Understanding the Trend of NO2, SO2 and CO over East Africa from 2005 to 2020

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Abstract: The atmospheric chemistry constituents of nitrogen dioxide (NO2), sulphur dioxide (SO2) and carbon monoxide (CO) are associated with air pollution and climate change. In sub-Saharan Africa, a lack of sufficient ground-based and aircraft observations has, for a long time, limited the study of these species. This study thus utilized satellite observations as an alternative source of data to study the abundance of these species over the East African region. The instruments used included the Ozone Monitoring Instrument (OMI), the Atmospheric InfraRed Sounder (AIRS), and the TROP-Ospheric Monitoring Instrument (TROPOMI). An investigation of trends in the data series from 2005 to 2020 was carried out using the sequential Mann-Kendall test while the Pearson correlation coefficient was used to compare the data records of the instruments. The analysis revealed no trend in NO2 (p > 0.05), a decreasing trend in SO2 (p < 0.05), a decreasing trend (p < 0.05) in CO closer to the surface (850 hPa to 500 hPa) and an increasing trend (p < 0.05) in CO higher up in the atmosphere (400 hPa to 1 hPa). There is likely a vertical ascent of CO. The correlation between the instrument records was 0.54 and 0.77 for NO2 and CO, respectively. Furthermore, seasonal fires in the savanna woodlands were identified as the major source of NO2 and CO over the region, while cities such as Kampala, Nairobi, and Bujumbura and towns such as Dar es Salaam and Mombasa were identified as important NO2 hotspots. Similarly, the active volcano at Mt. Nyiragongo near Goma was identified as the most important SO2 hotspot.

Keywords: nitrogen dioxide; sulphur dioxide; carbon monoxide; OMI; TROPOMI

1. Introduction

Air pollution is one of the outstanding environmental concerns in the world today, affecting both developed and developing regions [1,2]. In 2016 alone, it accounted for 4.2 million deaths globally, because of its association with stroke, lung cancer, ischemic heart disease, acute lower respiratory disease and chronic obstructive pulmonary disease [3]. This pollution often takes the form of particulate matter (PM2.5 and PM10), nitrogen oxide (NO), nitrogen dioxide (NO2), sulphur dioxide (SO2), carbon monoxide (CO), and ozone (O3). This study only focused on three species, that is, NO2, SO2 and CO.

In East Africa, the important sources of these pollutants include biomass burning in the savanna, which is mostly carried out to clear land and to destroy crop residue, although it is at times triggered naturally by lightning [4,5]. The other sources are volcanic eruptions (specifically for SO2) and incomplete fossil fuel combustion in activities tagged to the local economy. These include transportation, manufacturing, and electricity generation. Domestic activities such as rubbish burning, cooking with charcoal, briquettes or firewood also emit these gases locally [5–7]. In the dry seasons, December to February and June to August, long-range winds may transport these gases from neighbouring regions into East Africa, adding to the observed gas abundances over the region [5].
NO$_2$, SO$_2$, and CO are also categorized under short-lived climate forcers because of the radiative forcing they induce in the atmosphere through the formation of other chemical species. NO$_2$ and SO$_2$ are precursors for nitrate and sulphate aerosols, respectively, both of which induce a net cooling effect. CO is a precursor for tropospheric ozone, which induces a net warming effect. These species also have short atmospheric lifetimes, realistically, days to a few months. SO$_2$ resides in the atmosphere for 2 days while for NO$_2$ that time is longer at 1 to 10 days and CO has the longest residence time of 30 to 90 days. This lifetime can change depending on the species height in the atmosphere [8,9]. This is why these gases are increasingly gaining attention in climate discussions [10,11].

Since cities are a hub for economic activity, they are often associated with higher emissions of these short-lived climate forcers / air pollutants than their surrounding regions. Consequently, cities have been the target for most studies exploring the abundances of these gases. Over East Africa, these cities include Kampala [12,13], Nairobi [14,15] and Dar es Salaam [16], which have expanded over the past two decades [17,18]. Now, however, it might be important to also study the overall regional distribution of these gases, in addition to the city-scale studies. This is because long-range wind transportation can shift significant gas amounts out of cities and spread them into neighbouring areas [19]. Furthermore, given the increasing importance of these gas species in understanding regional climates, having regional-scale studies as well is more meaningful than having city-scale studies alone.

Understanding the long-term trends of these gas species over East Africa is useful for inferring the region’s emission patterns and trends in air pollution, for studying the regional climate change driven by such short-lived climate forcers and for informing both climate and air quality policy. However, there are limited studies that have explored the trends of East Africa’s NO$_2$, SO$_2$, and CO over a long period of time. The major hindrance has been a lack of consistent observational records covering the entire region. To fill this gap, some observation campaigns have been set-up. These include; the Rwanda Climate Observatory, which started making observations in May 2015 [5] and Clean Air Nairobi, whose observations started in May 2016 [14]. However, these data are not yet sufficient to infer the long-term, spatially representative trends of these gases over the East African region.

In this study, therefore, satellite observations are utilized to study the long-term trends of NO$_2$, SO$_2$, and CO over the region of East Africa. Records from two chemistry-focused satellite instruments are used for each gas species, these are the Ozone Monitoring Instrument (OMI) [20,21] for NO$_2$ and SO$_2$ observations from 2005 to 2020, the Atmospheric InfraRed Sounder (AIRS) [22] for CO observations from 2005 to 2020, and the TROPOspheric Monitoring Instrument (TROPOMI) [23] for observations of all the three gases from 2018 to 2020. OMI and AIRS were useful to ensure ample temporal coverage of the historical period, while TROPOMI records were included in the analysis as a complementary data source, especially since it has newer technology and makes observations at a higher spatial resolution than OMI and AIRS. These datasets have an advantage over the existing ground-based measurements because they cover the entire spatial extent of the region, which is currently not yet possible with the existing ground-based observation network.

The objectives of this study were to determine the trend in NO$_2$, SO$_2$, and CO over East Africa between 2005 and 2020 and to compare the observations of TROPOMI with those of the longer-serving instruments, OMI and AIRS. The association of these chemistry species with meteorological variables was also investigated. The rest of this paper is organized as follows: Section 2 is the methodology section which includes a description of the study region, the satellite instruments from which data were collected and the methods of analysis. In Section 3, the important findings of the study are presented, and the major conclusions are summarized in Section 4.
2. Methodology

2.1. Study Region

This study focused on five countries in the East African region: Uganda, Kenya, Tanzania, Rwanda and Burundi (Figure 1). Politically, however, South Sudan is considered part of this region. The five countries were selected because they have the same bimodal rainfall regime and South Sudan was excluded because it has a unimodal rainfall regime, which is distinct from the other five countries [24]. The combined population of the region is estimated at 177.2 million inhabitants, gaining annual increments at a rate of 2.9% [25]. Such regions of sub-Saharan Africa are projected to be one of the important emission sources of short-lived climate and air pollutants toward the end of the century [26], making it prudent to start investigating the existing abundances of such species over the region.

![Figure 1. Map of East Africa showing its vegetation cover along with the capital cities and other towns.](image)

2.2. Data

2.2.1. NO\(_2\) and SO\(_2\) from the Ozone Monitoring Instrument (OMI)

OMI is a spectrometer aboard the Aura satellite operated by the National Aeronautics and Space Administration (NASA). Since July 2004, OMI has provided near global coverage of critical trace gases and aerosols, such as NO\(_2\), SO\(_2\), HCHO, O\(_3\) and OCIO, on a 24-h basis. These observations are retrieved from measurements of solar irradiance scattered by atmospheric components via three wavelength channels consisting of both the ultraviolet and visible spectrum (\(\sim 270–500\) nm). The observations are made at a spatial resolution of \(13 \times 24\) km\(^2\) at nadir [20,21]. The NO\(_2\) product utilized in this study is version 4.0 of the
OMI tropospheric NO$_2$ vertical column density (VCD) with improved cloud (30% cloud screened) and surface treatments and comes in units of molecules/cm$^2$ [27]. These data have a slant column density (SCD) uncertainty of $\sim 0.8 \times 10^{15}$ molecules/cm$^2$ [28]. This study also utilized version 2 of the OMI anthropogenic SO$_2$ (OMISO2 v2) VCD which has been tested and shown to have $\sim 0.15$--0.3 Dobson Units (DU) margin of uncertainty in the slant column densities (SCD) over the region of interest [29]. Daily quantities of both NO$_2$ and SO$_2$ from OMI for the time range 2005 to 2020 were downloaded from the GIOVANNI data handle of NASA (https://giovanni.gsfc.nasa.gov/; accessed on 7 June 2021) at a spatial resolution of 0.25$^\circ$ (~27.5 km). The analysis was then made based on area averages over the entire region.

2.2.2. CO from the Atmospheric InfraRed Sounder (AIRS)

AIRS is both a spectrometer and radiometer aboard NASA’s Aqua satellite, which was launched in May 2002. It operates within a spectral range of 3.7–15.4 µm to make direct measurements of temperature and atmospheric chemistry species such as CH$_4$, O$_3$ and CO. It also contributes to atmospheric measurements of cloud properties and humidity [22]. This study utilized version 7.0 of the AIRS CO product in the daytime/ascending orbit which gives the CO mole fraction in the air in units of parts per billion by volume (ppbv). Data over the entire study region for the period 2005 to 2020 were downloaded from the GIOVANNI data handle of NASA at a daily temporal resolution and a spatial resolution of 1$^\circ$ × 1$^\circ$ (~110 km) [30]. Before analysis, the data were summed up for 22 atmospheric levels, from 850 hPa to 1 hPa to obtain the total column amount. The levels, 1000 hPa and 925 hPa that are associated with the boundary layer were excluded from the analysis because they had no data. AIRS has a limitation of not sensing the lowest part of the troposphere where biomass burning occurs. Consequently, the AIRS CO data used here is underestimated.

2.2.3. NO$_2$, SO$_2$ and CO data from the TROPOspheric Monitoring Instrument (TROPOMI)

TROPOMI is a spectrometer aboard the Sentinel-5 Precursor satellite operated by the European Space Agency (ESA). TROPOMI started operation in October 2017, and it makes global measurements of critical atmospheric trace gases such as NO$_2$, SO$_2$, O$_3$, CO, HCHO, and CH$_4$ along with cloud properties via four (4) wavelength bands: ultraviolet, visible, near infrared and shortwave infrared, all ranging from ~270–2385 nm. The measurements are made daily at a spatial resolution of 3.5 × 7 km$^2$ at nadir [23], but since 6th August 2019, the resolution has been improved to 3.5 × 5.5 km$^2$. This study utilized the level 3 offline products for NO$_2$, SO$_2$ and CO from 2018 to 2020. These were all downloaded from Google Earth engine [31] in units of mol/m$^2$ and a spatial resolution of 3.5 × 7 km$^2$ for NO$_2$ and SO$_2$ and 7 × 7 km$^2$ for CO. In the analysis of NO$_2$ and CO, the content downloaded were multiplied by the factor $6.02214 \times 10^{20}$ to obtain quantities in units of molecules/cm$^2$ while for SO$_2$, it was multiplied by 2241.15 to obtain a quantity in Dobson Units (DU). Quality control has already been applied to these data, and a quality assurance value (qa_value) ranging from 0 (very poor) to 1 (very good) appended. For the NO$_2$ data used, the qa_value was above 0.75, while for SO$_2$ and CO the qa_value was above 0.5. Since the SO$_2$ data observed over regions with low SO$_2$ emissions can have negative vertical column values in some instances, only SO$_2$ columns above $-0.001$ mol/m$^2$ were used as recommended by the data providers [32].

2.2.4. Meteorological Observations

The rainfall observations used in this study were based on the Climate Hazards Group Infrared Precipitation with Stations (CHIRPS). This dataset is a hybrid combination between satellite and ground-based rainfall observations at a spatial resolution of 0.05$^\circ$ (~5 km) [33]. These data were downloaded for the period 2005 to 2020 from Google earth
engine [31]. The temperature data on the other hand were based on version 6 of the MODIS land surface temperature observations, which are also available at a spatial resolution of 0.05° (~5 km) [34]. These data were downloaded from NASA’s data handle, GIOVANNI.

2.3. Methods

In this study, temporally averaged maps for each species were compared to each other for the two instruments used per species. On the other hand, the spatially averaged timeseries data were analysed using Pearson correlation to show the association in the instrument records. This was discussed along with the seasonal variations. Before investigating the trends, the data were decomposed in order to isolate the de-seasonalized component from the seasonal and residual components. The trends were then determined by applying the non-parametric sequential Mann-Kendall test to the de-seasonalized timeseries of the species. This test has been adequately described by Shikwambana et al. [35]. In this study, it was used to reveal the direction, strength and significance of the trends at a 95% confidence level.

3. Results and Discussion

3.1. Spatial Distribution of NO2, SO2 and CO and Comparison of Satellite Data Records

Figure 2 shows the atmospheric abundances of NO2, SO2 and CO averaged by time, along with the correlation coefficients calculated using records of the two instruments used per gas species. Data from matching time periods were used in this analysis. For NO2 this was 1 July 2018 to 31 December 2020, for SO2 this was 7 December 2018 to 31 December 2020 and for CO it was 1 July 2018 to 31 December 2020. It is evident that each of the two satellite instruments used per species observed a similar spatial distribution pattern and the major difference was in the observed column amounts. TROPOMI’s higher resolution offers an added advantage of isolating the emission hotspots.

For NO2, both instruments show that the concentrations were higher in Uganda and Tanzania compared to the other countries. More specifically, the higher NO2 amounts were in the areas of northern Uganda and western and southern Tanzania. OMI observed between $0.6 \times 10^{15}$ and $1.5 \times 10^{15}$ molecules/cm$^2$, while TROPOMI observed between $2.5 \times 10^{15}$ and $3.3 \times 10^{15}$ molecules/cm$^2$. This NO2 distribution is most likely associated with the savanna fires at the surface as the distribution matches the location of the savanna woodlands (Figure 1). TROPOMI also observed NO2 concentrated within the capital cities such as Kampala (Uganda), Nairobi (Kenya), and Bujumbura (Burundi), and towns like Dar es Salaam (Tanzania) and Mombasa (Kenya). The amounts were between $2.8 \times 10^{15}$ and $3.7 \times 10^{15}$ molecules/cm$^2$. A similar pattern was observed outside the region, in cities like Juba (South Sudan) and Kasama (Zambia). This is due to the concentration of fossil fuel combustion activities within these urban locations. Overall, regardless of the matching spatial distribution observed by both instruments, TROPOMI measures higher NO2 vertical column densities (VCD) than OMI, making the correlation, $R = 0.54$. This suggests that peak values are not occurring on the same days in the two datasets.

The concentration of SO2 was very low over most of the region, typically below 0.15 DU and 0 DU in OMI and TROPOMI measurements, respectively (Figure 2). The negative values denote clear sky measurements in areas with low SO2 amounts. The only outstanding emission source for SO2 was near Goma (Democratic Republic of Congo), a town neighbouring Rwanda in the north (Figure 1). This is primarily because of the existence of an active volcano at Mt. Nyiragongo which is about 15 km from Goma. It is an open-vent volcano with a lava lake at the top and it continuously emits SO2 [36,37]. Overall, TROPOMI observed higher SO2 column amounts than OMI.

Since TROPOMI makes CO measurements as totals of the entire atmospheric column, the AIRS CO data were summed up for 22 atmospheric levels, from 850 hPa at the bottom.
to 1 hPa at the top of the atmosphere. The summed data were then compared with TROPOMI. The instruments records showed a strong correlation; $R = 0.77$. The spatial distribution showed that CO is most concentrated on the western side of the region and least concentrated on the eastern side. Uganda, Rwanda, Burundi and Tanzania are the largest contributors of CO in the region. This pattern matches the location of the savanna woodlands (Figure 1) which implies that the savannah fires are the most important emission source of CO over the region. The results also suggest that NO$_2$ and CO are emitted from the same combustion processes of the savanna fires.
3.2. Seasonal Variation

Figure 3 shows the seasonal variation of the NO$_2$, SO$_2$, and CO, along with rainfall and temperature. These were plotted using the entire 16-year data record for OMI and AIRS and a 2-year data record for TROPOMI. The variations are based on spatial averages over the entire region, and since some parts of the East African region have very low abundances of these species, the spatial averages tend to be lower than temporal averages on which the maps in Figure 2 are based.

The abundances of NO$_2$ and CO are highest during the months of July to October and December to February, which coincides with the fire seasons over the African continent [5]. Their variation also shows a higher sensitivity to the meteorological parameters than is seen in SO$_2$. NO$_2$ and CO show an inverse response to rainfall. The abundances of these species are lowest during the first rain season, March to May. During that season, NO$_2$ does not exceed $5 \times 10^{14}$ molecules/cm$^2$, while CO is below 1150 ppbv (OMI) or $2.8 \times 10^{17}$ molecules/cm$^2$ (TROPOMI). During the dry season, June to September, the abundances of these species are higher. NO$_2$ reaches peak concentrations of about $11 \times 10^{14}$ molecules/cm$^2$ while CO reaches to about 1238 ppbv (OMI) or $3.06 \times 10^{17}$ molecules/cm$^2$ (TROPOMI). After this period, their abundances start declining as the second rain season sets in from October to December. This inverse association between these chemistry species and rainfall is because the prolonged occurrence of rainfall events during its peak seasons creates conditions that do not allow ample time for the woodlands and grasslands to dry-up enough, to the level that they can ignite [38].

Furthermore, NO$_2$ and CO have a trajectory that matches that of temperature. The high abundances occur during the months with higher temperatures, January to March and August to November, when temperatures are typically above 34 °C. On the other hand, low abundances occur during months with lower temperatures, that is, April to July, when temperatures are below 32.5 °C. This association with low temperatures is, however, more pronounced for CO than it is for NO$_2$.

SO$_2$ shows some variation over different months of the year. The OMI observations show the amounts to be between ~0.02 and 0.03 DU, while the TROPOMI observations show slightly higher amounts, between ~0.01 and 0.07 DU.
3.3. Long-Term Trend of NO$_2$, SO$_2$ and CO

Figure 4 shows the records of the three chemistry species based on observations from OMI for NO$_2$ and SO$_2$ and AIRS for CO. TROPOMI data were excluded because of their short temporal coverage, which were not sufficient to infer long-term trends. Since the daily data were too noisy, monthly averages were also plotted to give a clearer direction of the trend. For NO$_2$, the daily average was between a minimum of $\sim 2.3 \times 10^{14}$ molecules/cm$^2$ and a maximum of $22.7 \times 10^{14}$ molecules/cm$^2$, while the monthly average was between $\sim 4 \times 10^{14}$ and $10 \times 10^{14}$ molecules/cm$^2$. NO$_2$ peaks twice in each year, that is, August to September and in December. The latter peak extends into January of the following year.

For SO$_2$, the daily average for the majority of the period was up to a maximum of $\sim 0.2$ DU, however there were a few extreme events that ranged between $\sim 0.22$ DU and 0.55 DU. Coincidentally, some of them occurred within the months with overall higher SO$_2$ amounts that exceeded a monthly average 0.05 DU. These months included June and November of 2011 and February, June and July of 2014, and they caused notable spikes in the monthly timeseries (Figure 4). The overall monthly SO$_2$ average was between $\sim 0.01$ and 0.08 DU.

For CO, the daily average was between a minimum of 953 ppbv and a maximum of 1839 ppbv, while the monthly average was between 1064 ppbv and 1335 ppbv. CO also peaks twice a year, that is, January to February and September to October. Importantly, the peak seasons of NO$_2$ and CO overlap in September and January, both of which are fire seasons in the tropical savanna [4].

Figure 3. Seasonal variation of NO$_2$ (a), SO$_2$ (b), CO (c,d) along with rainfall (e) and temperature (f). The vertical bars show the monthly standard deviations.
Figure 5 shows the decomposed monthly timeseries for each of the species. For NO\(_2\), the two-peak pattern is re-echoed to be August to September and in December. The August peak had more than twice the gas abundance of the December peak. The de-seasonalized series shows that the three outstanding periods of NO\(_2\) abundance occur from June to October of 2010, March to December of 2012 and June to August of 2015. For SO\(_2\), the most outstanding period was from April to July of 2014 followed by November 2011 and May 2012. For CO, the two-peak pattern is clarified to occur in September–October and January to February, with the first peak having ~ 60 ppbv more CO than the second peak. The most outstanding period with high CO abundance was November 2015 to April 2016. Importantly, the most prominent peaks in NO\(_2\) and CO simultaneously occur in September, suggesting that it could be the most intense month for the savanna fires.

Table 1 is a summary of the results of the sequential Mann-Kendall test applied to the de-seasonalized timeseries of each species. The results show that there was no trend in NO\(_2\) (\(p > 0.05\)), while CO and SO\(_2\) had decreasing trends (\(p < 0.05\)). Since Mt. Nyiragongo is the dominant SO\(_2\) hotspot in the region, the overall decreasing trend is most likely associated with the processes going on within it. Arellano et al. [36] reported a decreasing trend in SO\(_2\) at this volcano and attributed it to degassed magma sinking down into the volcano’s conduit.
Figure 5. Decomposed time series of \(\text{NO}_2\) (\(\times 10^{14}\) molecules/cm\(^2\)), \(\text{SO}_2\) (DU) and CO (ppbv). All units labelled (a) are the monthly series, (b) are the de-seasonalized components, (c) are the seasonal components and (d) are the residual components.
A further analysis of the CO vertical profile revealed significant trends at specific heights (Figure 6 and Table 2). For levels closer to the surface, that is, 850 hPa to 500 hPa, which had the highest CO abundances, there was a decreasing trend ($p < 0.05$) which has been illustrated with the Mann-Kendall statistics for the 850 hPa level (Figure 6a and Table 2). For levels starting at 400 hPa to 70 hPa and those from 50 hPa to 1 hPa, there was an increasing trend ($p < 0.05$), and this has been illustrated using the Mann-Kendall statistics for the 150 hPa and 3 hPa levels, respectively (Figure 6b, c and Table 2). This trend is however stronger for the levels, 400 hPa to 70 hPa than it is for 50 hPa to 1 hPa. These CO trends could be due to the vertical uplift of CO from the boundary layer into the free atmosphere by the action of deep convective updrafts [39,40]. Figure 6d shows the average abundance of CO for the 22 atmospheric levels. The highest CO amounts are found closer to the surface because that is the most important location for biomass burning events.

**Figure 6.** CO trend for 22 pressure levels in the atmosphere (a–c) along with the CO vertical profile (d) for the period 2005 to 2020.
Table 2. A summary of the Mann-Kendall test statistic for CO at heights of 850 hPa, 150 hPa and 3 hPa.

|       | 850 hPa | 150 hPa | 3 hPa |
|-------|---------|---------|-------|
| p-value | 0.0     | 0.0     | 0.0   |
| Z-score | -13.3947 | 14.6096 | 14.8224 |
| S      | -10,828 | 11,810  | 11,982 |
| Tau    | -0.6721 | 0.7330  | 0.7437 |
| Slope  | -0.0737 | 0.0236  | 0.0028 |
| Intercept | 112.8257 | 56.9255 | 27.1086 |

3.4. Analysis of SO\(_2\) Extreme Years

For the SO\(_2\) timeseries which exhibited a significant trend (Table 1), further analysis was carried out considering the five months with the highest monthly average SO\(_2\) amounts. These were June and November of 2011 and February, June and July of 2014, which all had SO\(_2\) monthly averages exceeding 0.05 DU (Figure 5). Coincidentally, these months were within the years with the highest SO\(_2\) amounts, that is, 2011 and 2014. Since the beginning of the OMI observation campaign, in the East African region, these two years were recorded to have the highest SO\(_2\) amounts (Figure 7).

![Figure 7](image_url)

Figure 7. Relative change in SO\(_2\) over East Africa, normalized based on the year 2005.

The spatial distribution of SO\(_2\) during these months is presented in Figure 8 and it shows that the active volcano, Mt. Nyiragongo located near the border town of Goma stands out as an SO\(_2\) hotspot during all the months, but most especially in June 2014 when the SO\(_2\) amount exceeded 10 DU. The seasonal variation of SO\(_2\) emitted from this volcano (Figure 9) shows that the peak emissions occur during the months of May to August. Importantly, this volcano emits much higher SO\(_2\) amounts whenever it erupts, and such was the case in the most recent eruption that occurred on 22nd May 2021 [41]. Figure 8 shows that the region closest to the volcano had SO\(_2\) amounts between ~8 and 15 DU and the region of north western Tanzania had SO\(_2\) amounts of up to ~6 DU.

In June 2011, the regions of western and central Kenya also added to the observed SO\(_2\) amounts by ~1 to 5 DU. The underlying reason is not yet clear, however, since no such distribution is observed in the other months, then this could be due to a one-off SO\(_2\) related emission event that occurred during that month within this region. For example, fires set off by fuel leakages along the oil pipelines from Eldoret and Kisumu through Nairobi can emit SO\(_2\) due to combustion of the sulphur-rich fuel. Such an event occurred in 2011 [42]. In June 2014, Bujumbura, the capital city of Burundi also added to the observed SO\(_2\) amounts over the region by ~1.5 to 4.5 DU.
Figure 8. Spatial distribution of boundary layer SO\textsubscript{2} (in Dobson Units; DU) for the months with the highest monthly average SO\textsubscript{2} amounts, along with that observed by TROPOMI during the eruption of May 2021. The white areas are missing data.

Figure 9. Seasonal variation of SO\textsubscript{2} emitted from Mt. Nyiragongo.

4. Conclusions

In this study, satellite data records were used to investigate the trend of NO\textsubscript{2}, SO\textsubscript{2} and CO over the East African region. Along with this, the study also made comparisons between the satellite records and investigated the association that these three species have with rainfall and temperature. The major conclusions are summarized below.

(1) The major source of NO\textsubscript{2} and CO over the East African region are the fires that occur in the savanna woodlands. The observed spatial distribution of these species perfectly matches the spatial extent of the savanna. Cities and towns are also important emission sources of NO\textsubscript{2} in East Africa and its neighbouring regions. These cities include Kampala, Nairobi, Bujumbura, and Juba, and the towns include Dar es Salaam, Mombasa and Kasama. Furthermore, the active volcano at Mt. Nyiragongo near Goma is the single most important source of SO\textsubscript{2} in East Africa.
(2) There was no trend in NO\(_2\) over the 16-year period investigated. Perhaps there has not yet occurred any significant shift in the savanna fires. In addition to this, though there has been an expansion in the cities within East Africa over the past two decades, it seems not to have translated into a significant increase in the NO\(_2\) abundance over the region. There was an overall decreasing trend in SO\(_2\) over the region which is attributed to internal events within Mt. Nyiragongo. Degassed magma is reported to be sinking further down into the volcano’s conduit [37]. CO exhibited a decreasing trend from 850 hPa to 500 hPa and an increasing trend for the levels above, that is, 400 hPa to 1 hPa. The increasing trend was stronger from 400 hPa to 70 hPa than from 50 hPa to 1 hPa. There is likely to be vertical ascent of CO from the boundary layer into the free atmosphere. This uplift has been seen in other regions such as east Asia [39] and India [40].

(3) NO\(_x\) and CO are suppressed during rainfall seasons because the consecutive rain-fall events do not allow enough time for the savanna woods and grass to dry up sufficiently for ignition to happen.

(4) Finally, despite the time difference in the technology, the satellite observations had consistent spatial agreement in the distribution of the gases. TROPOMI’s higher resolution, however, gives it an added advantage of isolating the emission hotspots, and this cements its position as one of the most important instruments for observing atmospheric chemistry, both now and in the coming future.

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**Data Availability Statement**: The data used in this study are freely available for download from the GIOVANNI data handle of NASA (https://giovanni.gsfc.nasa.gov/); accessed on 7 June 2021) and Google Earth Engine (https://earthengine.google.com/); accessed on 8 June 2021).

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