formatted: Font: Not Italic

Formatted: Font: Not Italic

Formatted: Font: Not Italic Formatted: Font: Not Italic Formatted: Font: Not Italic 409 (Deeter et al., 2014), which may contribute to the higher rank of 2016 in the MOPITT upper tropospheric 410 data compared to MLS. It is therefore hard to argue that 2016 was outside of the normal IAV for this region. 411 2016 CO in the North Atlantic and Alaska regions was below average in both the MOPITT column 412 and the 700 hPa level, and is in fact the lowest August value in the MOPITT record for the 700 hPa level over Alaska. MLS also shows moderately low CO in the upper troposphere over Alaska in Aug. 2016. 413 414 Combined, this data suggests that the ATom-1 CO is not typical for the region. August 2016 CO column 415 values are also below the median over New Zealand and the eastern and central tropical Pacific, but the 416 relatively low variability of these regions makes this less of a concern for the representativeness of the ATom measurements. The IAV of these regions is larger for the MOPITT 700 hPa level, and 2016 lies slightly 417 below the 25<sup>th</sup> percentile for this level. 418 419 The regionally-averaged 500 nm AOT from MODIS (Fig. 11) shows similar features to the MOPITT 420 column. The highest values are found for the tropical Atlantic, followed by the Alaska and North Atlantic 421 regions. This similarity is consistent with the importance of biomass burning emissions for both CO and 422 aerosols. However, the difference between the tropical Atlantic and the other regions is larger in the aerosol 423 case, while the difference between the North Atlantic and the Pacific regions is smaller. There is also greater 424 relative year-to-year variability over the tropical Atlantic for the aerosols than for CO. The shorter lifetime 425 of aerosols compared to CO, as well as the large contribution from biomass burning, likely explains the greater prominence of the tropical Atlantic in the aerosol case. Furthermore, AOL (Fig. 12) shows a clear 426 427 peak in 2009 in several of the regions, whereas MOPITT data is missing for Aug. 2009, but MLS shows a minimum (Tropical Atlantic) or no anomaly (other regions). 428 429 In summary, the multi-year satellite record shows considerable variability in CO, particularly over the 430 North Atlantic and Alaska. Concentrations during August 2016 were on the low end of the distribution for 431 most regions, especially in the lower troposphere. Worden et al. (2013) showed negative trends in the 432 MOPITT CO column significant at the one-sigma level for both the Northern and Southern hemispheres for 433 2000-2012. In addition, Deeter et al. (2014) report a small negative bias drift in the MOPITT V6 TIR product 434 in the lower troposphere, although drift in the column is almost negligible. Decreasing MOPITT CO over 435 time is also visible in some regions in Fig. 10. Overall, the year 2016 shows anomalies for some regions, but 436 does not appear to be an extreme year.

#### 437 6 Conclusions

We place the observations from the ATom-1 campaign in the context of interannual variability and global source distributions using satellite observations and tagged tracers from GEOS-5, respectively. GEOS-5 gives a reasonable reproduction of the background CO levels for most flights despite the use of climatological fossil fuel and biofuel emissions, and captures the global distribution of CO observed during ATom-1. Simulations with both forecast and analysis winds capture the timing of many of the enhanced CO plumes encountered during the flights, although the magnitude of the enhancements was often underestimated, which is not unexpected given the difference in resolution between the observations and Formatted: Font: Not Italic

Deleted: D

Formatted: Font: Not Italic

Formatted: Font: Not Italic

**Deleted:** This negative trend may be contributing to the low values in 2016, although there is also substantial IAV in CO.

model. The strong performance of GEOS-5 with regards to the overall CO distribution and the timing of the
 enhancements gives us confidence in using tagged tracers to identify the sources affecting the air sampled in
 ATom-1.

452 We find that for most flights the dominant contribution to total CO is from non-biomass burning 453 sources, which include both fossil fuels and biofuels and oxidation of hydrocarbons including methane. An 454 exception to this is in the lower troposphere of the tropical Atlantic, where biomass burning from Africa 455 makes the largest contribution, exceeding 100 ppb in some locations. The non-BB source includes a large fraction from Asia for flights over the North Pacific and from both Asia and North America for the North 456 Atlantic and North American flights, while other regions dominate in the Southern Hemisphere. Plumes of 457 elevated CO from both biomass burning and non-BB sources led to observations of enhanced CO during 458 459 ATom-1

460 We use satellite observations of CO from MOPITT and MLS and AOT from MODIS to assess whether August 2016, the period sampled by ATom-1, is typical or atypical in the context of IAV in the satellite 461 record (2000-2016). MOPITT and MLS show that CO in the lower and upper troposphere, respectively, 462 were below average in August 2016 compared to the satellite record for August for most of the regions 463 464 sampled by ATom-1, but not usually the minimum year. CO concentrations in the North Atlantic and Alaskan 465 regions show a positive correlation with Siberian biomass burning and large interannual variability. In 466 contrast, both MODIS AOT and the MOPITT CO column show above average values for the Tropical Atlantic in 2016. This suggests that the high values of CO and aerosols from biomass burning encountered 467 during the tropical Atlantic portions of ATom may have been especially pronounced during this particular 468 469 vear.

470 The seasonality of biomass burning, the OH distribution, and atmospheric transport pathways can alter 471 the source contributions from season to season. Thus, the next three ATom campaigns, which occur in 472 different seasons, will likely show variations in the relative source contributions to each region.

473

# 474 Data Availability

The QCLS data version used in this paper and the corresponding tagged-CO model output will be available
from the ORNL DAAC at <u>https://doi.org/10.3334/ORNLDAAC/1604</u>. MOPITT data is available at
https://eosweb.larc.nasa.gov/datapool. MLS data is available from https://mls.jpl.nasa.gov/. MODIS aerosol
data are available from https://ladsweb.modaps.eosdis.nasa.gov/api/v1/productGroupPage/name=aerosol.
Acknowledgements

- 481 The authors thank the NASA GMAO for providing the GEOS-5 forecasts and analyses. We thank the NASA
- 482 Earth Venture Suborbital Program, ESPO, and the pilots, crew and support staff of the DC-8.
- 483
- 484
- 485 References

- 486 ATom Science Team (2017), Moffett Field, CA, NASA Ames Earth Science Project Office (ESPO),
- 487 Accessed at doi: 10.5067/Aircraft/ATom/TraceGas\_Aerosol\_Global\_Distribution
- 488 Bian, H., Colarco, P., Chin, M., Chen, G., Rodriguez, J., Liang, Q., Blake, D., Chu, D., da Silva, A.,
- 489 Darmenov, A., Diskin, G., Fuelberg, H., Huey, G., Kondo, Y., Nielsen, J., Pan, X., and Wisthaler, A.: Source
- 490 attributions of pollution to the Western Arctic during the NASA ARCTAS field campaign, Atmospheric
  491 Chemistry and Physics, 13, 4707-4721, 10.5194/acp-13-4707-2013, 2013.
- 492 e.d. Blunden, J., and D. S. Arndt: State of the Climate in 2015. In: Bull. Amer. Meteor. Soc., 97(8), 2016.
- 493 Darmenov, A. and A. da Silva, The Quick Fire Emissions Dataset (QFED): Documentation of versions 2.1,
- 494 2.2 and 2.4. NASA/TM-2015-104606, Vol. 38, 2015.
- 495 Deeter, M., Martinez-Alonso, S., Edwards, D., Emmons, L., Gille, J., Worden, H., Sweeney, C., Pittman, J.,
- Daube, B., and Wofsy, S.: The MOPITT Version 6 product: algorithm enhancements and validation,
  Atmospheric Measurement Techniques, 7, 3623-3632, 10.5194/amt-7-3623-2014, 2014.
- 498 Duncan, B., Bey, I., Chin, M., Mickley, L., Fairlie, T., Martin, R., and Matsueda, H.: Indonesian wildfires of
- 499 1997: Impact on tropospheric chemistry, Journal of Geophysical Research-Atmospheres, 108,
  500 10.1029/2002JD003195, 2003a.
- 501 Duncan, B. N., Martin, R. V., Staudt, A. C., Yevich, R., and Logan, J. A.: Interannual and seasonal variability
- 502 of biomass burning emissions constrained by satellite observations, Journal of Geophysical Research-
- 503 Atmospheres, 108, 10.1029/2002jd002378, 2003b.
- 504 Duncan, B. N., Logan, J. A., Bey, I., Megretskaia, I. A., Yantosca, R. M., Novelli, P. C., Jones, N. B., and
- Rinsland, C. P.: Global budget of CO, 1988-1997: Source estimates and validation with a global model, J.
  Geophys. Res. Atmos., 112, 10.1029/2007JD008459, 2007.
- 507 Duncan, B. N., and Logan, J. A.: Model analysis of the factors regulating the trends and variability of carbon
- 508 monoxide between 1988 and 1997, Atmospheric Chemistry and Physics, 8, 7389-7403, 2008.
- 509 Eastham, S. D. and Jacob, D. J.: Limits on the ability of global Eulerian models to resolve intercontinental
- 510 transport of chemical plumes, Atmos. Chem. Phys., 17, 2543-2553, doi:10.5194/acp-17-2543-2017, 2017.
- 511 Edwards, D., Petron, G., Novelli, P., Emmons, L., Gille, J., and Drummond, J.: Southern Hemisphere carbon
- 512 monoxide interannual variability observed by Terra/Measurement of Pollution in the Troposphere
- 513 (MOPITT), Journal of Geophysical Research-Atmospheres, 111, 10.1029/2006JD007079, 2006.
- 514 Edwards, D. P., Emmons, L. K., Hauglustaine, D. A., Chu, D. A., Gille, J. C., Kaufman, Y. J., Petron, G.,
- 515 Yurganov, L. N., Giglio, L., Deeter, M. N., Yudin, V., Ziskin, D. C., Warner, J., Lamarque, J. F., Francis, G.
- 516 L., Ho, S. P., Mao, D., Chen, J., Grechko, E. I., and Drummond, J. R.: Observations of carbon monoxide and
- 517 aerosols from the Terra satellite: Northern Hemisphere variability, Journal of Geophysical Research-
- 518 Atmospheres, 109, 10.1029/2004jd004727, 2004.
- 519 Emmons, L., Pfister, G., Edwards, D., Gille, J., Sachse, G., Blake, D., Wofsy, S., Gerbig, C., Matross, D.,
- 520 and Nedelec, P.: Measurements of Pollution in the Troposphere (MOPITT) validation exercises during
- 521 summer 2004 field campaigns over North America, Journal of Geophysical Research-Atmospheres, 112,
- 522 10.1029/2006JD007833, 2007.

- 523 Field, R., van der Werf, G., Fanin, T., Fetzer, E., Fuller, R., Jethva, H., Levy, R., Livesey, N., Luo, M., Torres,
- 524 O., and Worden, H.: Indonesian fire activity and smoke pollution in 2015 show persistent nonlinear
- 525 sensitivity to El Nino-induced drought, Proceedings of the National Academy of Sciences of the United States
- of America, 113, 9204-9209, 10.1073/pnas.1524888113, 2016. 526
- 527 Heald, C., Jacob, D., Fiore, A., Emmons, L., Gille, J., Deeter, M., Warner, J., Edwards, D., Crawford, J.,
- Hamlin, A., Sachse, G., Browell, E., Avery, M., Vay, S., Westberg, D., Blake, D., Singh, H., Sandholm, S., 528
- 529 Talbot, R., and Fuelberg, H.: Asian outflow and trans-Pacific transport of carbon monoxide and ozone 530 pollution: An integrated satellite, aircraft, and model perspective, Journal of Geophysical Research-
- Atmospheres, 108, 10.1029/2003JD003507, 2003. 531
- Hsu, J., Prather, M., Wild, O., Sundet, J., Isaksen, I., Browell, E., Avery, M., and Sachse, G.: Are the TRACE-532
- 533 P measurements representative of the western Pacific during March 2001?, Journal of Geophysical Research-
- 534 Atmospheres, 109, 10.1029/2003JD004002, 2004.
- Kasischke, E., Hyer, E., Novelli, P., Bruhwiler, L., French, N., Sukhinin, A., Hewson, J., and Stocks, B.: 535
- 536 Influences of boreal fire emissions on Northern Hemisphere atmospheric carbon and carbon monoxide,
- Global Biogeochemical Cycles, 19, 10.1029/2004GB002300, 2005. 537
- Langenfelds, R., Francey, R., Pak, B., Steele, L., Lloyd, J., Trudinger, C., and Allison, C.: Interannual growth 538
- 539 rate variations of atmospheric CO2 and its delta C-13, H-2, CH4, and CO between 1992 and 1999 linked to
- 540 biomass burning, Global Biogeochemical Cycles, 16, 10.1029/2001GB001466, 2002.
- Levy, R., Hsu, C., et al.: MODIS Atmosphere L2 Aerosol Product. NASA MODIS Adaptive Processing 541
- System, Goddard Space Flight Center, USA, doi:http://dx.doi.org/10.5067/MODIS/MYD04 L2.006, 2015. 542
- 543 Liu, J., J. A. Logan, L. T. Murray, H. C. Pumphrey, and I. A. Megretskaia, Transport analysis and source
- attribution of seasonal and interannual variability of CO in the tropical upper troposphere and lower 544 545 stratosphere. Atmos. Chem. Phys., 13: 129-146 doi: 10.5194/acp-13-129-2013, 2013.
- Livesey, N., Filipiak, M., Froidevaux, L., Read, W., Lambert, A., Santee, M., Jiang, J., Pumphrey, H., Waters, 546
- 547 J., Cofield, R., Cuddy, D., Daffer, W., Drouin, B., Fuller, R., Jarnot, R., Jiang, Y., Knosp, B., Li, Q., Perun,
- 548 V., Schwartz, M., Snyder, W., Stek, P., Thurstans, R., Wagner, P., Avery, M., Browell, E., Cammas, J.,
- 549 Christensen, L., Diskin, G., Gao, R., Jost, H., Loewenstein, M., Lopez, J., Nedelec, P., Osterman, G., Sachse,
- 550 G., and Webster, C.: Validation of Aura Microwave Limb Sounder O-3 and CO observations in the upper
- 551 troposphere and lower stratosphere, Journal of Geophysical Research-Atmospheres, 113, 552 10.1029/2007JD008805, 2008.
- 553 Logan, J., Megretskaia, I., Nassar, R., Murray, L., Zhang, L., Bowman, K., Worden, H., and Luo, M.: Effects
- 554 of the 2006 El Nino on tropospheric composition as revealed by data from the Tropospheric Emission Spectrometer (TES), Geophysical Research Letters, 35, 10.1029/2007GL031698, 2008.
- 555
- 556 Lucchesi, R, File Specification for GEOS-5 FP. GMAO Office Note No. 4 (Version 1.1), 61pp, 2017,
- 557 available from http://gmao.gsfc.nasa.gov/pubs/office\_notes.

- 558 Molod, A., Takacs, L., Suarez, M., and Bacmeister, J.: Development of the GEOS-5 atmospheric general
- circulation model: evolution from MERRA to MERRA2, Geoscientific Model Development, 8, 1339-1356,
- 560 10.5194/gmd-8-1339-2015, 2015.
- 561 Novelli, P., Masarie, K., Lang, P., Hall, B., Myers, R., and Elkins, J.: Reanalysis of tropospheric CO trends:
- 562 Effects of the 1997-1998 wildfires, Journal of Geophysical Research-Atmospheres, 108, 563 10.1029/2002JD003031, 2003.
- 564 Ott, L., Duncan, B., Pawson, S., Colarco, P., Chin, M., Randles, C., Diehl, T., and Nielsen, E.: Influence of
- the 2006 Indonesian biomass burning aerosols on tropical dynamics studied with the GEOS-5 AGCM,
- 566 Journal of Geophysical Research-Atmospheres, 115, 10.1029/2009jd013181, 2010.
- 567 Pfister, G., Emmons, L., Edwards, D., Arellano, A., Sachse, G., and Campos, T.: Variability of springtime
- 568 transpacific pollution transport during 2000-2006: the INTEX-B mission in the context of previous years,
- 569 Atmospheric Chemistry and Physics, 10, 1345-1359, 10.5194/acp-10-1345-2010, 2010.
- 570 Prather, M., Zhu, X., Flynn, C., Strode, S., Rodriguez, J., Steenrod, S., Liu, J., Lamarque, J., Fiore, A.,
- 571 Horowitz, L., Mao, J., Murray, L., Shindell, D., and Wofsy, S.: Global atmospheric chemistry which air
- 572 matters, Atmospheric Chemistry and Physics, 17, 9081-9102, 10.5194/acp-17-9081-2017, 2017.
- 573 Prather, M. J., Flynn, C. M., Zhu, X., Steenrod, S. D., Strode, S. A., Fiore, A. M., Correa, G., Murray, L. T.,
- and Lamarque, J.-F.: How well can global chemistry models calculate the reactivity of short-lived greenhouse
- 575 gases in the remote troposphere, knowing the chemical composition, Atmos. Meas. Tech., 11, 2653-2668,
- 576 https://doi.org/10.5194/amt-11-2653-2018, 2018.

577 Reinecker, M. M. et al., The GEOS-5 Data Assimilation System – Documentation of Versions 5.0.1, 5.1.0,

578 and 5.2.0, Technical Report Series on Global Modeling and Data Assimilation, Vol. 27, ed. M. J. Suarez,

579 <u>NASA/TM-2008-104606, 2008</u>

580 Rienecker, M. M., Suarez, M. J., Gelaro, R., Todling, R., Bacmeister, J., Liu, E., Bosilovich, M. G., Schubert,

- 581 S. D., Takacs, L., Kim, G.-K., Bloom, S., Chen, J., Collins, D., Conaty, A., da Silva, A., Gu, W., Joiner, J.,
- 582 Koster, R. D., Lucchesi, R., Molod, A., Owens, T., Pawson, S., Pegion, P., Redder, C. R., Reichle, R.,
- 583 Robertson, F. R., Ruddick, A. G., Sienkiewicz, M., and Woollen, J.: MERRA: NASA's Modern-Era
- Retrospective Analysis for Research and Applications, Journal of Climate, 24, 3624-3648, 10.1175/JCLI-D 11-00015.1, 2011.
- 586 Santoni, G., Daube, B., Kort, E., Jimenez, R., Park, S., Pittman, J., Gottlieb, E., Xiang, B., Zahniser, M.,
- 587 Nelson, D., McManus, J., Peischl, J., Ryerson, T., Holloway, J., Andrews, A., Sweeney, C., Hall, B., Hintsa,
- 588 E., Moore, F., Elkins, J., Hurst, D., Stephens, B., Bent, J., and Wofsy, S.: Evaluation of the airborne quantum
- 589 cascade laser spectrometer (QCLS) measurements of the carbon and greenhouse gas suite CO2, CH4, N2O,

590 and CO - during the CalNex and HIPPO campaigns, Atmospheric Measurement Techniques, 7, 1509-1526,

- 591 10.5194/amt-7-1509-2014, 2014.
- 592 Strode, S. A., and Pawson, S.: Detection of carbon monoxide trends in the presence of interannual variability,
- 593 Journal of Geophysical Research-Atmospheres, 118, 12257-12273, 10.1002/2013JD020258, 2013.

**Moved down [1]:** Reinecker, M. M. et al., The GEOS-5 Data Assimilation System – Documentation of Versions 5.0.1, 5.1.0, and 5.2.0, Technical Report Series on Global Modeling and Data Assimilation, Vol. 27, ed. M. J. Suarez, NASA/TM-2008-104606, 2008.¶

Moved (insertion) [1]

Deleted: 1

- 600 van der Werf, G., Randerson, J., Giglio, L., Collatz, G., Kasibhatla, P., and Arellano, A.: Interannual
- variability in global biomass burning emissions from 1997 to 2004, Atmospheric Chemistry and Physics, 6,
- 602
   3423-3441, 2006.

603 van der Werf, G. R., Randerson, J. T., Giglio, L., van Leeuwen, T. T., Chen, Y., Rogers, B. M., Mu, M., van

- 604 Marle, M. J. E., Morton, D. C., Collatz, G. J., Yokelson, R. J., and Kasibhatla, P. S.: Global fire emissions
- estimates during 1997–2016, Earth Syst. Sci. Data, 9, 697-720, https://doi.org/10.5194/essd-9-697-2017,
  2017.
- 607 Voulgarakis, A., Marlier, M., Faluvegi, G., Shindell, D., Tsigaridis, K., and Mangeon, S.: Interannual 608 variability of tropospheric trace gases and aerosols: The role of biomass burning emissions, Journal of
- variability of hopospheric face gases and acrosofs. The fore of biomass burning emissions, source
- 609 Geophysical Research-Atmospheres, 120, 7157-7173, 10.1002/2014JD022926, 2015.
- 610 Waters, J., Froidevaux, L., Harwood, R., Jarnot, R., Pickett, H., Read, W., Siegel, P., Cofield, R., Filipiak,
- 611 M., Flower, D., Holden, J., Lau, G., Livesey, N., Manney, G., Pumphrey, H., Santee, M., Wu, D., Cuddy, D.,
- 612 Lay, R., Loo, M., Perun, V., Schwartz, M., Stek, P., Thurstans, R., Boyles, M., Chandra, K., Chavez, M.,
- 613 Chen, G., Chudasama, B., Dodge, R., Fuller, R., Girard, M., Jiang, J., Jiang, Y., Knosp, B., LaBelle, R., Lam,
- 614 J., Lee, K., Miller, D., Oswald, J., Patel, N., Pukala, D., Quintero, O., Scaff, D., Van Snyder, W., Tope, M.,
- 615 Wagner, P., and Walch, M.: The Earth Observing System Microwave Limb Sounder (EOS MLS) on the Aura
- 616 satellite, Ieee Transactions on Geoscience and Remote Sensing, 44, 1075-1092,
  617 10.1109/TGRS.2006.873771, 2006.
- 618 Wofsy, S. C., Apel, E., Blake, D. R., Brock, C. A., Brune, W. H., Bui, T. P., Daube, B. C., Dibb, J. E., Diskin, G. S.,
- 619 Elkiins, J. W., Froyd, K., Hall, S. R., Hanisco, T. F., Huey, L. G., Jimenez, J. L., McKain, K., Montzka, S. A., Ryerson,
- 620 T. B., Schwarz, J. P., Stephens, B. B., Weinzierl, B., and Wennberg, P.: ATom: Merged Atmospheric Chemistry, Trace
- 621 Gases, and Aerosols. ORNL DAAC, Oak Ridge, Tennessee, USA, available at:
- 622 <u>https://doi.org/10.3334/ORNLDAAC/1581.</u>
- Wolter, K., and Timlin, M. S.: Monitoring ENSO in COADS with a seasonally adjusted principal component
   index, Proc. of the 17th Climate Diagnostics Workshop, 1993,
- 625 Worden, H. M., Deeter, M. N., Frankenberg, C., George, M., Nichitiu, F., Worden, J., Aben, I., Bowman, K.
- 626 W., Clerbaux, C., Coheur, P. F., de Laat, A. T. J., Detweiler, R., Drummond, J. R., Edwards, D. P., Gille, J.
- 627 C., Hurtmans, D., Luo, M., Martinez-Alonso, S., Massie, S., Pfister, G., and Warner, J. X.: Decadal record
- of satellite carbon monoxide observations, Atmospheric Chemistry and Physics, 13, 837-850, 10.5194/acp13-837-2013, 2013.
- 630 Yashiro, H., Sugawara, S., Sudo, K., Aoki, S., and Nakazawa, T.: Temporal and spatial variations of carbon
- 631 monoxide over the western part of the Pacific Ocean, Journal of Geophysical Research-Atmospheres, 114,
- 632 10.1029/2008JD010876, 2009.
- 633 Ziemke, J., and Chandra, S.: La Nina and El Nino-induced variabilities of ozone in the tropical lower
- atmosphere during 1970-2001, Geophysical Research Letters, 30, 10.1029/2002GL016387, 2003.
- 635

# 636 Table 1 : Regional August 2016 CO Emission Totals in the GEOS-5 FP Simulations

	Fossil Fuel <sup>1</sup>	<b>Biogenic</b> <sup>1</sup>	BB1
North America	6.7 (8.0)	5.8	2.4 (2.7)
Europe	4.9 (5.9)	2.4	
Asia	26 (31)	7.9	
Eurasia <sup>2</sup>			3.0 (3.3)
Africa			24 (27)
South America	13 (16)	17	11 (12)
Other <sup>3</sup>			2.9 (3.2)
Global	50 (60)	34	43 (48)

Global50 (60)3443 (48)637<sup>1</sup>Emissions are in units of Tg. Values in parentheses include the 20% and 11% scaling factors for fossil fuels

and biomass burning, respectively, to account for CO production from VOC oxidation.

<sup>639</sup> <sup>2</sup>The Eurasian tagged tracer for BB CO includes emissions from Europe and northern Asia, but excludes
 <sup>640</sup> southern Asia.

641 <sup>3</sup>Other fossil fuel emissions includes emissions from Africa and South America, while other BB emissions

642 excludes those regions since they are tagged separately. Other BB does include southern Asia as well

- 643 Australia.
- 644

645 Table 2: Mean and Standard Deviations in CO along Atlantic and Pacific Flight Tracks

Region	Flight	Obs Mean (ppb)	Obs Stdev (ppb)	Model Mean (ppb)	Model Stdev (ppb)
Eastern Pacific	1. Palmdale – Palmdale	75	14	77	19
	2. Palmdale - Anchorage	100	40	88	16
Pacific	3. Anchorage-Kona	85	36	81	18
	4. Kona –Pago Pago	61	5.1	63	5.5
	5. Pago Pago – Christchurch	55	11	57	6.1
Southern Ocean	6. Christchurch – Punta Arenas	56	6.4	54	4.7
Atlantic	7. Punta Arenas – Ascension	69	17	71	26
	8. Ascension – Azores	101	36	103	27

	9. Azores – Kangerlussuaq	88	32	87	19
N. America	10. Kangerlussuaq – Minneapolis	90	26	91	22
	11. Minneapolis – Palmdale	84	38	107	78

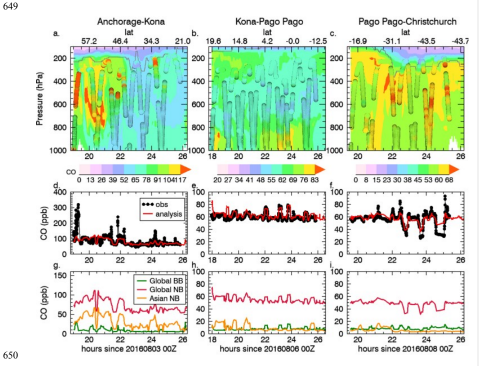




Figure 1: Curtain plot of CO (ppb) from the GEOS-5 analysis as a function of time and pressure overplotted with the model tropopause (gray line) and QCLS CO observations (circles) (top row) for the a) Anchorage to Kona 654 flight, b) Kona to Pago Pago flight, and c) Pago Pago to Christelturch flight. Axis ranges vary between panels due to the large range of concentrations encountered. The top x-axis indicates the latitudes of the flight track. d-f) The GEOS-5 CO interpolated to the flight track (red line) is compared to the observations (black circles). g-h) Tagged tracer contributions to the GEOS-5 CO.

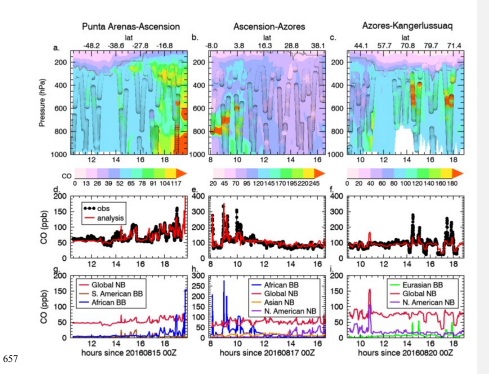
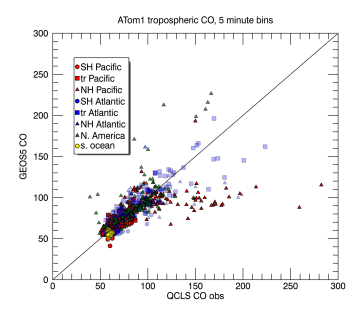
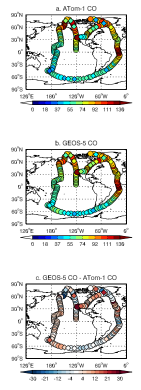


Figure 2: As in Fig. 1, but for the Atlantic flights: a,d,g) Punta Arenas-Ascension Island, b,e,h) Ascension Island
 to the Azores, and c,f,i) Azores to Kangerlussuaq.

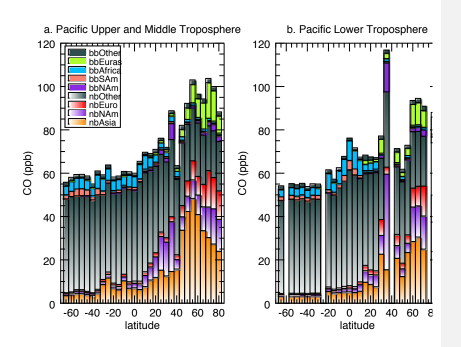


662 Figure 3: GEOS-5 simulated CO versus QCLS CO observations for all ATom-1 flights averaged into 5 minute bins. CO is in units of ppb. Pacific flights are shown in red, Atlantic flights in blue, N. American flights in green,

664 and southern ocean flights in yellow. Circles indicate Southern Hemisphere points, triangles indicate Northern Hemisphere points, and squares indicate tropical points. The one-to-one line is overplotted in black.



- Figure 4: CO (ppb) from the (a) QCLS observations, (b) GEOS-5 analysis, and (c) the GEOS-5 obs difference for the ATom-1 circuit including all 11 research flight segments. The GEOS-5 CO is taken from the analysis closest to the mid-point of the flight time and interpolated to the flight track following the longitude, latitude and pressure given in the observations. Both model forecast and ATom measurements are averaged into a sample rate of one per 360-second. Data in the troposphere are plotted in a circle, while data in the stratosphere are plotted in a diamond, based on the GEOS-5 tropopause.
- 671

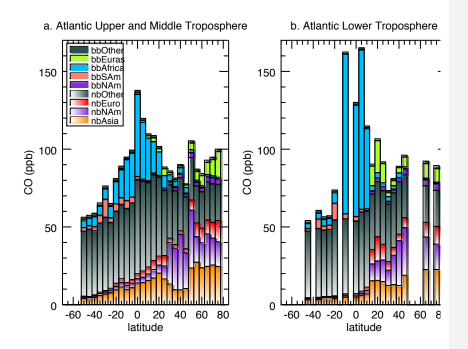


675 Figure 5: The contribution of each tagged CO tracer over the Pacific in the (a) upper and middle troposphere 676 677 (pressure <= 850 hPa) and (b) lower troposphere (pressure > 850 hPa). Data from multiple flights over the region

between 120°E and 110°W is included, with each bar representing data averaged over a 5 degree latitude bin.

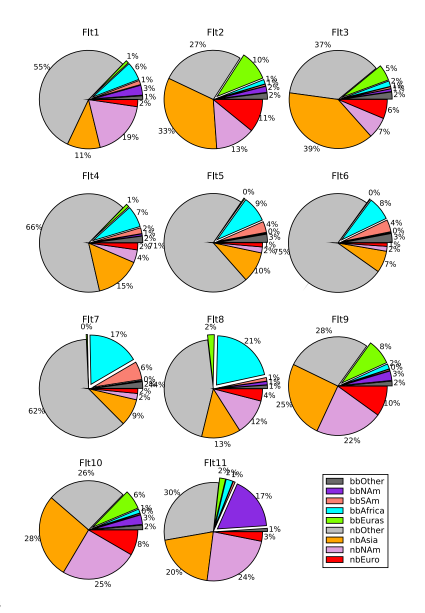
678 Shaded bars represent non-BB CO from Asia (orange), N. America (purple), Europe (red), and the rest of the 679 680 world (gray). Solid bars represent BB CO from N. America (purple), S. America (pink), Africa (cyan), Eurasia

(green), and the rest of the world (gray).





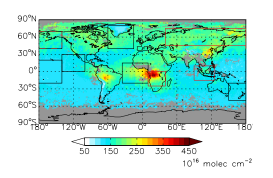
682Figure 6: As in Fig. 5, but for the Atlantic. Data from multiple flights over the region 0-60°W is included, with683each bar representing data averaged over a 5 degree latitude bin.



685Figure 7: Percent contributions of tagged tracers to total CO for each flight. Exploded slices represent the biomass686burning tracers: North American (purple), S. American (salmon), African (cyan), Eurasian (green), and Other

687 688 (dark gray). The non-biomass burning (nb) tracers are for Asia (orange), N. America (lavender), Europe (red), and other (light gray).

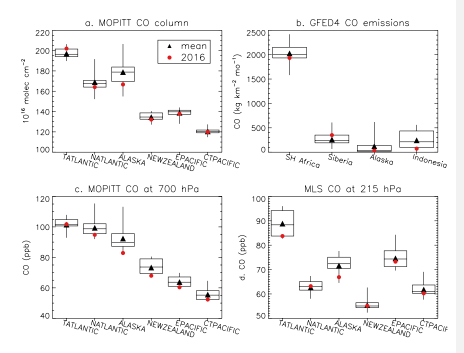




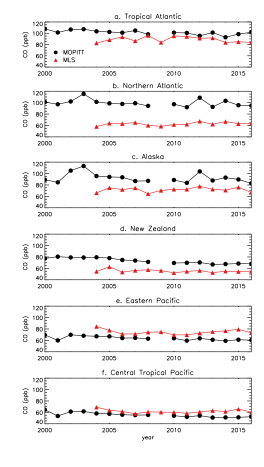


691 692 693 Figure 8: MOPITT CO column for August 2016 overplotted with the regions shown in Fig. 10. Black rectangles indicate the regions where we analyze CO concentrations, and red rectangles indicate the regions used for biomass

burning.



- Figure 9: Boxes show the 25th, 50th, and 75th percentile values; whiskers show the minimum and maximum values;
- 695 696 697 698 699 black triangles show the mean value, and 75 percentile values, winskers show the diminishing and maximum values, black triangles show the common value, and red circles show the 2016 value for a) the MOPITT CO column, b) the GFED4 CO emissions, c) MOPITT CO at 700 hPa, and d) MLS CO at 215 hPa. Statistics for MOPITT are for 2000-2016, statistics for GFED4 are for 2000-2015, and statistics for MLS are for 2004-2016. MOPITT and MLS values are for August, while the GFED4 emissions are averaged over June through August.



702 Figure 10: Time series of August MOPITT CO at the 700 hPa level (black circles) and MLS CO (red triangles) 703 at the 215 hPa level for the 6 regions shown in black in Fig. 8.

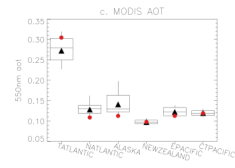


Figure 11: As in Figure 9, but for the August MODIS 550 nm AOT. Only values over oceans are included in the
 regional averages.

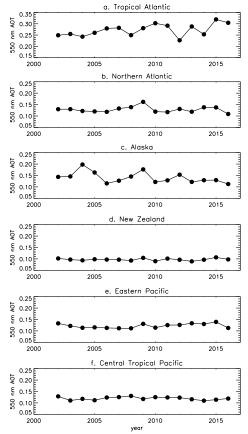




Figure 12: Time series of regionally averaged August MODIS 550 nm AOT. Only values over oceans are included

in the regional averages. The y-axis range for panel a differs from the other panels due to the higher AOT values
 in that region.