## Evaluation of stratospheric intrusions and biomass burning plumes on the vertical distribution of tropospheric ozone over the Midwestern U.S

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### Abstract

Naturally occurring ozone rich Stratosphere-to-Troposphere Transport (STT) intrusions and biomass burning (BB) plumes reaching the surface can contribute to exceedances of the U.S. National Ambient Air Quality Standards for ground-level ozone (70 ppbv implemented in 2015). Additionally, fires can inject significant pollution into the free troposphere where it can be transported long distances. The combined air quality impacts from these sources on ozone has only been analyzed in a few case studies for the Midwest U.S. Here we study ozone impacts in a Midwestern city, for the first time in St. Louis, Missouri, using a series of ozonesonde profiles taken during the SEAC4RS (Studies of Emissions and Atmospheric Composition, Clouds and Climate Coupling by Regional Surveys) field campaign in August-September 2013. All ozonesondes showed enhancements above the background profile levels (~55 ppbv) throughout each tropospheric column. Two models were used to estimate ozone origins in columns. A chemical transport model identified STT enhancements equivalent to 10 to 15 ppbv over the background with a 10 to 15% contribution overall to the column. Two FLEXPART-WRF simulations, one with smoke in the boundary layer and another with smoke above, identified BB enhancements equivalent to 10 to 80 ppbv. Overall, the total BB contribution is 15 to 30% of the total column. Five ozonesondes showed signatures of mixed BB plumes and STT intrusions. During this study period, BB in the western U.S. contributed 70% to ozone enhancements in the total column compared to 3% from the central U.S and 27% from other areas.

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Key Points:

• Stratospheric intrusions can contribute 10 to 25 ppbv to the background ozone.

- Biomass burning plumes can contribute 10 to 80 ppbv to the background ozone.
- Biomass burning in the western U.S. can significantly increase Midwest pollution levels.

### Abstract

Naturally occurring ozone rich Stratosphere-to-Troposphere Transport (STT) intrusions and biomass burning (BB) plumes reaching the surface can contribute to exceedances of the U.S. National Ambient Air Quality Standards for ground-level ozone (70 ppbv implemented in 2015). Additionally, fires can inject significant pollution into the free troposphere where it can be transported long distances. The combined air quality impacts from these sources on ozone has only been analyzed in a few case studies for the Midwest U.S. Here we study ozone impacts in a Midwestern city, for the first time in St. Louis, Missouri, using a series of ozonesonde profiles taken during the SEAC4RS (Studies of Emissions and Atmospheric Composition, Clouds and Climate Coupling by Regional Surveys) field campaign in August-September 2013. All ozonesondes showed enhancements above the background profile levels (~55 ppbv) throughout each tropospheric column. Two models were used to estimate ozone origins in columns. A chemical transport model identified STT enhancements equivalent to 10 to 15 ppbv over the background with a 10 to 15% contribution overall to the column. Two FLEXPART-WRF simulations, one with smoke in the boundary layer and another with smoke above, identified BB enhancements equivalent to 10 to 80 ppby. Overall, the total BB contribution is 15 to 30% of the total column. Five ozonesondes showed signatures of mixed BB plumes and STT intrusions. During this study period, BB in the western U.S. contributed 70% to ozone enhancements in the total column compared to 3% from the central U.S and 27% from other areas.

### Plain Language Summary

Because man-made emissions are decreasing, concentrations of harmful ozone pollution have also decreased in many areas of the USA. Not all sources can be easily controlled however. For example, biomass burning emits lots of pollutants to the atmosphere, and descending air from the stratosphere can bring with it high levels of ozone. These sources can pollute first the air above us, and then when the air is transported to the surface it can pollute the air we breathe. In this study we used balloons to measure ozone pollution as it changes from the surface to high in the atmosphere. We then used computer models to identify the sources responsible for higher pollution levels. Our study was part of a major field campaign that took place in the Midwest U.S. in the summer of 2013.

### 1 Introduction

Even moderate concentrations of ground-level ozone  $(O_3)$  can adversely affect human health and the environment across the United States (U.S. EPA, 2015). To lessen these impacts, the U.S. Environmental Protection Agency (U.S. EPA) has lowered the National Ambient Air Quality Standard (NAAQS) for  $O_3$  from 75 to 70 parts-per-billion-per-volume (ppbv) (U.S. EPA, 2015). Ozone is a key secondary air pollutant associated with a large number of health issues ranging from asthma to premature deaths (Fann et al., 2013, 2018; Reid et al., 2016; Rappold et al., 2017), leading to 12,300 to 52,000 annual premature deaths in North America alone (Silva et al., 2013). Due to more stringent regulations over the past twenty years, regional ozone levels have declined by 7 to 13% (Simon et al., 2015; Strode et al., 2015). New evidence has demonstrated that even at lower concentrations ozone can still be very toxic (U.S. EPA, 2013). Despite overall progress, ozone concentrations may still be increasing in some areas (Fishman et al., 2014; Cooper et al., 2015; Jaffe et al., 2018), particularly in fire prone regions (McClure & Jaffe 2018).

Regulating locally formed ozone is complicated by the fact that ozone has significant background levels in the troposphere typically ranging 30 to 50 ppbv (Jaffe et al., 2018). Background concentrations of ozone can be enhanced due to noncontrollable ozone sources (NCOS), defined by Jaffe et al. (2018) to include recent local pollution influence from Stratosphere-to-Troposphere Transport (STT), lingering biomass burning emissions, long-range transport from international sources, lightning, or photochemical production from precursor emissions (e.g. nitrogen oxides and volatile organic compounds). Therefore, in this study we define background ozone to include NCOS. While foreign sources of pollution and wildfires are theoretically controllable, these are beyond the control of any local jurisdiction, so for this discussion, we can justify including these as background ozone. Over the past decade, significant progress has been made in our efforts to understand aspects of the U.S. background ozone problem for example: episodic stratospheric sources (Lin et al., 2015), interannual variability (Strode et al., 2015), and wildfire contributions (Jaffe et al., 2013; Westerling, 2016). However, these efforts have lacked coordination, and are largely focused on the impacts to the western U.S. where NCOS are considered to be the greatest (Jaffe & Wigder, 2012; Jaffe et al., 2018). Despite a 15 to 33% reduction in ozone precursors from anthropogenic emissions across the Eastern and Midwestern U.S. (Strode et al., 2015), the Midwestern background ozone levels continue to increase ~0.23 ppbv per year (Fishman et al., 2014) on par with the trends in the intermountain western U.S.

The Midwestern U.S., like the intermountain west U.S., is frequently impacted by biomass burning (McCarty et al., 2007; Liu et al., 2016) and stratospheric intrusions (Lin et al., 2015; Langford et al., 2018). Although biomass burning in the Midwest consists mostly of smaller agricultural fields, their combined emitted pollution and higher frequency can result in emissions double that of wildfires (Larkin et al., 2014). Understanding stratospheric intrusions and biomass burning contributions for this region will be critical as they can confound NAAQS attainment (e.g., Jaffe, 2011; Lin et al., 2012; Hess and Zbinden, 2013), where summertime daily ozone maxima already range from 70 to 80 ppbv and occasionally reach mixing ratios higher than 140 ppbv (Fishman et al., 2014). The periodic nature of some NCOS makes it difficult to target these sources with dedicated field campaigns, but opportunistic measurements have been made during field studies with other objectives (Sullivan et al., 2015; Ott et al., 2016). For example, the influence of wildfires, long-range transport, and SST were the foci of the Las Vegas ozone Study (LVOS) where it was demonstrated that NCOS contributed ~30 ppbv to three ozone exceedances (Langford et al., 2015).

The SEAC4RS (Studies of Emissions and Atmospheric Composition, Clouds and Climate Coupling by Regional Surveys) field campaign in August-September 2013 presents a unique opportunity to study Midwest U.S. background ozone (Toon et al., 2016). During the flight mission several ozonesondes were launched nearly every day to measure vertical profiles of column ozone and meteorology over seven U.S. stations in a network known as SEACIONS (SouthEast American Consortium for Intensive Ozonesonde Network Study). The SEAC4RS campaign sparked a growing number of studies concerning ozone in the southeast U.S. (Travis et al., 2016; Wagner et al., 2015; Zhu et al., 2016). Although the research flights did not cover the Midwest U.S., this area is of interest because the time of the field campaign was an active period for NCOS with signatures for STT, large fires in the western U.S., and smaller prescribed fires in the central U.S. (~0.2 million hectares burned). During the 2013 ozone season (1 April through 31 October) fifteen total ozone exceedances occurred with six high-ozone days occurring during SEAC4RS. Additionally, a satellite-based detection method identified smoke plumes with confirmed injection heights above the boundary layer, initiated by smoke-infused thunderstorms known as pyrocumulonimbus or pyroCb (Fromm et al., 2010, 2019; Peterson et al., 2015, 2017a.b, 2018). The parametrization of plume rise from biomass burning can vary widely and cause undue biases and errors within model simulations of pollution transport (Paugam et al., 2016; Wilkins et al., 2018; Val martin et al., 2012, 2018). The detection algorithm used for the smoke plumes reduced the uncertainty of the injection heights and gave improved estimates based on remote sensing of each specific case.

Using a chemical transport model, it has been shown that fires can contribute ~14% to daily max ozone levels in the central U.S. (Wilkins et al., 2018). This has driven the need for developing accurate tropospheric ozone budgets for a typical summertime in the Midwest as suggested by Wilkins et al. (2018). Several approaches have been used to partition or identify the individual contribution of a naturally occurring ozone rich STT intrusions (Langford et al., 2018), a biomass burning plume (Westerling, 2016), or their combined impacts (Brioude et al., 2007) to elevated ozone levels. These approaches include the use of intensive balloonborne ozonesonde field campaigns (Cooper et al., 2007; Thompson et al., 2019); lidar and aircraft remote sensing (Brioude et al., 2007; Langford et al., 2018); and Lagrangian trajectory and chemical transport models (Morris et al., 2006; Baker et al., 2016, 2018). However, the determination of the contribution from individual sources to local ozone enhancement with a single point measurement remains difficult (Thompson et al., 2019), and it can be even more challenging to quantify accurately the contribution from multiple sources (Lin et al., 2015). In particular, the partitioning of these sources is critical for NAAQS attainment investigations along with long-term trend determinations, as they are two of the most critical NCOS imported (Parrish et al., 2012, 2014). Further, 30% of the present-day atmospheric ozone burden is attributable to human activity and these emissions are still rising (Granier et al., 2011; Parrish et al., 2010). For example, Li et al. (2019) using  $\tilde{1}$ ,000 surface ozone sites over a five year period (2013-2017) showed that East Asian concentrations are continuing at +1 to 3 ppbv per year. In this study, we use ozonesonde measurements from the SEAC4RS campaign and models to identify and characterize contributions from biomass burning and stratospheric intrusions to tropospheric ozone columns above the Midwest background ozone levels. Additionally, we evaluate the contribution from fires observed in the western and Midwest U.S. to ozone concentration in St. Louis.

### 2 Tropospheric Ozone and Meteorological Soundings

Vertical profiles of ozone, pressure, humidity, and temperature were measured using balloon-borne ozonesondes coupled with radiosondes. Each ozonesonde consists of a Teflon pump, an ozone sensing electrochemical concentration cell (ECC) (Komhyr et al., 1986, 1995), attached alongside a Vaisala RS-8015N radiosonde. The instruments were prepared in accordance with the Southern Hemisphere Additional Ozonesondes (SHADOZ) protocol (Thompson et al., 2011a,b, 2019). The ECC-type ozonesonde is currently the most widely used due to its well-characterized behavior and working capability under various sky conditions (Kuang et al., 2012). Comparisons with other  $O_3$  measuring instruments have demonstrated the ECC sonde precision to be ±5% near the ground and ±10% in the upper troposphere, although errors can reach 17% (Thompson et al., 2007; Stauffer et al., 2014). The radiosonde temperature and pressure sensors have accuracies below 20 km of ±0.3°C and ±0.5 hPa, respectively (Lal et al., 2014). The heights were calculated based on the observed pressure and are above sea level (asl). The humidity sensor has an accuracy of about ±2% near the ground which decreases to ±15-30% in the 5 to 15 km altitude range (Kley et al., 1997; Hurst et al., 2011).



Figure 1. Study area map with FLEXPART-WRF model domain. Rectangles mark the WRF domains D1 which has a horizontal resolution of 12 km and D2 (4 km resolution). SEACIONS locations are marked by black circles (St. Louis, Missouri station (STL) is a gray circle). Wildfire emissions from FLAMBE, August-September 2013, with eleven Pyro convective and five high-altitude injection elevated smoke plume activity areas are marked, red crosses indicate pyroCb plumes which transport smoke into the upper troposphere and lower stratosphere. Blue crosses indicate high-altitude injection of smoke in the absence of large pyroCb which transport smoke into the mid-troposphere).

Figure 1 displays the seven ozonesonde sites across the U.S. for the South East American Consortium for Intensive Ozone Network Study (SEACIONS) — archived at http://croc.gsfc.nasa.gov/seacions/. Here, we focus on the sole midwestern U.S. site located at the James S. McDonnell Planetarium in Forest Park (90.27W, 38.63N, 181 m ASL), 5 km west of downtown St. Louis, Missouri. Balloon soundings carrying an ozonesonde and a radiosonde were made near daily between 10:00 to 14:00 (UTC - 5 h). The exact launch time depended on clearance from the Air Traffic Control of the local airport and weather conditions. The first flight was made on 8 August 2013 and the last on 22 September 2013. There were breaks scheduled between the launches in order to cover the time period of the larger SEAC4RS flight mission (Toon et al., 2016). A total of 28 ozonesondes were launched during this period from St. Louis. The balloons reached a maximum altitude of 22 to 36 km with an average ascent rate of about 5 ms<sup>-1</sup>. The instruments fell mostly within 70 km of the launch site.

Vertical curtain plots of ozone concentrations are shown in Figure 2, displayed in 500 m bins up to 15 km height. The figure shows structures in the tropospheric ozone profile which will be analyzed for impacts of stratospheric air masses and biomass burning sources using the modeling techniques demonstrated by Thompson et al. (2007, 2011b).

### **3** Numerical Models

### 3.1. Stratospheric-to-Tropospheric Transport ozone tracer model

To simulate the stratospheric air masses entering the troposphere during the SEACIONS field campaign, the NASA Goddard Earth Observing System model (GEOS-5, Rienecker et al., 2008) was used. The model analysis was provided on 72 levels from the surface to 0.01 hPa (75 km), every 6 hours. Initial conditions for the atmospheric component were taken from uncoupled experiments forced by the observed sea surface temperature. The model defines a stratospheric intrusion as air masses with a high ozone composition, potential vorticity greater than 1.5 potential vorticity units (PVU; 1 PVU =  $10^{-6}$  m<sup>2</sup> s<sup>-1</sup> K kg<sup>-1</sup>), and a relative humidity (RH) of less than 20% (e.g. Holton et al., 1995; Waugh and Funatsu, 2003; Stohl, 2003). Here we categorize each STT by temporal duration, vertical extent, and meteorological cause and relate them to the contributions to the tropospheric ozone column. Previously, GEOS-5 has been used extensively for stratospheric intrusions and has been validated with ozonesonde observations (e.g. Stajner et al., 2008; Wargan et al., 2015). A review of data assimilation methodology applied to chemical constituents, including ozone, can be found in Lahoz et al. (2007).

### 3.2. Dispersion Model

### 3.2.1. FLEXPART-WRF Lagrangian Dispersion Model

The FLEXPART-WRF version 3.1 (Brioude et al., 2013) was used as a Lagrangian particle dispersion model using wind fields from the Weather Research and Forecasting (WRF version 3.4, Skamarock et al., 2008). Simulations were made for seven days to relate ozonesonde profiles to biomass burning events. FLEXPART-WRF was used to simulate carbon monoxide (CO) as a passive tracer for biomass burning plumes. The plume injection heights were adjusted to match the impacts with the vertical ozone profiles at a resolution of 100 m. The model does not include chemical loss or production in its trajectory calculations. The simulations can be used to estimate CO concentrations using multiple particles each representing a fixed amount of CO emitted and transported as a passive scalar. Therefore, trajectories aged 5 to 15 days can be considered as mixing ratios above background (Brioude et al., 2007). WRF was initialized using the North American Regional Reanalysis (NARR, Mesinger et al., 2006) wind fields, with hourly temporal resolution, two nested domains with a horizontal resolution of 27 km and 9 km, and 40 pressure levels. Figure 1 shows a map of the two nested WRF domains. Nine individual WRF simulations were performed to cover the entire time span of the SEACIONS mission. Each simulation lasted 162 h with 42 h of spin-up time and the remaining five days were used for the analysis (de Foy et al., 2014).

To determine the transport impacting each ozone measurement, a plume origin path was calculated. The plume path was determined based on several thousand forward trajectory particles released from a box surrounding the location and time of biomass burning present over the 44 day period. Plume origin path distributions were output in 1 day intervals on a 2.5@ x 2.5@ output grid at 500 m vertical resolution covering the U.S., with a 0.25@ x 0.25@ at 100 m vertical resolution nested grid over St. Louis. FLEXPART-WRF outputs the plume origin paths in units of s kg<sup>-1</sup> m<sup>3</sup>, which represents the residence time of the plume per grid cell divided by the air density. The residence of each plume was calculated for the surface to upper troposphere ( $^{0}$  to 15 km) layers of the atmosphere. This is known as a weighted plume residence time (Lal et al., 2014).

The dispersion of a plume origin path forward in time indicates the likely source regions contribution of the ozone precursors to the measured ozone but over the previous 7 days. For plumes passing through these layers, the residence time of the forward trajectories can interact with stratospheric intrusions allowing for the detection of mixed plumes. Hourly biomass burning emissions were provided by the Fire Locating and Modeling of Burning Emissions (FLAMBE) program (http://www.nrlmrv.navv.mil/flambe/) (Reid et al., 2009). The emission factors are outlined by Ferek et al. (1998). FLAMBE provides quasi-operational, with fire location, instantaneous estimates of fire size, and smoke emission flux in kg  $m^{-2}$  generated in near real time for the Western Hemisphere (Reid et al., 2009). The inventory uses active fire detection from the GOES's Wildfire Automated Biomass Burning Algorithm (WF-ABBA) and MODIS's Active Fire Products to detect biomass burning activity in near-real-time (Reid et al., 2009). The fire emissions are calculated in grams of CO from the inventories' burned area  $(m^2)$  and smoke aerosol emissions (CO in gm<sup>-2</sup>) variables. The fire emissions are mapped onto a 0.25<sup>(a)</sup> x 0.25<sup>(a)</sup> grid as input to FLEXPART-WRF. With this technique, the quantity of CO emitted into each plume from several fire source regions across the U.S. was tabulated. The CO tracer has no chemical or depositional removal processes and was treated as a passive tracer. The NARR analysis contains information that can be used to derive stratospheric ozone values above and within the troposphere. We used these values to calculate the quantity of ozone transported from the stratosphere to the location where a plume was released. NARR provides more information on the stratospheric intrusions structure with temporal coverage 8-times daily and a 32 km spatial grid. Stratospheric intrusions in NARR were calculated using PVU distribution on isentropic surfaces, 320 K to 410 K ([?] 80 hPa). Specifically, with NARR we can closely match ozonesonde launch times with meteorological data (~18:00 UTC); this allows for better comparison with model and observed stratospheric intrusions.

### 3.2.2. FLEXPART-WRF high-altitude smoke injection

To account for the vertical extent (or plume injection height) of varying biomass burning emissions in the simulations, we employ the FLEXPART-WRF model. The BASELINE simulation represents biomass burning emissions as a uniform release within the well-mixed boundary layer up to ~3.5 km (e.g. Liousse et al., 1996; Colarco, 2004; Davison, 2004; Brioude et al., 2007). To represent plumes that penetrate well above the boundary layer, we use the U.S. Naval Research Laboratory (NRL) satellite-based pyroCb detection algorithm and inventory (Peterson et al., 2017a,b) — referred to hereafter as the PYRO simulation. For a confirmed pyroCb, the plume tops are set at the model determined Upper Troposphere and Lower Stratosphere (UTLS) boundary (~9 to 15 km). For non-pyroCb, plume tops are set to be near the model determined middle troposphere (~5 km). In both cases, the plume bottoms are set above the PBL, with the emissions uniformly distributed within the tropospheric model layers. Lastly, a method is presented that is a combination of the two methods mentioned above (BASELINE + PYRO) referred to hereafter as the combined method. The combined method was run as a single simulation and compared to each individual method. The combined simulation included all the emissions from the Baseline, with addition of the elevated emissions from the fires with pyro convective plumes identified in Figure 1, see Table 1 for more details.

Peterson et al. (2017a) examined a variety of wildfires and pyroconvective events in western North America during the summer (June to August) of 2013. At least 26 intense pyroCb events were inventoried, injecting smoke particles well into the UTLS. Several fires produced smaller pyroCbs and others injected smoke above the boundary layer in the absence of significant pyroCb activity (Figure 1). The NRL pyroCb detection algorithm is designed to detect large plumes that impact the UTLS, and in general, is not designed to capture lower tropospheric transport. The algorithm only detects anvil cloud tops; therefore, neither the

exact quantitative plume height nor a plume bottom can be detected.

A variety of additional observations and methods exist to track the evolution of pyroCb smoke plumes in the UTLS after the initial injection (e.g., Fromm et al., 2010, 2019; Peterson et al., 2018). However, application of these methods is beyond the scope of this study. Evolution and transport of significant pyroCb plumes from the 2013 fire season are examined in greater detail by Peterson et al. (2015, 2017a,b) and Fromm et al. (2019).

### 4 Results and Discussions

### 4.1. The structure of tropospheric ozone columns



Figure 2. Ozonesondes in support of SEACIONS were launched at the Saint Louis University ozonesonde station located at the James S. McDonnell Planetarium in Forest Park (90.27@W, 38.63@N, 181 m asl), 5 km west of downtown St. Louis. Vertical tropospheric profiles at ~18:30 UTC (1:30 p.m. Local time) ozone profiles are averaged vertically every 500 m and shown below the thermal tropopause level (~15 km). Cases of ozone enhancements above the background (~55 ppbv) are shown by source origin: the gray box for Stratospheric-Tropospheric Transport (STT), the solid black line boxes for biomass burning (BB), and the dashed line black boxes for combined STT and BB.

The vertically averaged near-daily profiles of ozone are shown in Figure 2. The profiles are averaged every 500 m vertically from the surface to the tropopause (~15 km) with annotations for enhanced ozone rich plumes, above the tropospheric column mean (55.7 ppbv), from biomass burning and stratospheric intrusions. Ozone gradually increases with altitude in the lower troposphere (below about 3.5 km). Average ozone is about 55 ppbv near the surface but decreases to 45 ppbv near 3.5 km. Above 3.5 km, ozone increases slowly throughout the entire troposphere a rate of 3 ppbv km<sup>-1</sup>. In the middle troposphere (4 to 10 km, height), average ozone is highest in August and lowest in September. While above 10 km, ozone is similar for both months. August profiles averaged 10 to 15 ppbv higher than September. Ozone is lowest (~40 ppbv) during the month of September near the surface and remains low below 3.5 km. However, on 5 September the second highest recorded surface ozone day occurred at the surface (60 ppbv ozonesonde, 95 ppbv nearby surface monitor later in the day). The maximum measured ozone from a sonde at the surface occurred 30 August at 65 ppbv. Other notable enhancements (NAAQS exceedances) of O<sub>3</sub> (> 70 ppb near the surface) occurred on 23 August, and 4 to 6 and 10 to 11 September. A mean PBL of 2.4 km and a mean thermal

tropopause at 14.9 km could be clearly identified in the ozonesonde profiles. Note that these figures for biomass burning contributions from boreal fire contributions to ozone downwind are somewhat larger than the 5 to 15% estimates of this quantity by Thompson et al. (2011; see Figure 13) or Moeini et al. (2020) that are based on Canadian soundings. These last two studies attribute 20 to 25% of the ozone column amounts to stratospheric origins.

### 4.2. Stratospheric-tropospheric transport tracer impacts

Table 2 lists the GEOS-5 determined stratospheric instructions events that impacted St. Louis during August and September 2013, expressed as deep or shallow with the pressure level 600 hPa being the threshold. The 44 day study period consisted of 13 events. The intrusions recorded resulted from either a shortwave trough (6 days: 7, 10, 23, 28, 30 August and 6 September), frontal passage (7 days: 3, 12 August and 3, 12, 15, 22 September), or cut-off low (1 day: 14 August). Three frontal pass intrusions reached the lower troposphere and penetrated the boundary layer. About 40% of the intrusions reached the middle troposphere ( $^{500}$ hPa). Stratospheric intrusion generally lasted 1 to 3 days occurring either early morning before 3:00 (UTC - 5 h) or around noon local time. The intrusions where found to contribute on average 10 to 25 ppbv to tropospheric ozone columns which equates to around 5 to 15% of the total tropospheric ozone column.



### 4.3. Biomass Burning Emissions

**Figure 3.** Biomass burning emissions of carbon monoxide (CO) estimated by FLAMBE in units of Gg per km<sup>2</sup> per hour in North America during SEACIONS, 8 Aug to 22 Sept 2013. FLAMBE emissions are displayed on the model grid which has a resolution of 0.25@ per cell. Only cells with emissions above 2,000 kg are shown.



Figure 3 shows the hourly U.S. total mass of fire smoke emissions from FLAMBE in Gg-CO km<sup>-2</sup> hr<sup>-1</sup> gridded per 0.25@ cell with a minimum threshold of 2,000 kg. From 8 August to 22 September 2013 the calculated total emissions were 1,060 Gg of CO. FLAMBE emissions depict a regional distribution pattern in known fire regions: agricultural and prescribed fires in the southeast U.S. (e.g. the Carolinas, Florida, and Georgia) and southcentral U.S. (e.g. Louisiana, Mississippi, and Arkansas); and the wildfires in the central plains and intermountain west U.S. (e.g. Wyoming, Idaho/Montana area, and California). A time series of the hourly emissions indicates that August contained almost all the major fire emissions (Figure 4). The duration of the largest fire emissions are indicated in the figure by arrows. Between 14 and 24 August 2013, on average  $\tilde{}$  Gg-CO hr<sup>-1</sup> ( 1.2 x 10<sup>4</sup> FLEXPART particles where 1 particle = 500 kg = 0.5x10<sup>-3</sup> Gg) was being emitted daily with a maximum hourly total of 11 Gg-CO hr<sup>-1</sup> ( 2.2 x 10<sup>4</sup> FLEXPART particles). Regional fires emissions from the central U.S. were considerably lower than wildfires emissions in the intermountain west (3% compared to 70%).

Figure 4. A time series of FLAMBE emissions (mass/time) converted to particles released in FLEXPART-WRF (# of particles or particle count). Each 500 kg of mass is converted to 1 particle in FLEXPART-WRF simulations. Locations emitting less than 500 kg only emit a single particle. Arrows represent the duration of two of the largest wildfires (Rim Fire and Beaver Creek) with relevance to SEACIONS with other fires (Emigrant Fire and Pony/Elk Complex) contributing to those time frame emissions indicated. The dotted line is the California Rim Fire (37.85@N, 120.083@W) 17-31 August 2013, and the solid line is the Idaho Beaver Creek (43.593@N, 114.684@W) 7-21 August 2013. Note: Beaver Creek and Pony/Elk Complex occurred nearly at the same time and were fewer than 100 km apart in distance. While, the Emigrant Fire contributed a significant amount of emissions during its occurrence at the same time as the Rim Fire but was much smaller in magnitude.

Specifically, three large wildfires areas were identified that produced the most emissions and high-altitude smoke injection plumes. The Idaho Beaver Creek fire, caused by a lightning strike on 7 August 2013 in the Sawtooth National Forest, occurred in the lower southwestern edge of the Elk/Pony Complex (43.59@N, 114.68@W). This fire burned an area of 465 km<sup>2</sup> (114,900 acres), emitting 54.6 Gg-CO of smoke emissions, and accounted for 14% of the Idaho and Montana states combined total fire emissions (392 Gg-CO km<sup>-2</sup>hr<sup>-1</sup>) during the months of August and September. The second fire area, led by a fire named the Montana

Emigrant Fire, was a part of a small system of fires caused by multiple lightning strikes that spanned across Yellowstone National Park (45.20@N, 110.69@W) consuming  $< 81 \text{ km}^2$  (20,000 acres) starting 14 August emitting 9.4 Gg-CO of smoke emissions. Lastly, the largest of the three, the California Rim Fire, described in detail in Peterson et al. (2015), caused by an illegal campfire, occurred on 17 August 2013 in Yosemite National Park (37.85@N, 120.08@W). The Rim Fire burned an area of 1041 km<sup>2</sup> (257,314 acres), emitted 224 Gg-CO of smoke, and accounted for over 66% of the California biomass burning emissions (341 Gg-CO km<sup>-2</sup> hr<sup>-1</sup>) during SEAC4RS.

4.4. Residence time of particles over St. Louis



**Figure 5.** FLEXPART-WRF Biomass burning (BB) CO concentrations and age are simulated, binning trajectories over St. Louis within a 2.5@ x 2.5@ grid box and averaged vertically 500 m with daily temporal resolution. FLAMBE emissions of BB CO are converted into particle amounts and are released. (a) The base simulation, Boundary Layer Simulation, releases emissions within the PBL, (b) NRL's adjusted pyroconvention scheme is implemented, PyroCb Simulation, particles are released from pyroCbs and high-altitude injection as specified in Table 1, and (c). The combined (a) and (b). With panels (d-f), corresponding to plume ages in simulations. The average BB CO g cm<sup>-3</sup>per grid cell (top) lighter shading indicated higher concentration and transported CO plume ages (bottom) lighter shading indicates older air.

The FLEXPART-WRF plume origin path (forward trajectories) has been used to calculate residence times of air parcels above St. Louis (Figure 5). Figure 5a-5f show the average residence times of the biomass burning plume origin paths expressed as plume age and CO tracer particle amounts as concentrations. Results are displayed for the BASELINE simulation (Figure 5a and 5d), PYRO simulation (Figure 5b and 5e), and combined (Figure 5c and 5f). The lighter shading represents a higher concentration of CO particle counts (Figure 5a-5c) and for plume age the lighter shading indicates older air (Figure 5d-5f). The major contributions below 3 km are from the central U.S. fires. The contributions from the intermountain west U.S. fires dominates above 3 km, in the range of 80 to 90%. Other regions contribute in the range of 5 to 10%. Biomass burning plumes are indicated to have impacted the surface ozone each day but with varying magnitudes. The surface impacts were connected to mechanisms causing air parcels to move downward (e.g. shortwave on 30 August). Likewise, strong vertical motion was evident in bringing simulated air within the boundary to the middle and upper portions of the troposphere. For example, on 24-26 August at the 8 to 12 km layer, both simulations indicate enhancements of 0.74 ppm of CO that has been aged 7 days. This enhancement corresponded to 40 ppb of excess  $O_3$  at the 11 to 12 km layer in the ozonesondes. Ozone impacts for August from biomass burning, sharply increase after 3 km and vary daily above; up to 12 km where the biomass burning contributions to ozone decrease to near negligible amounts. The high-altitude smoke injection simulation indicates that plumes that get injected above the 4 km height can impact the surface. Evidence of aged lofted air, ranging from 4 to 15 km, impacting the surface is seen 22 August to 1

September 2013, where a constant 2 ppm of CO enhancement at the surface is present. Additional evidence of biomass burning emissions enhancing ozone within the lower tropopause and PBL (0 to 6 km) is present on 21-24 August over the St. Louis area — a greater than 15 ppbv enhancement of ozone within a 6 day aged plume containing an additional 9 ppm of CO.

The average plume age for the August month ranged between 4 to 7 days. Plumes reaching the lower troposphere (below 3.5 km) were  $\tilde{3}$  days fresher then plumes above. The simulated high-altitude smoke injection is typically  $\tilde{2}$  days fresher than the boundary layer injected smoke when above 3 km but can be 2 days older when below. The average aged plume lead to 15 ppbv increase in ozone. As plumes aged, they became more ozone enriched. Fresher plumes have higher levels of CO but lower amounts of ozone. An example of this occurred 23 August, where and ozone exceedance at the surface was indicated in both simulations to be from a polluted air mass layer with mixed ages. From the lower and middle troposphere (below 1.5 km and above 5.5 km) the plumes ages averaged 2 days older, with about 7 ppm fewer CO, and 20 ppbv more ozone than the air in the layers in-between.

### 4.5. High-altitude smoke injection analysis

The impacts of high-altitude smoke injection are demonstrated using two examples from commonly encountered summertime synoptic meteorological situations in the central U.S. (i) cyclonic flow or a system of low pressure (e.g. cut-off low 17-21 August) and (ii) anti-cyclonic flow or a system of high pressure (e.g. blocking high 25-30 August). Table 3 lists each episode's ozone enriched plume identified by an ozonesonde over St. Louis (Figure 2). For each identified ozone enhancement, the simulated source-contribution relationships from biomass burning (Table 1) and stratospheric intrusions (Table 2) are quantified. The corresponding travel time of the plume (or plume age) and concentrations are indicated by FLEXPART-WRF (Figure 5). St. Louis area for two selected days (21 and 30 August). Each plume was individually characterized with model simulations and ozonesonde observational data in addition to the meteorological conditions leading to the plume reaching St. Louis is discussed below.



Figure 6. Series of daily synoptic atmospheric plane slices at 500 hPa at 7:00 UTC for preceding conditions to individual transport cases (a-c) 21 August 2013 (d-f) 30 August 2013.



**Figure 7.** Atmospheric plane slices from NARR at 500 hPa at 18:00 UTC (a) 21 August 2013 (b) 30 August 2013. Vertical Cross sections at 90<sup>@</sup> W, 20-60<sup>@</sup> N 18:00 UT (c) 21 August 2013 (d) 30 August 2013. Blue contour lines show Potential Vorticity (PV) 10<sup>6</sup> PVU, Relative Humidity (RH) below 30% is shaded from light to dark, with darker shades representing the dryer air. The black lines on (a) and (b) are pressure height contours in m. While, the black lines on (c) and (d) represent potential temperature (isentropic surfaces) in K. Biomass burning sources tend to have low PV and can have moderate RH levels for pyroCbs, while stratospheric air masses have high PV and low RH. The blue dashed line PV contour (1.5 PV) shows the approximate boundary between stratospheric and tropospheric air masses. Arrows (green) show air-mass transport patterns from Stratospheric-Tropospheric Transport (STT) and from Biomass Burning (BB).



Figure 8. NOAA Hazard Management System (HMS) and Smoke Product analysis of smoke (a) 21 Aug 2013 and (b) 30 Aug 2013. Below, three-day backward FLEXPART-WRF trajectories corresponding to each episode is provided (c) 21 Aug 2013 and (d) 30 Aug 2013. Additional trajectories are placed in the data archive at https://tropo.gsfc.nasa.gov/seacions/.

4.5.1. 17-21 August 2013: Evidence of biomass burning advected by a cutoff low

The meteorological map provided in Figure 6a-c indicates biomass burning transported by a cut-off low originating on 17 August 2013 00:00 UTC at 500 hPa over central Missouri. In this case the cut-off low was created by a closed upper-level low pressure system that was cut off from a westerly current (14 August 06:30 UTC) and it began moving independently. The cut-off low remained nearly stationary until 19 August 00:00 UTC, at which time the trough filled in and moved eastward. Due to the nature of the quasi-stationary system, the westerly flow of air propagated the western fire plumes over the central U.S. and southeast U.S. Figure 7a and 7c present a cross section and plane view of the resulting low. Due to the low pressure system the trajectory analysis indicated that smoke flowed from various fire locations to the St. Louis area (see Figure 8). Figure 9a-c depicts the vertical ozone and meteorological profiles representative of this time period (21 August). Refer to Figure 2 for corresponding ozonesonde curtain plots.



Figure 9. Vertical tropospheric profiles over St. Louis at ~18:30 to 19:30 UTC for selected test cases 21 August 2013 and 30 August 2013. Labels P1 to P6 correspond to plume information in Table 3. Ozone measured from the ozonesonde in ppb are shown as a black line. Panel (a) and (d), the green line RH %. Panel (b) and (e), the blue line represents the GEOS-5 modeled Potential Vorticity  $10^6$  PVU. Panel (c) and (f), the FLEXPART-WRF modeled CO biomass burning g cm<sup>-3</sup>, for each simulation the PYRO simulation is red, and BASELINE simulation in gray. Refer to Figure 2 for corresponding ozonesonde curtain plots.

Three ozone enhanced layers are visible on 21 August (Table 3, Figure 9). None of these enhancements are deemed of stratospheric origins, as there was no significant PVU present in these layers. Plume P3 at 11 km was initially considered to be a stratospheric intrusion related but are ruled out as the NARR data and sonde thermal tropopause indicate that the tropopause was lowered (15 km to  $^{9}$  km) possibly by wave breaking from the cut-off low. The lower- and upper-middle tropospheric ozone enhancements at P1 (2 to 5.5 km) and P2 (7 to 9 km), respectively, are determined to be of biomass burning origin. The lower-tropospheric plume P1 was determined to be from the Idaho Beaver Creek plume  $^{6}$  days earlier. The upper-middle tropospheric plume P2 originated 2 days prior from high-altitude injection plumes from the Yellowstone National Park

Emigrant cluster fires. The contribution of 21 August tropospheric ozone layers from biomass burning was 10 to 25 ppbv. The pyroCb from the Idaho Pony Elk Fire (16 Aug at 23:41 UTC) reached the St. Louis area; But, the polluted air remained in the lower stratospheric air and did not mix with the lower layers. The incorporation of the high-altitude injection plumes allowed for the two middle- and upper- tropospheric polluted plumes to be characterized which contributed ~15 to 30% to layers in the total ozone column. This upper level pollution was later advected down to the surface and contributed to the exceedance on the 23 August.

4.5.2. 25-30 August 2013: Evidence of stratospheric air mixing with aged biomass burning during anticyclonic flow

The meteorological map provided in Figure 6d-f indicates aged biomass burning transported into a stratospheric intrusion by a high pressure system that became well-established over Missouri 25 August 00:00 UTC. The high pressure system began developing a week prior over the southcentral U.S. The high remained quasipermanent as it sat over Missouri reaching its maximum strength on 26 August 06:00 UTC until a mid-level (500 hPa) shortwave trough moved into the area on 29 August 12:00 UTC moving the system eastward. The near surface reflection of the shortwave reaches Eastern North Dakota 29 August 18:00 UTC as the trough extends south to Oklahoma where it penetrates deeper into the ridge. Figure 7b and 7d presents a cross section and plane view of the resulting high. Due to the high pressure system the trajectory analysis indicated that smoke flowed from various fire locations in additional to stratospheric intrusion impacts to the St. Louis area (see Figure 8). Figure 9d-f depicts the vertical ozone and meteorological profiles representative of this time period (30 August). Further evidence of this trough effects on elevated ozone aloft in the sonde measurements (see Figure 2) remained until 30 August 18:00 UTC (Figure 9d-f). Thereafter a shortwave passes the area slightly to the east and cleared out the excess ozone.

Three ozone enhanced layers are present during the high pressure event as evident on 30 August (Table 3, P4 to P6). Stratospheric air descended into the middle troposphere ~500 hPa on two occasions during this time period (28 Aug and 30 Aug) from shortwaves (Figure 7d). Stratospheric impacts are indicated at ~7 to 9 km and >12 km layers with high PVU values and low RH values (Figure 9d-f, P5 and P6). NARR PVU analysis provided further proof that these are in fact stratospheric in origin. Figure 7d depicts lowering dry stratospheric air and ascending moist tropospheric air. The near surface plume (0 to 3.5 km) was traced back to southeastern agricultural fires ~5 days prior (see Figure 8), indicating a likely recirculation of air by the high pressure system. The middle-tropospheric plume P5 was determined to be from an unknown source, potentially a recirculation of summertime ozone precursors (Cooper et al., 2006, 2007). Evidence of this layer is present in previous soundings for the week 25-29 August where all sondes show a single well-defined plume at 5 to 9 km. The GEOS-5 and NARR potential vorticity also give no indication of what the cause for this layer is. Our hypothesis is supported by the vertical wind profiles for the week being relatively stagnantthe ozonesonde launched that day landed within 50 m of the launch site — signifying air circulation over St. Louis for the entire week increasing photochemical effects. An alternative theory is that wind shear caused the layer and it's a part of the larger plume above. The layer between the unknown plume and above  $(^{\circ}6 \text{ to } 8 \text{ km})$  is a layer of moist and clean air (see Figure 7d). The upper-tropospheric layer plume P6 is a combination of both a biomass burning plume and a stratospheric intrusion air mass. The lower half of the plume P6 (8 to 10 km) is dominated by stratospheric air, while the above portion of P6 (10 to 12 km) is primarily biomass burning. There is another layer of clean air above prior to reaching the troppause. The upper-tropospheric plume P6 originated 5 days prior from high-altitude injection plumes from the California Rim Fire (25 August). The contribution of 30 August tropospheric ozone layers from Biomass burning was 10 to 80 ppbv and stratospheric air masses contributed 10 to 40 ppbv. The incorporation of high-altitude smoke injection allowed for the two mid- and upper- tropospheric polluted plumes to be characterized which contributed ~15 to 60% to layers in the total ozone column.

### 5 Summary and Conclusions

By incorporating balloon-borne ozonesonde observations with models, this study has quantitatively examined sources for tropospheric ozone enhancements due to Non-Controllable Ozone Sources (NCOS). In particular,

the pollution impacts from stratospheric intrusions and biomass burning contributions to background tropospheric and surface ozone levels in the Midwest United States were characterized. Additionally, emphasis is placed on partitioning the contribution from western U.S. fires, central U.S. fires, and other areas to St. Louis background ozone. A chemical transport model and trajectory model were run to quantify source contribution to ozone in St. Louis, Missouri. For the region and time period of this study, 10 to 15% of the ozone enhancements stems from a stratospheric airmass contribution and 15 to 30% from biomass burning. These NCOS contributions and ozonesonde profiles can be considered as baseline ranges for the Midwest U.S. area if direct ozone measurements (sondes, airplane, ground-based) are not available. Considering U.S. fires only, 70% of the biomass burning plumes originated from the western parts of the U.S. and only 3% came from the local central U.S. emissions. Moreover, it was demonstrated that a redistribution of the biomass burning emissions injection height, with part of the emissions above the boundary layer led to a reduction of model predicted surface ozone.

In agreement with earlier studies (e.g., Fishman et al., 2014), this study has identified a generally increasing relationship between background ozone and transported pollution. We followed the definition of background ozone described in Jaffe et al. (2018) as NCOS such as lingering biomass burning and long-range transported international sources. The major contributions below 3 km were from the central U.S. fires. While 80 to 90% of the high-altitude injection smoke (above 3.5 km) originated in the western U.S. During this campaign period only 5 to 10% of the biomass burning emissions reaching St. Louis originated from the southeast and other regions.

This study identified that biomass burning plumes in the western U.S. can have impacts on the daily atmospheric ozone column in the Midwest (10 to 80 ppbv of ozone) at a greater frequency and intensity than stratospheric intrusion (10 to 25 ppbv of ozone). We show the background ozone to be 55 ppbv, which was near the 30 to 50 ppbv range mentioned in Jaffe et al. (2018) which is typical for the U.S. We identified a relationship between smoke plume age and ozone enhancement where the high-altitude injection smoke plumes above 3.5 km generally were associated with higher amounts of CO concentrations but fresher smoke regarding ozone levels. In addition, we recognized that the high-altitude smoke had a higher tendency to mix with stratospheric intrusions, which together doubled the ozone enhancement in the tropospheric column.

An investigation of several individual smoke plumes has shown the importance of incorporating high-altitude smoke injection in model simulations in addition to ensuring that accurate biomass burning locations and temporal allocation of intensity are included in model emissions. Up to 60% of the smoke plume lies above 3.5 km, and this needs to be simulated as it can later be mixed down to the surface and lead to ozone exceedances days later. In addition, it was shown that the incorporation of satellite-based detections of high-altitude smoke injection (e.g., pyroCb activity, Peterson et al., 2014, 2017a,b) can be helpful for improve modeling results and explain ozone enhancements aloft and at the surface.

The individual cases were selected because they are associated with common meteorological situations in the Midwest leading to ozone exceedances. Evidence from the first case where biomass burning is advected by a cutoff low is an example of a common flow pattern that transports air masses from the west. Summertime occurrences of this synoptic situation in conjunction with large wildfires in the western U.S. can lead to increases in ozone in the Midwest of 10 to 80 ppbv or greater. In the second case a stratospheric air mass was shown to mix with an aged wildfire plume during anticyclonic flow, a pattern that was previously found to occur 40% of the time for the southcentral U.S. (Texas and the Gulf of Mexico area, Brioude et al., 2007). Additionally, the test cases showed that the surface impacts were connected to mechanisms causing air parcels to move downward (e.g. shortwave on 30 August). Likewise, strong vertical motion was evident in bringing simulated air within the boundary to the middle and upper portions of the troposphere.

While the results in this study highlight that Non-Controllable Ozone Sources can contribute significantly to local tropospheric ozone in the Midwest, future studies must combine satellite data and model integration techniques with meteorological information for a longer time period, both within and outside the Midwest to better characterize NCOS contribution to U.S. ozone. Good examples of how an air quality model could be used to assess NCOS has been shown in Baker et al. (2016, 2018). More specifically, Baker et al. (2016, 2018)

address the question of regional-scale pyrogenic ozone sources. In particular, Baker et al. (2018) investigated the Rim Fire, using non-sonde data, which occurred during our study period with a photochemical model and their findings were similar to ours. Furthermore, as shown by individual plume cases and several earlier studies (e.g. Morris et al., 2006; Brioude et al., 2007; Jaffe, 2011; Lin et al., 2012; Hess and Zbinden, 2013), high-altitude smoke injection from a fire can lead to long distance transport depending on weather conditions and hence impact surface ozone and NAAQS attainment. Plume rise and plume injection heights are a key source of uncertainty (Paugam et al., 2016), which can be reduced considerably using plume height information from remote sensing tools (e.g. Peterson et al., 2017a,b; Val Martin et al., 2018). Improved modeling techniques will be required to better characterize biomass burning transport and hence better simulate long range impacts and the possibility of unhealthy surface concentrations far from the biomass burning sources.

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**Figure 1**. Study area map with FLEXPART-WRF model domain. Rectangles mark the WRF domains D1 which has a horizontal resolution of 12 km and D2 (4 km resolution). SEACIONS locations are marked by black circles (St. Louis, Missouri station (STL) is a gray circle). Wildfire emissions from FLAMBE, August-September 2013, with eleven Pyro convective and five high-altitude injection elevated smoke plume activity areas are marked, red crosses indicate pyroCb plumes which transport smoke into the upper troposphere and lower stratosphere. Blue crosses indicate high-altitude injection of smoke in the absence of large pyroCb which transport smoke into the mid-troposphere).

**Figure 2.** Ozonesondes in support of SEACIONS were launched at the Saint Louis University ozonesonde station located at the James S. McDonnell Planetarium in Forest Park (90.27@W, 38.63@N, 181 m asl), 5 km west of downtown St. Louis. Vertical tropospheric profiles at ~18:30 UTC (1:30 p.m. Local time) ozone profiles are averaged vertically every 500 m and shown below the thermal tropopause level (~15 km). Cases of ozone enhancements above the background (~55 ppbv) are shown by source origin: the gray box for Stratospheric-Tropospheric Transport (STT), the solid black line boxes for biomass burning (BB), and the dashed line black boxes for combined STT and BB.

**Figure 3.** Biomass burning emissions of carbon monoxide (CO) estimated by FLAMBE in units of Gg per km<sup>2</sup> per hour in North America during SEACIONS, 8 Aug to 22 Sept 2013. FLAMBE emissions are displayed on the model grid which has a resolution of 0.25@ per cell. Only cells with emissions above 2,000 kg are shown.

Figure 4. A time series of FLAMBE emissions (mass/time) converted to particles released in FLEXPART-WRF (# of particles or particle count). Each 500 kg of mass is converted to 1 particle in FLEXPART-WRF simulations. Locations emitting less than 500 kg only emit a single particle. Arrows represent the duration of two of the largest wildfires (Rim Fire and Beaver Creek) with relevance to SEACIONS with other fires (Emigrant Fire and Pony/Elk Complex) contributing to those time frame emissions indicated. The dotted line is the California Rim Fire (37.85@N, 120.083@W) 17-31 August 2013, and the solid line is the Idaho Beaver Creek (43.593@N, 114.684@W) 7-21 August 2013. Note: Beaver Creek and Pony/Elk Complex occurred nearly at the same time and were fewer than 100 km apart in distance. While, the Emigrant Fire contributed a significant amount of emissions during its occurrence at the same time as the Rim Fire but was much smaller in magnitude.

**Figure 5.** FLEXPART-WRF Biomass burning (BB) CO concentrations and age are simulated, binning trajectories over St. Louis within a 2.5@ x 2.5@ grid box and averaged vertically 500 m with daily temporal resolution. FLAMBE emissions of BB CO are converted into particle amounts and are released. (a) The base simulation, Boundary Layer Simulation, releases emissions within the PBL, (b) NRL's adjusted pyroconvention scheme is implemented, PyroCb Simulation, particles are released from pyroCbs and high-altitude injection as specified in Table 1, and (c). The combined (a) and (b). With panels (d-f), corresponding to plume ages in simulations. The average BB CO g cm<sup>-3</sup>per grid cell (top) lighter shading indicated higher concentration and transported CO plume ages (bottom) lighter shading indicates older air.

Figure 6. Series of daily synoptic atmospheric plane slices at 500 hPa at 7:00 UTC for preceding conditions to individual transport cases (a-c) 21 August 2013 (d-f) 30 August 2013.

**Figure 7.** Atmospheric plane slices from NARR at 500 hPa at 18:00 UTC (a) 21 August 2013 (b) 30 August 2013. Vertical Cross sections at 90@ W, 20-60@ N 18:00 UT (c) 21 August 2013 (d) 30 August 2013. Blue contour lines show Potential Vorticity (PV) 10<sup>6</sup> PVU, Relative Humidity (RH) below 30% is shaded from light to dark, with darker shades representing the dryer air. The black lines on (a) and (b) are pressure height contours in m. While, the black lines on (c) and (d) represent potential temperature (isentropic surfaces) in K. Biomass burning sources tend to have low PV and can have moderate RH levels for pyroCbs, while stratospheric air masses have high PV and low RH. The blue dashed line PV contour (1.5 PV) shows the approximate boundary between stratospheric and tropospheric air masses. Arrows (green) show air-mass transport patterns from Stratospheric-Tropospheric Transport (STT) and from Biomass Burning (BB).

Figure 8. NOAA Hazard Management System (HMS) and Smoke Product analysis of smoke (a) 21 Aug 2013 and (b) 30 Aug 2013. Below, three-day backward FLEXPART-WRF trajectories corresponding to each episode is provided (c) 21 Aug 2013 and (d) 30 Aug 2013. Additional trajectories are placed in the data archive at https://tropo.gsfc.nasa.gov/seacions/.

**Figure 9.** Vertical tropospheric profiles over St. Louis at ~18:30 to 19:30 UTC for selected test cases 21 August 2013 and 30 August 2013. Labels P1 to P6 correspond to plume information in Table 3. Ozone measured from the ozonesonde in ppb are shown as a black line. Panel (a) and (d), the green line RH %. Panel (b) and (e), the blue line represents the GEOS-5 modeled Potential Vorticity  $10^6$  PVU. Panel (c) and (f), the FLEXPART-WRF modeled CO biomass burning g cm<sup>-3</sup>, for each simulation the PYRO simulation is red, and BASELINE simulation in gray. Refer to Figure 2 for corresponding ozonesonde curtain plots.

**Table 1.** Elevated smoke plumes escaping the planetary boundary layer identified by the Naval Research laboratory satellite detection algorithm for high-altitude injection during the Studies of Emissions and Atmospheric Composition, Clouds and Climate Coupling by Regional Surveys (SEAC4RS) in the summer 2013. Smoke plumes detected are categorized: 1. A pyroconvective plume containing smoke reaching the stratosphere (~10 to 15 km) or 2. A high-altitude plume containing smoke reaching into the middle troposphere (~5 to 9 km).

Date	Time (UTC)	Lat	Lon	State	Fire name	Plume type
7 Aug 2013	22:15	42.71	-123.66	OR	Big Windy Complex	High-altitude
8 Aug 2013	n/A	43.46	-114.55	ID	Beaver Creek	Pyroconvective
8 Aug 2013	n/A	43.42	-114.85	ID	McCan	Pyroconvective
9 Aug 2013	22:41	43.29	-115.55	ID	Pony/Elk	Pyroconvective
10 Aug 2013	21:00	43.36	-115.45	ID	Pony/Elk	Pyroconvective
12 Aug 2013	21:00	43.29	-115.55	ID	Pony/Elk	Pyroconvective
12 Aug 2013	23:00	43.62	-114.51	ID	Beaver Creek	Pyroconvective
13 Aug 2013	23:41	43.50	-115.43	ID	Pony/Elk	Pyroconvective
16 Aug 2013	21:41	45.66	-114.78	ID	Gold Pan Complex	Pyroconvective
16 Aug 2013	23:41	43.62	-114.51	ID	Pony/Elk	High-altitude
16 Aug 2013	23:41	43.46	-114.55	ID	Beaver Creek	Pyroconvective
19 Aug 2013	22:50	45.20	-110.69	$\mathbf{MT}$	Emigrant	High-altitude
19 Aug 2013	23:15	37.86	-120.09	CA	Rim	Pyroconvective
21 Aug 2013	22:41	37.86	-120.09	CA	Rim	Pyroconvective
22 Aug 2013	21:30	41.04	-123.47	CA	Corral Complex	High-altitude
25 Aug 2013	17:00	37.91	-120.00	CA	Rim	High-altitude

**Table 2.** Simulated stratospheric air masses entering the troposphere (stratospheric intrusion) over St. Louis, Missouri during August and September 2013.

Date	Time (UTC)	Duration (hr)	Height (hPa)	Extent	Meteorology
Date	Time (UTC)	Duration (hr)	Height (hPa)	Extent	Meteorology
3 Aug 2013	03:30	21	580	Shallow	Frontal passage
7 Aug 2013	13:30	28	690	Deep	Shortwave
10 Aug 2013	02:30	34	800	Deep	Shortwave
12 Aug 2013	18:30	28	500 to $850$	Deep	Frontal passage
14 Aug 2013	06:30	180	600	Shallow	Cutoff low
23 Aug 2013	16:30	9	500	Shallow	Shortwave
28 Aug 2013	21:30	18	480	Shallow	Shortwave
30 Aug 2013	08:30	94	500	Shallow	Shortwave
3 Sept 2013	12:30	30	700 to 1000	Deep	Frontal passage
6 Sept 2013	09:30	41	600	Shallow	Shortwave
12 Sept 2013	16:30	10	800	Deep	Frontal passage
15 Sept 2013	20:30	48	800 to 1000	Deep	Frontal passage
22 Sept 2013	00:30	94	900	Deep	Frontal passage

**Table 3.** Ozone  $(O_3)$  enriched plumes identified over St. Louis, Missouri using ozonesondes vertical profile measurements for two episodes. Included are the simulated biomass burning plumes and stratospheric air masses entering the troposphere (stratospheric intrusion (SI)) concentrations of a carbon monoxide (CO) tracer for each plume. Plume ID corresponds to labeling in Figure 9.

Plume ID	Date	Source	Plume height	Smoke age	BASELINE simulation	PYRO Simulation	So	
			AGL km	days	CO ppm	CO ppm	%	
P1	21 Aug	ID Beaver Creek	2  to  5.5	5.8	2.4	14	10	
P2	21 Aug	MT Emigrant	7 to 9	2.6	0	5.2	5	
P3	21 Aug	pyroCb and SI	11	1.2	0	0.63	20	
P4	30 Aug	SE US Ag. Fire	0  to  2.5	5.1	6.6	0.83	85	
P5	30 Aug	Unidentified	5  to  7	6.7	0	0.35	10	
P6	30 Aug	CA Rim Fire and SI	8 to 12	5.3	0	6.8	15	

# Evaluation of stratospheric intrusions and biomass burning plumes on the vertical distribution of tropospheric ozone over the Midwestern U.S.

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## 13 Key Points:

• Stratospheric intrusions can contribute 10 to 25 ppbv to the background ozone.

• Biomass burning plumes can contribute 10 to 80 ppbv to the background ozone.

• Biomass burning in the western U.S. can significantly increase Midwest pollution levels.

## 18 Abstract

19 Naturally occurring ozone rich Stratosphere-to-Troposphere Transport (STT) intrusions and

20 biomass burning (BB) plumes reaching the surface can contribute to exceedances of the U.S.

21 National Ambient Air Quality Standards for ground-level ozone (70 ppbv implemented in 2015).

Additionally, fires can inject significant pollution into the free troposphere where it can be transported long distances. The combined air quality impacts from these sources on ozone has

transported long distances. The combined air quality impacts from these sources on ozone has only been analyzed in a few case studies for the Midwest U.S. Here we study ozone impacts in a

Midwestern city, for the first time in St. Louis, Missouri, using a series of ozonesonde profiles

taken during the SEAC4RS (Studies of Emissions and Atmospheric Composition, Clouds and

27 Climate Coupling by Regional Surveys) field campaign in August-September 2013. All

ozonesondes showed enhancements above the background profile levels (~55 ppbv) throughout

29 each tropospheric column. Two models were used to estimate ozone origins in columns. A

30 chemical transport model identified STT enhancements equivalent to 10 to 15 ppbv over the

background with a 10 to 15% contribution overall to the column. Two FLEXPART-WRF

simulations, one with smoke in the boundary layer and another with smoke above, identified BB

enhancements equivalent to 10 to 80 ppby. Overall, the total BB contribution is 15 to 30% of the

total column. Five ozonesondes showed signatures of mixed BB plumes and STT intrusions.

35 During this study period, BB in the western U.S. contributed 70% to ozone enhancements in the

total column compared to 3% from the central U.S and 27% from other areas.

## 37 Plain Language Summary

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Because man-made emissions are decreasing, concentrations of harmful ozone pollution have 39 also decreased in many areas of the USA. Not all sources can be easily controlled however. For 40 example, biomass burning emits lots of pollutants to the atmosphere, and descending air from the 41 stratosphere can bring with it high levels of ozone. These sources can pollute first the air above 42 us, and then when the air is transported to the surface it can pollute the air we breathe. In this 43 study we used balloons to measure ozone pollution as it changes from the surface to high in the 44 atmosphere. We then used computer models to identify the sources responsible for higher 45 pollution levels. Our study was part of a major field campaign that took place in the Midwest 46 U.S. in the summer of 2013. 47

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## 50 **1 Introduction**

Even moderate concentrations of ground-level ozone  $(O_3)$  can adversely affect human 51 health and the environment across the United States (U.S. EPA, 2015). To lessen these impacts, 52 the U.S. Environmental Protection Agency (U.S. EPA) has lowered the National Ambient Air 53 Quality Standard (NAAQS) for  $O_3$  from 75 to 70 parts-per-billion-per-volume (ppbv) (U.S. EPA, 54 2015). Ozone is a key secondary air pollutant associated with a large number of health issues 55 ranging from asthma to premature deaths (Fann et al., 2013, 2018; Reid et al., 2016; Rappold et 56 al., 2017), leading to 12,300 to 52,000 annual premature deaths in North America alone (Silva et 57 al., 2013). Due to more stringent regulations over the past twenty years, regional ozone levels 58 have declined by 7 to 13% (Simon et al., 2015; Strode et al., 2015). New evidence has 59 demonstrated that even at lower concentrations ozone can still be very toxic (U.S. EPA, 2013). 60 Despite overall progress, ozone concentrations may still be increasing in some areas (Fishman et 61 62 al., 2014; Cooper et al., 2015; Jaffe et al., 2018), particularly in fire prone regions (McClure & Jaffe 2018). 63 64 Regulating locally formed ozone is complicated by the fact that ozone has significant

background levels in the troposphere typically ranging 30 to 50 ppby (Jaffe et al., 2018). 65 Background concentrations of ozone can be enhanced due to noncontrollable ozone sources 66 (NCOS), defined by Jaffe et al. (2018) to include recent local pollution influence from 67 Stratosphere-to-Troposphere Transport (STT), lingering biomass burning emissions, long-range 68 transport from international sources, lightning, or photochemical production from precursor 69 70 emissions (e.g. nitrogen oxides and volatile organic compounds). Therefore, in this study we define background ozone to include NCOS. While foreign sources of pollution and wildfires are 71 theoretically controllable, these are beyond the control of any local jurisdiction, so for this 72 discussion, we can justify including these as background ozone. Over the past decade, significant 73 progress has been made in our efforts to understand aspects of the U.S. background 74 ozone problem for example: episodic stratospheric sources (Lin et al., 2015), interannual 75 variability (Strode et al., 2015), and wildfire contributions (Jaffe et al., 2013; Westerling, 2016). 76 However, these efforts have lacked coordination, and are largely focused on the impacts to the 77 western U.S. where NCOS are considered to be the greatest (Jaffe & Wigder, 2012; Jaffe et al., 78 2018). Despite a 15 to 33% reduction in ozone precursors from anthropogenic emissions across 79 the Eastern and Midwestern U.S. (Strode et al., 2015), the Midwestern background ozone levels 80 continue to increase ~0.23 ppbv per year (Fishman et al., 2014) on par with the trends in the 81

82 intermountain western U.S.

The Midwestern U.S., like the intermountain west U.S., is frequently impacted by 83 biomass burning (McCarty et al., 2007; Liu et al., 2016) and stratospheric intrusions (Lin et al., 84 2015; Langford et al., 2018). Although biomass burning in the Midwest consists mostly of 85 smaller agricultural fields, their combined emitted pollution and higher frequency can result in 86 emissions double that of wildfires (Larkin et al., 2014). Understanding stratospheric intrusions 87 and biomass burning contributions for this region will be critical as they can confound NAAOS 88 attainment (e.g., Jaffe, 2011; Lin et al., 2012; Hess and Zbinden, 2013), where summertime daily 89 ozone maxima already range from 70 to 80 ppbv and occasionally reach mixing ratios higher 90 than 140 ppbv (Fishman et al., 2014). The periodic nature of some NCOS makes it difficult to 91 target these sources with dedicated field campaigns, but opportunistic measurements have been 92 made during field studies with other objectives (Sullivan et al., 2015; Ott et al., 2016). For 93

example, the influence of wildfires, long-range transport, and SST were the foci of the Las Vegas

ozone Study (LVOS) where it was demonstrated that NCOS contributed ~30 ppbv to three ozone
 exceedances (Langford et al., 2015).

The SEAC4RS (Studies of Emissions and Atmospheric Composition, Clouds and 97 Climate Coupling by Regional Surveys) field campaign in August-September 2013 presents a 98 99 unique opportunity to study Midwest U.S. background ozone (Toon et al., 2016). During the flight mission several ozonesondes were launched nearly every day to measure vertical profiles 100 of column ozone and meteorology over seven U.S. stations in a network known as SEACIONS 101 (SouthEast American Consortium for Intensive Ozonesonde Network Study). The SEAC4RS 102 campaign sparked a growing number of studies concerning ozone in the southeast U.S. (Travis et 103 al., 2016; Wagner et al., 2015; Zhu et al., 2016). Although the research flights did not cover the 104 Midwest U.S., this area is of interest because the time of the field campaign was an active period 105 for NCOS with signatures for STT, large fires in the western U.S., and smaller prescribed fires in 106 the central U.S. (~0.2 million hectares burned). During the 2013 ozone season (1 April through 107 31 October) fifteen total ozone exceedances occurred with six high-ozone days occurring during 108 SEAC4RS. Additionally, a satellite-based detection method identified smoke plumes with 109 confirmed injection heights above the boundary layer, initiated by smoke-infused thunderstorms 110 known as pyrocumulonimbus or pyroCb (Fromm et al., 2010, 2019; Peterson et al., 2015, 111 2017a,b, 2018). The parametrization of plume rise from biomass burning can vary widely and 112 cause undue biases and errors within model simulations of pollution transport (Paugam et al., 113 2016; Wilkins et al., 2018; Val martin et al., 2012, 2018). The detection algorithm used for the 114 smoke plumes reduced the uncertainty of the injection heights and gave improved estimates 115 based on remote sensing of each specific case. 116

Using a chemical transport model, it has been shown that fires can contribute  $\sim 14\%$  to 117 daily max ozone levels in the central U.S. (Wilkins et al., 2018). This has driven the need for 118 developing accurate tropospheric ozone budgets for a typical summertime in the Midwest as 119 120 suggested by Wilkins et al. (2018). Several approaches have been used to partition or identify the individual contribution of a naturally occurring ozone rich STT intrusions (Langford et al., 121 2018), a biomass burning plume (Westerling, 2016), or their combined impacts (Brioude et al., 122 2007) to elevated ozone levels. These approaches include the use of intensive balloon-borne 123 124 ozonesonde field campaigns (Cooper et al., 2007; Thompson et al., 2019); lidar and aircraft remote sensing (Brioude et al., 2007; Langford et al., 2018); and Lagrangian trajectory and 125 126 chemical transport models (Morris et al., 2006; Baker et al., 2016, 2018). However, the determination of the contribution from individual sources to local ozone enhancement with a 127 single point measurement remains difficult (Thompson et al., 2019), and it can be even more 128 challenging to quantify accurately the contribution from multiple sources (Lin et al., 2015). In 129 particular, the partitioning of these sources is critical for NAAQS attainment investigations along 130 with long-term trend determinations, as they are two of the most critical NCOS imported (Parrish 131 et al., 2012, 2014). Further, 30% of the present-day atmospheric ozone burden is attributable to 132 human activity and these emissions are still rising (Granier et al., 2011; Parrish et al., 2010). For 133 example, Li et al. (2019) using ~1,000 surface ozone sites over a five year period (2013-2017) 134 showed that East Asian concentrations are continuing at +1 to 3 ppbv per year. In this study, we 135 use ozonesonde measurements from the SEAC4RS campaign and models to identify and 136 characterize contributions from biomass burning and stratospheric intrusions to tropospheric 137 ozone columns above the Midwest background ozone levels. Additionally, we evaluate the 138

contribution from fires observed in the western and Midwest U.S. to ozone concentration in St.Louis.

## 141 2 Tropospheric Ozone and Meteorological Soundings

Vertical profiles of ozone, pressure, humidity, and temperature were measured using 142 balloon-borne ozonesondes coupled with radiosondes. Each ozonesonde consists of a Teflon 143 pump, an ozone sensing electrochemical concentration cell (ECC) (Komhyr et al., 1986, 1995), 144 145 attached alongside a Vaisala RS-8015N radiosonde. The instruments were prepared in accordance with the Southern Hemisphere Additional Ozonesondes (SHADOZ) protocol 146 (Thompson et al., 2011a,b, 2019). The ECC-type ozonesonde is currently the most widely used 147 due to its well-characterized behavior and working capability under various sky conditions 148 (Kuang et al., 2012). Comparisons with other  $O_3$  measuring instruments have demonstrated the 149 ECC sonde precision to be  $\pm 5\%$  near the ground and  $\pm 10\%$  in the upper troposphere, although 150 151 errors can reach 17% (Thompson et al., 2007; Stauffer et al., 2014). The radiosonde temperature and pressure sensors have accuracies below 20 km of  $\pm 0.3$  °C and  $\pm 0.5$  hPa, respectively (Lal et 152 al., 2014). The heights were calculated based on the observed pressure and are above sea level 153 (asl). The humidity sensor has an accuracy of about  $\pm 2\%$  near the ground which decreases to 154  $\pm 15$ -30% in the 5 to 15 km altitude range (Kley et al., 1997; Hurst et al., 2011). 155



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158 Figure 1. Study area map with FLEXPART-WRF model domain. Rectangles mark the WRF

domains D1 which has a horizontal resolution of 12 km and D2 (4 km resolution). SEACIONS

- locations are marked by black circles (St. Louis, Missouri station (STL) is a gray circle).
- 161 Wildfire emissions from FLAMBE, August-September 2013, with eleven Pyro convective and

162 five high-altitude injection elevated smoke plume activity areas are marked, red crosses indicate

pyroCb plumes which transport smoke into the upper troposphere and lower stratosphere. Bluecrosses indicate high-altitude injection of smoke in the absence of large pyroCb which transport

crosses indicate high-altitude injection of smoke in the absence o smoke into the mid-troposphere).

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Figure 1 displays the seven ozonesonde sites across the U.S. for the South East American 167 Consortium for Intensive Ozone Network Study (SEACIONS) - archived at 168 http://croc.gsfc.nasa.gov/seacions/. Here, we focus on the sole midwestern U.S. site located at 169 the James S. McDonnell Planetarium in Forest Park (90.27W, 38.63N, 181 m ASL), ~5 km west 170 of downtown St. Louis, Missouri, Balloon soundings carrying an ozonesonde and a radiosonde 171 were made near daily between 10:00 to 14:00 (UTC - 5 h). The exact launch time depended on 172 clearance from the Air Traffic Control of the local airport and weather conditions. The first flight 173 was made on 8 August 2013 and the last on 22 September 2013. There were breaks scheduled 174 between the launches in order to cover the time period of the larger SEAC4RS flight mission 175 176 (Toon et al., 2016). A total of 28 ozonesondes were launched during this period from St. Louis. The balloons reached a maximum altitude of 22 to 36 km with an average ascent rate of about ~5 177 ms<sup>-1</sup>. The instruments fell mostly within 70 km of the launch site. 178

Vertical curtain plots of ozone concentrations are shown in Figure 2, displayed in 500 m bins up to 15 km height. The figure shows structures in the tropospheric ozone profile which will be analyzed for impacts of stratospheric air masses and biomass burning sources using the modeling techniques demonstrated by Thompson et al. (2007, 2011b).

## 183 **3 Numerical Models**

## 184 3.1. Stratospheric-to-Tropospheric Transport ozone tracer model

To simulate the stratospheric air masses entering the troposphere during the SEACIONS 185 field campaign, the NASA Goddard Earth Observing System model (GEOS-5, Rienecker et al., 186 2008) was used. The model analysis was provided on 72 levels from the surface to 0.01 hPa (~75 187 km), every 6 hours. Initial conditions for the atmospheric component were taken from uncoupled 188 experiments forced by the observed sea surface temperature. The model defines a stratospheric 189 intrusion as air masses with a high ozone composition, potential vorticity greater than 1.5 190 potential vorticity units (PVU; 1 PVU =  $10^{-6}$  m<sup>2</sup> s<sup>-1</sup>K kg<sup>-1</sup>), and a relative humidity (RH) of less 191 than 20% (e.g. Holton et al., 1995; Waugh and Funatsu, 2003; Stohl, 2003). Here we categorize 192 each STT by temporal duration, vertical extent, and meteorological cause and relate them to the 193 contributions to the tropospheric ozone column. Previously, GEOS-5 has been used extensively 194 for stratospheric intrusions and has been validated with ozonesonde observations (e.g. Stajner 195 et al., 2008; Wargan et al., 2015). A review of data assimilation methodology applied to 196 chemical constituents, including ozone, can be found in Lahoz et al. (2007). 197

1983.2. Dispersion Model

199 3.2.1. FLEXPART-WRF Lagrangian Dispersion Model

The FLEXPART-WRF version 3.1 (Brioude et al., 2013) was used as a Lagrangian particle dispersion model using wind fields from the Weather Research and Forecasting (WRF version 3.4, Skamarock et al., 2008). Simulations were made for seven days to relate

ozonesonde profiles to biomass burning events. FLEXPART-WRF was used to simulate carbon 203 204 monoxide (CO) as a passive tracer for biomass burning plumes. The plume injection heights were adjusted to match the impacts with the vertical ozone profiles at a resolution of 100 m. The 205 model does not include chemical loss or production in its trajectory calculations. The simulations 206 can be used to estimate CO concentrations using multiple particles each representing a fixed 207 amount of CO emitted and transported as a passive scalar. Therefore, trajectories aged 5 to 15 208 days can be considered as mixing ratios above background (Brioude et al., 2007). WRF was 209 initialized using the North American Regional Reanalysis (NARR, Mesinger et al., 2006) wind 210 fields, with hourly temporal resolution, two nested domains with a horizontal resolution of 27 km 211 and 9 km, and 40 pressure levels. Figure 1 shows a map of the two nested WRF domains. Nine 212 individual WRF simulations were performed to cover the entire time span of the SEACIONS 213 mission. Each simulation lasted 162 h with 42 h of spin-up time and the remaining five days 214 were used for the analysis (de Foy et al., 2014). 215

To determine the transport impacting each ozone measurement, a plume origin path was 216 calculated. The plume path was determined based on several thousand forward trajectory 217 particles released from a box surrounding the location and time of biomass burning present over 218 the 44 day period. Plume origin path distributions were output in 1 day intervals on a 2.5° x 2.5° 219 output grid at 500 m vertical resolution covering the U.S., with a 0.25° x 0.25° at 100 m vertical 220 resolution nested grid over St. Louis. FLEXPART-WRF outputs the plume origin paths in units 221 of s kg<sup>-1</sup> m<sup>3</sup>, which represents the residence time of the plume per grid cell divided by the air 222 density. The residence of each plume was calculated for the surface to upper troposphere (~0 to 223 15 km) layers of the atmosphere. This is known as a weighted plume residence time (Lal et al., 224 2014). 225

The dispersion of a plume origin path forward in time indicates the likely source regions 226 contribution of the ozone precursors to the measured ozone but over the previous 7 days. For 227 plumes passing through these layers, the residence time of the forward trajectories can interact 228 229 with stratospheric intrusions allowing for the detection of mixed plumes. Hourly biomass burning emissions were provided by the Fire Locating and Modeling of Burning Emissions 230 (FLAMBE) program (http://www.nrlmry.navy.mil/flambe/) (Reid et al., 2009). The emission 231 factors are outlined by Ferek et al. (1998). FLAMBE provides quasi-operational, with fire 232 location, instantaneous estimates of fire size, and smoke emission flux in kg m<sup>-2</sup> generated in 233 near real time for the Western Hemisphere (Reid et al., 2009). The inventory uses active fire 234 235 detection from the GOES's Wildfire Automated Biomass Burning Algorithm (WF-ABBA) and MODIS's Active Fire Products to detect biomass burning activity in near-real-time (Reid et al., 236 2009). The fire emissions are calculated in grams of CO from the inventories' burned area  $(m^2)$ 237 and smoke aerosol emissions (CO in  $gm^2$ ) variables. The fire emissions are mapped onto a 0.25° 238 x 0.25° grid as input to FLEXPART-WRF. With this technique, the quantity of CO emitted into 239 each plume from several fire source regions across the U.S. was tabulated. The CO tracer has no 240 chemical or depositional removal processes and was treated as a passive tracer. The NARR 241 analysis contains information that can be used to derive stratospheric ozone values above and 242 within the troposphere. We used these values to calculate the quantity of ozone transported from 243 the stratosphere to the location where a plume was released. NARR provides more information 244 on the stratospheric intrusions structure with temporal coverage 8-times daily and a 32 km spatial 245 grid. Stratospheric intrusions in NARR were calculated using PVU distribution on isentropic 246 surfaces, 320 K to 410 K ( $\approx$  80 hPa). Specifically, with NARR we can closely match ozonesonde 247

launch times with meteorological data (~18:00 UTC); this allows for better comparison with
model and observed stratospheric intrusions.

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## 3.2.2. FLEXPART-WRF high-altitude smoke injection

To account for the vertical extent (or plume injection height) of varying biomass burning 251 emissions in the simulations, we employ the FLEXPART-WRF model. The BASELINE 252 simulation represents biomass burning emissions as a uniform release within the well-mixed 253 boundary layer up to ~3.5 km (e.g. Liousse et al., 1996; Colarco, 2004; Davison, 2004; Brioude 254 et al., 2007). To represent plumes that penetrate well above the boundary layer, we use the U.S. 255 Naval Research Laboratory (NRL) satellite-based pyroCb detection algorithm and inventory 256 (Peterson et al., 2017a,b) — referred to hereafter as the PYRO simulation. For a confirmed 257 pyroCb, the plume tops are set at the model determined Upper Troposphere and Lower 258 Stratosphere (UTLS) boundary (~9 to 15 km). For non-pyroCb, plume tops are set to be near the 259 260 model determined middle troposphere ( $\sim 5$  km). In both cases, the plume bottoms are set above the PBL, with the emissions uniformly distributed within the tropospheric model layers. Lastly, a 261 method is presented that is a combination of the two methods mentioned above (BASELINE + 262 PYRO) referred to hereafter as the combined method. The combined method was run as a single 263 simulation and compared to each individual method. The combined simulation included all the 264 emissions from the Baseline, with addition of the elevated emissions from the fires with pyro 265 convective plumes identified in Figure 1, see Table 1 for more details. 266

Peterson et al. (2017a) examined a variety of wildfires and pyroconvective events in 267 western North America during the summer (June to August) of 2013. At least 26 intense pyroCb 268 events were inventoried, injecting smoke particles well into the UTLS. Several fires produced 269 smaller pyroCbs and others injected smoke above the boundary layer in the absence of 270 significant pyroCb activity (Figure 1). The NRL pyroCb detection algorithm is designed to detect 271 large plumes that impact the UTLS, and in general, is not designed to capture lower tropospheric 272 transport. The algorithm only detects any cloud tops; therefore, neither the exact quantitative 273 plume height nor a plume bottom can be detected. 274

A variety of additional observations and methods exist to track the evolution of pyroCb smoke plumes in the UTLS after the initial injection (e.g., Fromm et al., 2010, 2019; Peterson et al., 2018). However, application of these methods is beyond the scope of this study. Evolution and transport of significant pyroCb plumes from the 2013 fire season are examined in greater detail by Peterson et al. (2015, 2017a,b) and Fromm et al. (2019).

## 280 **4 Results and Discussions**





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Figure 2. Ozonesondes in support of SEACIONS were launched at the Saint Louis University 283 ozonesonde station located at the James S. McDonnell Planetarium in Forest Park (90.27°W, 284 38.63°N, 181 m asl), 5 km west of downtown St. Louis. Vertical tropospheric profiles at ~18:30 285 UTC (1:30 p.m. Local time) ozone profiles are averaged vertically every 500 m and shown 286 below the thermal tropopause level ( $\sim$ 15 km). Cases of ozone enhancements above the 287 background (~55 ppby) are shown by source origin: the gray box for Stratospheric-Tropospheric 288 Transport (STT), the solid black line boxes for biomass burning (BB), and the dashed line black 289 boxes for combined STT and BB. 290

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The vertically averaged near-daily profiles of ozone are shown in Figure 2. The profiles 292 are averaged every 500 m vertically from the surface to the tropopause ( $\sim$ 15 km) with 293 annotations for enhanced ozone rich plumes, above the tropospheric column mean (55.7 ppbv), 294 from biomass burning and stratospheric intrusions. Ozone gradually increases with altitude in the 295 lower troposphere (below about 3.5 km). Average ozone is about 55 ppbv near the surface but 296 decreases to 45 ppbv near 3.5 km. Above 3.5 km, ozone increases slowly throughout the entire 297 troposphere a rate of 3 ppbv km<sup>-1</sup>. In the middle troposphere (4 to 10 km, height), average ozone 298 is highest in August and lowest in September. While above 10 km, ozone is similar for both 299 months. August profiles averaged 10 to 15 ppbv higher than September. Ozone is lowest (~40 300 ppbv) during the month of September near the surface and remains low below 3.5 km. However, 301 on 5 September the second highest recorded surface ozone day occurred at the surface (60 ppbv 302 ozonesonde, 95 ppbv nearby surface monitor later in the day). The maximum measured ozone 303

- from a sonde at the surface occurred 30 August at 65 ppbv. Other notable enhancements (NAAQS exceedances) of  $O_3$  (> 70 ppb near the surface) occurred on 23 August, and 4 to 6 and 10 to 11 September. A mean PBL of 2.4 km and a mean thermal tropopause at 14.9 km could be clearly identified in the ozonesonde profiles. Note that these figures for biomass burning contributions from boreal fire contributions to ozone downwind are somewhat larger than the 5 to 15% estimates of this quantity by Thompson et al. (2011; see Figure 13) or Moeini et al. (2020) that are based on Canadian soundings. These last two studies attribute 20 to 25% of the
- 311 ozone column amounts to stratospheric origins.
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4.2. Stratospheric-tropospheric transport tracer impacts

Table 2 lists the GEOS-5 determined stratospheric instructions events that impacted St. 313 Louis during August and September 2013, expressed as deep or shallow with the pressure level 314 600 hPa being the threshold. The 44 day study period consisted of 13 events. The intrusions 315 recorded resulted from either a shortwave trough (6 days: 7, 10, 23, 28, 30 August and 6 316 September), frontal passage (7 days: 3, 12 August and 3, 12, 15, 22 September), or cut-off low (1 317 day: 14 August). Three frontal pass intrusions reached the lower troposphere and penetrated the 318 boundary layer. About 40% of the intrusions reached the middle troposphere (~500 hPa). 319 Stratospheric intrusion generally lasted 1 to 3 days occurring either early morning before 3:00 320 (UTC - 5 h) or around noon local time. The intrusions where found to contribute on average 10 321 to 25 ppbv to tropospheric ozone columns which equates to around 5 to 15% of the total 322 tropospheric ozone column. 323



4.3. Biomass Burning Emissions



Figure 3. Biomass burning emissions of carbon monoxide (CO) estimated by FLAMBE in units of Gg per km<sup>2</sup> per hour in North America during SEACIONS, 8 Aug to 22 Sept 2013. FLAMBE emissions are displayed on the model grid which has a resolution of 0.25° per cell. Only cells

with emissions above 2,000 kg are shown.

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Figure 3 shows the hourly U.S. total mass of fire smoke emissions from FLAMBE in Gg-CO km<sup>-</sup> 331 <sup>2</sup> hr<sup>-1</sup> gridded per 0.25° cell with a minimum threshold of 2,000 kg. From 8 August to 22 332 September 2013 the calculated total emissions were 1,060 Gg of CO. FLAMBE emissions depict 333 a regional distribution pattern in known fire regions: agricultural and prescribed fires in the 334 southeast U.S. (e.g. the Carolinas, Florida, and Georgia) and southcentral U.S. (e.g. Louisiana, 335 Mississippi, and Arkansas); and the wildfires in the central plains and intermountain west U.S. 336 (e.g. Wyoming, Idaho/Montana area, and California). A time series of the hourly emissions 337 338 indicates that August contained almost all the major fire emissions (Figure 4). The duration of the largest fire emissions are indicated in the figure by arrows. Between 14 and 24 August 2013, 339 on average ~6 Gg-CO hr<sup>-1</sup> (~1.2 x 10<sup>4</sup> FLEXPART particles where 1 particle =  $500 \text{ kg} = 0.5 \times 10^{-3}$ 340 Gg) was being emitted daily with a maximum hourly total of 11 Gg-CO hr<sup>-1</sup> ( $\sim 2.2 \times 10^4$ 341 FLEXPART particles). Regional fires emissions from the central U.S. were considerably lower 342 than wildfires emissions in the intermountain west (3% compared to 70%). 343



**Figure 4.** A time series of FLAMBE emissions (mass/time) converted to particles released in

346 FLEXPART-WRF (# of particles or particle count). Each 500 kg of mass is converted to 1

particle in FLEXPART-WRF simulations. Locations emitting less than 500 kg only emit a single

particle. Arrows represent the duration of two of the largest wildfires (Rim Fire and Beaver
 Creek) with relevance to SEACIONS with other fires (Emigrant Fire and Pony/Elk Complex)

contributing to those time frame emissions indicated. The dotted line is the California Rim Fire

351 (37.85°N, 120.083°W) 17-31 August 2013, and the solid line is the Idaho Beaver Creek

352 (43.593°N, 114.684°W) 7-21 August 2013. Note: Beaver Creek and Pony/Elk Complex occurred

nearly at the same time and were fewer than 100 km apart in distance. While, the Emigrant Fire

contributed a significant amount of emissions during its occurrence at the same time as the Rim

355 Fire but was much smaller in magnitude.

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Specifically, three large wildfires areas were identified that produced the most emissions 357 and high-altitude smoke injection plumes. The Idaho Beaver Creek fire, caused by a lightning 358 strike on 7 August 2013 in the Sawtooth National Forest, occurred in the lower southwestern 359 edge of the Elk/Pony Complex (43.59°N, 114.68°W). This fire burned an area of 465 km<sup>2</sup> 360 (114,900 acres), emitting 54.6 Gg-CO of smoke emissions, and accounted for 14% of the Idaho 361 and Montana states combined total fire emissions (392 Gg-CO km<sup>-2</sup> hr<sup>-1</sup>) during the months of 362 August and September. The second fire area, led by a fire named the Montana Emigrant Fire, 363 was a part of a small system of fires caused by multiple lightning strikes that spanned across 364 Yellowstone National Park (45.20°N, 110.69°W) consuming <81 km<sup>2</sup> (20,000 acres) starting 14 365 August emitting 9.4 Gg-CO of smoke emissions. Lastly, the largest of the three, the California 366 Rim Fire, described in detail in Peterson et al. (2015), caused by an illegal campfire, occurred on 367 17 August 2013 in Yosemite National Park (37.85°N, 120.08°W). The Rim Fire burned an area 368 of 1041 km<sup>2</sup> (257,314 acres), emitted 224 Gg-CO of smoke, and accounted for over 66% of the 369 California biomass burning emissions (341 Gg-CO km<sup>-2</sup> hr<sup>-1</sup>) during SEAC4RS. 370

#### 4.4. Residence time of particles over St. Louis 371

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Figure 5. FLEXPART-WRF Biomass burning (BB) CO concentrations and age are simulated, 374 binning trajectories over St. Louis within a 2.5° x 2.5° grid box and averaged vertically 500 m 375 with daily temporal resolution. FLAMBE emissions of BB CO are converted into particle 376 amounts and are released. (a) The base simulation, Boundary Layer Simulation, releases 377 emissions within the PBL, (b) NRL's adjusted pyro-convention scheme is implemented, PyroCb 378 379 Simulation, particles are released from pyroCbs and high-altitude injection as specified in Table 1, and (c). The combined (a) and (b). With panels (d-f), corresponding to plume ages in 380 simulations. The average BB CO g cm<sup>-3</sup> per grid cell (top) lighter shading indicated higher 381 concentration and transported CO plume ages (bottom) lighter shading indicates older air. 382

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The FLEXPART-WRF plume origin path (forward trajectories) has been used to 384 calculate residence times of air parcels above St. Louis (Figure 5). Figure 5a-5f show the average 385 residence times of the biomass burning plume origin paths expressed as plume age and CO tracer 386 particle amounts as concentrations. Results are displayed for the BASELINE simulation (Figure 387 388 5a and 5d), PYRO simulation (Figure 5b and 5e), and combined (Figure 5c and 5f). The lighter shading represents a higher concentration of CO particle counts (Figure 5a-5c) and for plume age 389 the lighter shading indicates older air (Figure 5d-5f). The major contributions below 3 km are 390 from the central U.S. fires. The contributions from the intermountain west U.S. fires dominates 391 above 3 km, in the range of 80 to 90%. Other regions contribute in the range of 5 to 10%. 392 Biomass burning plumes are indicated to have impacted the surface ozone each day but with 393 394 varying magnitudes. The surface impacts were connected to mechanisms causing air parcels to move downward (e.g. shortwave on 30 August). Likewise, strong vertical motion was evident in 395 bringing simulated air within the boundary to the middle and upper portions of the troposphere. 396

For example, on 24-26 August at the 8 to 12 km layer, both simulations indicate enhancements 397 of 0.74 ppm of CO that has been aged ~7 days. This enhancement corresponded to ~40 ppb of 398 excess O<sub>3</sub> at the 11 to 12 km layer in the ozonesondes. Ozone impacts for August from biomass 399 burning, sharply increase after 3 km and vary daily above; up to 12 km where the biomass 400 burning contributions to ozone decrease to near negligible amounts. The high-altitude smoke 401 injection simulation indicates that plumes that get injected above the 4 km height can impact the 402 surface. Evidence of aged lofted air, ranging from 4 to 15 km, impacting the surface is seen 22 403 August to 1 September 2013, where a constant ~2 ppm of CO enhancement at the surface is 404 present. Additional evidence of biomass burning emissions enhancing ozone within the lower 405 tropopause and PBL (0 to 6 km) is present on 21-24 August over the St. Louis area — a greater 406 than 15 ppby enhancement of ozone within a 6 day aged plume containing an additional 9 ppm 407 of CO. 408

The average plume age for the August month ranged between 4 to 7 days. Plumes 409 reaching the lower troposphere (below 3.5 km) were  $\sim$ 3 days fresher then plumes above. The 410 simulated high-altitude smoke injection is typically  $\sim 2$  days fresher than the boundary layer 411 injected smoke when above 3 km but can be 2 days older when below. The average aged plume 412 lead to 15 ppbv increase in ozone. As plumes aged, they became more ozone enriched. Fresher 413 plumes have higher levels of CO but lower amounts of ozone. An example of this occurred 23 414 August, where and ozone exceedance at the surface was indicated in both simulations to be from 415 a polluted air mass layer with mixed ages. From the lower and middle troposphere (below 1.5 km 416 and above 5.5 km) the plumes ages averaged 2 days older, with about 7 ppm fewer CO, and 20 417 ppbv more ozone than the air in the layers in-between. 418

## 419 4.5. High-altitude smoke injection analysis

The impacts of high-altitude smoke injection are demonstrated using two examples from 420 commonly encountered summertime synoptic meteorological situations in the central U.S. (i) 421 cyclonic flow or a system of low pressure (e.g. cut-off low 17-21 August) and (ii) anti-cyclonic 422 flow or a system of high pressure (e.g. blocking high 25-30 August). Table 3 lists each episode's 423 ozone enriched plume identified by an ozonesonde over St. Louis (Figure 2). For each identified 424 ozone enhancement, the simulated source-contribution relationships from biomass burning 425 (Table 1) and stratospheric intrusions (Table 2) are quantified. The corresponding travel time of 426 the plume (or plume age) and concentrations are indicated by FLEXPART-WRF (Figure 5). St. 427 Louis area for two selected days (21 and 30 August). Each plume was individually characterized 428 with model simulations and ozonesonde observational data in addition to the meteorological 429 conditions leading to the plume reaching St. Louis is discussed below. 430

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Figure 6. Series of daily synoptic atmospheric plane slices at 500 hPa at 7:00 UTC for preceding conditions to individual transport cases (a-c) 21 August 2013 (d-f) 30 August 2013.



Figure 7. Atmospheric plane slices from NARR at 500 hPa at 18:00 UTC (a) 21 August 2013 (b) 445 30 August 2013. Vertical Cross sections at 90° W, 20-60° N 18:00 UT (c) 21 August 2013 (d) 30 446 August 2013. Blue contour lines show Potential Vorticity (PV) 10<sup>6</sup> PVU, Relative Humidity 447 (RH) below 30% is shaded from light to dark, with darker shades representing the dryer air. The 448 black lines on (a) and (b) are pressure height contours in m. While, the black lines on (c) and (d) 449 represent potential temperature (isentropic surfaces) in K. Biomass burning sources tend to have 450 low PV and can have moderate RH levels for pyroCbs, while stratospheric air masses have high 451 PV and low RH. The blue dashed line PV contour (1.5 PV) shows the approximate boundary 452 between stratospheric and tropospheric air masses. Arrows (green) show air-mass transport 453 454 patterns from Stratospheric-Tropospheric Transport (STT) and from Biomass Burning (BB). 455



Figure 8. NOAA Hazard Management System (HMS) and Smoke Product analysis of smoke (a)
21 Aug 2013 and (b) 30 Aug 2013. Below, three-day backward FLEXPART-WRF trajectories
corresponding to each episode is provided (c) 21 Aug 2013 and (d) 30 Aug 2013. Additional
trajectories are placed in the data archive at <a href="https://tropo.gsfc.nasa.gov/seacions/">https://tropo.gsfc.nasa.gov/seacions/</a>.

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## 4.5.1. 17-21 August 2013: Evidence of biomass burning advected by a cutoff low

The meteorological map provided in Figure 6a-c indicates biomass burning transported 464 by a cut-off low originating on 17 August 2013 00:00 UTC at 500 hPa over central Missouri. In 465 this case the cut-off low was created by a closed upper-level low pressure system that was cut off 466 from a westerly current (14 August 06:30 UTC) and it began moving independently. The cut-off 467 low remained nearly stationary until 19 August 00:00 UTC, at which time the trough filled in 468 and moved eastward. Due to the nature of the quasi-stationary system, the westerly flow of air 469 propagated the western fire plumes over the central U.S. and southeast U.S. Figure 7a and 7c 470 present a cross section and plane view of the resulting low. Due to the low pressure system the 471 472 trajectory analysis indicated that smoke flowed from various fire locations to the St. Louis area





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Figure 9. Vertical tropospheric profiles over St. Louis at ~18:30 to 19:30 UTC for selected test
cases 21 August 2013 and 30 August 2013. Labels P1 to P6 correspond to plume information in
Table 3. Ozone measured from the ozonesonde in ppb are shown as a black line. Panel (a) and
(d), the green line RH %. Panel (b) and (e), the blue line represents the GEOS-5 modeled
Potential Vorticity 10<sup>6</sup> PVU. Panel (c) and (f), the FLEXPART-WRF modeled CO biomass
burning g cm<sup>-3</sup>, for each simulation the PYRO simulation is red, and BASELINE simulation in

484 gray. Refer to Figure 2 for corresponding ozonesonde curtain plots.

Three ozone enhanced layers are visible on 21 August (Table 3, Figure 9). None of these 486 enhancements are deemed of stratospheric origins, as there was no significant PVU present in 487 these layers. Plume P3 at 11 km was initially considered to be a stratospheric intrusion related 488 but are ruled out as the NARR data and sonde thermal tropopause indicate that the tropopause 489 490 was lowered (15 km to ~9 km) possibly by wave breaking from the cut-off low. The lower- and upper-middle tropospheric ozone enhancements at P1 (2 to 5.5 km) and P2 (7 to 9 km), 491 respectively, are determined to be of biomass burning origin. The lower-tropospheric plume P1 492 was determined to be from the Idaho Beaver Creek plume ~6 days earlier. The upper-middle 493 tropospheric plume P2 originated 2 days prior from high-altitude injection plumes from the 494 Yellowstone National Park Emigrant cluster fires. The contribution of 21 August tropospheric 495 ozone layers from biomass burning was 10 to 25 ppbv. The pyroCb from the Idaho Pony Elk Fire 496 (16 Aug at 23:41 UTC) reached the St. Louis area; But, the polluted air remained in the lower 497 stratospheric air and did not mix with the lower layers. The incorporation of the high-altitude 498 injection plumes allowed for the two middle- and upper- tropospheric polluted plumes to be 499 characterized which contributed ~15 to 30% to layers in the total ozone column. This upper level 500 pollution was later advected down to the surface and contributed to the exceedance on the 23 501 August. 502

503 504 4.5.2. 25-30 August 2013: Evidence of stratospheric air mixing with aged biomass burning during anticyclonic flow

The meteorological map provided in Figure 6d-f indicates aged biomass burning 505 transported into a stratospheric intrusion by a high pressure system that became well-established 506 over Missouri 25 August 00:00 UTC. The high pressure system began developing a week prior 507 over the southcentral U.S. The high remained quasi-permanent as it sat over Missouri reaching 508 its maximum strength on 26 August 06:00 UTC until a mid-level (500 hPa) shortwave trough 509 moved into the area on 29 August 12:00 UTC moving the system eastward. The near surface 510 reflection of the shortwave reaches Eastern North Dakota 29 August 18:00 UTC as the trough 511 extends south to Oklahoma where it penetrates deeper into the ridge. Figure 7b and 7d presents a 512 cross section and plane view of the resulting high. Due to the high pressure system the trajectory 513 analysis indicated that smoke flowed from various fire locations in additional to stratospheric 514 intrusion impacts to the St. Louis area (see Figure 8). Figure 9d-f depicts the vertical ozone and 515 meteorological profiles representative of this time period (30 August). Further evidence of this 516 trough effects on elevated ozone aloft in the sonde measurements (see Figure 2) remained until 517 30 August 18:00 UTC (Figure 9d-f). Thereafter a shortwave passes the area slightly to the east 518 and cleared out the excess ozone. 519

520 Three ozone enhanced layers are present during the high pressure event as evident on 30 August (Table 3, P4 to P6). Stratospheric air descended into the middle troposphere ~500 hPa on 521 two occasions during this time period (28 Aug and 30 Aug) from shortwaves (Figure 7d). 522 Stratospheric impacts are indicated at  $\sim$ 7 to 9 km and >12 km layers with high PVU values and 523 low RH values (Figure 9d-f, P5 and P6). NARR PVU analysis provided further proof that these 524 are in fact stratospheric in origin. Figure 7d depicts lowering dry stratospheric air and ascending 525 526 moist tropospheric air. The near surface plume (0 to 3.5 km) was traced back to southeastern agricultural fires ~5 days prior (see Figure 8), indicating a likely recirculation of air by the high 527 pressure system. The middle-tropospheric plume P5 was determined to be from an unknown 528

source, potentially a recirculation of summertime ozone precursors (Cooper et al., 2006, 2007). 529 Evidence of this layer is present in previous soundings for the week 25-29 August where all 530 sondes show a single well-defined plume at ~5 to 9 km. The GEOS-5 and NARR potential 531 vorticity also give no indication of what the cause for this layer is. Our hypothesis is supported 532 by the vertical wind profiles for the week being relatively stagnant— the ozonesonde launched 533 that day landed within 50 m of the launch site — signifying air circulation over St. Louis for the 534 entire week increasing photochemical effects. An alternative theory is that wind shear caused the 535 layer and it's a part of the larger plume above. The layer between the unknown plume and above 536 (~6 to 8 km) is a layer of moist and clean air (see Figure 7d). The upper-tropospheric layer 537 plume P6 is a combination of both a biomass burning plume and a stratospheric intrusion air 538 mass. The lower half of the plume P6 (8 to 10 km) is dominated by stratospheric air, while the 539 above portion of P6 (10 to 12 km) is primarily biomass burning. There is another layer of clean 540 air above prior to reaching the tropopause. The upper-tropospheric plume P6 originated 5 days 541 prior from high-altitude injection plumes from the California Rim Fire (25 August). The 542 contribution of 30 August tropospheric ozone layers from Biomass burning was 10 to 80 ppbv 543 and stratospheric air masses contributed 10 to 40 ppby. The incorporation of high-altitude smoke 544 injection allowed for the two mid- and upper- tropospheric polluted plumes to be characterized 545 which contributed  $\sim 15$  to 60% to layers in the total ozone column. 546

## 547 **5 Summary and Conclusions**

548 By incorporating balloon-borne ozonesonde observations with models, this study has quantitatively examined sources for tropospheric ozone enhancements due to Non-Controllable 549 Ozone Sources (NCOS). In particular, the pollution impacts from stratospheric intrusions and 550 551 biomass burning contributions to background tropospheric and surface ozone levels in the Midwest United States were characterized. Additionally, emphasis is placed on partitioning the 552 contribution from western U.S. fires, central U.S. fires, and other areas to St. Louis background 553 554 ozone. A chemical transport model and trajectory model were run to quantify source contribution to ozone in St. Louis, Missouri. For the region and time period of this study, 10 to 15% of the 555 ozone enhancements stems from a stratospheric airmass contribution and 15 to 30% from 556 557 biomass burning. These NCOS contributions and ozonesonde profiles can be considered as baseline ranges for the Midwest U.S. area if direct ozone measurements (sondes, airplane, 558 ground-based) are not available. Considering U.S. fires only, 70% of the biomass burning 559 plumes originated from the western parts of the U.S. and only 3% came from the local central 560 U.S. emissions. Moreover, it was demonstrated that a redistribution of the biomass burning 561 emissions injection height, with part of the emissions above the boundary layer led to a reduction 562 of model predicted surface ozone. 563

In agreement with earlier studies (e.g., Fishman et al., 2014), this study has identified a 564 generally increasing relationship between background ozone and transported pollution. We 565 followed the definition of background ozone described in Jaffe et al. (2018) as NCOS such as 566 lingering biomass burning and long-range transported international sources. The major 567 contributions below 3 km were from the central U.S. fires. While 80 to 90% of the high-altitude 568 injection smoke (above 3.5 km) originated in the western U.S. During this campaign period only 569 5 to 10% of the biomass burning emissions reaching St. Louis originated from the southeast and 570 other regions. 571

This study identified that biomass burning plumes in the western U.S. can have impacts 572 on the daily atmospheric ozone column in the Midwest (10 to 80 ppbv of ozone) at a greater 573 frequency and intensity than stratospheric intrusion (10 to 25 ppbv of ozone). We show the 574 575 background ozone to be 55 ppbv, which was near the 30 to 50 ppbv range mentioned in Jaffe et al. (2018) which is typical for the U.S. We identified a relationship between smoke plume age 576 and ozone enhancement where the high-altitude injection smoke plumes above 3.5 km generally 577 were associated with higher amounts of CO concentrations but fresher smoke regarding ozone 578 579 levels. In addition, we recognized that the high-altitude smoke had a higher tendency to mix with stratospheric intrusions, which together doubled the ozone enhancement in the tropospheric 580 column. 581

An investigation of several individual smoke plumes has shown the importance of 582 583 incorporating high-altitude smoke injection in model simulations in addition to ensuring that accurate biomass burning locations and temporal allocation of intensity are included in model 584 emissions. Up to 60% of the smoke plume lies above 3.5 km, and this needs to be simulated as it 585 can later be mixed down to the surface and lead to ozone exceedances days later. In addition, it 586 was shown that the incorporation of satellite-based detections of high-altitude smoke injection 587 (e.g., pyroCb activity, Peterson et al., 2014, 2017a,b) can be helpful for improve modeling results 588 and explain ozone enhancements aloft and at the surface. 589

The individual cases were selected because they are associated with common 590 meteorological situations in the Midwest leading to ozone exceedances. Evidence from the first 591 case where biomass burning is advected by a cutoff low is an example of a common flow pattern 592 that transports air masses from the west. Summertime occurrences of this synoptic situation in 593 conjunction with large wildfires in the western U.S. can lead to increases in ozone in the 594 Midwest of 10 to 80 ppby or greater. In the second case a stratospheric air mass was shown to 595 mix with an aged wildfire plume during anticyclonic flow, a pattern that was previously found to 596 occur 40% of the time for the southcentral U.S. (Texas and the Gulf of Mexico area, Brioude et 597 598 al., 2007). Additionally, the test cases showed that the surface impacts were connected to mechanisms causing air parcels to move downward (e.g. shortwave on 30 August). Likewise, 599 strong vertical motion was evident in bringing simulated air within the boundary to the middle 600 and upper portions of the troposphere. 601

While the results in this study highlight that Non-Controllable Ozone Sources can 602 contribute significantly to local tropospheric ozone in the Midwest, future studies must combine 603 satellite data and model integration techniques with meteorological information for a longer time 604 period, both within and outside the Midwest to better characterize NCOS contribution to U.S. 605 606 ozone. Good examples of how an air quality model could be used to assess NCOS has been shown in Baker et al. (2016, 2018). More specifically, Baker et al. (2016, 2018) address the 607 question of regional-scale pyrogenic ozone sources. In particular, Baker et al. (2018) 608 investigated the Rim Fire, using non-sonde data, which occurred during our study period with a 609 photochemical model and their findings were similar to ours. Furthermore, as shown by 610 individual plume cases and several earlier studies (e.g. Morris et al., 2006; Brioude et al., 2007; 611 Jaffe, 2011; Lin et al., 2012; Hess and Zbinden, 2013), high-altitude smoke injection from a fire 612 can lead to long distance transport depending on weather conditions and hence impact surface 613 ozone and NAAQS attainment. Plume rise and plume injection heights are a key source of 614 uncertainty (Paugam et al., 2016), which can be reduced considerably using plume height 615 information from remote sensing tools (e.g. Peterson et al., 2017a,b; Val Martin et al., 2018). 616

- 617 Improved modeling techniques will be required to better characterize biomass burning transport
- and hence better simulate long range impacts and the possibility of unhealthy surface
- 619 concentrations far from the biomass burning sources.

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- 628 <u>https://www.nasa.gov/mission\_pages/seac4rs/index.html</u> (mission data e.g., fire emissions and
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**Figure 1**. Study area map with FLEXPART-WRF model domain. Rectangles mark the WRF

- domains D1 which has a horizontal resolution of 12 km and D2 (4 km resolution). SEACIONS
- locations are marked by black circles (St. Louis, Missouri station (STL) is a gray circle). Wildfire
- emissions from FLAMBE, August-September 2013, with eleven Pyro convective and five high-

altitude injection elevated smoke plume activity areas are marked, red crosses indicate pyroCb

- plumes which transport smoke into the upper troposphere and lower stratosphere. Blue crosses
- 921 indicate high-altitude injection of smoke in the absence of large pyroCb which transport smoke
- into the mid-troposphere).
- **Figure 2.** Ozonesondes in support of SEACIONS were launched at the Saint Louis University
- ozonesonde station located at the James S. McDonnell Planetarium in Forest Park (90.27°W,
- 925 38.63°N, 181 m asl), 5 km west of downtown St. Louis. Vertical tropospheric profiles at ~18:30
- 926 UTC (1:30 p.m. Local time) ozone profiles are averaged vertically every 500 m and shown
- 927 below the thermal tropopause level ( $\sim$ 15 km). Cases of ozone enhancements above the
- background (~55 ppbv) are shown by source origin: the gray box for Stratospheric-Tropospheric
   Transport (STT), the solid black line boxes for biomass burning (BB), and the dashed line black
- <sup>929</sup> Transport (511), the solid black line boxes for blomass burning (BB), and the dashed line black <sup>930</sup> boxes for combined STT and BB
- boxes for combined STT and BB.
- Figure 3. Biomass burning emissions of carbon monoxide (CO) estimated by FLAMBE in units
  of Gg per km<sup>2</sup> per hour in North America during SEACIONS, 8 Aug to 22 Sept 2013. FLAMBE
  emissions are displayed on the model grid which has a resolution of 0.25° per cell. Only cells
- with emissions above 2,000 kg are shown.
- Figure 4. A time series of FLAMBE emissions (mass/time) converted to particles released in 935 FLEXPART-WRF (# of particles or particle count). Each 500 kg of mass is converted to 1 936 particle in FLEXPART-WRF simulations. Locations emitting less than 500 kg only emit a single 937 particle. Arrows represent the duration of two of the largest wildfires (Rim Fire and Beaver 938 Creek) with relevance to SEACIONS with other fires (Emigrant Fire and Pony/Elk Complex) 939 contributing to those time frame emissions indicated. The dotted line is the California Rim Fire 940 (37.85°N, 120.083°W) 17-31 August 2013, and the solid line is the Idaho Beaver Creek 941 (43.593°N, 114.684°W) 7-21 August 2013. Note: Beaver Creek and Pony/Elk Complex occurred 942 nearly at the same time and were fewer than 100 km apart in distance. While, the Emigrant Fire 943 contributed a significant amount of emissions during its occurrence at the same time as the Rim 944 Fire but was much smaller in magnitude. 945
- Figure 5. FLEXPART-WRF Biomass burning (BB) CO concentrations and age are simulated, 946 binning trajectories over St. Louis within a 2.5° x 2.5° grid box and averaged vertically 500 m 947 with daily temporal resolution. FLAMBE emissions of BB CO are converted into particle 948 amounts and are released. (a) The base simulation, Boundary Layer Simulation, releases 949 emissions within the PBL, (b) NRL's adjusted pyro-convention scheme is implemented, PyroCb 950 Simulation, particles are released from pyroCbs and high-altitude injection as specified in Table 951 1. and (c). The combined (a) and (b). With panels (d-f), corresponding to plume ages in 952 simulations. The average BB CO g cm<sup>-3</sup> per grid cell (top) lighter shading indicated higher 953 concentration and transported CO plume ages (bottom) lighter shading indicates older air. 954
- Figure 6. Series of daily synoptic atmospheric plane slices at 500 hPa at 7:00 UTC for preceding
  conditions to individual transport cases (a-c) 21 August 2013 (d-f) 30 August 2013.

- **Figure 7.** Atmospheric plane slices from NARR at 500 hPa at 18:00 UTC (a) 21 August 2013 (b)
- 958 30 August 2013. Vertical Cross sections at 90° W, 20-60° N 18:00 UT (c) 21 August 2013 (d) 30
- August 2013. Blue contour lines show Potential Vorticity (PV) 10<sup>6</sup> PVU, Relative Humidity
- (RH) below 30% is shaded from light to dark, with darker shades representing the dryer air. Theblack lines on (a) and (b) are pressure height contours in m. While, the black lines on (c) and (d)
- black lines on (a) and (b) are pressure height contours in m. While, the black lines on (c) and (d) represent potential temperature (isentropic surfaces) in K. Biomass burning sources tend to have
- low PV and can have moderate RH levels for pyroCbs, while stratospheric air masses have high
- PV and low RH. The blue dashed line PV contour (1.5 PV) shows the approximate boundary
- between stratospheric and tropospheric air masses. Arrows (green) show air-mass transport
- patterns from Stratospheric-Tropospheric Transport (STT) and from Biomass Burning (BB).
- **Figure 8.** NOAA Hazard Management System (HMS) and Smoke Product analysis of smoke (a)
- 969 21 Aug 2013 and (b) 30 Aug 2013. Below, three-day backward FLEXPART-WRF trajectories
- corresponding to each episode is provided (c) 21 Aug 2013 and (d) 30 Aug 2013. Additional
- 971 trajectories are placed in the data archive at <u>https://tropo.gsfc.nasa.gov/seacions/</u>.
- **Figure 9.** Vertical tropospheric profiles over St. Louis at ~18:30 to 19:30 UTC for selected test
- cases 21 August 2013 and 30 August 2013. Labels P1 to P6 correspond to plume information in
- Table 3. Ozone measured from the ozonesonde in ppb are shown as a black line. Panel (a) and
- (d), the green line RH %. Panel (b) and (e), the blue line represents the GEOS-5 modeled
- Potential Vorticity 10<sup>6</sup> PVU. Panel (c) and (f), the FLEXPART-WRF modeled CO biomass
- 977 burning g cm<sup>-3</sup>, for each simulation the PYRO simulation is red, and BASELINE simulation in
- 978 gray. Refer to Figure 2 for corresponding ozonesonde curtain plots.

**Table 1.** Elevated smoke plumes escaping the planetary boundary layer identified by the Naval

Research laboratory satellite detection algorithm for high-altitude injection during the Studies of

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983 (SEAC4RS) in the summer 2013. Smoke plumes detected are categorized: 1. A pyroconvective

plume containing smoke reaching the stratosphere ( $\sim 10$  to 15 km) or 2. A high-altitude plume

containing smoke reaching into the middle troposphere ( $\sim$ 5 to 9 km).

Date	Time	Lat	Lon	State	Fire name	Plume type
	(UTC)					
7 Aug 2013	22:15	42.71	-123.66	OR	Big Windy Complex	High-altitude
8 Aug 2013	n/A	43.46	-114.55	ID	Beaver Creek	Pyroconvective
8 Aug 2013	n/A	43.42	-114.85	ID	McCan	Pyroconvective
9 Aug 2013	22:41	43.29	-115.55	ID	Pony/Elk	Pyroconvective
10 Aug 2013	21:00	43.36	-115.45	ID	Pony/Elk	Pyroconvective
12 Aug 2013	21:00	43.29	-115.55	ID	Pony/Elk	Pyroconvective
12 Aug 2013	23:00	43.62	-114.51	ID	Beaver Creek	Pyroconvective
13 Aug 2013	23:41	43.50	-115.43	ID	Pony/Elk	Pyroconvective
16 Aug 2013	21:41	45.66	-114.78	ID	Gold Pan Complex	Pyroconvective
16 Aug 2013	23:41	43.62	-114.51	ID	Pony/Elk	High-altitude
16 Aug 2013	23:41	43.46	-114.55	ID	Beaver Creek	Pyroconvective
19 Aug 2013	22:50	45.20	-110.69	MT	Emigrant	High-altitude
19 Aug 2013	23:15	37.86	-120.09	CA	Rim	Pyroconvective
21 Aug 2013	22:41	37.86	-120.09	CA	Rim	Pyroconvective
22 Aug 2013	21:30	41.04	-123.47	CA	Corral Complex	High-altitude
25 Aug 2013	17:00	37.91	-120.00	CA	Rim	High-altitude

Table 2. Simulated stratospheric air masses entering the troposphere (stratospheric intrusion)
 over St. Louis, Missouri during August and September 2013.

Date	Time	Duration	Height	Extent	Meteorology
	(UTC)	(hr)	(hPa)		
3 Aug 2013	03:30	21	580	Shallow	Frontal passage
7 Aug 2013	13:30	28	690	Deep	Shortwave
10 Aug 2013	02:30	34	800	Deep	Shortwave
12 Aug 2013	18:30	28	500 to 850	Deep	Frontal passage
14 Aug 2013	06:30	180	600	Shallow	Cutoff low
23 Aug 2013	16:30	9	500	Shallow	Shortwave
28 Aug 2013	21:30	18	480	Shallow	Shortwave
30 Aug 2013	08:30	94	500	Shallow	Shortwave
3 Sept 2013	12:30	30	700 to 1000	Deep	Frontal passage
6 Sept 2013	09:30	41	600	Shallow	Shortwave
12 Sept 2013	16:30	10	800	Deep	Frontal passage
15 Sept 2013	20:30	48	800 to 1000	Deep	Frontal passage
22 Sept 2013	00:30	94	900	Deep	Frontal passage

- **Table 3.** Ozone (O<sub>3</sub>) enriched plumes identified over St. Louis, Missouri using ozonesondes
- vertical profile measurements for two episodes. Included are the simulated biomass burning
- plumes and stratospheric air masses entering the troposphere (stratospheric intrusion (SI))
- concentrations of a carbon monoxide (CO) tracer for each plume. Plume ID corresponds to
- 993 labeling in Figure 9.

Plume	Date	Source	Plume	Smoke	BASELINE	PYRO	Sonde	Sonde	Excess
ID			height	age	simulation	Simulation	RH	$O_3$	$O_3$
			AGL km	days	CO ppm	CO ppm	%	ppb	ppb
P1	21 Aug	ID Beaver Creek	2 to 5.5	5.8	2.4	14	10	65	10
P2	21 Aug	MT Emigrant	7 to 9	2.6	0	5.2	5	85	25
Р3	21 Aug	pyroCb and SI	11	1.2	0	0.63	20	65	10
P4	30 Aug	SE US Ag. Fire	0 to 2.5	5.1	6.6	0.83	85	65	10
Р5	30 Aug	Unidentified	5 to 7	6.7	0	0.35	10	100	45
P6	30 Aug	CA Rim Fire and SI	8 to 12	5.3	0	6.8	15	140	80