Greenhouse gas column observations from a portable spectrometer in Uganda

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Abstract. The extensive terrestrial ecosystems of tropical Africa are a significant store of carbon, and play a key but uncertain role in the atmospheric budgets of carbon dioxide and methane. As ground-based observations in the tropics are scarce compared with other parts of the world, recent studies have instead made use of satellite observations assimilated into atmospheric chemistry and transport models to conclude that methane emissions from this geographical region have increased since 2010 as a result of increased wetland extent, accounting for up to a third of global methane growth, and that the tropical Africa region dominates net carbon emission across the tropics. These studies rely critically on the accuracy of satellite datasets such as those from OCO-2, GOSAT, and Sentinel-5P TROPOMI, along with results from atmospheric transport models, over a geographical region where there are few independent data to test the robustness of published results.

In this paper we present the first ground-based observations of greenhouse gas column concentrations over East Africa, obtained using a portable Bruker EM27/SUN FTIR spectrometer during a deployment covering the first few months of 2020 in Jinja, Uganda. We operated the instrument near-autonomously by way of an automated weatherproof enclosure, and observed total atmospheric column concentrations of the greenhouse gases carbon dioxide and methane, as well as carbon monoxide, a useful proxy for emissions from incomplete combustion processes in the region. We discuss the performance of the combined enclosure and spectrometer system that we deployed in Jinja to obtain this data, and show comparisons of our ground-based observations with satellite datasets from OCO-2 and OCO-3 for carbon dioxide, and TROPOMI for methane and carbon monoxide, whilst also comparing our results with concentration data from the GEOS-Chem and CAMS atmospheric inversions that provide a means of increasing spatial and temporal coverage where satellite data are not available. For our measurement period, we find mean differences in $X_{\rm CO_2}$ between OCO-2 and the EM27/SUN of $-0.29\,\%$, and between OCO-3 and the EM27/SUN of $-0.28\,\%$. In the case of TROPOMI, the mean difference in $X_{\rm CH_4}$ that we find between TROPOMI and the EM27/SUN is $-0.44\,\%$, whilst for $X_{\rm CO}$ the mean difference is $-5.65\,\%$. In each of these cases, the mean difference observed between the

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satellite and ground-based column concentrations is either close to or within the precision and accuracy requirements for the respective missions. With regards to the model and reanalysis comparisons with the EM27/SUN column concentrations, we see mean differences from the EM27/SUN of a global GEOS-Chem inversion for $X_{\rm CO_2}$ of -0.08%, a regional high-resolution GEOS-Chem inversion for $X_{\rm CH_4}$ of -0.22%, and the CAMS global reanalysis $X_{\rm CO}$ of -9.79%. Our results demonstrate the value of ground-based observations of total column concentrations, and show that the combined EM27/SUN and enclosure system employed would be suitable for acquisition of the longer-term observations needed to rigorously evaluate satellite observations, and model and reanalysis calculations over tropical Africa.

1 Introduction

Gaps in our understanding of the global carbon cycle add uncertainty to our predictions of future climate change, including how the future climate will respond to different carbon emissions scenarios (IPCC, 2021; Friedlingstein et al., 2022). One part of the carbon cycle which still requires further investigation is that of carbon fluxes from terrestrial tropical ecosystems, which store large quantities of carbon in vegetation and soil whilst being sensitive to changes in the climate (Pan et al., 2011; Crowther et al., 2015). Carbon dioxide is released by these ecosystems into the atmosphere through a combination of respiration and fire, and is removed from the atmosphere by photosynthesis and subsequent conversion into plant biomass. The tropics are also home to extensive areas of wetlands, which are the most significant natural source of methane in the atmosphere via the decomposition of organic matter in anaerobic conditions (Kayranli et al., 2010; Mitsch et al., 2013). Further microbial sources of methane in tropical regions include agricultural practices, particularly the farming of ruminants, and waste disposal. Looking at Africa in particular, an additional factor having an increasingly significant impact on the tropical African carbon cycle is the recent increase in population in many African countries resulting in increasing demand for energy (Ayompe et al., 2021), as reflected in the rapid projected growth of cities such as Kampala (Uganda), Nairobi (Kenya) and Kinshasa (Democratic Republic of Congo). This combination of natural and anthropogenic fluxes that contribute to the atmospheric carbon budget in tropical Africa is challenging to accurately represent in climate and atmospheric chemistry models, so we need to make use of atmospheric composition measurements to evaluate our understanding.

Compared with other parts of the world, however, ground-based measurements of atmospheric composition are scarce in tropical Africa, placing an upper limit on how well we can understand the carbon cycle in this region (López-Ballesteros et al., 2018; Nickless et al., 2020). This measurement gap is partially addressed by satellites such as the JAXA GOSAT (Kuze et al., 2009), NASA OCO-2 (Eldering et al., 2017) and OCO-3 (Eldering et al., 2019), and Copernicus Sentinel-5P (Veefkind et al., 2012) missions, although the measurement technique employed usually requires cloud-free and low aerosol conditions to retrieve molecular concentrations from the observed radiance spectra, resulting in relatively poor coverage over the tropics where conditions are often cloudy. The satellite data that are obtained can however be used as an input for atmospheric chemistry models, which use prior estimates of surface fluxes and meteorological fields to calculate a most-likely state for the atmosphere constrained by the observations available (e.g. Basu et al. (2013); Deng et al. (2014); Feng et al. (2017); Chevallier et al. (2019); Crowell et al. (2019); Chen et al. (2022); Peiro et al. (2022)). The atmospheric chemistry models can then be

used in an inversion framework to produce *a posteriori* estimates of emissions where satellite data are not available, since increased greenhouse gas concentrations remain in the atmospheric column for some time downwind of where they are originally emitted. In addition, these models (such as GEOS-Chem: Turner et al. (2015); Feng et al. (2017); Lunt et al. (2019)) are a useful means for estimating atmospheric concentrations where observations are not available, and have underpinned a number of studies that address the tropical African carbon cycle (Palmer et al., 2019; Lunt et al., 2019, 2021; Pandey et al., 2021; Qu et al., 2022; Feng et al., 2022, 2023; Drinkwater et al., 2023). It is therefore important to validate model output with independent observations, to confirm how well the models represent the atmosphere and add weight to the conclusions of the studies which use them.

In this study, we describe observations of the total column concentrations of greenhouse gases in Uganda in the first few months of 2020, obtained using a portable spectrometer with a built-in solar tracker. We used an automated enclosure to provide a weatherproof environment for the spectrometer, and to allow us to operate the spectrometer remotely. This setup allowed us to produce for the first time a dataset of ground-based total column concentrations of carbon dioxide, methane, and carbon monoxide for a tropical East African location. In Section 2 we outline the measurement site and describe the instrument, enclosure, and retrieval algorithm used to obtain the dataset. Section 3 covers the satellite and model datasets that we then compare with our ground-based observations in Section 4. We finally conclude and consider the implications of this study in Section 5.

70 2 The measurement site at NaFIRRI in Jinja, Uganda

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For this study, we established our measurement site at the headquarters of the Ugandan National Fisheries Resources Research Institute (NaFIRRI) in Jinja (0.4165°N, 33.2070°E, 1157 metres above sea level). Jinja is located on the northern shore of Lake Victoria, approximately 70 km to the east of Kampala, Uganda's capital city with a population of approximately 3.5 million people across its wider urban area. The source of the White Nile is in Jinja, from which it flows northwards out of Lake Victoria, through Lakes Kyoga and Albert, and onwards into South Sudan. The Nile feeds the neighbouring wetlands, which are amongst the main sources of methane emissions in the East Africa through the anaerobic decomposition of organic matter. There is a strong link between CH₄ emissions and water table depth in tropical regions, such that anomalies in precipitation can lead to wetland CH₄ emissions anomalies (Bloom et al., 2010). The hydrological flow from increased precipitation over Lake Victoria to higher water table depth in Ugandan and South Sudanese wetlands, as a result of increased volumes of water transported along the Nile, is covered in more detail by Lunt et al. (2019). Precipitation in Uganda is driven by the annual northsouth movement of the inter-tropical convergence zone, resulting in two main wet seasons during the year: these are known as the 'long rains' which occur from March to May, and the 'short rains' occurring from October to December (Herrmann and Mohr, 2011). The amount of precipitation over East Africa during these wet seasons is in turn partly influenced by ocean temperatures in the Indian Ocean (Palmer et al., 2023), where an unusually high contrast in temperatures (greater than 0.4° C) between the warmer western Indian Ocean and cooler Eastern Indian Ocean (defined quantitatively as the Indian Ocean Dipole, Saji et al. (1999)) in 2019 resulted in one of the wettest short rains seasons on record (Wainwright et al., 2021).

In the wider region beyond Uganda, there are a number of environmental factors which can potentially affect the column of air that we measure at Jinja. To the north, the Sudd wetlands in South Sudan represent a significant natural source of CH₄ as discussed and investigated by Lunt et al. (2019) and Lunt et al. (2021). To the west, atmospheric CO₂ signals are dominated by the biospheric influence of the Congo rainforest (Palmer et al., 2019). This part of the world is also subject to a high frequency of biomass burning events, evidence of which can be seen in TROPOMI observations of carbon monoxide (a product of incomplete combustion, see Section 3.2).

To help understand which sources influence the composition of the air columns we observe, we can use a Lagrangian dispersion model to calculate the history of the air masses arriving over Jinja during the measurement period (Fleming et al., 2012; Panagi et al., 2020). We use the UK Met Office's (UKMO) Numerical Atmospheric Modelling Environment (NAME) to perform this task. NAME, along with other Lagrangian dispersion models, works by releasing a large number of inert particles from a specific location in the atmosphere and then tracking their pathways backwards in time using meteorological model data (which in the NAME modelling framework comes from the UKMO Unified Model). We then count the number of released particles that pass within 100 m of the surface over each spatial grid point, to determine where and to what extent the back trajectories are influenced by surface emissions from that location within a certain period of time. To account for our measurements being sensitive to the whole atmospheric column, we perform the particle releases from heights throughout the vertical grid of the model domain, and weight the contributions from each release height according to the pressure weighting function of the EM27/SUN observations. We perform the calculation for each day of the measurement period (performing the particle release at 1030 UTC each day, equivalent to 1330 local time – the time at which TROPOMI and OCO-2 pass overhead - and tracking it back in time for one day and five days) to obtain a daily column footprint. Figure 1 shows the mean daily footprint for the whole measurement period (panels A and B showing the results from back trajectories going back one and five days, respectively), giving us an estimate of where the surface has influenced the measured column. Although the highest contribution arrives via a region directly to the south, coinciding with Lake Victoria, the footprint of influence also covers regions to the north and east, reaching as far as South Sudan and Kenya respectively, where emissions from wetlands and agriculture can potentially have an impact on the observed atmospheric column.

2.1 The EM27/SUN portable spectrometer

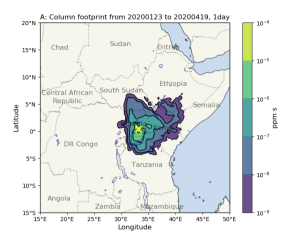
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Ground-based remote sensing of the atmospheric column has proved to be an invaluable tool in the validation of atmospheric composition data from satellite observations. The global network of Bruker 125HR spectrometers that form TCCON (Total Carbon Column Observing Network, Wunch et al. (2011)), for example, is now routinely used in the validation of greenhouse gas column observations from GOSAT, OCO-2, Sentinel-5P and others, allowing those working on the retrieval algorithms to identify, and correct for, systematic biases in their data (Inoue et al., 2016; Wunch et al., 2017; Sha et al., 2021). The standard configuration for a TCCON site is, however, both expensive and logistically challenging to set up and maintain. As a result, there are certain regions around the world – South America, Africa, and Central/Southern Asia – which do not currently have the resources and infrastructure in place to host TCCON sites, leaving significant gaps in the validation of GHG column data products, often in geographical areas of great scientific interest (e.g. the Amazon rainforest, sub-Saharan Africa).



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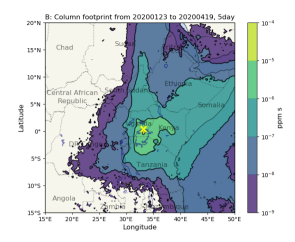


Figure 1. Mean column footprint of 1030 UTC (1330 local time) column observations, calculated from 1 day (panel A) and 5 day (panel B) NAME back trajectories for each day within the measurement period (23rd January to 19th April 2020). The yellow cross indicates the location of our EM27/SUN instrument in Jinja. The colour scale indicates the calculated contribution of the surface in that location to the observed atmospheric column, in ppm, integrated over the duration of the back trajectory.

The Bruker EM27/SUN FTIR (Fourier Transform InfraRed) spectrometer concept (Gisi et al., 2012) was developed at the Karlsruhe Institute of Technology (KIT), in part to address this problem. It comprises a portable Fourier transform spectrometer with built-in solar tracker, which trades off a reduced spectral resolution compared with the Bruker 125HR used at TCCON sites in favour of being less expensive, and much easier to transport to and operate in different locations. A number of previous studies have demonstrated comparable stability and precision when operated side-by-side with the higher resolution 125HR (Frey et al., 2015; Hedelius et al., 2016; Hase et al., 2016; Sha et al., 2020; Alberti et al., 2022). At the time of writing, over 150 EM27/SUNs have been purchased by research groups around the world, and operated in a variety of locations. Prior to shipment, the instruments are first calibrated at KIT to obtain the instrument line shape parameters, and are operated side-by-side with a reference instrument in Karlsruhe to derive instrument-specific scaling factors, which can be applied by the user to their retrieved GHG column data to maintain consistency between all EM27/SUN data sets, regardless of who operates the instrument and where. This work is done under the COCCON project (COllaborative Carbon Column Observing Network: Frey et al. (2019); Alberti et al. (2022)), which also develops and maintains the PROFFAST retrieval software used to calculate atmospheric column concentrations from the measured interferograms. Information on the spectral ranges used in the retrievals, along with some example spectra and vertical column sensitivities, can be found in Appendix A.

As well as being used for validation studies in various locations (Jacobs et al., 2020; Tu et al., 2020; Frey et al., 2021), the portability and relatively low cost of the EM27/SUN has lead to a variety of other scientific applications. In the city of Munich, Germany, a permanent network of five EM27/SUNs has been established to observe the city's carbon emissions using the differential column observation method (Chen et al., 2016; Dietrich et al., 2021), and provide a means of validating

spatial gradients in OCO-2 target mode observations of $X_{\rm CO_2}$ (Rißmann et al., 2022). Similar city-focused studies using EM27/SUNs have taken place in Berlin (Hase et al., 2015), St. Petersburg (Makarova et al., 2021), Beijing (Che et al., 2022; Zhou et al., 2022), and Indianapolis (Jones et al., 2021) amongst others. Further studies have taken advantage of the instrument's portability in another way, adapting the instrument with a specially designed solar tracker for operation on board a cargo ship to provide a unique opportunity for validation of satellite and model data over the ocean (Klappenbach et al., 2015; Knapp et al., 2021). Some of the studies listed here make use of various designs of weatherproof enclosure to operate the EM27/SUN more effectively; the enclosure we use here, developed at TU Munich, is described in Section 2.2.

2.2 An automated enclosure for the EM27/SUN

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The EM27/SUN, whilst very useful for greenhouse gas column observations, is not suitable for unattended operations 'out of the box'. Firstly the instrument itself is not weatherproof, so the user has to keep a close eye on the weather forecast when deciding whether to set up for a day of measurements, and be in close attendance to move it indoors in case of rain. In addition, the user has to manually start the solar tracker and then the spectrometer at the beginning of each day, before powering down and moving it indoors once the day's observations are complete. This labour intensive mode of operations works well for short term measurement campaigns, but is less suitable if the goal is to obtain long term observations in a single location.

To make the EM27/SUN suitable for use on longer term deployments, the Environmental Sensing and Modelling Group at the Technical University of Munich have developed an automated enclosure (Heinle and Chen, 2018; Dietrich et al., 2021) which provides weatherproofing, environmental control, and automation of the observations. The main components of the TU Munich enclosure are labelled in Figure 2, and are described in detail by Dietrich et al. (2021).

A modified Zarges K470 aluminium box is used for the main body of the enclosure system. On top of the housing, a rotating cover closes to protect the contents of the enclosure system when rain is detected by an optical rain sensor, and overnight when no measurements are taking place. When conditions are dry during the day time, the cover rotates to track the azimuth angle of the solar tracker and allow sunlight into the system.

The user controls and monitors the enclosure system by remotely accessing the enclosure computer, which also controls the EM27/SUN spectrometer and solar tracker, and stores the measured interferograms. The automated features of the system are controlled by a programmable logic controller (PLC), ensuring that critical safety features protecting the system (detection of rain or power failure, control of the cover motor, temperature control) are not dependent on the enclosure computer. An additional challenge posed by the location of these measurements very close to the equator is that of very high solar zenith angles, which at times are beyond the normal operating range of both the solar tracker and the protective cover. A pair of car-jacks attached to the side of the enclosure (see Figure 2) allow the entire enclosure system to be tilted, such that the sun can be tracked throughout the middle of the day.

Control and automation of the enclosure system is achieved by two software programs, both developed in-house at TU Munich (Dietrich et al., 2021). The first of these (Enclosure Control, or ECon: see Heinle and Chen (2018)) controls the enclosure itself – moving the rotating cover into the correct position, maintaining internal temperature using the thermoelectrical coolers, monitoring the rain sensor data and the UPS, and powering the spectrometer. ECon also checks that the



Figure 2. Left: internal view of the EM27/SUN spectrometer and solar tracker housed within the TU Munich enclosure system; Right: the enclosure system in operation at NaFIRRI, with the two car-jacks used to tilt the system to enable tracking of the sun at very high solar zenith angles (see text in Section 2.2). The bricks attached to the car-jacks anchor the enclosure down in case of strong winds.

Ethernet connections linking the different components of the enclosure system are working correctly, and performs automatic restarts or specific components if a malfunction is detected. Alongside ECon, control of the spectrometer and solar tracker are automated using a Python program called Pyra. Pyra effectively acts as a wrapper for the software provided by Bruker that controls the spectrometer and the solar tracker (OPUS and CamTracker, respectively), providing the means to start, stop, and control them automatically. For these measurements we used Pyra in a semi-automated mode which started and stopped the observations when the solar zenith angle passed a minimum threshold; a more detailed description of Pyra can be found in Appendix A of Dietrich et al. (2021), whilst the latest version is described in full by Aigner et al. (2023).

2.3 Total column concentrations over Jinja from the EM27/SUN and automated enclosure system

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The data processing method we use, taking us from the raw interferograms measured by the EM27/SUN spectrometer to the column averaged greenhouse gas abundances over Jinja presented in this paper, is described in more detail in Section 2.2 of Frey et al. (2021). The method comprises two parts, both written in FORTRAN: PREPROCESS, which performs Fast Fourier Transforms on the interferograms (that are first corrected for intensity fluctuations and apodised) to obtain solar absorption spectra; and PROFFAST, which then retrieves the column averaged greenhouse gas abundances from the solar absorption spectra. Several quality filters, summarised in Table 1 of Frey et al. (2021), are applied to each interferogram by the PREPROCESS routine. The *a priori* profiles we use for trace gas concentrations, pressure, and temperature, are those generated for use in the TCCON GGG2014 data version (Wunch et al., 2015). The profiles of pressure, temperature, geopotential height,

and water vapour come from the NCEP/NCAR reanalysis (National Centres for Environmental Prediction/National Centre for Atmospheric Research, Kalnay et al. (1996)), and are then used to generate the trace gas profiles by a set of empirical functions optimised to fit a range of *in situ* profile measurements as described in Wunch et al. (2011).

We then apply the PROFFAST retrieval algorithm to those spectra whose interferograms have passed the quality filters applied during the PREPROCESS stage. PROFFAST is a non-linear least squares algorithm which scales *a priori* trace gas profiles to fit forward modelled atmospheric spectra to the measured spectra, and then calculates the retrieved total column abundances from the scaled profiles. These are finally converted into column-averaged dry-air mole fractions X_{gas} , given by

$$X_{\rm gas} = \frac{\rm VC_{\rm gas}}{\rm VC_{\rm O_2}} \times 0.2095,\tag{1}$$

where VC_{gas} is the retrieved total column abundance for that gas. Taking the ratio of the total column abundances has the benefit of at least partially cancelling out any spectroscopic errors which affect both VCs in a similar way (Wunch et al., 2010), whilst also reducing the dependence on the measured ground pressure (Frey et al., 2021).

To monitor the stability of the spectrometer, we use the column-averaged amount of dry air (X_{air}) . This is the ratio of the total column abundance of dry air calculated from the retrieved total column abundance of oxygen, VC_{O_2} , to the total column abundance of dry air calculated from the measured surface pressure P_S , and is given by

$$X_{\text{air}} = \frac{g}{P_{\text{S}}} \cdot \left(\frac{\text{VC}_{\text{O}_2} \cdot \overline{\mu}}{0.2095} + \text{VC}_{\text{H}_2\text{O}} \cdot \mu_{\text{H}_2\text{O}} \right),$$
 (2)

where the molecular masses of dry air and water vapour are given by $\overline{\mu}$ and $\mu_{\rm H_2O}$ respectively, g is the column-averaged gravitational acceleration, and ${\rm VC_{H_2O}}$ is the retrieved water vapour total column (this correction is required to allow for the measured surface pressure including the whole air column, whereas it is the dry air column that we retrieve using the oxygen absorption band). As long as the spectrometer is working nominally, $X_{\rm air}$ should remain close to 1.0 and stable over time. We therefore use $X_{\rm air}$ as a final quality filter on the retrieved column data, by removing any data points where the difference from the daily median value of $X_{\rm air}$ is greater than 0.002, and then removing any further data points which deviate from the rolling hourly mean $X_{\rm air}$ by more than 0.0005.

A final step in the data processing is to apply calibration factors to the retrieved column concentrations, which bring the results into line with the rest of the EM27/SUNs involved in COCCON (as discussed earlier in this Section, a full list of calibration factors is given in Table 6 of Frey et al. (2019)). For the spectrometer used here (serial number 059), the calibration factors with respect to the reference EM27/SUN operated by Karlsruhe Institute of Technology (serial number 037) are 0.9998, 0.9991 and 1.0019 for $X_{\rm CO_2}$, $X_{\rm CH_4}$ and O_2 , respectively. The column concentrations retrieved by following the procedure described above are shown in Figure 3, along with the number of quality-controlled soundings obtained on each measurement date. The daily count of measurements leading to a valid retrieval is determined by a combination of weather conditions (cloudy vs. cloud-free) and the availability of mains power on the NaFIRRI site, during the times of day when the Sun is at least 20° above the horizon.

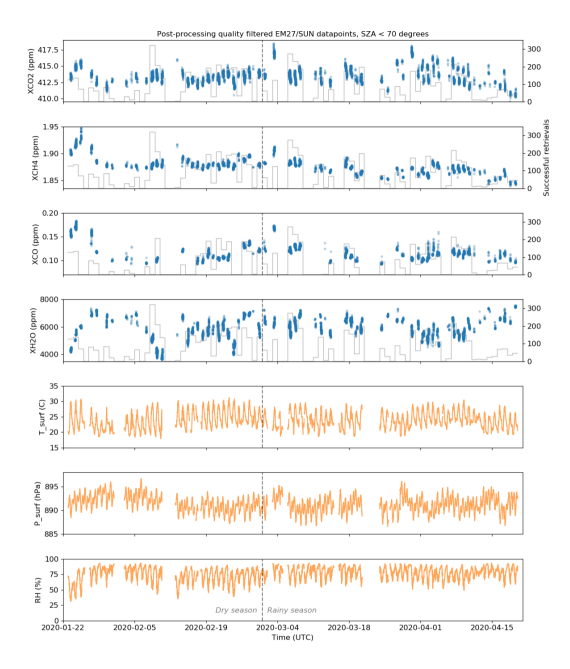


Figure 3. From top to bottom: column concentrations of CO_2 , CH_4 , CO, and H_2O retrieved from the EM27/SUN measurements using the PROFFAST algorithm as described in Section 2.3; surface air temperature; surface pressure; and relative humidity. The solid line in the upper four panels indicates the number of successful retrievals on each day, for each gas. The vertical dashed line marks the onset of the 'long rains' as described in Section 2.

220 3 Satellite and model datasets used in this study

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In this section, we introduce the satellite and model datasets that we compare with our Jinja EM27/SUN column concentration data.

3.1 Orbiting Carbon Observatory (OCO-2 and OCO-3) $X_{\rm CO_2}$ retrievals

The Orbiting Carbon Observatory-2 (OCO-2) was launched in 2014, and was specifically designed by NASA to have the precision required to detect the changes in $X_{\rm CO_2}$ that correspond to surface emissions and uptake of ${\rm CO_2}$, on a regional scale with global coverage (Eldering et al., 2017). The sole OCO-2 payload comprises a three-band grating spectrometer, which measures the radiance spectra of sunlight reflected back into space by the Earth's surface. Of the three spectral bands, two coincide with carbon dioxide absorption features (the so-called 'weak' and 'strong' ${\rm CO_2}$ bands, centred at wavelengths of 1.6 and 2.0 μm respectively), whilst the third band at 0.76 μm is used to measure absorption by molecular oxygen. The instrument samples eight spatial footprints across-track, which are each nominally 1.25 km in width at the surface. Along-track, each footprint is around 2.4 km in length owing to the distance travelled by the satellite during the instrument's 0.33 s integration time. The orbit track and the narrow swath width (approximately 10 km wide) mean that the same ground location is resampled once every 16 days. A full-physics retrieval algorithm based on an optimal estimation technique is used to retrieve $X_{{\rm CO}_2}$ from the OCO-2 measured spectra (O'Dell et al., 2012, 2018), taking into account multiple scattering and polarisation effects. The retrieved column concentrations are validated against the TCCON ground-based network of Bruker 125HR spectrometers (Wunch et al., 2017). For this study, we use Version 10r of the OCO-2 data (Taylor et al., 2023) – the spatially gridded mean ${\rm CO}_2$ column concentrations from this dataset observed over East Africa during our measurement period are shown in Figure 4A.

In 2019 NASA also integrated the flight spare for OCO-2, under the name Orbiting Carbon Observatory-3 (OCO-3, Eldering et al. (2019); Taylor et al. (2020)), onto the International Space Station (ISS). The low-inclination orbit occupied by the ISS introduces significant differences to the sampling pattern, with the main implication being that overpasses of a particular location do not take place at the same local time each day. The overpass time instead shifts about 20 minutes earlier from one day to the next, such that all times of day are eventually observed. In contrast to the observations obtained from the sunsynchronous orbit followed by OCO-2, this means that OCO-3 can provide information on how X_{CO_2} varies with time of day. In addition, the Pointing Mirror Assembly (PMA) allows pointing towards the ocean glint spot to maximise the observed signal over water (for OCO-2 the whole spacecraft is rotated to achieve a similar goal), or towards stationary ground targets such as validation sites. Uniquely to OCO-3, snapshot area maps (SAMs) can also be acquired. These involve sweeping the PMA fore-optics back and forth across an area approximately 85 km \times 85 km in size, effectively producing spatially resolved 2D images of X_{CO_2} over areas of interest. The same full-physics retrieval algorithm is used on OCO-3 measurements as for OCO-2 to obtain the column concentrations of CO_2 . Here we use Version 10.4r of the OCO-3 data (Taylor et al., 2023). The gridded mean X_{CO_2} over East Africa for our whole measurement period is shown in Figure 4B, which clearly illustrates the different spatial sampling pattern employed by OCO-3 compared with OCO-2 (Figure 4A). Note that Jinja is included on the

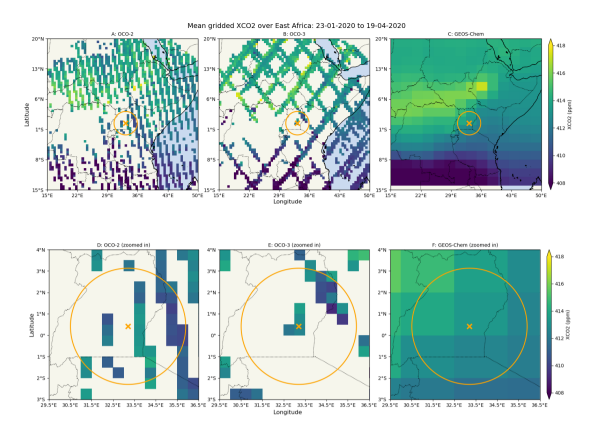


Figure 4. $X_{\rm CO_2}$ over East Africa retrieved from OCO-2 (panel A) and OCO-3 (panel B) observations, averaged over the EM27/SUN measurement period ($23^{\rm rd}$ January to $19^{\rm th}$ April 2020) and spatially binned into a 1x1 degree grid. The orange cross shows the location of the measurement site in Jinja, and the circle indicates the co-location criteria ($300 \, \rm km$ radius) used in the comparisons described in Section 4.1. Panel C shows the $X_{\rm CO_2}$ output from the global GEOS-Chem inversion described in Section 3.3. Panels D, E and F show the same data as panels A, B and C respectively, zoomed in on the co-location region.

list of targets for the SAM mode, such that the OCO-3 soundings taken as the ISS passes over Uganda tend to be concentrated within a short distance of our measurement site.

3.2 TROPOMI X_{CH_4} and X_{CO} retrievals

The Copernicus Sentinel-5 Precursor (S5P) mission was launched in October 2017 to measure atmospheric composition, specifically air quality and climate change indicators, with daily global coverage and moderately high (up to $5.5 \text{ km} \times 3.5 \text{ km}$ at nadir) spatial resolution. The sole payload of the S5P mission is the TROPOspheric Monitoring Instrument (TROPOMI, Veefkind et al. (2012)), a grating spectrometer with four spectral bands covering ultraviolet, visible, near-infrared (NIR), and shortwave infrared (SWIR) wavelength ranges, respectively. The S5P operational CH_4 retrieval algorithm uses the NIR and

SWIR spectral channels in a full-physics, optimal estimation method to derive the column-averaged dry mixing ratio of methane (X_{CH₄}) from the TROPOMI measurements (Hu et al., 2016; Hasekamp et al., 2021). The X_{CH₄} data used in this study has been processed using version 01.03.02 of the TROPOMI CH₄ processor, which has been shown to perform well within the mission requirements through comparison with ground-based observations from 28 TCCON stations (Sha et al., 2021). This version includes a surface albedo dependent *a posteriori* bias correction, based on comparisons between co-located TROPOMI and GOSAT X_{CH₄} data (Hasekamp et al., 2021). We show the gridded mean X_{CH₄} from this dataset over East Africa, averaging over our whole measurement period, in Figure 5A. More recent TROPOMI X_{CH₄} data (measurements from 1st July 2021 onwards) use version 2 of the processor, which incorporates a number of improvements including updated CH₄, CO and H₂O spectroscopic cross sections, and an updated *a posteriori* bias correction that is independent of external reference data (Lorente et al., 2021).

The TROPOMI SWIR band is also used to retrieve the total column abundances of carbon monoxide, $X_{\rm CO}$ (Landgraf et al., 2016). The retrieval uses a two-step process: firstly, a non-scattering retrieval of the total amount of ${\rm CH_4}$ is performed, and then compared with modelled methane abundances to act as a cloud filter (if the retrieved ${\rm CH_4}$ assuming no-scattering differs significantly from the model value, this indicates that the impact of scattering from high or optically thick cloud is too great to perform a useful carbon monoxide retrieval). The second step then retrieves the CO column using a profile scaling approach, along with the ${\rm H_2O}$ abundance and effective cloud parameters using the *a priori* knowledge of methane acquired during the first step. Validation of the operational TROPOMI ${\rm X_{CO}}$ against ground-based TCCON observations has demonstrated that the requirements for systematic and random uncertainties in the data are being met (Sha et al., 2021). Figure 6A shows the gridded mean ${\rm X_{CO}}$ from the operational TROPOMI product for our measurement period, over the East Africa region. The less strict requirements on accuracy and precision for ${\rm X_{CO}}$ compared with those for ${\rm X_{CH_4}}$ allows retrievals to be made over land and ocean scenes, under both clear-sky and (with the exception of high or optically thick clouds) cloudy conditions. This is reflected in the comprehensive spatial coverage shown in Figure 6A that is achieved by the ${\rm X_{CO}}$ retrieval compared with that of ${\rm X_{CH_4}}$ (as seen in Figure 5A), which requires cloud-free conditions and minimal scattering for a successful retrieval.

3.3 GEOS-Chem and CAMS concentration data

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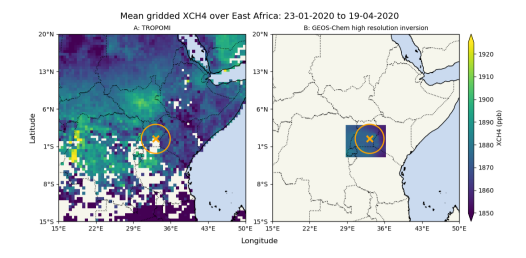
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GEOS-Chem is an atmospheric chemistry transport model that is used here to simulate the emissions, sinks, chemistry, and transport of carbon dioxide and methane (Turner et al., 2015; Feng et al., 2017; Lunt et al., 2019, 2021), and produce three-dimensional fields of their concentrations. This can provide a useful extension of satellite data in spatial regions and at times of day where the satellite data are not available. For a more detailed description of the GEOS-Chem model and the ensemble Kalman filter inverse method used, we refer the reader to the papers cited below.

For carbon dioxide, we use a global GEOS-Chem model run on a $2.0^{\circ} \times 2.5^{\circ}$ latitude-longitude grid with 47 vertical levels. We use emissions inventories for our *a priori* flux estimates, taking into account CO_2 emissions from biomass burning (van der Werf et al., 2010), fossil fuels (Oda et al., 2018), ocean fluxes (Takahashi et al., 2009) and biosphere fluxes (Olsen and Randerson, 2004). An ensemble Kalman Filter approach is then used to estimate the CO_2 fluxes, with either *in-situ* or satellite



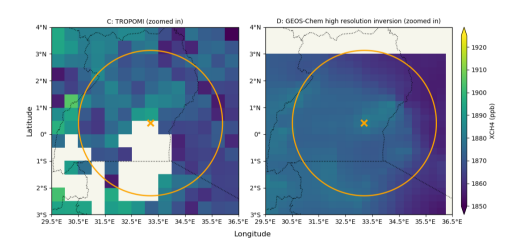
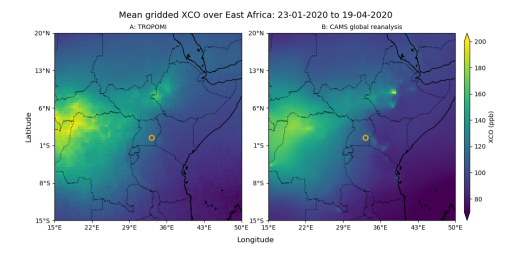


Figure 5. $X_{\rm CH_4}$ over East Africa retrieved from TROPOMI observations (Panel A, see Section 3.2) and calculated from a high resolution GEOS-Chem inversion (Panel B, see Section 3.3 for details), averaged over the EM27/SUN measurement period ($23^{\rm rd}$ January to $19^{\rm th}$ April 2020). The TROPOMI data are spatially binned into a 1x1 degree grid. The orange cross shows the location of the measurement site in Jinja, and the circles indicate the co-location criteria (300 km radius for $X_{\rm CH_4}$) used in the comparisons described in Section 4.2. Panels C and D show the same data as panels A and B respectively, zoomed in on the co-location region.

measurements of atmospheric CO_2 used as prior information on concentration (Feng et al., 2009; Palmer et al., 2019). The mean X_{CO_2} values for the measurement period calculated from the output of this global inversion are shown in Figure 4C.

In the case of methane, we run GEOS-Chem in a nested configuration at high spatial resolution $(0.25^{\circ} \times 0.3125^{\circ})$ over a latitude-longitude box covering sub-Saharan Africa $(-36.0 \text{ to } + 20.0^{\circ}\text{N}, -20.0 \text{ to } 55.0^{\circ}\text{E})$, using the setup described in detail



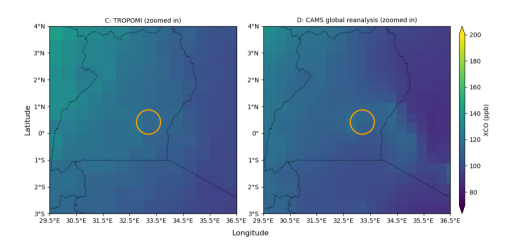


Figure 6. $X_{\rm CO}$ over East Africa retrieved from TROPOMI observations (Panel A, see Section 3.2) and calculated from a global CAMS reanalysis (Panel B, see Section 3.3 for details), averaged over the EM27/SUN measurement period ($23^{\rm rd}$ January to $19^{\rm th}$ April 2020.) The TROPOMI data are spatially binned into a 1x1 degree grid. The circles indicate the co-location criteria (50 km radius for $X_{\rm CO}$, centred on the measurement site in Jinja) used in the comparisons described in Section 4.3. Panels C and D show the same data as panels A and B respectively, zoomed in on the co-location region.

by Lunt et al. (2021). The inversion analysis we show here is an extension of the inversion presented in Lunt et al. (2021), from the end of 2019 through the first four months of 2020. For the *a priori* methane emissions inside the nested domain we use the EDGAR v4.3.2 database for anthropogenic emissions (Janssens-Maenhout et al., 2019), the WetCHARTs dataset for emissions from wetlands (Bloom et al., 2017), and the GFAS database for daily biomass burning emissions (Global Fire Assimilation

System, Kaiser et al. (2012)). The boundary conditions for the nested domain come from a global GEOS-Chem model run at lower spatial resolution ($2.0^{\circ} \times 2.5^{\circ}$). An ensemble Kalman Filter system (Hunt et al., 2007) is then used to perform the inversion, taking into account column CH₄ concentrations from TROPOMI (Lunt et al., 2021), which gives us estimates of the methane emissions within the nested domain along with the model-derived atmospheric concentrations. A subset of the mean $X_{\rm CH_4}$ values for the measurement period calculated from the output of this high-resolution regional inversion, covering the region surrounding the Jinja site (-3.0 to $+3.0^{\circ}$ N, +28.0 to $+36.0^{\circ}$ E), are shown in Figure 5B.

The non-satellite dataset we use in this study for carbon monoxide is the Copernicus Atmosphere Monitoring Service (CAMS) global reanalysis dataset (Inness et al., 2019), which covers the period from January 2003 to December 2021 with a spatial resolution of approximately $80 \,\mathrm{km}$ (interpolated onto a regular $0.75^\circ \times 0.75^\circ$ grid) and 60 vertical levels. A 4D-VAR assimilation framework (Rabier et al., 2000; Hollingsworth et al., 2008) is used to produce the reanalysis, which is described in detail for carbon monoxide by Flemming et al. (2017). Total column carbon monoxide data retrieved from the MOPITT instrument on board the NASA Terra satellite is used as input for the reanalysis (Deeter et al., 2014). Figure 6B shows the mean column concentration of carbon monoxide over East Africa for the measurement period, as calculated from the CAMS reanalysis output.

315 4 Comparisons of the EM27/SUN total column data with satellite, model and reanalysis datasets

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In this section we show how the total column concentrations observed using the EM27/SUN in Jinja compare with both satellite and model datasets, considering each species in turn. Before comparing the EM27/SUN and satellite data, we need to take into account that each retrieval algorithm used provides an estimate of the total column concentration that is based on different *a priori* information. Following the theory underpinning the optimal estimation retrieval method as described by Rodgers (2000), we correct for the different *a priori* profiles used in PROFFAST (see Section 2.3) and the respective satellite algorithms following Equation 3 in Dils et al. (2014), which assumes the ground-based *a priori* as the common *a priori* profile when validating satellite GOSAT data and ground-based TCCON total column concentrations of carbon dioxide and methane:

$$x_{\text{cor}} = x + \frac{1}{m_0} \sum_{i} m_i (A_i - 1) (ap_{i,sat} - ap_{i,EM27}).$$
(3)

Here, x_{cor} and x are the *a priori*-corrected and uncorrected dry-air total column concentrations; i is the vertical layer index, with corresponding mass of dry air m_i contained within the layer (derived from $\Delta p_i/g_i$, where Δp_i is the dry air pressure difference over layer i and g_i is the acceleration due to gravity at that height). m_0 is the total dry air mass of the atmospheric column, obtained by taking the sum of m_i over all layers. A_i is the column averaging kernel used by the satellite retrieval algorithm, and finally $ap_{i,sat}$ and $ap_{i,EM27}$ are the *a priori* dry air concentrations in layer i assumed by the satellite and EM27/SUN retrieval algorithms respectively.

For these comparisons, we use co-location criteria which represent a trade-off between ensuring there are enough measurement days to be able to make meaningful conclusions from the observed EM27/SUN vs. satellite/model differences, whilst also

ensuring that we are spatially comparing like with like. In the case of carbon dioxide and methane retrievals from OCO-2/3 and TROPOMI respectively, we employ a wider co-location radius ($300\,\mathrm{km}$) than used in the validation study of Sha et al. (2021) for example, as the cloudy conditions commonly encountered in the tropics limit the number of successful satellite retrievals. The close proximity of Lake Victoria to the south of the measurement site also has an impact here, since the low albedo of the lake surface at shortwave infrared wavelengths reduces the intensity of the observed signal below the level where a successful X_{CO_2} or X_{CH_4} retrieval is possible. We also split the time period in two, to check whether there is a notable difference in the comparisons as a result of the onset of the 'long rains' in March (see Section 2). In the following sections the two subsets are labelled 'dry', corresponding to January and February, and 'rainy', corresponding to the long rains period from March onwards.

340 4.1 Comparison of EM27/SUN $X_{\rm CO_2}$ with OCO-2/-3 and GEOS-Chem

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Here, we compare our EM27/SUN $X_{\rm CO_2}$ values with those retrieved from OCO-2 and OCO-3 observations (*a priori*-corrected as described above), and obtained from a global GEOS-Chem ${\rm CO_2}$ inversion which assimilates OCO-2 v10r data (see Section 3.3 for details). For the comparison we take OCO-2 and OCO-3 soundings (see Section 3.1), and GEOS-Chem grid points, within a 300 km radius of the EM27/SUN location, and calculate the median $X_{\rm CO_2}$ for each day (see Table B1 for a summary of how the choice of co-location radius affects the comparison). For $X_{\rm CO_2}$ we use all EM27/SUN data points, regardless of the time of day, in order to maximise the number of days of coincident OCO-2 and OCO-3 observations, and to take into account the varying OCO-3 overpass time. We also limit the OCO-2/3 vs. EM27/SUN comparison to days where there are at least ten $X_{\rm CO_2}$ soundings of sufficient quality that meet the co-location criteria described here. Figure 7 shows time series of these data, along with scatter plots directly comparing the EM27/SUN daily $X_{\rm CO_2}$ with the satellite and model datasets. The statistics of the $X_{\rm CO_2}$ comparisons are summarised in Table 1. The mean (standard deviation) *a priori* profile corrections given by Equation 3 are $-0.238\,(0.013)\,{\rm ppm}$ and $-0.373\,(0.082)\,{\rm ppm}$ for OCO-2 and OCO-3 respectively.

Whilst acknowledging that there are only a few days during the measurement period where OCO-2 and OCO-3 data can be compared with our EM27/SUN measurements, the data that we have available suggests that during this period the $X_{\rm CO_2}$ from OCO-2 is biased low with respect to that from the EM27/SUN, by 1.20 ppm (standard deviation is 1.05 ppm). The $X_{\rm CO_2}$ data from OCO-3 are also lower on average than what we observed with the EM27/SUN, however there are insufficient days of coincident observations during the measurement period to conclude that there is a statistically significant difference between the two. In both cases, the percentage difference from the EM27/SUN measurements (-0.29% for OCO-2, -0.28% for OCO-3) falls just outside the OCO mission precision requirement of 0.25%.

The GEOS-Chem model columns are also biased low on average with respect to the EM27/SUN data, and for our measurement period the inversion is insensitive to whether both satellite and *in-situ* or only *in-situ* observational data are assimilated. We see in both cases that these biases are primarily observed during the 'dry' period, where there are statistically significant differences from the EM27/SUN columns of $-0.86\,\mathrm{ppm}$ and $-0.72\,\mathrm{ppm}$ respectively (standard deviations are $0.76\,\mathrm{ppm}$ and $0.79\,\mathrm{ppm}$). The differences are most clear during the last week of February 2020 (see Figure 7B), and suggest the possibility of local sources not captured by the relatively coarse $(2.0^{\circ} \times 2.5^{\circ})$ latitude-longitude grid) global configuration of GEOS-Chem used here.

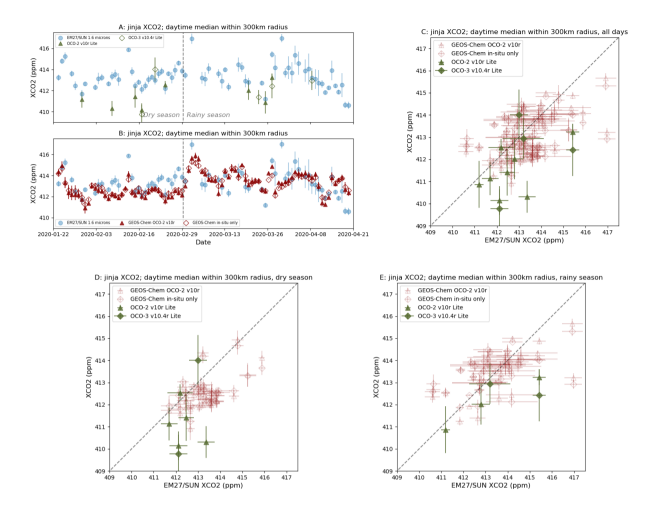


Figure 7. Panel A: daily median values of X_{CO_2} retrieved from EM27/SUN (blue circles), OCO-2 (olive filled triangles) and OCO-3 (olive open diamonds) measurements, error bars show the inter-quartile range. Panel B: same as Panel A, but with global GEOS-Chem model data (red filled triangles are with OCO-2 data assimilated, red open diamonds use in-situ observations only). Panel C: scatter plot showing EM27/SUN daily median observations vs. OCO-2 (olive filled triangles), OCO-3 (olive filled diamonds) and global GEOS-Chem (red open triangles with OCO-2 assimilated, red open diamonds with in-situ data only), error bars show the inter-quartile range. We use a co-location radius of 300 km. Panel D: same as Panel C, but including data from the 'dry' season only (January and February 2020). Panel E: same as Panel C, but including data from the 'rainy' season only (March and April 2020).

4.2 Comparison of EM27/SUN X_{CH4} with TROPOMI and GEOS-Chem

For X_{CH_4} we compare the EM27/SUN column concentrations with *a priori*-corrected data from TROPOMI observations (see Section 3.2), and column concentrations calculated using *a priori* and *a posteriori* emissions from the high resolution GEOS-Chem inversion (see Section 3.3 for details of the model run). A 300km co-location radius is used for both satellite and

Table 1. Mean and standard deviation of the differences (ΔX_{CO_2} in ppm) between daily median EM27/SUN and satellite/model X_{CO_2} . The 'dry' subset includes data from January and February 2020, whilst the 'rainy' subset covers data from March and April 2020.

Satellite/model	Subset	Number of days	Total number of satellite re- trievals	Mean ΔX_{CO_2} [ppm]	Std $\Delta { m X_{CO_2}}$ [ppm]	$\begin{array}{c} \text{Mean} \\ \frac{\Delta X_{\rm CO_2}}{X_{\rm CO_2,EM27}} \\ [\%] \end{array}$	$\begin{array}{c} \textbf{Std} \\ \frac{\Delta X_{\text{CO}_2}}{X_{\text{CO}_2, \text{EM27}}} \\ [\%] \end{array}$
OCO-2 v10r	All	8	1725	-1.20	1.05	-0.29	0.25
	Dry	5	1173	-1.26	1.17	-0.31	0.28
	Rainy	3	552	-1.10	0.79	-0.27	0.19
OCO-3 v10.4r	All	4	324	-1.15	1.61	-0.28	0.39
	Dry	2	248	-0.67	1.68	-0.16	0.41
	Rainy	2	76	-1.62	1.37	-0.39	0.33
GEOS-Chem including OCO-2	All	68	n/a	-0.35	1.08	-0.08	0.26
	Dry	31	n/a	-0.86	0.76	-0.21	0.18
	Rainy	37	n/a	0.077	1.13	0.02	0.27
GEOS-Chem in-situ only	All	68	n/a	-0.28	1.12	-0.07	0.27
	Dry	31	n/a	-0.72	0.79	-0.18	0.19

model data, and we only use EM27/SUN data and GEOS-Chem time steps within ± 2 hours of the TROPOMI overpass time, to calculate the median X_{CH_4} for each day (see Table B2 for a summary of how the choice of co-location radius affects the comparison). We also restrict the TROPOMI vs. EM27/SUN comparison to days where there are at least ten X_{CH_4} soundings of sufficient quality meeting the spatial and temporal co-location criteria described here. Figure 8 shows time series of these data, along with scatter plots directly comparing the EM27/SUN daily X_{CH_4} with the satellite and model datasets. The statistics of the X_{CH_4} comparisons are summarised in Table 2. The mean (standard deviation) *a priori* profile correction applied to the TROPOMI data, given by Equation 3, is +1.56 (0.15) ppb.

The short measurement period limits the number of days where comparisons can be made between the ground-based and satellite retrievals of $X_{\rm CH_4}$. In the data we have, the TROPOMI retrievals are lower than the EM27/SUN columns by a mean of $8.33\,\rm ppb$, albeit within the standard deviation ($10.5\,\rm ppb$) in the data. The mean percentage difference from the EM27/SUN measurements of $-0.44\,\%$ falls within the $1.5\,\%$ bias requirement on the TROPOMI $X_{\rm CH_4}$ data product.

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The data from the GEOS-Chem high resolution inversions show better agreement with the EM27/SUN data in terms of the mean differences. The difference is slightly greater ($-3.80\,\mathrm{ppb}$ compared with $-1.15\,\mathrm{ppb}$) when the *a posteriori* emissions incorporating TROPOMI $X_{\mathrm{CH_4}}$ are used, though the difference between the two is well within their respective standard deviations. The only comparison where there is a statistically significant difference from the EM27/SUN columns is that with the

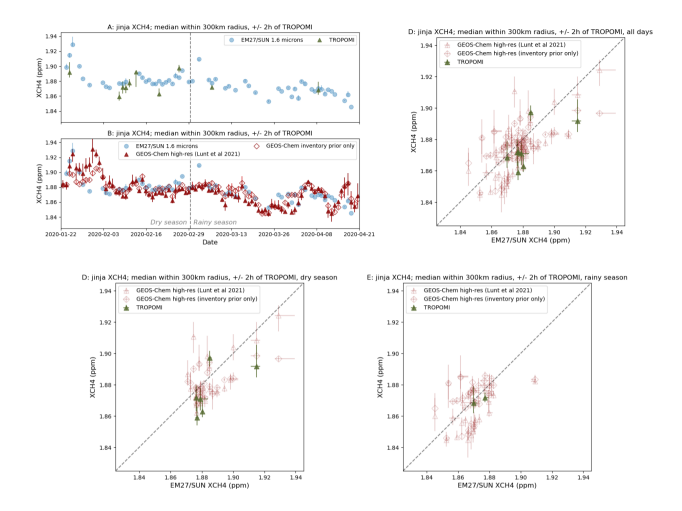


Figure 8. Panel A: daily median values of X_{CH_4} retrieved from EM27/SUN (blue) and TROPOMI (dark green) measurements, bars show the inter-quartile range. Panel B: same as Panel A, but with high resolution regional GEOS-Chem model data (darker shade of red is with TROPOMI data assimilated, lighter shade uses inventory emissions only). Panel C: scatter plot showing EM27/SUN observations vs. TROPOMI (dark green filled circles) and high resolution regional GEOS-Chem (red open circles with TROPOMI assimilated, blue diamonds with inventory emissions only), error bars show the inter-quartile range. We use a co-location radius of 300 km, and only consider data and model output within ± 2 hours of the TROPOMI overpass at 1030 UTC. Panel D: same as Panel C, but including data from the 'dry' period only (January and February 2020). Panel E: same as Panel C, but including data from the 'rainy' period only (March and April 2020).

385 GEOS-Chem simulation using *a posteriori* emissions, though there is not sufficient data to attribute this to either the 'dry' or the 'rainy' periods considered here.

There are a couple of possible explanations for differences seen between the EM27/SUN and GEOS-Chem $X_{\rm CH_4}$ data. Firstly, the posterior scale factors which are applied to the prior emission fields have an exponential correlation length scale of

Table 2. Mean and standard deviation of the differences (ΔX_{CH_4} in ppb) between daily median EM27/SUN and satellite/model X_{CH_4} . The 'dry' subset includes data from January and February 2020, whilst the 'rainy' subset covers data from March and April 2020.

Satellite/model	Subset	Number of days	Total number of satellite re- trievals	Mean $\Delta { m X}_{ m CH}_4$ [ppb]	Std $\Delta { m X}_{ m CH_4}$ [ppb]	$\begin{array}{c} \text{Mean} \\ \frac{\Delta X_{\mathrm{CH_4}}}{X_{\mathrm{CH_4,EM27}}} \\ [\%] \end{array}$	$\begin{array}{c} \textbf{Std} \\ \frac{\Delta X_{\text{CH}_4}}{X_{\text{CH}_4, \text{EM27}}} \\ [\%] \end{array}$
TROPOMI	All	8	1674	-8.39	10.5	-0.44	0.56
	Dry	6	1610	-9.97	11.7	-0.53	0.62
	Rainy	2	64	-3.66	1.98	-0.19	0.11
GEOS-Chem							
HR including	All	58	n/a	-3.80	12.5	-0.22	0.66
TROPOMI data							
	Dry	27	n/a	-2.85	11.9	-0.16	0.62
	Rainy	31	n/a	-4.53	13.0	-0.24	0.70
GEOS-Chem							
HR inventory	All	58	n/a	-1.15	11.6	-0.06	0.62
only							
	Dry	27	n/a	-3.59	10.7	-0.19	0.55
	Rainy	31	n/a	0.90	12.2	0.05	0.65

 $50\,\mathrm{km}$, meaning that smaller scale variations in the emissions that influence the EM27/SUN measurements may not be reflected in the differences between the posterior and prior inversions. It is also worth noting that the *a posteriori* inversion minimises the residual to all TROPOMI $X_{\mathrm{CH_4}}$ data within the larger inversion domain, rather than this specific grid box centred on Jinja. We can see from the EM27/SUN vs. TROPOMI comparison in Figure 8 that there are only a limited number of TROPOMI data available to constrain emissions during the measurement period, such that emissions local to the site are unlikely to be well represented in the inversion.

4.3 Comparison of EM27/SUN X_{CO} with TROPOMI and CAMS

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In the final part of this section we compare $X_{\rm CO}$ retrieved from the EM27/SUN ground-based observations with $X_{\rm CO}$ from TROPOMI data (see Section 3.2), and from the output of the global CAMS reanalysis (see Section 3.3 for details). The greater number of soundings with successful retrievals of $X_{\rm CO}$ allows us to apply a tighter 50 km co-location radius to the satellite and reanalysis data (see Table B3 for a summary of how the choice of co-location radius affects the comparison). As for $X_{\rm CH_4}$, we only use EM27/SUN data and CAMS time steps within $\pm 2\,{\rm hours}$ of the Sentinel-5P overpass time, to calculate the median $X_{\rm CO}$ value for each day. In addition we further restrict the TROPOMI vs. EM27/SUN comparison to days where there are at least ten $X_{\rm CO}$ soundings of sufficient quality meeting these spatial and temporal co-location criteria. Figure 9 shows time series of these data, along with scatter plots directly comparing the EM27/SUN daily $X_{\rm CO}$ with the satellite and reanalysis datasets.

The statistics of the X_{CO} comparisons are summarised in Table 3. The mean (standard deviation) *a priori* profile correction applied to the TROPOMI data, given by Equation 3, is -2.84(3.27) ppb.

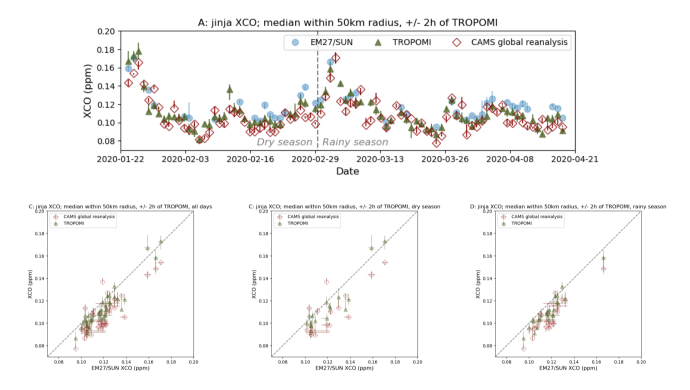


Figure 9. Panel A: daily median values of X_{CO} retrieved from EM27/SUN (blue filled circles) and TROPOMI (dark green filled triangles) measurements, and obtained from CAMS global reanalysis output (open red diamonds), bars show the inter-quartile range. Panel B: scatter plot showing EM27/SUN observations vs. TROPOMI (dark green filled triangles) and global CAMS reanalysis data (red open diamonds), error bars show the inter-quartile range. We use a co-location radius of $50 \, \mathrm{km}$, and only consider data and model output within 2 hours of the TROPOMI overpass at 1030 UTC. Panel C: same as Panel B, but including data from the 'dry' period only (January and February 2020). Panel D: same as Panel B, but including data from the 'rainy' period only (March and April 2020).

As discussed in Section 3.2, the availability of X_{CO} data from TROPOMI in partially cloudy conditions means that we have a greater number of measurement days suitable for comparison compared with X_{CO_2} and X_{CH_4} . We find a statistically significant difference between our ground-based observations and the TROPOMI satellite data, with the X_{CO} from TROPOMI being biased lower than that from the EM27/SUN by a mean value of $6.62\,\mathrm{ppb}$, which falls just outside the standard deviation $(6.25\,\mathrm{ppb})$ of the bias. These values are still well within the mission data requirements for the TROPOMI carbon monoxide data product, which stipulate that the magnitude of the bias $(-5.65\,\%$ for the whole measurement period, with standard deviation of $4.99\,\%$) should be less than 15% and the random error less than 10% (Landgraf et al., 2016).

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The X_{CO} values from the CAMS global reanalysis are also significantly low with respect to the ground-based EM27/SUN data, by a mean value of 11.7 ppb. This mean bias is greater than the standard deviation (8.94ppb) both throughout our

Table 3. Mean and standard deviation of the differences (ΔX_{CO} in ppb) between daily median EM27/SUN and satellite/reanalysis X_{CO} . The 'dry' subset includes data from January and February 2020, whilst the 'rainy' subset covers data from March and April 2020.

Satellite/model	Subset	Number of days	Total number of satellite retrievals	Mean ΔX_{CO} [ppb]	$egin{aligned} \mathbf{Std} \ \mathbf{\Delta X_{CO}} \ \mathbf{[ppb]} \end{aligned}$	$\frac{\Delta X_{\rm CO}}{X_{\rm CO,EM27}}$ [%]	Std $\frac{\Delta X_{CO}}{X_{CO,EM27}}$ [%]
TROPOMI	All	41	4738	-6.62	6.25	-5.65	4.99
	Dry	17	2323	-6.28	7.89	-5.30	6.03
	Rainy	24	2415	-6.85	4.74	-5.90	4.07
CAMS global reanalysis	All	43	n/a	-11.7	8.94	-9.79	7.45
	Dry	18	n/a	-11.8	11.4	-9.71	9.35
	Rainy	25	n/a	-11.6	6.65	-9.85	5.69

measurement period, and when separating into 'dry' and 'rainy' periods, suggesting that the CAMS global model may not be taking into account all local sources of carbon monoxide. In addition, recent work by Inness et al. (2022) has shown that assimilating TROPOMI carbon monoxide data into the CAMS system (in addition to the satellite data from MOPITT and IASI that is already assimilated in this version) increases the CAMS carbon monoxide columns by 8% on average, which would bring the CAMS reanalysis output into closer agreement with our EM27/SUN observations.

420 5 Conclusions and outlook

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In this paper, we describe the first ground-based remote sensing observations of total column greenhouse gas concentrations to have been performed in the Tropical East Africa region. We set up a Bruker EM27/SUN spectrometer at the headquarters of the National Fisheries Resources Research Institute in Jinja, Uganda, in January 2020. An automated enclosure for the instrument, designed and built by the Environmental Sensing and Modelling Group at the Technical University of Munich, allowed us to operate the instrument remotely and autonomously for a period of three months, providing a temporal density of greenhouse gas column data over this period that would have been challenging to achieve manually. The combined performance of the instrument and enclosure shown in this paper demonstrates the possibility to deploy EM27/SUN instruments as validation sites for satellite greenhouse gas retrievals, in parts of the world where it would be logistically difficult to establish new sites to extend established ground-based validation networks such as TCCON.

The ground-based measurements of carbon dioxide, methane, and carbon monoxide column concentrations that we have acquired using the EM27/SUN and automated enclosure allow us, for the first time, to see how well satellite and model datasets have performed in observing or calculating these concentrations over Uganda during our measurement period. For carbon dioxide, we find OCO-2 $X_{\rm CO_2}$ to be lower than our EM27/SUN measurements by a mean of 1.20 ppm, with standard deviation 1.05 ppm. Given the lack of days of coincident observations during the measurement period, we do not observe a

statistically significant difference between EM27/SUN and OCO-3 $X_{\rm CO_2}$ for this dataset (for the days we do have coincident observations, OCO-3 was lower by a mean of $1.15\,(1.61)\,\rm ppm$). In both cases, the percentage difference from the EM27/SUN measurements ($-0.29\,\%$ for OCO-2, $-0.28\,\%$ for OCO-3) falls just outside the OCO mission precision requirement of $0.25\,\%$. The global GEOS-Chem inversion we use for this study – irrespective of whether OCO-2 data has been assimilated – is also generally biased low with respect to the EM27/SUN measurements. We find GEOS-Chem $X_{\rm CO_2}$ to be lower by a mean of $0.35(1.08)\,\rm ppm$ with, and $0.28\,(1.12)\,\rm ppm$ without OCO-2 included in the inversion.

In the case of $X_{\rm CH_4}$, we find that TROPOMI is lower than our EM27/SUN by a mean of $8.33\,(10.5)\,\rm ppb$), albeit for a limited number of days where we had coincident observations. The mean percentage difference from the EM27/SUN measurements of $-0.44\,\%$ does however fall within the $1.5\,\%$ bias requirement on the TROPOMI $X_{\rm CH_4}$ data product. The high resolution GEOS-Chem inversion we use in this study, which incorporates TROPOMI data, is also biased low with respect the EM27/SUN observed $X_{\rm CH_4}$ (GEOS-Chem lower by a mean of $3.80\,(12.5)\,\rm ppb$). This may be a result of the *a posteriori* inversion being set up to minimise the residual to all TROPOMI $X_{\rm CH_4}$ data within the larger inversion domain, rather than just data within the specific grid box centred on Jinja where TROPOMI methane soundings are relatively scarce. This means that emissions local to the measurement site are unlikely to be well represented in the inversion.

When we consider carbon monoxide from TROPOMI, the quality flagging of the column concentration retrievals is much less sensitive to cloud cover, such that there were many more days with coincident observations that we could compare with our EM27/SUN measurements. Even over a three month period, there was sufficient data to be able to conclude that the carbon monoxide columns from TROPOMI were biased low with respect to the EM27/SUN data by a mean value of $3.68\,\mathrm{ppb}$ (standard deviation $7.00\,\mathrm{ppb}$). This is still well within the mission data requirements for the TROPOMI carbon monoxide data product, which stipulate that the bias ($-5.65\,\%$ for the whole measurement period, with standard deviation of $4.99\,\%$) should be less than 15% and the random error less than 10% (Landgraf et al., 2016). We also see a statistically significant difference between our EM27/SUN measurements and the CAMS global reanalysis (CAMS $X_{\rm CO}$ lower by a mean of $11.7\,\mathrm{ppb}$, with standard deviation $8.94\,\mathrm{ppb}$), suggesting that the CAMS global reanalysis may not be taking into account all local sources of carbon monoxide. Recent work by Inness et al. (2022) has, however, shown that assimilating TROPOMI carbon monoxide data into the CAMS system (in addition to the satellite data from MOPITT and IASI that is already assimilated in this version) would increase the CAMS carbon monoxide columns by 8% on average, which if applicable to tropical East Africa would bring the CAMS reanalysis output into closer agreement with our EM27/SUN observations.

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An important aspect of this work is the comparison with atmospheric chemistry and transport model output. Models and reanalyses such as GEOS-Chem and CAMS provide a means of studying atmospheric processes where observations are not available. This is of particular relevance in tropical Africa (e.g. Lunt et al. (2019, 2021); Palmer et al. (2019); Feng et al. (2022)), where ground-based observations of greenhouse gases are scarce and the data coverage provided by satellites is often limited by cloud cover. Ground-based column concentration observations such as those presented in this study provide data that can be used to evaluate these models and reanalyses which, unlike in-situ measurements, are not overly sensitive to emission sources on a local scale. Our results show that only three months of measurements can be sufficient to demonstrate the effectiveness of these models and reanalyses at this time of year, whilst also highlighting short periods where there are discrepancies to be

investigated further. A comprehensive validation of the models would require at least a whole calendar year of observations. Figure 10 shows radial histograms of the wind direction for each month of the year 2020. These demonstrate how the typical wind direction at 800 hPa, in the lower troposphere where the retrieved column concentrations are generally most sensitive, varies throughout the year. During the measurement period for this study, as summarised by the radial histogram in Figure 10A, the wind has most frequently blown from the north (particularly in January and February) and from the south east (from late February to April). A full year of measurements would be more representative of the variety of atmospheric conditions we would expect to observe from satellites or estimate from models, and would give us greater confidence in the performances of the retrieval algorithms and the model and reanalysis calculations respectively. The period from October to December would be particularly interesting to focus on in the future, as the typically northerly winds we see at that time of year (Figure 10B) coincide with the 'short rains', the intensity of which Lunt et al. (2021) have linked to changes in methane emissions from the Sudd wetlands, located in South Sudan to the north of the measurement site.

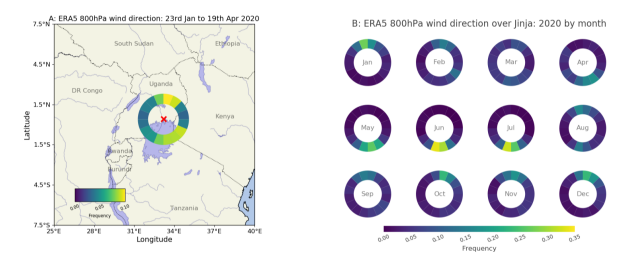


Figure 10. Panel A: radial histogram showing frequency of wind direction in the ERA5 reanalysis at 800 hPa above Jinja during the measurement period for this study. Panel B: radial histograms showing the frequency of wind direction at 800 hPa in ERA5 over Jinja for each month in 2020. The angles of the histogram segments correspond to the direction that the wind is coming from. Note that we use different colour scales in Panels A and B.

In summary, this study demonstrates the feasibility of a longer-term, autonomous deployment of the EM27/SUN instrument in a tropical environment, through the use of an automated weatherproof enclosure. This EM27/SUN plus enclosure system allows us to meet the goal of seasonal observations in support of studies focusing on the tropical carbon cycle, and the validation of greenhouse gas column concentration data from satellite retrievals, and from model and reanalysis calculations in the tropical East Africa region.

Code and data availability. The EM27/SUN column data and GEOS-Chem data will be made available on the Centre for Environmental Data Analysis archive. The latest version of the Pyra software used to control the automated weatherproof enclosure is available on GitHub (https://github.com/tum-esm/pyra). The latest version of the PROFFAST interferogram processing and analysis code is available from https://www.imk-asf.kit.edu/english/3225.php. The L2 column carbon dioxide data from OCO-2 and OCO-3 are available from the Goddard Earth Sciences Data and Information Services Centre (https://disc.gsfc.nasa.gov/datasets). The TROPOMI column methane and carbon monoxide data are available from the Sentinel-5P Pre-Operations Data Hub (https://s5phub.copernicus.eu/). The CAMS global reanalysis carbon monoxide concentration data are available from the CAMS Atmosphere Data Store (see https://www.ecmwf.int/en/research/climate-reanalysis/cams-reanalysis). The ERA5 reanalysis wind data are available from the Copernicus Climate Data Store (https://cds.climate.copernicus.eu/).

495 Appendix A: Further characteristics of the EM27/SUN instrument

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Here, we provide some further details on the EM27/SUN instrument and PROFFAST retrieval outputs for the reader's information. The spectral windows used by the PROFFAST retrieval code described in Section 2.3 are summarised in Table A1, and in Figure A1 we show example spectra and spectral residuals which can be used to check how well the retrieval has performed. Finally, Figure A2 shows example column sensitivities for carbon dioxide, methane, and carbon monoxide, presented as a function of pressure and solar zenith angle.

Table A1. Summary of the spectral windows used in the PROFFAST retrieval algorithm. The primary detector window near $1.6 \, \mu \mathrm{m}$ is used for the X_{CH_4} results shown in this study.

Gas	Wavenumber range (cm^{-1})	Wavelength range (μm)	Detector
$X_{\mathrm{CH_4}}$	5897 - 6145	1.627 – 1.696	Primary
X_{CO_2}	6173 – 6390	1.565 – 1.620	Primary
X_{O_2}	7765 – 8005	1.249 – 1.288	Primary
$\rm X_{\rm H_2O}$	8353 – 8463	1.182 - 1.197	Primary
$X_{\rm CO}, X_{{ m CH}_4}$	4210 – 4320	2.315 - 2.375	Secondary

Appendix B: Effect of spatial co-location criteria on EM27/SUN vs. satellite $X_{\rm gas}$ comparison

In these tables, we show how the mean and standard deviation of the differences between the EM27/SUN and satellite $X_{\rm gas}$ vary with increasing co-location radius. In each case, when choosing the co-location radius we make a compromise between expanding the radius enough to include enough data for a meaningful comparison across the measurement period, whilst also keeping the radius narrow enough that we minimise the potential influence of geo-spatial biases.

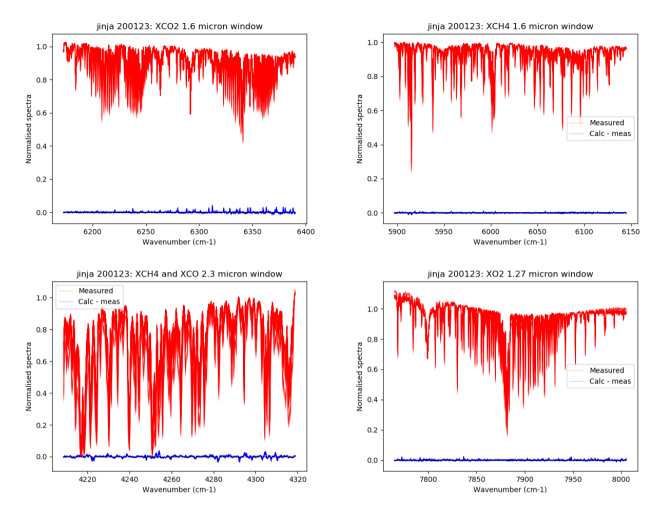


Figure A1. Example EM27/SUN spectra (shown in red) and spectral residuals (shown in blue) output by PROFFAST for measurements taken on January 23rd 2020.

Author contributions. NH set up the measurements, remotely monitored the instrument performance during the study period, performed the EM27/SUN data processing and analysis, prepared the figures, and wrote the manuscript. HB conceived the study, advised on interpretation of the results, and reviewed the manuscript. WO provided and prepared the measurement site, helped set up the measurements, and oversaw the site once measurements were up and running. JC and FD developed and built the automated weatherproof enclosure for the EM27/SUN used in this study. MFL performed the GEOS-Chem methane model runs, advised on interpretation of the results, and reviewed the manuscript. LF performed the GEOS-Chem carbon dioxide model runs and advised on their interpretation. PIP advised on the wider context and interpretation of the results, and reviewed the manuscript. FH devised the EM27/SUN instrument concept, and developed the PROFFAST retrieval code used in this study.

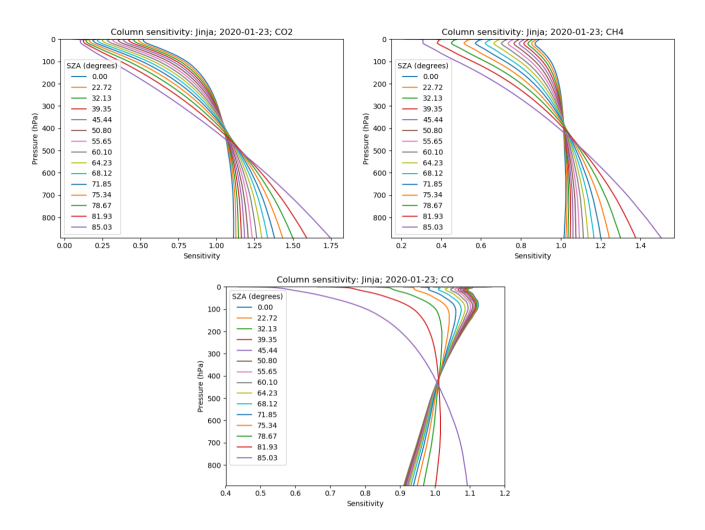


Figure A2. Column sensitivities calculated by the PROFFAST algorithm for January 23rd 2020 for carbon dioxide, methane, and carbon monoxide, presented as a function of pressure and solar zenith angle (SZA).

Competing interests. FH is a member of the editorial board of Atmospheric Measurement Techniques. The authors have no other competing interests to declare.

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Table B1. Mean and standard deviation of the differences (ΔX_{CO_2} in ppm) between daily median EM27/SUN and satellite X_{CO_2} , assuming a range of different spatial co-location criteria. The co-location radius chosen for this study is highlighted in bold. The $100 \, \text{km}$ and $200 \, \text{km}$ rows are identical for OCO-3 because of the sampling pattern used, which targets the measurement site whilst the ISS is passing overhead (see Figure 4).

Satellite	Radius (km)	Number of days	Total number of satellite re- trievals	Mean ΔX_{CO_2} [ppm]	Std ΔX_{CO_2} [ppm]	$\frac{\Delta X_{\rm CO_2}}{X_{\rm CO_2,EM27}}$ [%]	$\begin{array}{c} \text{Std} \\ \frac{\Delta X_{\text{CO}_2}}{X_{\text{CO}_2, \text{EM27}}} \\ [\%] \end{array}$
OCO-2 v10r	100	4	305	-0.39	1.05	-0.09	0.25
	200	5	752	-0.66	0.90	-0.15	0.21
	300	8	1725	-1.20	1.05	-0.29	0.25
	400	11	2894	-1.27	0.87	-0.31	0.21
	500	13	4464	-1.31	0.78	-0.32	0.19
	600	17	5969	-1.35	0.97	-0.33	0.23
OCO-3 v10.4r	100	2	76	-1.62	1.37	-0.39	0.33
	200	2	76	-1.62	1.37	-0.39	0.33
	300	4	324	-1.15	1.61	-0.28	0.39
	400	4	336	-1.13	1.59	-0.27	0.39
	500	13	938	-1.31	0.78	-0.32	0.19
	600	17	3980	-1.35	0.97	-0.33	0.23

Table B2. Mean and standard deviation of the differences (ΔX_{CH_4} in ppb) between daily median EM27/SUN and satellite X_{CH_4} , assuming a range of different spatial co-location criteria. The co-location radius chosen for this study is highlighted in bold.

Satellite	Radius (km)	Number of days	Total number of satellite re- trievals	Mean $\Delta { m X}_{ m CH}_4$ [ppb]	Std $\Delta { m X}_{ m CH_4}$ [ppb]	$\begin{array}{c} \text{Mean} \\ \frac{\Delta X_{\text{CH}_4}}{X_{\text{CH}_4, \text{EM27}}} \\ [\%] \end{array}$	$\begin{array}{c} \textbf{Std} \\ \frac{\Delta X_{\text{CH}_4}}{X_{\text{CH}_4, \text{EM27}}} \\ [\%] \end{array}$
TROPOMI	100	2	96	-3.52	2.86	-0.19	0.15
	200	6	711	-7.59	10.8	-0.40	0.57
	300	8	1674	-8.39	10.5	-0.44	0.56
	400	16	3079	-5.38	16.8	-0.28	0.88
	500	25	5312	-0.89	16.8	-0.04	0.88
	600	31	9258	0.07	18.6	0.009	0.98

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Table B3. Mean and standard deviation of the differences ($\Delta X_{\rm CO}$ in ppb) between daily median EM27/SUN and satellite $X_{\rm CO}$, assuming a range of different spatial co-location criteria. The co-location radius chosen for this study is highlighted in bold.

Catallita	Doding (km)	Number of days	Total number of satellite re-	Mean	Std	Mean $_{\Delta X_{CO}}$	Std $_{\Delta X_{CO}}$
Satellite	Radius (km)	Number of days	trievals	$\Delta m X_{CO}$ [ppb]	$\Delta m X_{CO}$ [ppb]	X _{CO,EM27} [%]	X _{CO,EM27} [%]
TROPOMI	25	32	1217	-6.31	6.04	-5.63	5.11
	50	41	4738	-6.62	6.25	-5.65	4.99
	75	41	9822	-5.98	6.30	-5.06	5.04
	100	42	16667	-5.66	5.95	-4.73	4.76
	125	42	25133	-5.44	5.75	-4.51	4.56

were produced by the OCO-2 project at the Jet Propulsion Laboratory, California Institute of Technology, and obtained from the OCO-2 data archive maintained at the NASA Goddard Earth Science Data and Information Services Center.

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