Measurement report: Combined use of MAX-DOAS and AERONET ground-based measurements in Montevideo, Uruguay, for the detection of distant biomass burning

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Abstract. Biomass burning releases large amounts of aerosols and chemical species into the atmosphere, representing a major source of air pollutants. Emissions and by-products can be transported over long distances, presenting challenges in quantification. This is mainly done using satellites, which offer global coverage and data acquisition for places that are difficult to access. In this study, ground-based observations are used to assess the abundance of trace gases and aerosols. On 24 November 2020, a significant increase in formaldehyde was observed with a Multi-AXis Differential Optical Absorption Spectroscopy (MAX-DOAS) instrument located in Montevideo (Uruguay), and its vertical column densities reached values of $2.4 \times 10^{16}$ molec. cm$^{-2}$, more than twice the values observed during the previous days. This was accompanied by an increase in the aerosol levels measured by an AErosol RObotic NETwork (AERONET) photometer located at the same site. The aerosol optical depth (AOD) at 440 nm reached values close to 1, an order of magnitude larger than typical values in Montevideo.

Our findings indicate that the increase was associated with the passage of a plume originating from distant biomass burning. This conclusion is supported by TROPOspheric Monitoring Instrument (TROPOMI) satellite observations as well as HYbrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) simulations. The profiles of the gases and aerosols retrieved from the MAX-DOAS observations are consistent with the HYSPLIT analysis, showing the passage of a plume over Montevideo on 24 November located at a height of $\sim 1.5$ km. This corroborates the finding that biomass burning events occurring about 800 km north of Montevideo can affect the local atmosphere through long-distance emissions transport. This study underscores the potential of ground-based atmospheric monitoring as a tool for detection of such events. Furthermore, it demonstrates greater sensitivity compared to satellite when it comes to detection of relatively small amounts of carbonyls like glyoxal and formaldehyde.
1 Introduction

Biomass burning (BB) events significantly impact air quality, both regionally and globally, due to the direct emissions of various substances into the atmosphere and the production of species during the subsequent long-distance transport. This issue and its impact on cities have been addressed in several studies using satellite observations, which offer the advantage of broad spatial coverage compared with ground-based measurements (Wittrock et al., 2006; Zarzana et al., 2017; Alvarado et al., 2020; Schutgens et al., 2021; Lerot et al., 2023).

In addition to producing aromatic hydrocarbons, nitrogen oxides (NOx), carboxyls, and organic carbon, fires are the main source of aerosols containing particulate matter (PM), generating peaks of very high PM10 concentrations during the dry seasons (de Oliveira Alves et al., 2015). Furthermore, the pollutants produced during these fires can be transported over long distances, affecting areas far away from their source (Hsu et al., 1996; Freitas et al., 2005; Ravindra et al., 2008).

Aerosols have adverse effects on human health in exposed populations (Chen et al., 2017; Contini et al., 2021), leading to respiratory diseases. Additionally, aerosols originating from BB contain soot, which absorbs solar radiation, thus altering the radiative budget of the atmosphere, cloud formation processes, and the albedo of snow- and ice-covered areas (Seinfeld and Pandis, 2006; Bond et al., 2013; Bellouin et al., 2020).

In South America, thousands of square kilometres of the Amazon is affected by burning to carry out agricultural, livestock, mining, and infrastructure projects each year. Uncontrolled fires of other types of vegetation also occur, for example, in Argentina, in the Paraná Delta, and in the border region between Argentina, Brazil, and Paraguay (Gassmann and Ulke, 2008).

Uruguay is located south of Brazil and east of Argentina, with its capital, Montevideo, positioned about 450 km from the Paraná Delta (Argentina) and approximately 800 km from the tri-border region between Argentina, Paraguay, and Brazil. Large fires occur frequently in both regions, affecting air quality in Montevideo. These fires can be identified in satellite images due to the presence of large amounts of smoke. Air quality in Uruguay is generally good (Ministerio de Ambiente, República Oriental del Uruguay, 2024; Intendencia de Montevideo, 2024); however, these fires induce significant short-term changes in the lower atmosphere, altering its composition.

Since 2020, a CIMEL sun photometer of the AERosol RObotic NETwork (AERONET, led by NASA – Goddard Space Flight Center) and a Multi-AXis Differential Optical Absorption Spectroscopy (MAX-DOAS) system of the Applied Optics Group of the Institute of Physics at the Engineering Faculty (UDELAR) have been operational to perform ground-based continuous monitoring of atmospheric constituents over Montevideo.

DOAS has become a widely used technique to study the atmosphere remotely, due to its versatility and adaptability, allowing a broad range of configurations (Platt and Stutz, 2008; Rivera et al., 2010; Ibrahim et al., 2010; Shaiganfar et al., 2017). Applying DOAS, the main sources of SO2 and NO2 emissions from industrial activities in Montevideo have been studied, such as the “La Teja” oil refinery (Frins et al., 2012; Osorio et al., 2017) and also the “Central Batlle y Ordóñez” power plant (Frins et al., 2014). Despite these local sources of emissions, the air quality in Montevideo remains at good levels, due to its location at the coast of the La Plata River and the prevailing winds. Although the combination of ground-based instruments such as DOAS with AERONET is widely used in the Northern Hemisphere for detecting and analysing emissions from biomass burning events, its application is quite rare in the Southern Hemisphere, especially in South America.

The objective of this study is to explore the use of ground-based observations for the detection of aerosols and trace gases produced by distant BB emissions transported over long distances to Montevideo. In particular, we investigated an event that occurred in November 2020 on the border between Paraguay and Argentina, situated approximately 800 km north of Montevideo. This relatively strong event resulted in the detection of a plume passing over Montevideo, by means of multiple instruments covering ground-based and satellite observations.

This article is organized as follows: in Sect. 2 we describe the methods used for monitoring the event and the additional tools employed to support our conclusions, in Sect. 3 we present the DOAS observations and additional results used for detecting the event in November 2020, in Sect. 4 we present a discussion of our findings, and in Sect. 5 we provide some general remarks and the conclusions from this work.

2 Instruments and methods

2.1 Site description

In this study, a MAX-DOAS system and a sun photometer from the AERONET programme were used for remote sensing of the atmosphere. Both instruments are installed on the roof of the Faculty of Engineering building located in Montevideo, Uruguay (34.9175° S, 56.1669° W). The MAX-DOAS system has been providing data since November 2020, and the sun photometer since January 2020. Figure 1a indicates the location of the instruments and the MAX-DOAS observation directions on the map. Figure 1b shows a picture of both instruments on the roof of the building. The main characteristics of each instrument are described below.
2.2 MAX-DOAS instrument

A MAX-DOAS instrument manufactured by AirYX GmbH operates continuously and acquires diffuse solar radiation spectra at several elevation angles in a vertical plane from east to west. The spectra are acquired by a thermo-controlled Avantes spectrometer that operates in the spectral range of 301–463 nm and has a spectral resolution of approximately 0.6 nm. The total field of view of the system is 0.3° in the vertical direction and 1° in the horizontal. The MAX-DOAS instrument includes a mercury lamp for wavelength calibration. Dark-current and offset measurements are acquired at night to correct the spectra. The temperature of the spectrometer was set to 20 °C.

For this study we considered elevation angles of 1, 2, 3, 5, 10, 20, 40, and 90° for both western and eastern viewing directions of observation (see Fig. 1a). The acquisition time of each spectrum was 1 min, and the duration of each vertical plane scanning sequence was approximately 20 min.

2.3 AERONET sun photometer

The AERONET node at Montevideo uses a CIMEL CE3128-T multispectral photometer (Holben et al., 1998) installed adjacent to the MAX-DOAS instrument. This photometer measures direct and diffuse solar radiation at different bands ranging from 340 to 1640 nm to retrieve aerosol data from the observations. AERONET provides several data products derived from these measurements, for example, aerosol optical depth (AOD), single scattering albedo (SSA), and phase function (Holben et al., 2001; Dubovik et al., 2000; Dubovik and King, 2000). In this study we used AERONET V3 level 2.0 data, ensuring good quality of data by applying radiometric and instrumental corrections, as well as cloud-cover removal algorithms (Sinyuk et al., 2020).

2.4 Satellite observations

The TROPOspheric Monitoring Instrument (TROPOMI), carried by the Copernicus Sentinel-5 Precursor satellite, was successfully launched on 13 October 2017. It operates within the ultraviolet (UV), visible (VIS), near-infrared (NIR), and short-wave infrared (SWIR) spectral range, covering wavelengths from 270 to 500 nm in the UV–VIS, from 675 to 775 nm in the NIR, and from 2305 to 2385 nm in the SWIR band.

TROPOMI provides near-global daily coverage, achieving a spatial resolution of 3.5 km × 5.5 km (7 km × 7 km in the SWIR) since August 2018. It crosses the Equator at 13:30 local time (ascending node). Similar to the Ozone Monitoring Instrument (OMI), TROPOMI uses a nadir-viewing imaging spectograph with a two-dimensional charged-couple device (CCD). One dimension captures spectral information, while the other dimension covers the spatial information. The TROPOMI instrument has been delivering data since November 2017 (Veefkind et al., 2012). Its spectral bands enable the observation of relevant atmospheric species such as CHOCHO, HCHO, NO₂, and CO.

We employed TROPOMI observations of NO₂ and CO in our analysis to detect long distance transport events. We also used UV Aerosol Index (UVAI) data as a reference for the presence of aerosols in the plume, which likely coincides with the location of the trace gases (see for instance Torres et al., 2020; Alvarado et al., 2020). These products have been operational since March 2018, and they include the reprocessed (RPRO) and offline-mode (OFFL) data versions OFFL V01.03.02, RPRO V02.04.00, and OFFL V01.03.02, respectively. For the retrieval of formaldehyde (HCHO) and glyoxal (CHOCHO), volatile organic compounds (VOCs), the products considered in this study are based on Alvarado et al. (2020). Glyoxal has been retrieved in the wavelength range from 433 to 465 nm, which is slightly larger than windows used in previous investigations (Vrekoussis et al., 2010; Alvarado et al., 2014). HCHO is retrieved in the fitting window from 323.5 to 361 nm, which results in a reduction of the noise of slant column density. The main interfering absorbers are included in the respective wavelength ranges as described in Alvarado et al. (2020). To compute vertical column densities (VCDs), air mass factors based on trace gas profiles simulated with the Tracer Model 5 – Massively Parallel version (TM5-MP) global chemistry transport model are applied to the retrieved slant column density (SCD) data (Myriokefalitakis et al., 2020). For an individual CHOCHO measurement, the detection limit is of the order of 5 × 10¹⁴ molec. cm⁻². For HCHO, the detection limit is of the order of 4.5 × 10¹⁴ molec. cm⁻².

2.5 HYSPLIT model

The Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model, developed by the National Oceanic and Atmospheric Administration (NOAA), is used to track emissions of gases and aerosols into the atmosphere. It is used to model trajectories of air mass parcels, as well as to simulate transport, dispersion, chemical processes, and deposition (Stein et al., 2015). HYSPLIT uses a stratified atmosphere and relies on three-dimensional weather data, such as wind speed and direction, temperature, and humidity. These weather data can be obtained from ground-based weather stations, weather balloons, satellites, or numerical weather models.

In this study, we used HYSPLIT to simulate the trajectories of air parcel originating from distant biomass burning sites. This approach allowed the assessment of long-distance plume transport (see Sect. 3.2).

2.6 BOREAS

The Bremen Optimal estimation REtrieval for Aerosols and trace gases (BOREAS; Bösch et al., 2018) is an inversion algorithm designed for retrieving vertical profiles of trace
gases and aerosol from MAX-DOAS measurements of differential slant column densities (dSCDs). This algorithm was developed at the Institute for Environmental Physics in Bremen, Germany (Bösch et al., 2018), and has been validated in various studies (e.g. Frieß et al., 2019; Tirpitz et al., 2021). Each input vector includes dSCD measurements taken at different lines of sight for either the trace gas of interest or the oxygen collision complex $\text{O}_4$. The latter has a well-known vertical profile in the atmosphere, and its slant columns can therefore be used to retrieve the vertical distribution of aerosols.

In this study, aerosols have been retrieved from the $\text{O}_4$ absorption band at 360 nm. Using the mean Ångström exponent 440–870 nm from AERONET Inversion data (Version 3, Almucantar, Level 1.5), the retrieval results have been extrapolated to the matching spectral windows of HCHO and CHOCHO, as the AERONET station only retrieves data at 440 nm or higher wavelengths. AERONET Inversion data were also used for the aerosol parametrization within BOREAS. The AERONET 440 nm single scattering albedo (SSA) and phase function obtained at the time closest to the corresponding slant column measurement scan have been used within the profile inversion (440 nm products are the closest available to 360 nm wavelength products).

Atmospheric profiles over Montevideo were also simulated using the Community Atmosphere Model – Chemistry model (Buchholz et al., 2019; Emmons et al., 2020) and served as a priori meteorological data sets within BOREAS. Each trace gas profile was retrieved by applying a pre-scaled exponential a priori profile (Bösch et al., 2018) with an exponential scale height of 1 km for all species. In general, the sensitivity of MAX-DOAS profiling algorithms is highest for the lowest retrieval layers and decreases strongly for altitudes above 2–3 km. However, if an elevated trace gas layer is dominant – with no significant trace gas concentration present below the elevated layer – the retrieval is possible, but the vertical extent of the layer might be retrieved with slightly decreased accuracy (Bösch, 2019; Tirpitz et al., 2021).

3 Detection of a distant biomass burning event

3.1 Signals of the biomass burning event

AERONET data provided early indications of the presence of a biomass burning plume over Montevideo in November of 2020. Aerosol optical depth (AOD) values usually range around $0.09 \pm 0.03$ in the 440 nm band when measured at this station. However, on 24 November, AOD values reached up to 0.96 from the photometer measurements, indicating the presence of large amounts of aerosols in the atmosphere.

Figure 2a shows AOD values for a period around 24 November at wavelengths 380, 440, and 500 nm, and Fig. 2b shows the contribution of the fine and coarse modes to the AOD at 440 nm (O’Neill et al., 2003). The prevalence of fine particles (those with radius smaller than 1 µm) is interpreted as an indicator of a biomass burning plume passing above Montevideo (Agesta, 2023). Notably, the AOD due to fine particles increased on 24 November with respect to the average of the period ($0.46 \pm 0.06$ versus $0.03 \pm 0.01$). This increase is a characteristic of biomass burning smoke (Shi et al., 2019).

Analysing RGB satellite images from the VIIRS instrument (NOAA-20) for dates around 24 November, the presence of an extensive plume over a large region in the central-eastern Uruguayan territory was detected. The origin of this plume can be attributed to several distant emission sources primarily related to biomass burning, as shown in Fig. 3. Some fire spots appear near the border between Paraguay and Argentina, approximately 800 km north of Montevideo,
around 22 November. The number of foci observed increased towards 23 November, and a distinguishable plume was transported southwards over Uruguay on 24 November. As a proxy for identifying fire location and fire intensity, the Fire Radiative Power (FRP) data product from Copernicus Atmosphere Service Information is superimposed in Fig. 3, indicating the locations of the fires (Kaiser et al., 2012).

3.2 Plume transport simulations with the HYSPLIT model

To investigate possible transport pathways, several simulations were conducted using the HYSPLIT trajectory model around the event date to track the transport of air mass parcels from the fires and verify if the emitted plume was expected to appear over Montevideo. We estimated the location and height of the main fire spots appearing around 23 November at 17:00 UTC using satellite data (see Fig. 3).

The trajectory simulations start on 23 November at 17:00 UTC from a total of nine points, using wind field data from the Global Forecast System at 0.25° horizontal resolution to drive the HYSPLIT model, which also provides the boundary layer height. Data from the Weather Research and Forecasting model were selected as input for the simulation. This also defines the height interpolation, with a maximum height set to 15 km. Figure 4 shows an example of these simulations, for which the height of each source was set at 100 m above ground level. Additional initial conditions were investigated with starting heights set to 500 and 1000 m altitude to ensure that the air parcels appear well within the boundary layer, allowing them to be transported hundreds of kilometres. The results are similar, and, in all cases, air parcels from the areas surrounding the fires reached Montevideo at

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**Figure 2.** AOD detected at the AERONET station in Montevideo for the period 18 to 30 November 2020. Panel (a) shows the AOD for the wavelength bands of 380, 440, and 500 nm. Panel (b) shows the AOD at 500 nm for fine and coarse particles separately.
Figure 3. RGB satellite images from the VIIRS instrument (NOAA-20) for (a) 22 November, (b) 23 November, and (c) 24 November 2020. The corresponding Fire Radiative Power (FRP) data product was overlaid to visualize the location of the principal burning sources.

Figure 4. HYSPLIT-simulated trajectories for fire sources located 800 km north of Montevideo detected on 23 November. The line markers indicate 3 h intervals. Several trajectories are passing near Montevideo around the time of the measurements (labels e, f, and g).

the time corresponding to the detection through our observations.

These trajectories indicate that a portion of the emitted plume is transported from north to south and then towards east of Montevideo on 24 November (trajectories e, f, and g in Fig. 4), in agreement with the rest of the observations for the same date. Other parts of the plume were transported towards the west of Uruguay, at distances exceeding 100 km from Montevideo (trajectories a–d, h, and i in Fig. 4).

The modelled plume remained below an altitude of 2500 m with respect to sea level (bottom panel in Fig. 4). Here, we assume that aerosols and the principal derived products of the biomass burning are transported at the same altitude.

3.3 DOAS retrieval

The principles of the DOAS analysis are described for example in Höninger et al. (2004) and in Platt and Stutz (2008). We performed the DOAS analysis of the spectra recorded by the MAX-DOAS instrument, with primary focus on retrieving the dSCD of trace gases commonly associated with wildfire events, such as formaldehyde (HCHO), glyoxal (CHOCHO), and nitrogen dioxide (NO₂). Additionally, the analysis addressed the dSCD of the oxygen dimer O₄, which is used to quantify the presence of aerosols in the atmosphere (Wagner et al., 2004; Frieß et al., 2006). The evaluations were performed using the QDOAS software (Danckaert et al., 2017). Table 1 summarizes the settings used for the retrievals of each trace gas of interest in three fitting intervals: 333 to 359 nm for HCHO, 352 to 387 nm for O₄, and 432.5 to 459 nm for CHOCHO and NO₂. The high-resolution absorp-
tion cross sections required for the trace gas retrieval were taken from the MPI-Mainz UV/VIS Spectral Atlas (Keller-Rudek et al., 2013). The reference spectra used in the retrievals were the zenith measurements closest in time to each scanning sequence. This choice is intended to enhance the sensitivity towards tropospheric absorptions. In the DOAS retrieval, a synthetic Ring spectrum calculated with QDOAS was also included (Grainger and Ring, 1962; Chance and Spurr, 1997). A polynomial of degree 5 was used to fit broadband structures, along with allowance for a second-order shift and stretch of the wavelengths of the spectra.

Figure 5 shows an example of the DOAS analysis for each fitting interval, performed on a spectrum acquired on 24 November 2020 at 13:04 local time (LT). The first row shows the fitting of the target gases, HCHO, O₂, and CHOCHO. Other trace gases exhibiting absorption in the same spectral interval, as well as the synthetic Ring spectrum, are shown in the second and third rows. The residual of the analysis is shown in the last row.

In this example, the DOAS fits reveal a strong formaldehyde signal, in contrast to what usually appears in Montevideo. In addition, glyoxal was also observed, which had never been detected previously in the same location. A strong signal was also observed for NO₂. However, this cannot necessarily be attributed to the influence of the event over Montevideo. A detailed discussion of this point is presented in Sect. 4.

Cloudy days were screened out from our spectral data set using an algorithm based on the diurnal cycle of the colour index given by the ratio between intensities at 370 and 440 nm (Gielen et al., 2014; Wagner et al., 2014). This ratio is strongly affected by cloudy conditions, and its variations provide a way to identify clear days. The days of 25 and 28 November were tagged as completely cloudy and removed from the analysis as a result of this first-order data screening. We used data from weather stations to manually confirm this result.

4 Results and discussion

4.1 MAX-DOAS

The results of the DOAS retrievals for the period around the detection of the event are shown in Fig. 6 for the trace gases HCHO, O₂, CHOCHO, and NO₂ (first to fourth rows, respectively). The coloured circles correspond to the dSCD values for elevation angles of 5° (red), 10° (green), and 20° (blue) in the east and west observation directions. The O₂ dSCDs obtained in the period from 20 to 28 November have a behaviour typical of days with clear skies and low aerosol load, except for 24 November. On this day, O₂ results are unusually low, showing approximately the same values at all elevation angles. This can be interpreted as the result of multiple scattering in the atmosphere, which alters light path lengths and indicates a high aerosol load (Wagner et al., 2004; Sinreich et al., 2013).

The main interest of our DOAS analysis is to assess simple carbonyls such as formaldehyde and glyoxal that will be used as plume tracers from biomass burning events (Hoque et al., 2018; Kluge et al., 2020). Formaldehyde dSCDs showed a clear increase on 24 November compared to earlier dates. Following 24 and 25 November (the latter was a cloudy day), HCHO levels returned to the values observed before the arrival of air masses from biomass burning. A difference is also observed between the dSCDs obtained from the spectra acquired in the east and west directions. This may arise because the measurements obtained in the westerly direction may be influenced by industrial and port activities around Montevideo Bay, while eastbound observations were only affected by city activity, primarily traffic.

A slight increase in retrieved glyoxal dSCDs is observed on 24 November, although it is not as pronounced as for formaldehyde. The difference of this signal compared to the background is more noticeable for the spectra obtained looking in the easterly viewing direction. NO₂ dSCDs show a more regular behaviour throughout the study period in the westerly viewing direction but have some unusually large values at 20° elevation on 24 November. When analysing the spectra obtained from observations in the easterly viewing direction, the effect is larger, and again the 20° measurements are unusually high. The DOAS retrieval uncertainties range from 3% to 11% for HCHO and from 6% to 17% for glyoxal on 24 November. On other days, these uncertainties vary between 15% and 50% respectively, as tropospheric columns are much smaller.

The detection limit for HCHO and glyoxal columns was estimated to be 4.0 × 10^{15} and 5.6 × 10^{14} molec. cm⁻², respectively (Platt and Stutz, 2008).

4.2 Vertical column densities and profiles

We used the dSCD to derive the VCD with the BOREAS profile retrieval algorithm (Bösch et al., 2018). The VCDs are related to the dSCD via the air mass factor (AMF), which depends on the local time, surface albedo, aerosol profile, trace-gas profiles, and observation geometry (Sect. 2.6). The BOREAS output was screened to keep points with solar zenith angles less than 85° and with relative azimuth angles greater than 20°.

Figure 7 shows the retrieved VCD of the trace gases and AOD considered in this study for the given period. We calculated percentile values for the corresponding TROPOMI-S5P products in a circular region with a radius of 25 km centred on Montevideo. These are shown as horizontal lines in panels (a), (b), and (c). AERONET values for the same period are also displayed in panel (d). Figure 8a shows the AOD on 24 November, comparing the BOREAS retrieval with AERONET values (at 360 and 380 nm, respectively), and Fig. 8b shows the scatter plot and a linear fit between

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Figure 5. Example of DOAS fits for a spectrum measured during the event on 24 November at 13:04 LT, recorded at 5° elevation angle pointing towards west. Blue lines are the optical densities associated with each trace gas, and the orange lines are the fitted differential absorption cross sections. The retrieved dSCDs and their fitting errors are displayed in molec. cm$^{-2}$ for HCHO, CHOCHO, NO$_2$, and H$_2$O and in molec.$^2$ cm$^{-5}$ for O$_4$.

AERONET and BOREAS AOD values. The following remarks can be made regarding these results:

- The vertical column densities obtained from the BOREAS retrieval show a pronounced increase for formaldehyde and glyoxal on 24 November compared to the values of the other days, which is consistent with the passage of the plume over Uruguay found in satellite images (see Figs. 2 and 3). The variability in the median values of TROPOMI-S5P CHOCHO, HCHO, and NO$_2$ generally aligns with the corresponding MAX-DOAS measurements for most days compared. However, we note that there is a considerable variability in CHOCHO values – similar to MAX-DOAS observations – likely arising from the inherent difficulties in its retrieval.

- On this particular day, the AOD values derived from BOREAS are also above those from the days before and after 24 November, qualitatively following the behaviour of the AERONET values (see Fig. 7d). However, the latter are approximately 30% higher, as shown in Fig. 8a and b. This discrepancy could be attributed to one of the following reasons: (1) an inhomogeneous spatial distribution of the aerosols within the plume, potentially causing a mismatch between MAX-DOAS and AERONET measurements, which are taken in different directions, or (2) the use of an a priori profile during the inversion procedure that does not completely match the atmospheric conditions for 24 November. Previous studies have suggested that scaling factors for the O$_4$ dSCD are sometimes needed to solve this issue.
Figure 6. Differential slant column densities for elevation angles 5° (red), 10° (green), and 20° (blue), for the trace gases HCHO, O₄, CHOCHO, and NO₂ (first to fourth row, respectively) for the study period. Results for east and west directions are separated into the left and right panels, respectively. An increase in the dSCD values of HCHO is observed on 24 November. The O₄ dSCD values observed for that date are lower and coalesce for the different elevation angles, which indicates a high aerosol load. The glyoxal signal also increases on 24 November although not as clearly as that of formaldehyde. The observed changes in NO₂ dSCD values for that day with respect to the background are relatively small and may not be attributed to the event.
Table 1. Retrieval settings and cross sections used for each DOAS analysis.

<table>
<thead>
<tr>
<th>Target trace gas</th>
<th>Fitting interval</th>
<th>Fitted absorber</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCHO (294 K)</td>
<td>333.0–359.0 nm</td>
<td>HCHO</td>
<td>Meller and Moortgat (2000)</td>
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<td></td>
<td></td>
<td>NO2 (294 K)</td>
<td>Vandaele et al. (1998)</td>
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<tr>
<td></td>
<td></td>
<td>O3 (223 K, 243 K)</td>
<td>Serdyuchenko et al. (2014)</td>
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<tr>
<td></td>
<td></td>
<td>O4</td>
<td>Thalman and Volkamer (2013)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>BrO</td>
<td>Fleischmann et al. (2004)</td>
</tr>
<tr>
<td>O4 (294 K)</td>
<td>352.0–387.0 nm</td>
<td>NO2 (294 K)</td>
<td>Vandaele et al. (1998)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>O3 (223 K)</td>
<td>Serdyuchenko et al. (2014)</td>
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<td></td>
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<td>O4</td>
<td>Thalman and Volkamer (2013)</td>
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<td></td>
<td></td>
<td>BrO</td>
<td>Fleischmann et al. (2004)</td>
</tr>
<tr>
<td>CHOCHO and NO2</td>
<td>432.5–459.0 nm</td>
<td>NO2 (220 K, 294 K)</td>
<td>Vandaele et al. (1998)</td>
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<td></td>
<td></td>
<td>CHOCHO</td>
<td>Rothman et al. (2013)</td>
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<td></td>
<td></td>
<td>O4</td>
<td>Thalman and Volkamer (2013)</td>
</tr>
</tbody>
</table>

(Wang et al., 2016; Wagner et al., 2019). In Tirpitz et al. (2021), a similar behaviour was detected during the second Cabauw Intercomparison campaign of Nitrogen Dioxide measuring Instruments (CINDI). However, the atmospheric conditions there were rather different from the ones present over Montevideo.

In this study, only aerosols up to 4 km above mean sea level were considered in the BOREAS inversion model. It is possible that aerosols above the highest retrieval altitude or in altitudes with lower sensitivity were retrieved by the AERONET station but could not be retrieved by BOREAS. Wang et al. (2016) report a similar behaviour in their study.

On 24 November, the VCD values of formaldehyde increased by approximately a factor of 4 in comparison to the previous days, further supporting the conclusion that part of a plume, caused by a biomass burning event, was detected over Montevideo. No unusual activities were identified during that week in Montevideo. Atypical NO2 values are not apparent for that day, compared to previous days. However, there is a good agreement when comparing VCD values derived from both viewing directions, which is unusual because of the difference in NO4 emitters in the two directions. This could be explained by the fact that the transported plume covered all of Montevideo on that day. For days prior to the detection, the NO2 diurnal cycle can be seen with its characteristic peak near noon, and the signal is stronger in the values computed from the west viewing direction, coinciding with the main industrial activity of the city such as the harbour and the oil refinery.

Glyoxal VCD values show an increase on 24 November. This suggests the transport of glyoxal by a plume of distant origin. However, caution must be taken because the DOAS fitting error for glyoxal is relatively high for the period under study. The reason is that the CHOCHO dSCD is close to the detection limit, since Montevideo does not have strong sources of glyoxal. Hence, the majority of the glyoxal signal would likely originate from the transported plume.

Figure 9 shows examples of the retrieved gas and aerosol profiles for 22 November (top row), as reference, and for 24 November (bottom row).

The profiles show a change in the concentration and aerosol extinction during later hours of 24 November. Glyoxal and formaldehyde show maxima in their profiles at approximately the same altitude, ranging between 1 and 2 km, which is consistent with the results obtained by the HYSLIT model and supports our initial hypothesis, suggesting that gases and aerosols are transported at a similar altitude.

The aerosol profiles show a clean air situation on 22 November, with the highest concentration close to the surface, indicating emissions primarily at or near the surface. On 24 November, the aerosol profiles confirm the presence of a transported plume, characterized by atypically high aerosol loads in elevated layers over Montevideo. However, it is worth noting that accurately retrieving the upper boundary of the aerosol is not possible due to the lower sensitivity of MAX-DOAS inversion algorithms for higher altitudes, especially in the presence of high aerosol loads. This issue is more pronounced for aerosols, primarily due to the non-linearity of the dependence of extinction to O4 dSCD compared to trace gases.

4.3 Satellite observations of glyoxal and formaldehyde from TROPOMI

In order to evaluate the event on a larger scale, the products derived from the TROPOMI instrument were analysed for the days around 24 November. Figure 10 shows the different products considered for this analysis. On 23 November, TROPOMI detects the beginning of a significant burning event near the south of Paraguay, which can be asso-
Figure 7. Panels (a)–(c) show the time series of vertical column densities of glyoxal, formaldehyde, and nitrogen dioxide. Percentile values for the corresponding products from TROPOMI-S5P over Montevideo are also shown as horizontal lines. Panel (d) shows the time series for the retrieved AOD by BOREAS from the MAX-DOAS observations and AERONET AOD values (at 360 and 380 nm, respectively).

Figure 8. (a) AOD values obtained by AERONET at 380 nm and retrieved by BOREAS at 360 nm for 24 November in Montevideo. The latter is based on the observed $O_4$ dSCD for western and eastern viewing directions using MAX-DOAS. (b) Scatter plot and linear fit between AERONET and BOREAS AOD values for 24 November.
Figure 9. Retrieved gas and aerosol profiles for 22 November (top row) and 24 November (bottom row) at different hours during the day for the eastern direction.

Associated with various fire sources. Strong signals of CO and NO$_2$ result from incomplete biomass combustion and the result of oxidation processes. The UV aerosol index also shows high values in northern Argentina, associated with the movement of the plume to the south. Formaldehyde and glyoxal are detected near the origin of the plume. Glyoxal appears to have a less dispersed plume, most likely due to direct emission of glyoxal from fires, but there is also a contribution from oxidation processes within the plume. Formaldehyde is mainly produced through atmospheric oxidation processes. In TROPOMI data, glyoxal is less visible than formaldehyde because the VCD is closer to the detection limit.

On 24 November, new fire sources became active in the same region, leading to an increase in the size of the detected plume. Observations of the UV aerosol index and CO show the plume in the region of northern Argentina and southern Brazil, along with several NO$_2$ hotspots. These signals are attributed to emissions from the active fires since there are no other identifiable sources in this region, such as large cities, capable of producing these signals. Formaldehyde and glyoxal levels also show an increase compared to the previous day, mainly due to the mixing with emissions from the new fire sources. The strongest signals of glyoxal and formaldehyde correlate with the RGB images of a dense plume within the Argentinian territory (see Fig. 3). This can be associated with the well-known fact that glyoxal is formed in the early stages of a wildfire, influenced by factors such as the intensity of the fire, its extension, and the number of ignition sources (Alvarado et al., 2020; Lerot et al., 2023) and references therein).

On this particular day, the aerosol index and CO columns, RGB images and the trajectories simulated with HYSPLIT, suggest that the plume moved across Uruguayan territory in a southeast direction. However, there are no clear indications that the plume extends over the city of Montevideo since there are no other identifiable sources in this region, such as large cities, capable of producing these signals. Formaldehyde and glyoxal values are near to the detection limit, making it challenging to infer the overall pattern of the plume. Consequently, there are no significant changes compared to previous days. This could also be in part the result of cloudiness and the substantial aerosol load in the atmosphere over Montevideo, since aerosols are not accounted for in the glyoxal and formaldehyde retrievals (see Sect. 2.4). Typical mechanisms for the removal of both gases from the troposphere, such as the reaction with the OH radical and photolysis, may explain why the gases are not transported so far from the source (Atkinson, 2000). The disagreement between satellite and ground-based measurements over Montevideo is probably the result of the much better detection limit of the MAX-DOAS observations. It is also possible that the plume had not yet arrived at its highest concentrations at the satellite overpass time around noon.

A NO$_2$ spot over Montevideo is simply showing the typical pollution of the city itself. NO$_2$ produced by the biomass burning event is rapidly removed from the troposphere most
likely through conversion to HNO₃. The evidence that part of the transport of the initial emissions is what is detected by the ground-based instruments is mostly provided by the CO columns product used as a tracer of biomass burning (Shi et al., 2015), since a small part of the border of the plume can effectively be seen over Montevideo (Fig. 10).

On 25 November, the Uruguayan territory experienced cloudy conditions, resulting in a gap in both satellite and terrestrial data. Despite the continued presence of sources in Paraguay and Argentina in the RGB images, their potential effect on Montevideo could not be detected after 25 November.

Figure 10. Vertical column densities (in molec. cm⁻²) for glyoxal, formaldehyde, nitrogen dioxide, and carbon monoxide and the aerosol index (in arbitrary units) from TROPOMI on 23, 24, and 25 November. The location of Montevideo is indicated by the red squares in the maps.
5 Conclusions

A plume of aerosols and trace gases from biomass burning in the Paraguayan–Argentinian border region on 22 November 2020 was detected over Montevideo 2 d later. On 24 November, AERONET measured very high AOD values compared to the usual Montevideo background in all bands. For example, there were AOD values at 380 nm greater than 1.0. Furthermore, ground-based MAX-DOAS measurements for that period showed an increase in formaldehyde and glyoxal vertical column densities, accompanied with an increase in the aerosol optical depth derived from the \( \text{O}_4 \) observations. Formaldehyde is known to be a useful tracer for biomass burnings. The observed increase in glyoxal levels was not so pronounced, probably due to the location of the ground-based instrument relative to the plume and the magnitude of the fires, so it is not clear that this gas can be used for plume detection in this scenario.

To confirm that the ground-based instruments detected this biomass burning plume, we used additional tools to ensure that the transport of the air masses to Uruguay was indeed possible. Examination of satellite RGB images shows the plume travelling from its source towards the vicinity of the city. Satellite data from TROPOMI show a vast carbon monoxide plume covering the entire south-east of Uruguay on 24 November. The formaldehyde, nitrogen dioxide, and glyoxal signals are relatively weak and not conclusive. Cloud coverage on 25 November prevented further investigation.

Trajectory simulations we conducted using HYSPLIT also support the conclusion that the plume reached Uruguay at the expected time. The air parcels are shown to be transported from the fire sources to the region near Montevideo, reaching a height of about 1.5 km above the ground, even when variations on the initial conditions are introduced. This agrees well with vertical profiles retrieved from the MAX-DOAS data using the BOREAS inversion algorithm for all species.

Atypical high trace gas concentrations and aerosol extinction levels confirm the ability of detecting transported plumes with ground-based instruments. The comparison of different trace gas and aerosol profiles shows possible temporal and spatial inhomogeneities within the plume for different species.

The above points can be interpreted as a partial validation of the ground-based detection of the plume, and we think that, in the future, ground-based atmospheric monitoring in Montevideo can be used as a tool for the detection of biomass burning events on its own, especially when satellite data are inconclusive for certain species such as HCHO, NO\(_2\), and CHOCHO.

As shown in this case study, ground-based remote sensing measurements of trace gases and aerosols are useful tools to detect the impact of biomass burning emissions at long distances from the source. Such measurements are much more sensitive than satellite data and therefore complement space-borne observations which provide full spatial coverage. Continuous measurements of this type of stations at several locations in the South American continent would provide valuable information on the regional- and continental-scale effects of wild fires. They also would deliver the data needed to validate atmospheric models to investigate the chemical processes in the plumes.

Appendix A: ICESat-2 observations of the biomass burning event

Data from the Ice, Cloud and land Elevation Satellite-2 (ICESat-2) were analysed to search for additional information about the BB plume passing over Uruguay. ICESat-2 is part of the Earth Observing System from NASA, and its main mission is to measure ice sheet and glacier elevation, as well as to measure and study ice thickness and vegetation characteristics in forests worldwide. It is equipped with the Advanced Topographic Laser Altimeter System (ATLAS) instrument, which times the travel of 532 nm laser pulses for high-resolution altimetry to perform surface and atmospheric measurements (Palm et al., 2021).

The data product analysed in this study was ATL09 Level 3A, which contains calibrated attenuated backscatter profiles and layer-integrated attenuated backscatter from the data acquired by ATLAS (Palm et al., 2022).

Between 23 and 24 November, the satellite performed two overpasses over Uruguay with a time stamp close to the event and with a trajectory almost over the BB plume. Figures A1 and A2 show the layer data for 23 and 24 November respectively. Only cloud and aerosol layers are detected over Uruguay in these passes.

Figure A1 reveals that the plume has not yet reached Uruguayan territory at the time of the overpass on 23 November, as no aerosol layers were detected. A low-altitude aerosol signal appears at northern latitudes when the satellite passes over the regions of Argentina and Paraguay, close to the location of the fires. However, it was not possible to obtain independent confirmation with additional satellite imagery (such as the VIIRS instrument) since the ICESat-2 overpass occurs at nighttime.

For 24 November, an aerosol layer was detected in the eastern part of Uruguay, approximately 200 to 375 km from Montevideo, at heights mostly between 1500 and 4000 m, as shown in Fig. A2. This result is in agreement with the profiles obtained from the BOREAS inversion analysis and with the VIIRS RGB images (see Fig. 3).
Figure A1. (a) ICESAT-2 ascending track above Uruguay for 23 November 2020. The colour scale represents the height of the topmost layer detected. The red square indicates the location of Montevideo. (b) Distribution of clouds and aerosol layers along the trajectory, from ATLAS ATL09 product data.

Figure A2. (a) ICESAT-2 descending track above Uruguay for 24 November 2020. The colour scale represents the height of the topmost layer detected. The red square indicates the location of Montevideo. (b) Distribution of clouds and aerosol layers along the trajectory, from ATLAS ATL09 product data.

Code and data availability. QDOAS software can be downloaded from the Royal Belgian Institute for Space Aeronomy web page (https://uv-vis.aeronomie.be/software/QDOAS/, Danckaert et al., 2017). Access to BOREAS software can be requested to the Institute of Environmental Physics, University of Bremen.

Data availability. AERONET data for Montevideo_FING station can be found at https://aeronet.gsfc.nasa.gov (Holben et al., 1998). BOREAS data are available upon request from Tim Bösch (tim.boesch@iup.physik.uni-bremen.de). TROPOMI data are available upon request from Leonardo M. A. Alvarado (leonardo.alvarado@awi.de). The ATL09 data from ICESat-2 satellite used in the present study can be freely obtained from https://doi.org/10.5067/ATLAS/ATL09.006, by searching for the ATL09 product (Palm et al., 2019).
**Author contributions.** The project was conceptualized by EF and LMAA. The analysis and discussion of results were carried out by MO, EF, AA, TB, NC, AR, and LMAA. The original draft was written by MO. All authors contributed to the final paper.

**Competing interests.** At least one of the (co-)authors is a member of the editorial board of *Atmospheric Chemistry and Physics*. The peer-review process was guided by an independent editor, and the authors also have no other competing interests to declare.

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