Exploring the Potential of Using Carbonyl Sulfide to Track the Urban Biosphere Signal

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Abstract  Cities are implementing additional urban green as a means to capture CO₂ and become more carbon neutral. However, cities are complex systems where anthropogenic and natural components of the CO₂ budget interact with each other, and the ability to measure the efficacy of such measures is still not properly addressed. There is still a high degree of uncertainty in determining the contribution of the vegetation signal, which furthermore confounds the use of CO₂ mole fraction measurements for inferring anthropogenic emissions of CO₂. Carbonyl sulfide (OCS) is a tracer of photosynthesis which can aid in constraining the biosphere signal. This study explores the potential of using OCS to track the urban biosphere signal. We used the Sulfur Transport and deposition Model (STEM) to simulate the OCS concentrations and the Carnegie Ames Stanford Approach ecosystem model to simulate global CO₂ fluxes over the Bay Area of San Francisco during March 2015. Two observation towers provided measurements of OCS and CO₂: The Sutro tower in San Francisco (upwind from the area of study providing background observations), and a tower located at Sandia National Laboratories in Livermore (downwind of the highly urbanized San Francisco region). Our results show that the STEM model works better under stable marine influence, and that the boundary layer height and entrainment are driving the diurnal changes in OCS and CO₂ at the downwind Sandia site. However, the STEM model needs to better represent the transport and boundary layer variability, and improved estimates of gross primary productivity for characterizing the urban biosphere signal are needed.

1. Introduction

Cities are home to 50% of the world population and potentially 70% by 2050 (UN/DESA, 2018). Consequently, cities are major energy consumers and contributors of anthropogenic greenhouse gas (GHG) emissions, representing almost two-thirds of the global energy demand and 70% of carbon emissions from the energy sector (IEA, 2016). Aware of their critical role in mitigating GHG emissions, many cities are adopting measures to reduce the impact of energy-intensive activities, such as urban passenger travel, residential heating, and waste and waste-water treatment through initiatives such as the Covenant of Mayors (Covenant of Mayors, 2009) and the Local Governments for Sustainability (ICLEI, http://wwwICLEI.org). According to the International Energy Agency, cities must take the lead to limit the global temperature increase to no more than 2°C outlined by the Paris Agreement (United Nations/Framework Convention on Climate Change, 2015). To measure the efficiency of reduction strategies and track progress over time to support policy intervention, cities must establish ways of tracking GHG emissions that are consistent, repeatable and reliable.

The most widely used method for GHG emissions accounting at city level is a bottom-up approach based on energy consumption multiplied by emissions factors specific for each type of fuel and process (GPC, 2014;
ICLEI, 2012; Kennedy et al., 2009). There are several standardized methodologies such as those presented by Kennedy et al., but in general most methods involve determining the CO₂ emissions from electricity, heating and industrial fuels, ground transportation fuels, air and marine fuels, industrial processes and waste. Thousands of cities have committed to report their emissions this way by signing the covenant of mayors agreement (Covenant of Mayors, 2009). However, given the vast amount of time and resources required to carry out such bottom-up inventories, generic emissions factors and data assumptions are often used to calculate the carbon footprint potentially resulting in highly inaccurate estimates of GHG emissions. CO₂ emissions estimated this way at the urban scale have been shown to have an uncertainty of 50%–200% (Asefi-Najafabady et al., 2014; J. C. Turnbull et al., 2011). The bottom-up approach also fails to accurately determine the contribution of urban CO₂ capture by green infrastructures, which are currently being promoted as a means to reduce the carbon footprint (Byrne et al., 2020). Some of the limitations include uncertainty in the time scales for respiration (RESP) of carbon previously taken up through photosynthesis and the effect of urban climate and topography on the urban biosphere. Furthermore, equations that predict biomass generation (carbon sequestration) derived from measurements are not applicable to urban green because of its unique form and maintenance (Pataki et al., 2011).

In recent years, “top-down” simulations of GHG at the urban scale have used bottom-up anthropogenic GHG emission inventories to determine urban GHG concentrations using atmospheric and chemical transport models (Feng et al., 2016; J. C. Turnbull et al., 2014). In other cases, atmospheric measurements of the CO₂ concentrations are combined with atmospheric chemical transport models to determine the CO₂ emissions through inverse simulations (McKain et al., 2012; Newman et al., 2013; Turner et al. 2016) and flux estimation methods (Mays et al., 2009). These modeling efforts require biogenic emissions as input which are still highly uncertain because there is significant divergence among the various biosphere models that provide this biospheric component of the carbon budget (Pugh et al., 2016; Sitch et al., 2008). The biogenic emissions are further confounded at the urban level because anthropogenic heat fluxes, impervious surfaces, phenological shifts, and air quality can alter spatial and temporal patterns of biogenic carbon fluxes compared to natural systems (Krupa & Manning, 1988; Melaas et al., 2016; Zhao et al., 2016).

Determining the urban biosphere contribution to the urban CO₂ budget from measurements also has its complexity. One approach is to constrain the anthropogenic CO₂ instead by using CO or C¹⁴ CO₂ as gas tracers of fossil-derived CO₂ (Campbell et al., 2008; LaFranchi et al., 2013; J. C. Turnbull et al., 2011; J. Turnbull et al., 2009; Levin et al. 2011) (Miller et al., 2012). These studies have served to quantify discrepancies between bottom-up CO₂ inventories with observations, but have not provided much information in terms of the urban biosphere contribution to the overall carbon budget. While anthropogenic tracers exist, biosphere tracers have not yet been applied in urban analysis.

Thus quantifying the contribution of urban vegetation to the overall CO₂ budget in urban areas is a challenge. Several studies have pursued this endeavor, not only to determine the effectiveness of mitigating GHG by incrementing urban green, but also to accurately monitor, report, and verify such strategies. Hardiman et al. (2017) adapts the Vegetation Photosynthesis and Respiration Model (VPRM) to urban phenology for the cities of Massachusetts, and estimates that urban vegetation absorbs 14% of the anthropogenic emissions, but that this benefit is significantly offset by the urban heat island effect and the asynchrony between temporal patterns of biogenic and anthropogenic carbon fluxes (Hardiman et al., 2017). Park et al. also use an optimized VPRM together with Hestia fossil fuel CO₂ emission data and estimate higher biospheric contribution in the Southern California Air Basin, uptaking 20%–24% of the total anthropogenic CO₂ during the daytime (Park et al., 2018). Using a combination of Light Detection and Ranging (LiDAR) and field techniques, another study was able to quantify that the trees of the city of Meran, Italy, were able to sequester only 0.61% of the traffic emissions (Speak et al., 2020). Eddie co-variance flux estimates performed by Velasco et al., however, show that when considering vegetation and soil together, the biogenic component was found to actually add and extra 4.4% to the total CO₂ flux of a neighborhood in Singapore (Velasco et al., 2016).

To further improve understanding of the influence of urban vegetation, here we explore the potential of using the atmospheric tracer carbonyl sulfide (OCS or COS) to provide information about the contribution of the urban biosphere to the overall carbon budget. Observations of OCS concentrations have recently been used to estimate gross carbon uptake of the biosphere at regional and global scales (Campbell et al., 2017;
This technique is based on the close relationship of CO$_2$ and OCS uptake by plants (Campbell et al., 2008; Sandoval-Soto et al., 2005; Stimler et al., 2011). In contrast to CO$_2$, plants can take up OCS but do not release it during respiration; this phenomenon allows OCS to be used to estimate the gross primary productivity (GPP) (i.e., the total amount of carbon fixed by photosynthesis in the ecosystem). OCS has been demonstrated to be a good predictor for GPP in forests (Rastogi et al., 2018; Spielmann et al., 2019) and orchards (Yang et al., 2018); although some studies also reported that there is OCS production from marsh vegetation (Whelan et al., 2013), crops (Bloem et al., 2012), and paddy fields (Yi et al., 2008). Previous studies that use OCS as a tracer for GPP have focused on highly vegetated areas at the canopy-scale (Kooljmans et al., 2017) or continental and global scales (Campbell et al., 2017; Hilton et al., 2017).

More recently, high-resolution temporal and spatial surveys of OCS and CO$_2$ mixing ratios confirm that OCS may be a proxy for CO$_2$ uptake in homogenous ecosystems on a local scale (Belviso et al., 2016; Campbell et al., 2017). However, less is known about the usefulness of using OCS as a GPP tracer in urban environments, where the complexity of urban and natural activity could include additional sources and sinks of OCS. In this respect, Belviso et al. (2020) conclude that long-range transport of OCS from anthropogenic sources located in Benelux, Eastern France and Germany impacts the Paris area during wintertime, limiting the use of OCS to assess regional-scale CO$_2$ uptake because of such strong anthropogenic influence. The present study reduces the potential influence of anthropogenic OCS sources by performing the measurements and simulations in the coastal area of San Francisco with frequent westerly winds from the ocean and where the anthropogenic OCS sources have been deemed insignificant (Zumkehr et al., 2018).

On this premise, and with the overall goal of learning how to best quantify the biosphere contribution using OCS as a tracer, we developed a 9-km resolution observation-model framework of the densely populated San Francisco region in California. We used regional atmospheric chemical transport simulations to interpret observed OCS concentrations from an urban tower and explored the drivers of atmospheric OCS variation in the densely populated area.

### 2. Methods

We use the Sulfur Transport and dEposition Model (STEM) (Carmichael et al., 1991) to simulate the OCS concentrations over the densely populated San Francisco Bay Area at 9-km resolution during March 2015 at one hour temporal resolution. The domain, shown in Figure 1, is a 50 by 50 horizontal grid with 59
pressure-based vertical layers centered around the city of Livermore and is characterized by urban land use, croplands of the San Joaquin Valley, and forests of coastal areas and the Sierra Nevada. There is photosynthetic activity all year in this domain, with the most active period being February through June, which is the wet season. The study captures the month of March, which is the beginning of the most active time of year for photosynthetic activity, peaking in April.

There are two observation towers that provided measurements of OCS and CO2 for this study. One is the Sutro tower located in San Francisco (37°45′18.72″N, 122°27′10.08″W), which stands 298 m above ground and is 70 km upwind from Livermore. Given itsproximity to the ocean and the predominance of westerly winds, Sutro tower provided a representation of background concentrations most of the time. The second tower is located at Sandia National Laboratories in Livermore (Lon: 121.71°, Lat: 37.67°) (hereafter referred to as Sandia tower, which is downwind of the highly urbanized San Francisco region with an inlet height of 27 m a.g.l.). OCS and CO2 samples were collected using 0.7-liter flasks filled to 40 psia over a few minutes at Sutro tower twice a day, which shipped to Boulder, Colorado, for analysis in NOAA's gas chromatography and mass spectrometry laboratory by the methods described by Montzka et al. (2007); at Sandia tower air samples were analyzed hourly, using a Los Gatos Research cavity-enhanced laser absorption spectrometer (Rastogi et al., 2018). CO2 measurements at Sandia tower were referred to 3 NOAA GMD standards via synthetic air transfer standards. The OCS analyzer was calibrated by a CO2-referenced method and was thus limited by the uncertainty of the OCS/CO2 ratio in the standard, estimated to be ±2.8% based on the quadrature sum of the uncertainties in OCS and CO2 (both ±2%). This included uncertainty in the WMO scale of ±0.07 ppm and repeatability of the transfer of the WMO calibration (based on repeatability during transfer standard calibrations) to the synthetic reference gases of ±0.11 ppm (in total ±0.13 ppm). The median replicate precision for OCS flask samples from the NOAA network was 0.4% (~2 ppt), and 95% of all pairs had a replicate precision of less than 1.3% (<6.3 ppt).

We used the Weather Research and Forecasting model (WRF) version 3.7.1 (Skamarock et al., 2008) using the same domain, vertical profile and time period described above for STEM. The configuration of WRF included the planetary boundary layer (PBL) scheme Mellor-Yamada-Janjic, the Noah land surface model, and the Grell-Devenyi ensemble cumulus scheme (Yver et al., 2013). We ran WRF with reanalysis data provided by the Global Forecast System of NOAA's National Center for Environmental Information at 28-km, 6-hour resolution (NOAA, 2017) to generate the meteorological fields needed to run STEM for March 5–25, 2015 (discarding the first 5 days of the run as spin-up period). The hourly WRF output was validated for surface temperature at 2 m and wind speed and direction at 10 meters observed at the Lawrence Livermore National Laboratory weather station, and provided by https://weather.llnl.gov (see Supporting Information S1).

In addition to the meteorological data, we generated hourly surface fluxes of OCS and CO2 as input to STEM for the same time period March 5–25, 2015. We used a global OCS hourly inventory of the biospheric surface-flux at 1.25° × 1° resolution (approximately 87 km by 87 km for this latitude) that was produced by the Simple Biosphere Model SIB3 (Baker et al., 2009; Berry et al., 2013). This inventory was regridded to 1-km resolution using normalized difference vegetation index (NDVI) as a spatial proxy for the STEM regional domain (NASA and USGS, 2017). The steps were to first download modis terra NDVI values in text format at the desired resolution from http://modis-land.gsfc.nasa.gov/vi.html. Next, we ran a fortran script to match the NDVI values to the STEM domain based on the longitude and latitude of each pixel. The result was a map of NDVI values for each cell (1 km by 1 km) of the entire STEM domain. Then we used the OCS flux data from SIB and the following simple equation to determine the OCS value for each cell (or each pixel) of the STEM domain:

\[
\frac{\text{NDVI value of 1 STEM pixel}}{\text{NDVI value of entire STEM domain}} \times \text{OCS flux value of entire STEM domain} = \text{OCS value of 1 STEM pixel}
\]

According to this method, the hourly average OCS flux field generated for the entire study period is presented in Figure 2 at 1 km resolution. The supporting information summarizes the input required for running STEM in Figure S3.

Global CO2 fluxes for GPP and RESP are available from the Carnegie-Ames-Stanford-Approach (CASA) ecosystem model at 1° by 1.25° resolution (NACP, 2015). We regridded each flux from its native grid and resolution to the 50 × 50 grid at 9 km resolution used in this study using the mass-conservative algorithm of the
MTXCLE program of the Models-3/EDSS Input/Output Applications Programming Interface (Coates, 2014). Figure 3a (GPP) and Figure 3b (Respiration) show the regridded fluxes. We used the U.S. high resolution inventory for the fossil-fuel derived anthropogenic CO$_2$ emissions (Gurney et al., 2009; NACP, 2010). This emissions inventory is available hourly at a 10-km resolution, and was regridded to the 9-km resolution as well (see Figure 3c). Unlike the other emissions inventories that were available for the study period, the Vulcan inventory was only available for 2008. Net ecosystem exchange (NEE) is the net exchange of C between the ecosystem and the atmosphere (Kramer et al., 2002). Here we define NEE as the CO$_2$ released to the atmosphere by RESP minus the CO$_2$ sequestered by GPP; a sum of all these fluxes results in total CO$_2$ anomaly from the background (total CO$_2$ anomaly = ANTHRO CO$_2$ + NEE) (Figure 3d).

3. Results and Discussion

3.1. Background Climatology

Figure 4 shows long term observation from the NOAA measurements of flasks collected on the Sutro tower, which is usually upwind from the study area of Livermore and provides background mixing ratios for both OCS and CO$_2$. Comparisons to results from other surface sites in the NOAA network (Montzka et al., 2007, and updates available at ftp://aftp.cmdl.noaa.gov/hats/carbonyl_sulfide/) suggest that the data from Sutro are typically representative of the OCS mole fractions in background air:

(a) The mean mixing ratio measured between 2007 and 2015 of 493 ppt is very similar to the mean measured for Niwot Ridge, Colorado, (494 ppt), which is relatively close in latitude in the Northern Hemisphere (40°N, 106°W); (b) the mean Sutro value is only slightly higher than the mean value of 475 ppt at Trinidad Head (California), which is a coastal site north of Sutro and close to the boarder to Oregon (41°N, 124°W), which is consistent with the observation that OCS mole fractions decrease with higher latitudes in the Northern Hemisphere; (c) the OCS mixing ratios at Sutro do not have strong long term trends, which is also suggested for air at more remote, background sites in the NOAA record; and (d) the seasonality of OCS (phase and amplitude) at Sutro is similar in magnitude to other Northern Hemisphere sites in the NOAA network. The observed average OCS concentration during the study period of March 5–25, 2015 is 505 ppt. This value is used as the background value to initialize the OCS in the STEM model runs. A constant initial condition of OCS seems adequate for this study because our interest is in determining the enhancements rather than absolute concentrations. Furthermore, STEM simulations of OCS fluxes in North America run under different initial conditions showed that GPP variability dominates STEM boundary conditions for OCS concentrations (Hilton et al., 2017).

Sutro tower CO$_2$ observations also show the seasonal variation established by other studies with higher values of CO$_2$ in winter and lower values in summer (Sweeney et al., 2015). The enhanced mole fraction in the winter indicates an urban influence, mainly attributed to the urban biospheric component acting as a source in winter, rather than a sink as it does in the summer (Miller et al., 2012). The average CO$_2$ concentration during the study period of March 5–25 is 405 ppm, similar to background concentrations reported by other monitoring stations such as Mauna Loa, Hawaii, United States (NOAA, 2015). We use this value of 405 ppm as background for the CO$_2$ model runs.

3.2. Urban Observations

We compared background observations at Sutro tower with downwind concentrations at Sandia Tower in Livermore. Background CO$_2$ measured at Sutro are relatively stable, while at Sandia, CO$_2$ concentrations follow a strong diurnal cycle. The result is consistent with the expectation that Sutro air typically represents a relatively stable clean marine boundary layer without recent anthropogenic input, whereas the Sandia
samples were taken in a continental boundary layer (BL) downwind of an urban region and from a continental boundary layer with high diurnal variability in thickness.

CO$_2$ mixing ratios are higher at Sandia than Sutro, reflecting the larger impact of anthropogenic sources within the San Francisco region on air downwind of this urban region. The CO$_2$ observations at Sandia follow a clear diurnal pattern. Concentrations are higher at night and early morning, which is typical of urban areas that are characterized by dominant anthropogenic sources and a diurnally varying boundary layer that is shallowest at night time. The lowest CO$_2$ concentrations at the Sandia tower location occur at noon, when vertical mixing is enhanced at photosynthetic rates are elevated. Highest CO$_2$ enhancements relative to background occur at nighttime when plant CO$_2$ assimilation, anthropogenic activity, and PBL height are all at a minimum. The enhancement maxes despite anthropogenic emissions being at a minimum because the low PBL height and no C uptake by plants is more than enough to compensate for the drop in anthropogenic emissions. Similarly, a study performed for the city of Salt Lake City by McKain et al. (2012) found that the highest concentrations of CO$_2$ occurred during nighttime and early morning (up to 60 ppm above background values), notably out of phase with anthropogenic emissions, which are elevated during the day. The study by McKain also found that CO$_2$ concentrations were lowest from 12 to 8 p.m, with enhancements
relative to background of 0–20 ppm. In natural ecosystems, where there are no other sources of CO$_2$ other than soil and day-to-night differences in CO$_2$ are attributed to GPP and RESP (Knox et al., 2018). This explanation may not apply to urban areas, where anthropogenic CO$_2$ emissions are expected to dominate local mole fractions. In an attempt to separate anthropogenic from biogenic sources of urban CO$_2$, Wei et al. (2020) attributed the highest CO$_2$ at early morning (7:00) to the unusual high traffic during this time, with a peak-to-peak amplitude of 12.5 ppm compared to lower values occurring at midday (14:00), when atmospheric boundary-layer mixing and biogenic uptake (i.e., photosynthetic peak) maximize reducing atmospheric CO$_2$ mole fractions.

The lower panel of Figure 5 shows the observed concentrations of OCS. At Sutro, OCS mixing ratios showed variability between 450 and 550 ppt until March 21, at which point the concentrations stabilized around 550 ppt. One potential explanation for this change in OCS concentration is that continental air can sometimes be transported offshore in the residual layer and then entrained in the marine boundary layer, as was found by Riley (2005). This hypothesis is further validated by the fact that, after March 21, there were stable westerly winds that transported constant OCS (background) concentrations from above the ocean. Before March 21, the wind vectors indicate that the air was from the continent and thus OCS mole fractions may vary according to the degree to which the different air masses were exposed to biosphere OCS sinks over land (see animation available at https://ddd.uab.cat/record/232963). Another potential explanation could be that changes in the boundary layer depth could cause Sutro tower to sometimes sample in the continental boundary layer and some other times in the free troposphere.

The mixing ratios of OCS at the Sandia location were on average 40 ppt lower than at the Sutro tower location, which may reflect the strong OCS plant sink (Campbell et al. 2008) existing between the two towers. Typically OCS concentrations peaked at midday/early afternoon and were lowest at midnight. This variability is similar with other studies of the continental boundary layer (Berkelhammer et al., 2014), which show that OCS mixing ratios are highest in the middle of the day because of deep boundary layer depths, causing increased mixing with the free troposphere, which has a greater effect on the diurnal variation in the mixing ratio than the capture of OCS by photosynthesis. This mixing effect in the middle of the day is also true for CO$_2$, as can be seen by the lower mixing ratios at noon in Figure 5a.
3.3. Simulated OCS

Simulated OCS concentrations were compared to observed concentrations at the Sandia location, as shown in Figure 6. After March 20, both the observed and simulated mixing ratios exhibited lower day-to-day variability than the earlier period. The earlier part of the period had a synoptic system causing irregular flow, but after March 20 wind patterns returned to a typical summer regime for this region of northwesterly flow. The observations reflect this better than the simulations, which are much more consistent due to the fixed lateral boundary conditions used in the simulations, limiting the capability of our model for interpreting...
this trend. An animation of OCS fluxes (micromole/m²/s) and drawdown (ppt) is available at https://ddd.uab.cat/record/232963.

Figure 6 shows that the simulation result includes a significant bias at night, which may be due to typical deficiencies in modeling the PBL depth during this time. This bias is most apparent prior to March 21, at which point there was a shift from continental to marine air (please refer to Figure S4 in the supporting information for a graphical representation of the bias). After March 20, observed mixing ratios were higher, and the modeled mixing ratios enter a more standard diurnal pattern (hitting 500 ppt every night and dropping around noon every day). This latter pattern is due to the fact that after March 20 there are strong westerly winds and a consistent marine influence, as opposed to the continental influence occurring before March 20, which resulted in less stable patterns. Figure 6 also shows that the model underpredicts measured OCS mole fractions after March 20 indicating an overestimation of OCS uptake by urban vegetation by the STEM model or underestimated OCS fluxes from the SiB model. Wind direction is decisive in the interpretation of OCS values in regions with heterogeneously dispersed sources and sinks; in a recent study, (Belviso et al., 2016) described nighttime transport of anthropogenic sources of OCS related to changes in winds directions.

Modeled and observed average daily OCS mixing ratios at the Sandia location for the times 12:00 to 15:00 are shown in Figure 7a. We concentrate the study on this period because it is when the biosphere sink is expected to be the most significant. Additionally, the PBL height as simulated by WRF is lower than the height

Figure 7. (a) 3-Hour average mixing ratios of carbonyl sulfide observed (blue) and modeled (red) at the Sandia location. The averages are calculated between 12:00 and 15:00 for March 9th-March 24th at local time PST; (b) the observed mole fraction enhancement (blue) is calculated as the difference between the mixing ratios observed at the Sandia tower and Sutro tower at 14:00 local time PST. The modeled enhancement (red) calculated as the difference between the mixing ratios of the two towers averaged over a three-hour period in the afternoon (12:00, 13:00, 14:00, and 15:00), local time PST.
derived from radiosonde measurements from NOAA/ESRL Radiosonde Database for Oakland (NOAA/ESRL, 2015) area and the Heffter (1980) method during evening and early hours, while more agreement was found at midday—a known limitation in WRF simulations (Jia & Zhang, 2020). (See Supporting Information S2 for PBL simulations, calculations, and method description). The big discrepancy between modeled and observed OCS values at maximum photosynthetic rates suggest that the model is not well suited for determining GPP. The agreement between the modeled and observed enhancements improves considerably toward the end of the study period after the synoptic system has passed, as shown in Figure 7b, where the enhancement of OCS concentrations is calculated as the difference between the concentrations at Sandia minus the concentration at Sutro for the same hour period.

3.4. Simulated CO$_2$

Figure 8 compares modeled (initialized with Sutro tower data) and observed CO$_2$ at the Sandia tower location. The modeled CO$_2$ includes NEE and anthropogenic emissions of CO$_2$. During the continental influence on air sampled at Sandia during the earlier period, there is a large bias between 10 p.m. and 7 a.m., in which observed CO$_2$ mole fractions were higher than modeled CO$_2$, ranging from 20 to 72 ppm. The model does a better job during midday (11 a.m.–3 p.m.), when the bias is reduced to 0–15 ppm. During the more typical marine influence after March 20th, the bias is improved overall, but the model performs worst at midday (11 a.m.–3 p.m., bias ranges from 15 to 45 ppm). There are several potential reasons for this discrepancy. One possible answer is that the biosphere uptake signal is being overestimated by the model; in fact we can observe during the whole period that modeled CO$_2$ is lower in the model than in the observed values. The anthropogenic signal (represented by orange triangles in Figure 8) seems to be underestimated by the model, and it is more notable during nighttime when CO$_2$ values are much greater than observed values at midday. This explanation is difficult to confirm because there could be model deficiencies in simulating the boundary layer depth that could also be affecting modeled mixing ratios (Hu et al., 2010). In Figure 9, we plot concentrations of CO$_2$ only for the time period 12–3 p.m., which has been documented as being the time period with best model performance in terms of boundary layer height; this result is consistent with other studies that show a better performance during daytime for modeling boundary layer and wind (Lee et al., 2011).

Figure 9 shows 3-hour averages (between 12 and 3 p.m.) of CO$_2$ mole fractions from simulations that include different CO$_2$ fluxes (NEE CO$_2$ and anthropogenic CO$_2$ from Vulcan), and they are compared to observed CO$_2$ concentrations. A comparison of modeled values in Figure 9 show that the biosphere signal is an important contributor to simulated CO$_2$ mole fractions. This apparent influence could be attributed to two hypotheses: (a) That the biotic signal is overestimated in the model resulting in higher capture of CO2 than observed and/or (b) the anthropogenic flux used as input to the model is perhaps underestimated, also resulting lower values of CO$_2$ than the observed. Normally, the difference between modeled NEE CO$_2$ and
observed CO$_2$ is attributed to anthropogenic CO$_2$ (McKain et al., 2012; Wei et al., 2020). However, Figure 9 shows that there is a significant difference between the modeled anthropogenic CO$_2$ concentrations (orange bars) and the sum of NEE and anthropogenic CO$_2$ concentrations (gray bars). This difference, which ranges between 9 and 29 ppm, is in the same order of magnitude as the overall model bias. On average, the model bias is 35 ppm. The average contribution from the biosphere is 17 ppm. These results show that the contribution of the urban biosphere is significant, and it is important to quantify it with a certain degree of confidence. The use of OCS as biosphere tracer should improve our estimations of NEE by improving our capacity to fine-tune the GPP signal.

Modeled mixing ratios of OCS and CO$_2$ present the same problem: They both have bias at night due to deficiencies in modeling the PBL. OCS is biased in the opposite direction to CO$_2$ because OCS is a small sink at night (Belviso et al., 2020; Maseyk et al., 2014), while CO$_2$ is a source at night due to respiration. Figure 10 shows the correlation between OCS model error and CO$_2$ model mismatch. This correlation is a strong indicator that model mismatch is due to transport, not due to model input of sinks and sources.

Figure 9. 3-hour averages (between 12:00 and 15:00) of observed, anthropogenic (modeled Vulcan emissions), and modeled total (net ecosystem exchange plus anthropogenic) CO$_2$ concentrations.

Figure 10. Model mismatch between CO$_2$ and carbonyl sulfide from Sulfur Transport and dEposition Model. The vertical lines represent midnight local time PST.
4. Conclusions

This study evaluates and describes the present limitations of using OCS as a tracer of the urban biosphere to determine the contribution of the urban ecosystem to CO$_2$ uptake. We used the STEM to simulate the OCS concentrations and the CASA ecosystem model to simulate global CO$_2$ fluxes over the Bay Area of San Francisco during March 2015. Our results show that the STEM model works better under stable marine influence, and that the BL height and entrainment are driving the diurnal changes in CO$_2$ at the downwind Sandia site. However, the model needs a better characterization of transport and BL variability and improvements in the estimation of GPP for characterizing the urban biosphere signal. A synoptic event in the middle of our run clearly shows a shift in the background concentrations, evidencing a shift from continental to marine influence. The transport model does a better job simulating OCS concentrations during the marine influence, when Sutro tower can be assumed to be representative of background concentrations. BL depth is more important in driving diurnal variability in mixing ratios than CO$_2$ emissions in the urban area. This conclusion is supported by the fact that we found more CO$_2$ emissions but lower mixing ratios in the afternoon than in the evening, due to the reduction in the the BL depth at night.

The chemical transport model STEM is not correctly representing the diurnal cycle we see with observations reflecting deficiencies in its transport feature (how wind and boundary layer depth affect fluxes) or in the quantification of fluxes affecting CO$_2$ and COS. Furthermore, we identify the need to improve the model in terms of daytime mixing ratios of biogenic emissions to be able to refine GPP predictions. However, we also show that the “enhancement” of urban OCS concentrations (compared to the background) agrees with modeled enhancement when the observed background OCS values are stable during the period of marine influence.

Data Availability Statement

The data on which this article is based are publically available at the following web sites: The reanalysis meteorological data used for the WRF runs are available in https://www.ncdc.noaa.gov/data-access/model-data/model-datasets/ climate-forecast-system-version2-cfsv2; temperature at 2 m and wind speed and direction at 10 meters from the Lawrence Livermore National Laboratory weather station are available at https://weather.llnl.gov; hourly OCS flux data for the STEM runs were extracted from the OCS hourly inventory of the biospheric surface-flux at 1.25° × 1° resolution (Baker et al., 2009; Berry et al., 2013); NDVI data used in this study was downloaded from http://modis-land.gsfc.nasa.gov/vi.html; Global CO$_2$ fluxes for GPP and RESP are available at https://nacp-files.nacarbon.org/nacp-kawa-01/; U.S. high resolution inventory for the fossil-fuel derived anthropogenic CO$_2$ emissions were downloaded from the Vulcan inventory https://daacornl.gov/NACP/guides/Vulcan_V3_Annual_Emissions.html; Radiosonde measurements were obtained from https://ruc.noaa.gov/raobs/.

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