Article

Three-Dimensional Distribution of Biomass Burning Aerosols from Australian Wildfires Observed by TROPOMI Satellite Observations

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Abstract: We present a novel passive satellite remote sensing approach for observing the three-dimensional distribution of aerosols emitted from wildfires. This method, called AEROS5P, retrieves vertical profiles of aerosol extinction from measurements of the TROPOMI satellite sensor onboard the Sentinel 5 Precursor mission. It uses a Tikhonov-Phillips regularization which iteratively fits cloud-free near-infrared and visible selected reflectances to simultaneously adjust the vertical distribution and abundance of aerosols. The information on the altitude of the aerosol layers is provided by TROPOMI measurements of the reflectance spectra at the oxygen A-band near 760 nm. In the present paper, we use this new approach for observing the daily evolution of the three-dimensional distribution of biomass burning aerosols emitted by Australian wildfires on 20-24 December 2019. Aerosol optical depths (AOD) derived by vertical integration of the aerosol extinction profiles retrieved by AEROS5P are compared with MODIS, VIIRS and AERONET coincident observations. They show a good agreement in the horizontal distribution of biomass burning aerosols, with a correlation coefficient of 0.86 and a mean absolute error of 0.2 with respect to VIIRS. A fair agreement is found between coincident transects of vertical profiles of biomass burning aerosols derived from AEROS5P and from the CALIOP spaceborne lidar. The mean altitude of these aerosols derived from these two measurements show a good agreement, with a small mean bias (185 m) and a correlation coefficient of 0.83. Moreover, AEROS5P observations reveal the height of injection of the biomass burning aerosols in 3D. The highest injection heights during the period of analysis are coincident with the largest fire radiative power derived from MODIS. Consistency is also found with respect to the vertical stability of the atmosphere.

Keywords: fine particulate matter; biomass burning; black carbon; smoke; aerosol extinction vertical profile; 3D distribution of aerosols; Australian fire; TROPOMI; aerosol injection height

1. Introduction

Biomass burning (BB) aerosols have a major impact on the environment. Large-scale emissions of these particles have large impacts on the radiation balance and climate. Fresh and aged biomass burning plumes produce diabatic radiative heating e.g. [1] then can

lead to self-lofting of the plume e.g. [2]. They can produce a warming or cooling radiative forcing, depending on their optical properties and then their atmospheric lifetime [3]. If vertically and horizontally co-located with clouds, they may modify their properties and precipitation by acting as cloud condensation nuclei e.g. [4], depending on the age and the composition of the particles e.g. [5]. When transported near the surface, BB aerosols degrade air quality and cause negative long-term health effects e.g. [6]. The characterization of BB aerosols and their environmental impacts is hampered by the large spatiotemporal variability of wildfires and the difficulty to predict their occurrence and intensity.

The altitudes at which BB aerosol plumes are injected in the atmosphere during emission by wildfires is also important for the estimation of their impacts, and is very uncertain. Chemistry-transport models often predict them using semi-empirical methods based on satellite measurements of fire radiative power, burned surface areas, fire temperature, etc. e.g. [7,8]. The limitations of these methods are shown by comparisons with observations of the fire plumes [9,10]. Their uncertainties are linked to the lack of sufficient knowledge on the characteristics of the fires, pyro-convection processes, its effects on atmospheric stability and feedback. Accurate knowledge on the smoke injection heights is important for studying the regions that these particles may reach and thus their impacts. This is linked to the fact that horizontal transport strongly varies depending on the altitude of the atmospheric layers.

A description of both the horizontal and vertical distributions of atmospheric constituents within BB plumes is required to study their evolution in time and space and their impacts on the environment. Passive remote sensing allows the observation of the abundance of several atmospheric gaseous species, such as ozone or water vapor, in terms of vertical profiles. This is done by using the dependence of the spectral absorption signatures of these gases with respect to their vertical distribution. On the other hand, classic approaches of aerosol passive remote sensing, using sensors such as MODIS (Moderate Resolution Imaging Spectroradiometer), have provided for several decades total atmospheric columns of aerosol optical depth (AOD) which only describe their horizontal distributions. This is partly linked to the fact that the spectral signature of aerosols only depends on its altitude for specific parts of the radiation spectrum, such as the oxygen absorption bands in the near infrared e.g.[11] and the thermal infrared but only for coarse particles as desert dust e.g.[12].

Satellite high spectral resolution measurements in the near infrared, with strong spectral signatures of both fine and coarse aerosols are only performed by a few space-borne sensors, such as Global Ozone Monitoring Experiment-2 (GOME-2) [13] and TROP-ospheric Ozone Monitoring Instrument (TROPOMI) [14], respectively onboard MetOp-A/B/C since 2007 and Sentinel 5 Precursor (S5P) since 2017. Their analysis is challenging since the interactions between radiation and aerosols simultaneously depend on their abundance, altitude, size distribution [15] and composition [16]. Another difficulty for observing aerosols from space comes from the fact that the composition and thermodynamic state of the atmosphere are constantly evolving, and so are the optical properties of aerosols [17,18], which hampers tracing of the path of aerosol plumes in the atmosphere.

Several approaches allow the observation of the vertical profiles of particulate matter in the atmosphere. Airborne in situ sensors provide direct and accurate measurements of aerosol vertical profiles [19], but their availability is limited in space and time as they are mainly performed during field campaigns. Active remote sensing based on lidar instruments like the CALIOP lidar onboard the Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observation (CALIPSO) satellite can retrieve latitudinal transects of aerosol vertical profiles [20] but only for a very narrow field of view (90 m wide footprints). On a given day, these measurements are spaced longitudinally by about ~2000 km at mid-latitudes (for either daytime or nighttime overpasses). Recently, passive remote sensing of the reflectance spectrum of the oxygen absorption A-band has allowed the retrieval of mean aerosol heights over dark targets, from GOME-2 [11,20] and TROPOMI [21,22] measurements with horizontal resolutions of respectively 40 km x 80 km and 3.5 km x 5.5

km. Using nine viewing angles, aerosol mean heights are also derived from the Multi Angle Imaging Spectroradiometer (MISR) [23] on bright and dark surfaces at 4.4 km horizontal resolution [24,25]. The vertical profile of desert dust coarse particles is retrieved from space using measurements of the thermal infrared spectrum from the Infrared Atmospheric Sounding Interferometer (IASI) instrument [26–28]. The latter observations are horizontally spaced by 25 km (at nadir), so that they allow the observation of the 3D distribution of coarse particles having a strong radiative signature in the thermal infrared (such as desert dust). However, this is not the case for fine particles such as those originated from fires and most urban pollution.

In this paper, we present a new approach called AEROS5P for observing the 3D distribution of BB fine aerosols based on TROPOMI measurements. This method derives vertical profiles of aerosol extinction at 550 nm (linearly interpolated between 496 and 673 nm) for cloud-free pixels of TROPOMI. AEROS5P thus provides the daily observation of the full 3D distribution of fine BB particles, for cloud-free conditions. The determination of the vertical profile of aerosol extinction from passive spectral measurements is based on a Tikhonov-Philips-like auto-adaptive regularization. This technique was originally developed for desert dust aerosols and IASI measurements [12], lately applied also to volcanic sulphate aerosol [29], but it is here adapted for BB particles and TROPOMI data. For simultaneously constraining the aerosol vertical distribution and AOD, the method uses both near infrared and visible spectra from TROPOMI (instrument details are given in section 2). The vertical profiles of BB aerosols derived from AEROS5P can also be used to calculate AODs and mean aerosol layer heights.

We use AEROS5P for describing the 3D distribution of BB aerosols emitted during the extreme large-scale wildfires occurring over Australia in late 2019 and the beginning of 2020. Our approach provides an unprecedented 3D description of the smoke injection height and transport pathways. We mainly focus on the period 20-24 December 2019, which is particularly characterized by cloud-free conditions over most of the Australian continent during several days. The record-breaking Australian fires [30] caused a catastrophic impact on the economy, the population health, and biodiversity [31]. Massive amounts of smoke were injected into the upper-troposphere and the stratosphere. The subsequent in-plume radiative heating contributed to the formation of a smoke-charged confined ascending vortex that reached 35 km of altitude [2]. The plumes were transported very long distances and dominated the stratospheric aerosol layer of the whole Southern Hemisphere for several months, producing the most important impact on the climate system linked to stratospheric aerosol perturbations since the eruption of Pinatubo in 1991.

This paper provides a description of the AEROS5P algorithm backbone, and all additional datasets used in this work (Section 2). Section 3 shows comparisons of AEROS5P retrievals with respect to observations of the AOD from AERONET network stations, its horizontal distribution from MODIS and VIIRS (Visible Infrared Imaging Radiometer Suite) and vertical profiles derived from the CALIOP spaceborne lidar. The following section describes the 3D distribution of smoke plumes and their injection heights for the period of analysis, and the link with respect to the wildfire activity and meteorological conditions. Section 4 draws conclusions and discusses perspectives of this work.

2. Materials and Methods

2.1. The AEROS5P method

An overview scheme of the AEROS5P algorithm is presented in Fig. 1. The approach simulates radiative transfer in the surface-atmosphere system and calculates reflectance spectra in the visible and near infrared. The simulated reflectances are then compared with those measured by TROPOMI. The retrieval algorithm estimates the vertical profile of aerosol content in the atmosphere with a regularized iterative process, as done for the profiles of desert dust by [12,26,32] and ozone by e.g. [33,34]. As typically done in the Tikhonov-Philips' regularization [32] and the optimal estimation method [35], the cost

function that is minimized is the sum of the residuals of the measurement vector and the difference between the state vector and a priori values. These last ones are weighted by respectively radiometric noise and the regularization constraint (as described in [32]). The final products of AEROS5P are vertical profiles of aerosol extinction at 550 nm (α 550), aerosol top layer height (htop), mean aerosol height (hmean) and AODs for cloud-free TROPOMI pixels both over land and ocean, at daily scale. The following paragraphs describe more in detail the inputs, RT calculations and inversion procedure of our method.

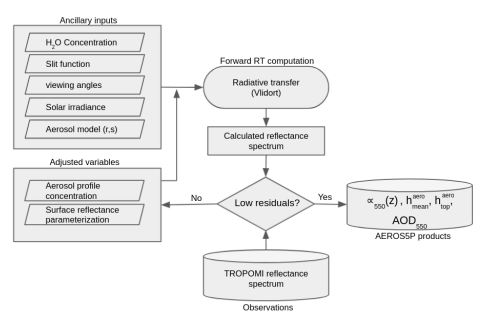


Figure 1. Scheme of the AEROS5P algorithm for retrieving the 3D distribution of fine BB aerosols. r : aerosol size distribution modal radius; s: aerosol size distribution standard deviation; h_{mean} : aerosol layer mean height; α_{550} : aerosol extinction profile at 550 nm; h_{top} : aerosol layer top height; AOD₅₅₀: aerosol optical depth at 550 nm.

2.1.1. Inputs

AEROS5P uses TROPOMI measurements in the visible and near infrared. This instrument is a spectrometer onboard the Copernicus S5P mission. Since its deployment in 2017, the sensor has been tested and calibrated extensively [36], and its quality is periodically monitored [37]. It has a swath width of around 2600 km, offering daily global coverage with a ground pixel resolution of 3.5×5.5 km2 (after 6 August 2019) and an overpass time of 13h30 local time. The instrument observes the spectrum of radiation emitted by the sun and backscattered by the Earth and the atmosphere towards the sensor, from the ultraviolet to the short-wave infrared (8 bands). The sensor also observes the sun to measure the solar irradiance spectrum.

AEROS5P uses reflectance measurements, derived as the ratio between radiance backscattered by the Earth and solar irradiance at each wavelength. This quantity is less dependent on calibration and aging of the instrument than absolute measurements of backscattered radiance alone. For each pixel, our approach uses reflectance measured at 194 wavelengths, corresponding to all channels within 12 spectral microwindows (see the central wavelength of these microwindows in Table 1). Radiation at these wavelengths is mainly affected by aerosols, surface properties and oxygen absorption, while avoiding the spurious influence of other gasses. They sample the visible spectrum from violet to red (specific wavelengths are taken from the list suggested by [38]) and the longest wavelength microwindow covers the whole oxygen absorption A-band (from 758 to 775 nm). These measurements are taken from the instrument bands 3, 4, 5 and 6 and are downloaded from [39]. For pixel co-localization, two pixels on the west side of the swath of bands 5 and 6 are not used. The present paper only shows results for 1 out of 13 pixels of TROPOMI for reducing computation time, although AEROS5P can process all cloud-free

pixels. The effective horizontal resolution is approximately 0.12° x 0.2° (near 12 x 20 km2). Only TROPOMI cloud-free pixels are processed. Cloudy pixels are screened out using the VIIRS level 2 cloud mask data CLDMSK_L2 [40] downloaded from [41] and co-localized with TROPOMI pixels. Pixels flagged as "cloudy" and "probably cloudy" are removed for robust cloud filtering.

In addition, we apply a so-called "soft correction" for improving the calibration of TROPOMI measurements. This correction is an adaptation from the work of [42,43] developed for the OCRA cloud retrieval for GOME-2 and TROPOMI measurements. It reduces the effects of parasite light sources, and it depends on the across-track position, the month of the year, land/ocean surface and instrument aging. These corrections improve the consistency of AOD and altitude estimation of successive scans.

Ancillary inputs of the algorithm describe the state of the atmosphere and geophysical properties. Meteorological conditions are taken from ERA-Interim reanalysis from the European Center for Medium Weather Forecast [44] in terms of temperature, pressure, and water vapor profiles. A vertically constant oxygen mixing ratio of 0.209 is assumed and oxygen absorption spectroscopic parameters from HITRAN 2012 [45] are used. Other geophysical inputs are a high spectral resolution spectrum of solar irradiance measured by [46] and monthly directionally dependent Lambertian-equivalent reflectivity (DLER) of the surface derived from TROPOMI measurements within the Sentinel-5p+ Innovation project [47] at 19 discrete wavelengths and 0.125° x 0.125° spatial resolution. Satellite boresight and relative azimuth angles between TROPOMI and the sun are also considered.

While several aerosol types may be found in Australia, most of the particulate matter responsible for the high AOD is fine BB aerosols or smoke during the period we analyze (see supplementary data, figure S1). Thus, we assume the microphysical properties of the latter in the retrieval. BB aerosols are assumed to have a monomodal lognormal size distribution with modal radius of $0.1~\mu m$ and a width of 1.43 (taken from AERONET retrieval of the event at Birdsville) and refractive indexes from [48]. All retrievals consider a unique a priori aerosol vertical profile of background concentrations, homogenous up to 9 km of altitude and decreasing above. A priori total column aerosol concentration is set to consider a background AOD of 0.03 at 550 nm.

Table 1. AEROS5P inputs and their corresponding sources.

| | Variable | Characteristics | | |
|-------------------------------|-----------------------------------|---|--|--|
| Meteorological | Reflectance spectra | 12 microwindows at 406, 416, 425, 436, 442, 451.5, 463, 477, 483, | | |
| state | | 494.5, 674.5 and 766.5 nm | | |
| Meteorological state | Temperature profile | ERAI reanalyzes | | |
| | Pressure profile | ERAI reanalyzes | | |
| | Water vapor profile | ERAI reanalyzes | | |
| Atmospheric species | A unique a priori aerosol concen- | - Homogeneous concentration up to 9 km corresponding to a back- | | |
| | tration profile | ground AOD of 0.03 at 550 nm | | |
| | BB aerosol refractive indices | From [48] | | |
| Gas absorption cross sections | Aerosol size distribution | A modal radius of 0.1 μm and width of 1.43 taken from AERONET | | |
| | | retrievals at Birdsville on 22 December 2019 | | |
| | Oxygen vertical profile | A constant mixing ratio of 0.209 | | |
| | | Calculation using HITRAN 2012 [45] spectroscopic parameters, | | |
| | Oxygen absorption cross sections | s line mixing, H2O broadening and collision-induced absorption fol- | | |
| | | lowing [49] | | |
| | H2O absorption cross sections | From HITRAN 2012 [45] | | |
| Geophysical state | Solar spectrum | High spectral measurements from [46] | | |
| | Surface albedo | DLER at 19 wavelengths, 0.125x0.125°, version 0.3 from the TRO- | | |
| | | POMI s5p+ Innovation project [47] | | |
| Instrumental calibration | Soft correction | Adapted from the soft correction for the OCRA cloud retrieval | | |
| | | [42,43] | | |
| | Instrumental spectral response | Slit function provided for each swath position and wavelengths | | |
| | | from [50] | | |

^{*}Central wavelengths indicated

2.1.2. Forward calculations of reflectance spectra

Radiative transfer calculations are performed with the VLIDORT (Vectorised Linear Discrete Ordinate Radiative Transfer) model version 2.7 [51,52]. It performs plane-parallel scalar computations accounting for light multiple scattering and absorption in the atmosphere at each wavelength. VLIDORT configuration is chosen for providing acceptable accuracy while reducing computation time. It is set for using a linearized Mie code (assuming spherical particle shape), 6 aerosol moments, 2 discrete radiative streams and no vector calculations. Computations are performed line by line for 262 wavelengths spaced by 0.2 nm covering the 12 spectral microwindows used in the approach.

Oxygen absorption cross sections at the A-band are derived for each RT calculation considering line mixing, water vapor broadening and collision-induced absorption following [49]. They are calculated at a high spectral resolution of 0.01 nm and then convolved by the instrument response function for deriving effective cross sections at the RT coarser spectral resolution. This technique significantly speeds up computations while keeping sufficient accuracy, as done for ozone retrievals by [33,34]. Sensitivity tests based on the retrieval of one day of measurements over Australia (20 december 2019) suggest that retrievals of the mean altitude of the aerosol layers using effective oxygen cross sections are highly correlated (correlation coefficient R of 0.96) with those based on high spectral resolution radiative transfer calculations. A mean bias of -270 m (lower altitudes with effective cross sections) and a root-mean-squared (RMS) difference of 560 m is also seen between the two datasets. This indicates that this source of error is not negligible but we consider it acceptable for analyzing the 3D distribution of BB aerosol plumes in the present study.

Assuming that most of the aerosols are located within the troposphere during the period of analysis, the atmosphere is divided from the surface to 12 km of altitude into

layers of equal depth of 1 km. Each layer contains water vapor, oxygen, and an initial background concentration of aerosols.

Surface albedo ω_0 is spectrally resolved and directionally dependent. For each wavelength λ of the RT calculations, it is derived as a linear combination of high spectral resolution empirical orthogonal functions (EOF) according to the following equation:

$$\omega_0(\lambda) = \sum_{i=1}^{n} c_i \cdot EOF_i(\lambda)$$
 (1)

the EOFs are derived from 400 to 800 nm at 1 nm resolution from a principal component analysis of a large database of high spectral resolution measurements of land reflectance for several surface types (vegetation, soils, man-made and ocean from [53]) and the United States Geological Survey (USGS) digital spectral library (version splib06a, [54]). We consider 6 and 3 spectrally resolved EOFs respectively over land and ocean. The a priori values of the coefficients c are calculated using the TROPOMI-derived DLER database mentioned in Table 1 for a given location and pointing geometry. The latter database is provided for 19 specific wavelengths. As done by [53], the coefficients are obtained by solving the linear system of equation (1) evaluated at the 19 wavelengths of the DLER data. This is performed with the following operation:

$$C^{a \text{ priori}} = (A^{T} A)^{-1} A^{T} \Omega_{DLER}$$
 (2)

where $C_{a priori}$ is a vector with the a priori values of the coefficients c, A a matrix where each column is a EOF interpolated at the 19 wavelengths and the vector Ω_{DLER} the DLER values from the TROPOMI-derived database. The values of the coefficients c are also jointly adjusted with the vertical profile of aerosols for better fitting TROPOMI reflectance spectra during the AEROS5P inversion process.

Once the reflectance spectra are simulated by VLIDORT, they are convolved by the instrument spectral response function (or slit function) of TROPOMI for enhancing the resemblance with measurements. These instrumental functions are provided for each swath position and wavelengths from [50].

2.1.3. Inversion procedure

The variables that are jointly adjusted for reducing the differences between measured and simulated reflectance spectra are given by the state vector $x_{AEROSSP}$:

$$\mathbf{x}_{\mathsf{AEROS5P}} = [\mathbf{N}_{\mathsf{log\,aero}}^{\mathsf{T}} \, \mathbf{C}^{\mathsf{T}}]^{\mathsf{T}} \tag{3}$$

Thus, the vertical profile of BB aerosol concentrations is obtained while simultaneously adjusting the weighting coefficients c in the vector C of the spectrally resolved EOFs of surface albedo. For avoiding negative values, the vector N_{togsero} contains the logarithm of aerosol number concentrations at each vertical level (spaced by 1 km). Moreover, only the four most significant coefficients c defining the surface albedo are adjusted for limiting the size of the state vector.

As TROPOMI measurements do not provide enough information to derive independent values of aerosol abundance at each vertical level, the approach constrains the solution for describing a single aerosol layer with a variable vertical distribution. This is done with Tikhonov-Philips constraints [55] that vary with altitude and auto-adapt to the convergence conditions as done in [32,56]. The constraint strengths are empirically derived from numerical simulations as done by [27] for desert dust but allowing for BB aerosols more variability at the upper atmospheric levels. During the iterative procedure, the constraints of the aerosol concentrations are automatically scaled by a Levenberg-Marquardt-type damping factor. Following [27], this method releases (enhances) the constraints when converging to smaller (diverging to higher) differences between measured and simulated spectra. The number of iterations is empirically set to 10 as convergence is usually found and the state vector varies at most by 5%. Using these constraints, the number of independent pieces of information (or degrees of freedom DOF) in the retrieval of

BB aerosols profiles varies during the iterative procedure. For the results shown in section 3 and 4, a typical value of ~1.5 is reached during the first iterations which allows the determination of the shape of the aerosol profile.

Once the iterative procedure is finished, the AEROS5P retrieval results for each pixel are taken from the iteration with the least difference between measurements and simulations. Quality assurance post-processing screens out pixels where the spectral differences remain too large (>10%) or too large DOFs (>3).

The final product of AEROS5P are aerosol extinction profiles at 550 nm, which are calculated with the adjusted concentration profiles and the assumed size distribution and refractive indexes, for cloud-free TROPOMI pixels over land and ocean. We consider that the aerosol profile in terms of extinction is better constrained by AEROS5P than that in terms of concentrations, since this last one directly depends on the accuracy of the assumed particle size distribution and refractive index. AODs are directly calculated by vertical integration of the aerosol extinction profiles, the mean aerosol height as the altitude at which half of the AOD is above and the other below, and the top aerosol height as the highest altitude with an aerosol extinction larger than a given threshold (we use here 0.1 km⁻¹). AODs and aerosol extinction at 550 nm are derived from linear interpolation between retrievals at 496 and 673 nm.

2.2. Other aerosol observations

For assessing the quality of AEROS5P aerosol retrievals, we compare them with several widely used and validated aerosol products. First, a comparison is conducted with respect to AOD derived from ground-based and satellite measurements. We consider sun photometer AOD measurements from eight AErosol RObotic NETwork (AERONET) stations [57] in Australia (Birdsville, Fowlers Gap, Lake Argyle, Lake Lefroy, Learmonth, Lucinda, Tumbarumba and Adelaide). We use level 1.5 data downloaded from [58]. They are measurements taken around noon at 440 nm, which we extrapolated to 550 nm using the corresponding Angstrom exponent measurements between 440 and 870 nm.

The horizontal distribution of AOD derived from AEROS5P is compared to that from satellite products derived from MODIS Dark Target and Deep Blue algorithms [59]. For a better horizontal coverage, we use the Multi-angle Implementation of Atmospheric Correction (MAIAC), which includes data from two MODIS sensors, as well as VIIRS using Dark Target. We consider the MYD04_L2 product from MODIS onboard the Aqua platform, MCD19A2 combining MODIS/Aqua+Terra (MAIAC), and the AERDB_L2 product from VIIRS onboard the Suomi National Polar-orbiting Partnership (NPP) satellite. We under-sampled these products at the AEROS5P grid resolution. For comparison of the horizontal distribution of AOD, we use MODIS/MAIAC data as it offers an enhanced horizontal coverage compared to the MODIS approaches that use one sensor. Co-localized comparisons of AOD are done with respect to MODIS Aqua as the overpass time is similar to that of TROPOMI.

Aerosol vertical profiles and mean heights retrieved from AEROS5P are validated against those derived from co-located measurements from CALIOP spaceborne lidar. This instrument measures latitudinal transects of total attenuated backscatter profiles at 532 and 1064 nm, with a 30 m vertical resolution below 8 km and 60 m above. These profiles qualitatively depict the vertical distribution of aerosols. For the cases analyzed here, the operational CALIOP product (v4.2) misclassifies the aerosols emitted from fires as "polluted dust" and "marine". Since the operational level 2 product of aerosol extinction is largely biased, and their spatial coverage is poor, we derive here aerosol extinction profiles using a classic Fernald lidar inversion procedure [60,61] (also called Klett inversion). The particle lidar ratio was adjusted for obtaining a rough match with respect to colocated measurements of AOD from MODIS. We use these aerosol extinction profiles to derive AODs and mean aerosol heights from CALIOP. It is worth noting that the operational TROPOMI product of aerosol layer height is not included in the comparison due to

limited spatial coverage. Indeed, almost no data is available in coincidence with the transects of CALIOP and AEROS5P observations analyzed in this paper.

In addition to aerosol products, we analyze the results derived from AEROS5P in terms of the 3D distribution of BB aerosols with other datasets describing the fire activity and the atmospheric conditions. The distribution of carbon monoxide (CO) is examined with the TROPOMI operational product [62,63]. Fire activity is surveyed using fire detections and estimates of Fire Radiative Power (FRP) from merged MODIS data onboard Aqua and Terra (MYD14 and MOD14 respectively) products [64] gridded to $0.2^{\circ} \times 0.2^{\circ}$ resolution.

Meteorological conditions are described with ERA5 reanalysis [65] from ECMWF at 0.25° x 0.25° resolution and 37 pressure levels (downloaded from the ESPRI IPSL mesocentre https://mesocentre.ipsl.fr). In addition to wind fields, we describe atmospheric static stability in terms of Brunt-Vaisala frequencies. This quantity describes the buoyancy of the air parcels, according to the surrounding meteorological conditions. It is calculated using ERA5 potential temperature, geopotential, and wind fields with formulas adapted from [66]. This variable does not consider processes as pyro-convection, whose explicit consideration needs a dedicated dynamical model which is out of the scope of the present study.

3. Comparison of AEROS5P retrievals with other aerosol products

3.1. Aerosol optical depth

The horizontal distribution of the AOD observed by AEROS5P is compared to that from MODIS on 20 and 21 December 2019 (Figure 2). It can be seen that the two approaches show very similar structures of AODs for the two days, both in variability and absolute values. On 20 December, a large and dense BB aerosol plume of an AOD reaching ~1 was depicted over eastern Australia extending from 150°E to 140°E, clearly seen with a similar shape from both satellite data. South and west of the plume some intermediate AODs of up to 0.4 are also seen in both cases. On the following day, two BB aerosol plumes are similarly depicted in two AOD maps. A plume over Central Australia of AODs of ~0.9 and a second one over Eastern Australia of AODs above 1, their shape and AOD values are very similar. We note a larger horizontal coverage of MODIS/MAIAC data, provided the use of two sensors and a less conservative detection of clouds as compared to AEROS5P. AOD maps from MODIS/Aqua show overall similar features (not shown for avoiding redundancy) and agreement with AEROS5P, but with a more limited horizontal coverage than MODIS/MAIAC.

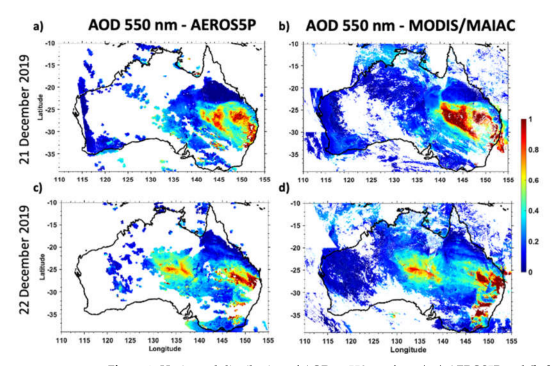


Figure 2. Horizontal distribution of AOD at 550 nm from **(a,c)** AEROS5P and **(b,d)** MODIS using the MAIAC algorithm on **(a,b)** 21 and **(c,d)** 22 December 2019 over Australia. This period is characterized by intense wildfires with large emission of BB aerosols.

Figure 3 presents a comparison of the AOD at 550 nm measured at eight AERONET stations with the three co-located satellite products (AEROS5P, MODIS/Aqua and VIIRS). AOD from AEROS5P is in good agreement with VIIRS and MODIS/Aqua on most of the days of the analyzed intense fire episode on 20-24 December 2021. The agreements with respect to AERONET ground based AOD observations are fairly good and similar for the 3 satellite products, including AEROS5P. Globally, the largest AODs are seen for all measurements near the fire regions (at Tumbarumba over the South-East of Australia) and medium values at Central Australia (at Birdsville) downwind from fires. Locations far from the BB aerosol sources clearly show the lowest AODs (Learmonth and Lucinda on the Western and North-eastern coasts respectively). Some differences in AOD may come from the time of the measurements, between around noon for AERONET and 13h30 local time for the satellite products (MODIS, VIIRS, and AEROS5P).

We also notice 3 cases where AEROS5P AOD deviates from the AOD measured by AERONET stations. First, on 20 December on the station Tumbarumba, we see a substantial difference in the AOD observed by AEROS5P and measured by the AERONET station (around 1.2 and 0.4 respectively), we notice that MODIS AOD also overestimate the AOD on that case (around 1). The difference may be caused by co-localization issues as the dense and heterogeneous smoke plume emitted from very intense fires located in the vicinity of this region (see supplementary data Figure S2 and S3). The second case where AEROS5P AOD is different from AERONET measurement is on 21 december, the satellite depicts an AOD around 0.9 and the sun photometer from Birdsville station measures 0.6. The slight discrepancy may be caused by the presence of desert dust in the center of the continent which induces positive biases in the AEROS5P BB retrievals. The last case of the mismatch between the AOD of AERONET and that of AEROS5P is on 23 december on the station Lake Lefroy; the AEROS5P AOD is then around 0.6 while the AERONET station measures 0.1 (as MODIS and VIIRS). This difference is probably caused by the cloud contamination not filtered by the VIIRS cloud mask used by AEROS5P. This seems to also be the reason for the overestimation of AEROS5P AOD at Adelaide, (particularly on 22 December). A better cloud filtering mask may improve the retrieval quality in the pixels neighboring cloudy situations.

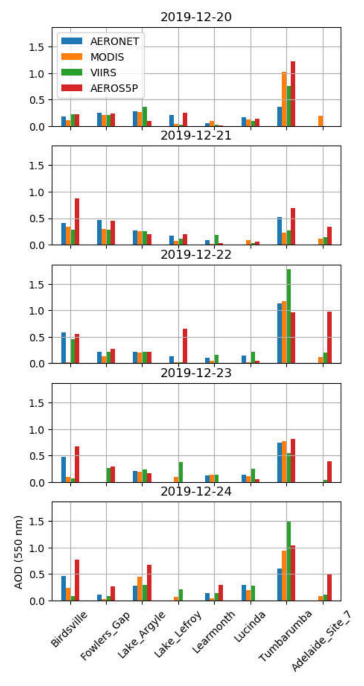


Figure 3. Comparison of AOD at 550 nm measured at 8 AERONET stations in Australia by sun photometer with respect to three satellite products: AEROS5P, MODIS/Aqua and VIIRS on 20 to 24 December 2019 (one panel per day from the upper to lower one).

A more systematic comparison of all co-located retrievals of AOD from AEROS5P, MODIS/Aqua and VIIRS is shown in Figure 4. It covers the period 20-24 December 2019. A good overall agreement is found between the 3 datasets, whereas the match is better between AEROS5P and VIIRS. This is shown by the fact that 65 % of AEROS5P/VIIRS co-located pixels show differences smaller than 0.2, a correlation coefficient of 0.87 and RMS differences of 0.3. These values are roughly similar to those found from the comparisons of AOD between MODIS and AEROIASI from the desert dust over the Sahara, with a correlation coefficient of 0.90 and an RMS difference of 0.2 [27]. The large value of our RMS difference is likely linked to the large AOD values of the comparison (up to 5 here, where it is at most 1 for [27]). With respect to MODIS Aqua, these indicators are moderately smaller (respectively 58 %, 0.63 and 0.36). A mean absolute error around 0.2 is seen

in both cases. The range of values of comparison for VIIRS is clearly greater (AOD up to 5) than for the comparison with MODIS, explaining partly the larger correlation in the first case. A comparison by the aerosol type detected by VIIRS shows that larger AOD values are mostly identified as "smoke" as expected (Table 2). Intermediate values with AOD around 0.6 are detected as "non-smoke fine aerosols". For those two aerosol types (smoke and non-smoke fine aerosols), the match between AEROS5P and VIIRS is much better (correlation coefficient larger than 0.82). Most of the points of AOD below 0.5 correspond to pixels identified by VIIRS as "mixed" and "background" (Table 2). For those cases, the AOD derived from AEROS5P that assumes the presence of only BB aerosols shows as expected larger differences (mainly overestimating) with respect to the retrievals from VIIRS and MODIS.

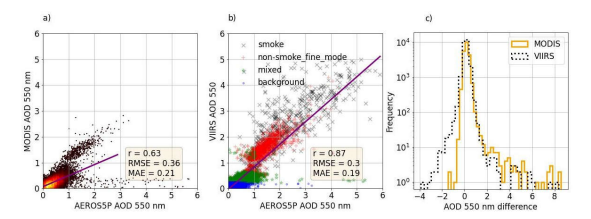


Figure 4. Scatterplot comparison of AOD at 550 nm derived from AEROS5P with respect to colocated retrievals from **(a)** MODIS/AQUA and **(b)** VIIRS for the period 20-24 December 2019. **(c)** Frequency histogram of differences of AOD at 550 nm of MODIS/AQUA and VIIRS with respect to AEROS5P. Symbols and colors in panel (b) indicate the aerosol type derived from the VIIRS approach.

Table 2. Comparison of AOD from AEROS5P and VIIRS for each aerosol type detected by VIIRS. r : pearson correlation coefficient; RMSE : Root Mean Squared Error; MAE : Mean Absolute Error.

| Aerosol type | r | RMSE | MAE | Number of co-located pixels |
|---------------------|------|------|------|-----------------------------|
| smoke | 0.82 | 0.69 | 0.49 | 667 |
| non-smoke_fine_mode | 0.86 | 0.30 | 0.24 | 3168 |
| mixed | 0.23 | 0.35 | 0.21 | 5868 |
| background | 0.24 | 0.20 | 0.13 | 8595 |

3.2. Aerosol extinction profiles

A comparison of two transects aerosol extinction profiles derived from CALIOP and AEROS5P on 21 and 22 December 2019 are shown in Figure 5. The vertical distribution of aerosols by the two datasets is in fairly good agreement. On 21 December, both retrievals show a denser aerosol layer extending from 1.0 to about 2.5 km in the southern part, approaching the surface at 28-30°S and progressively increasing in altitude up to 1-3 km around 24°S. The transect of the following day shows in both cases a rather homogenous aerosol layer from the surface up to ~4 km. The aerosol layer top gradually varies from 4 km of altitude at the southern part up to 2 km of altitude near 16°S. Quantitatively, values of aerosol extinction profiles are quite similar between CALIOP and AEROS5P transects. This is particularly remarked on 22 December 2019 (Fig. 5b,d), when the vertically integrated AOD for AEROS5P and CALIOP is very similar to that derived from MODIS (Fig. 5f). The transect of 21 December 2019 shows moderately larger values of extinction for AEROS5P than those derived from inversion of CALIOP measurements, while AOD from AEROS5P is in better agreement with MODIS than that from CALIOP (it underestimates

the AOD south of 28°S in Fig. 5e). Some other differences may partly come from the presence of clouds and the difference in overpass times of CALIOP and TROPOMI of about 1.5 h. The mean altitude of the aerosol layers is depicted as dashed lines in the figure. There is a good agreement between the two datasets. Considering all coincidences in the two transects, they show a correlation of 0.83, a mean bias of 180 m and a RMS difference of about 500 m. The uncertainty of the mean aerosol layer height derived from AEROS5P is moderately weaker and in similar order of magnitude than that of the TROPOMI operational product (between 500 and 1000 m according to [67].

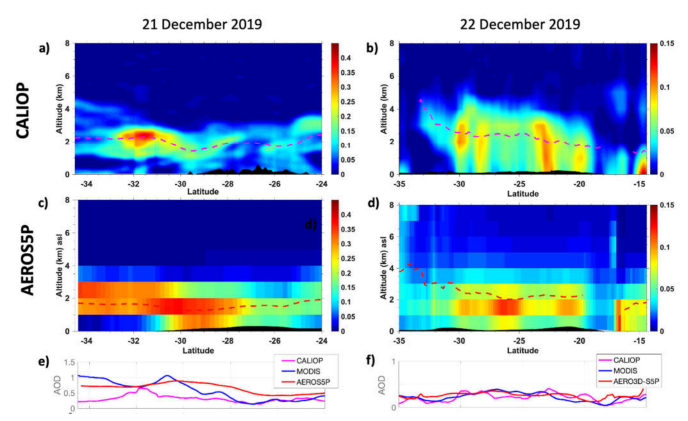


Figure 5. Comparison of transects of aerosol extinction profiles (in km¹) derived from **(a,b)** the CALIOP spaceborne lidar and **(c,d)** the AEROS5P passive approach on **(a,c)** 21 December and **(b,d)** 22 December 2019. Dashed lines indicate the mean altitude of the aerosol layers defined as the ones with half of the AOD above and below. The corresponding AOD at 550 nm from MODIS MAIAC and AEROS5P, and CALIOP at 532 nm is shown in panels (e) and (f).

4. Daily evolution of the 3D distribution of biomass burning aerosol plumes

During the period of analysis, exceptionally hot and dry conditions over Australia are responsible for numerous fires in a wide variety of locations. However, the most intense fires leading to the emission of massive amounts of smoke are seen principally located over Eastern and Southeastern Australia (principally in the New South Wales region [68]). This is seen in the example of fire detections from MODIS on 20 December, where the highest density of fire hotspots is located near the South-eastern coast (Figure 6c). As shown in Figure 2 in terms of AOD and also CO retrievals derived from TROPOMI (see supplementary Figure S2 and S3), the densest smoke plumes are seen over the eastern part of Australia (east of 140°E). The 3D distribution of biomass burning aerosols on 20 December is presented in terms of iso-aerosol extinction surface in Figure 6a. It shows that this large BB aerosols plume located at 25-37°S 140-150°E mainly extends vertically from the surface up to 3-4 km of altitude, with the core plume section reaching 5 km of altitude (seen for some pixels, see the aerosol top heights in Fig. 6b). According to estimations of the atmospheric stability in terms of Brunt-Vaisala frequency *N*² (calculated at 0.25° degree resolution from ERA5 reanalysis) meteorological conditions at these locations lead to a

relatively stable atmosphere (values of 1.5 10⁴ s²). Winds at the northern sector of this aerosol plume blow towards the northwest direction (Fig. 6b). The southern and western parts of this aerosol plume reach higher altitudes (5-7 km, seen for some pixels between clouds), co-located with a less stable atmosphere (lower values of N²). Strong winds at 600 hPa (Fig. 6d) are expected to transport these aerosols in the southeastern direction over the Pacific.

20 December 2019

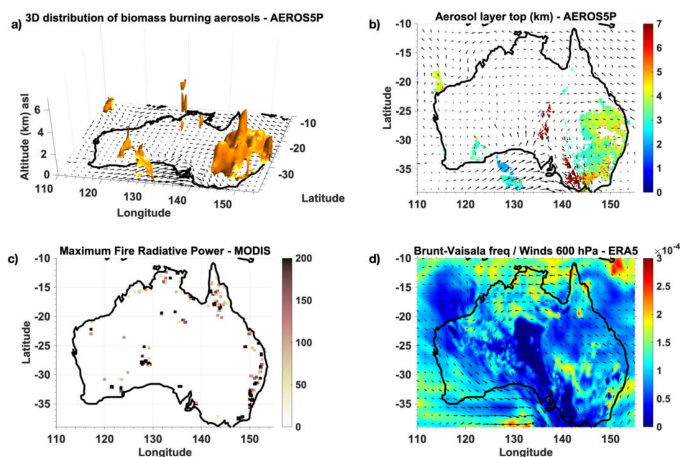


Figure 6. (a) Three-dimensional distribution of biomass burning aerosols on 20 December 2019 derived from TROPOMI measurements using the AEROS5P method, depicted as iso-aerosol extinction surface of 0.1 km³. **(b)** Aerosol layer top derived from AEROS5P as the aerosol highest layer with aerosol extinction larger than 0.1 km³. **(c)** Maximum fire radiative power derived from MODIS sensors onboard Aqua and Terra within each 0.2°x0.2° grid pixel. **(d)** Static atmospheric stability depicted by the Brunt-Vaisala frequency derived from ERA5 reanalysis. Winds from ERA5 reanalysis are overlaid on panels (a,b) at 850 hPa and (d) at 600 hPa.

On 21 December, the density of intense fires at the south-east hotspot (35°S 150°E) enhances while it diminishes elsewhere (Fig. 7c). Atmospheric stability is rather low above this region and within a band in the northwestern direction, according to low values of N² (Fig. 7d). As clearly depicted by AEROS5P, the 3D distribution of injections of biomass burning aerosol plumes forms a consistent structure extending from the surface near the fire region and ascending up to 7 km altitude at the northwest fringe of the plume. The slanted shape of the aerosol plume is consistent with a northwest direction of the wind at 600 hPa. The eastern part of this large aerosol plume (east of 145°E) extends vertically up to 3-4 km at maximum, which is concomitant with a clearly more vertically stable atmosphere (see higher N² values in Fig. 7d). Other aerosol plumes extending vertically up to 3-4 km (at the south-center part of the continent) are also co-located with rather stable conditions.



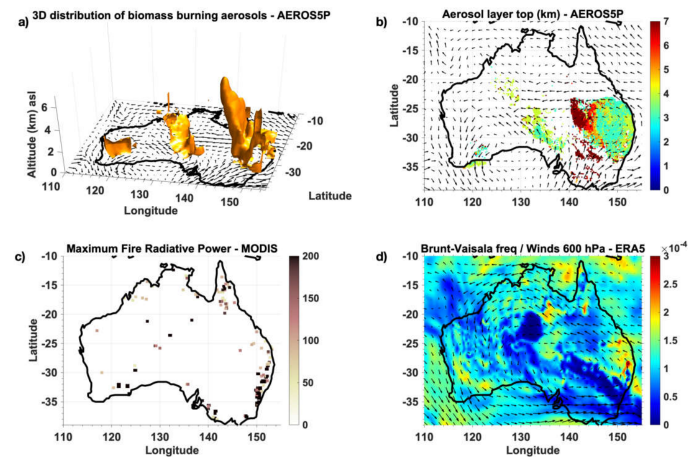


Figure 7. Same as Figure 6 but for 21 December 2019.

On the following day (22 December), the fire activity diminishes over the Eastern coast of Australia (Fig. 8c). Two aerosol plumes are observed by AEROS5P (Fig. 2b,d). The aerosols located at Central Australia are likely those emitted the previous day over Eastern Australia and then transported northeastward by the winds (see clear and steady wind direction at 850 and 600 hPa in Figs. 8b,d). This aerosol plume distributes vertically from the surface up 4-5 km, while located far from pyro-convective activity of the fire hotspots. The other aerosol plume over Eastern Australia is likely freshly emitted by fires. It is injected up to 6 km of altitude (Fig. 8a,b), in a region with mostly low atmospheric stability.

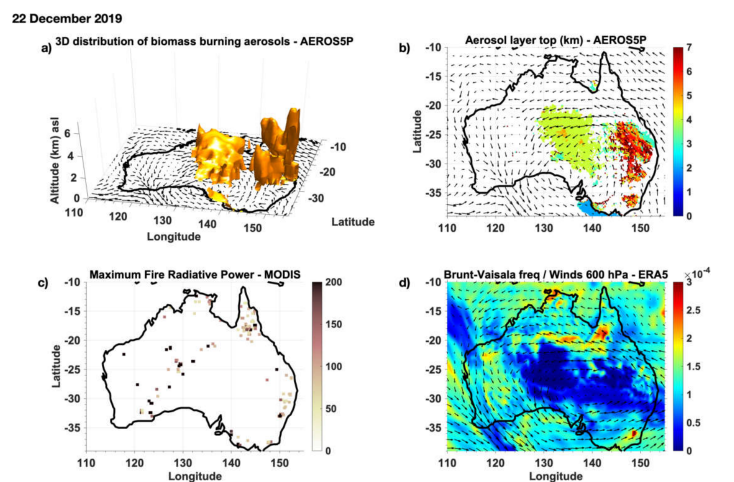


Figure 8. Same as Figure 6 but for 22 December 2019.

On 23 December, wildfire activity is almost absent over the Eastern coast of Australia. The two aerosol plumes observed in central Australia by AEROS5P extend vertically up to 4 km of altitude. This limited vertical extent and their geographical location west from the fire regions suggest that they are likely smoke plumes emitted the previous days. This is consistent with the reduction of new fire emissions on this day. In absence of direct link with fire emissions, low atmospheric stability (Fig. 9d) does not seem to favor the uplift of these aerosol plumes into higher atmospheric layers. A similar situation is remarked the following day (24 December 2019).

Additionally, we notice that the fire activity detected by MODIS in some regions does not necessarily imply the emission of large amounts of aerosols. This is the case for example of the Cape York Peninsula located northeast the continent, where fires are detected consistently every day (20-23 December in Figs. 6-9c) but dense aerosol plumes are not clearly seen as being emitted from this area. This might be linked to the kind of vegetation in this area (see Fig. S4). Grasslands and savanna are the main vegetation over the Cape York Peninsula, where fire events are more frequent but less intense [69,70]. Over the South-east coast of the continent, the climate is more temperate and evergreen forests are present. This may explain the emission of large amounts of biomass burning aerosols (e.g. Fig. 2) and carbon monoxide (Figure S2) from this area.

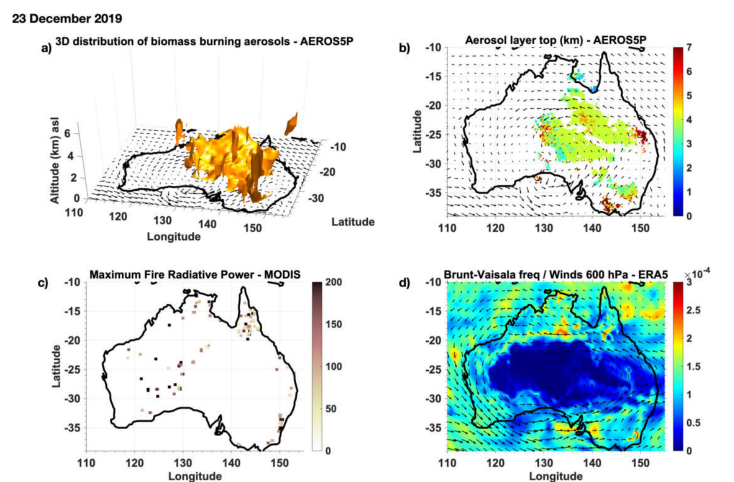


Figure 9. Same as Figure 6 but for 23 December 2019.

5. Conclusions

We present the new satellite-based AEROS5P approach, which enables the observation of the daily evolution of the 3D distribution of biomass burning aerosols. It is used here to characterize the 3D structure of aerosol plumes emitted by Australian wildfires during late December 2019. This approach derives vertical profiles of aerosol extinction at 550 nm by analyzing the reflectance spectra in the near infrared and visible measured by TROPOMI at $3.5 \times 5.5 \text{ km}^2$ resolution and cloud-free conditions. The spectral signature at the oxygen A-band near 760 nm provides information on the altitude of the aerosols while that in the visible spectrum from 406 to 674 nm about their total column abundance. These two quantities are derived simultaneously.

Aerosol optical depths retrieved by AEROS5P show good agreement with those derived from MODIS and VIIRS, with similar horizontal distribution. These datasets present correlations of up to 0.86 and root-mean-squared differences of 0.3. Transects of vertical profiles of aerosol extinction derived by AEROS5P show similar structure of aerosol plumes as those derived from measurements of the CALIOP spaceborne lidar. Mean altitudes of aerosol layers derived by AEROS5P and CALIOP are also in good agreement. These two datasets show a small mean bias (185 m), root-mean-squared differences of 500 m and a correlation coefficient of 0.83, for the mean plume altitude, for this event.

Moreover, AEROS5P provides an unprecedented daily description of 3D structure of the vertical injection of smoke plumes. The 3D distribution of these plumes is consistent with fire radiative power hotspots detected by MODIS and atmospheric stability described by a buoyancy frequency diagnostic. For the period of analysis (20-24 December 2019), we remark that aerosol plumes linked with intense fire activities and relatively low atmospheric stability are injected into the atmosphere up to 6 and 7 km of altitude. On the

other hand, aerosol plumes extend vertically up to 3-4 km for relatively higher atmospheric stability even near the emission regions. Aerosol plumes reach these altitudes once they are transported far away from sources.

Future work will be dedicated to the analysis of the 3D distribution of BB aerosol using AEROS5P in more diverse situations, such as other periods when BB aerosols from the Australian fires reached the stratosphere or other large fire events in other regions (e.g. Central and South Africa, California, Siberia, etc). Moreover, new versions of the AEROS5P will aim at characterizing other types of aerosols such as those from anthropogenic pollution.

Author Contributions: Investigation, software writing, data curation, formal analysis, writing original draft, performing and analyzing the retrievals, FL; Conceptualization, project administration, investigation, methodology, supervision, software writing, analysis, writing—review and editing, funding acquisition, JC; Project administration, partial funding acquisition, CD; software writing, methodology, ME; Project administration, writing—review, GD; providing datasets for analysis, GS; writing—review, analysis PS; writing-review, providing surface albedo principal components, XL and PZ; writing—review, analysis, fire power data providing, ST; Oxygen A-band cross section calculation, DT; providing swath dependent soft correction for TROPOMI and writing—review, RL and DL.

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Data Availability Statement: The data used/generated in this study are available the following plateforms: MODIS, CALIOP, and VIIRS data are downloadable from:https://search.earthdata.nasa.gov/; TROPOMI L1B data from: https://s5phub.copernicus.eu; TROPOMI Instrument Spectral Response Function from: http://www.tropomi.eu; AERONET AOD from: https://aeronet.gsfc.nasa.gov; AEROS5P data are available on request from the corresponding author.

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References

- 1. Yu P, Davis SM, Toon OB, Portmann RW, Bardeen CG, Barnes JE, et al. Persistent stratospheric warming due to 2019–2020 Australian wildfire smoke. Geophys Res Lett. 2021;48. doi:10.1029/2021gl092609
- 2. Khaykin S, Legras B, Bucci S, Sellitto P, Isaksen L, Tencé F, et al. The 2019/20 Australian wildfires generated a persistent smoke-charged vortex rising up to 35 km altitude. Communications Earth & Environment. 2020;1: 1–12.
- 3. Sellitto P, Belhadji R, Kloss C, Legras B. Radiative impacts of the Australian bushfires 2019–2020 Part 1: Large-scale radiative forcing. doi:10.5194/egusphere-2022-42

- 4. Chen L, Li Q, Wu D, Sun H, Wei Y, Ding X, et al. Size distribution and chemical composition of primary particles emitted during open biomass burning processes: Impacts on cloud condensation nuclei activation. Sci Total Environ. 2019;674: 179–188.
- 5. Li Y. Cloud Condensation Nuclei Activity and Hygroscopicity of Fresh and Aged Biomass Burning Particles. Pure and Applied Geophysics. 2019. pp. 345–356. doi:10.1007/s00024-018-1903-0
- Kennedy IM. The health effects of combustion-generated aerosols. Proc Combust Inst. 2007;31: 2757–2770.
- Freitas SR, Longo KM, Chatfield R, Latham D, Silva Dias MA, Andreae MO, et al. Including the sub-grid scale plume rise of vegetation fires in low resolution atmospheric transport models. Atmospheric Chemistry and Physics. 2007. pp. 3385–3398. doi:10.5194/acp-7-3385-2007
- 8. Turquety S, Menut L, Siour G, Mailler S, Hadji-Lazaro J, George M, et al. APIFLAME v2.0 biomass burning emissions model: impact of refined input parameters on atmospheric concentration in Portugal in summer 2016. Geoscientific Model Development. 2020. pp. 2981–3009. doi:10.5194/gmd-13-2981-2020
- Kahn RA, Chen Y, Nelson DL, Leung F-Y, Li Q, Diner DJ, et al. Wildfire smoke injection heights: Two perspectives from space. Geophysical Research Letters. 2008. doi:10.1029/2007gl032165
- 10. Sofiev M, Ermakova T, Vankevich R. Evaluation of the smoke-injection height from wild-land fires using remote-sensing data. Atmospheric Chemistry and Physics. 2012. pp. 1995–2006. doi:10.5194/acp-12-1995-2012
- 11. Sanders AFJ, de Haan JF, Sneep M, Apituley A, Stammes P, Vieitez MO, et al. Evaluation of the operational Aerosol Layer Height retrieval algorithm for Sentinel-5 Precursor: application to O2 A band observations from GOME-2A. Atmospheric Measurement Techniques. 2015. pp. 4947–4977. doi:10.5194/amt-8-4947-2015
- 12. Cuesta J, Eremenko M, Flamant C, Dufour G, Laurent B, Bergametti G, et al. Three-dimensional distribution of a major desert dust outbreak over East Asia in March 2008 derived from IASI satellite observations. Journal of Geophysical Research: Atmospheres. 2015. pp. 7099–7127. doi:10.1002/2014jd022406
- 13. Callies J, Corpaccioli E, Eisinger M, Lefebvre A, Munro R, Perez-Albinana A, et al. GOME-2 ozone instrument onboard the European METOP satellites. Weather and Environmental Satellites. SPIE; 2004. pp. 60–70.
- 14. Veefkind JP, Aben I, McMullan K, Förster H, de Vries J, Otter G, et al. TROPOMI on the ESA Sentinel-5 Precursor: A GMES mission for global observations of the atmospheric composition for climate, air quality and ozone layer applications. Remote Sens Environ. 2012;Complete: 70–83.
- 15. Bond TC, Doherty SJ, Fahey DW, Forster PM, Berntsen T, DeAngelo BJ, et al. Bounding the role of black carbon in the climate system: A scientific assessment. J Geophys Res. 2013;118: 5380–5552.
- Singh S, Fiddler MN, Bililign S. Measurement of size-dependent single scattering albedo of fresh biomass burning aerosols
 using the extinction-minus-scattering technique with a combination of cavity ring-down spectroscopy and nephelometry. Atmospheric Chemistry and Physics. 2016. pp. 13491–13507. doi:10.5194/acp-16-13491-2016
- 17. Konovalov IB, Golovushkin NA, Beekmann M, Andreae MO. Insights into the aging of biomass burning aerosol from satellite observations and 3D atmospheric modeling: evolution of the aerosol optical properties in Siberian wildfire plumes. Atmospheric Chemistry & Physics. 2021;21: 357–392.
- 18. Hosseini S, Li Q, Cocker D, Weise D, Miller A, Shrivastava M, et al. Particle size distributions from laboratory-scale biomass fires using fast response instruments. Atmospheric Chemistry and Physics. 2010. pp. 8065–8076. doi:10.5194/acp-10-8065-2010
- 19. Sandvik OS, Friberg J, Martinsson BG, van Velthoven PFJ, Hermann M, Zahn A. Intercomparison of in-situ aircraft and satellite aerosol measurements in the stratosphere. Sci Rep. 2019;9: 15576.
- 20. Winker DM, Tackett JL, Getzewich BJ, Liu Z, Vaughan MA, Rogers RR. The global 3-D distribution of tropospheric aerosols as characterized by CALIOP. Atmospheric Chemistry and Physics. 2013. pp. 3345–3361. doi:10.5194/acp-13-3345-2013
- 21. Chen X, Wang J, Xu X, Zhou M, Zhang H, Castro Garcia L, et al. First retrieval of absorbing aerosol height over dark target using TROPOMI oxygen B band: Algorithm development and application for surface particulate matter estimates. Remote Sensing of Environment. 2021. p. 112674. doi:10.1016/j.rse.2021.112674
- 22. Rao L, Xu J, Efremenko DS, Loyola DG, Doicu A. Hyperspectral Satellite Remote Sensing of Aerosol Parameters: Sensitivity Analysis and Application to TROPOMI/S5P. Front Environ Sci Eng China. 2022;9. doi:10.3389/fenvs.2021.770662
- 23. Diner DJ, Beckert JC, Reilly TH, Bruegge CJ, Conel JE, Kahn RA, et al. Multi-angle Imaging SpectroRadiometer (MISR) instrument description and experiment overview. IEEE Trans Geosci Remote Sens. 1998;36: 1072–1087.
- 24. Limbacher JA, Kahn RA. Updated MISR dark water research aerosol retrieval algorithm Part 1: Coupled 1.1 km ocean surface chlorophyll a retrievals with empirical calibration corrections. Atmos Meas Tech. 2017;10: 1539–1555.
- 25. Garay MJ, Witek ML, Kahn RA, Seidel FC, Limbacher JA, Bull MA, et al. Introducing the 4.4 km Spatial Resolution MISR Aerosol Product. doi:10.5194/amt-2019-340
- 26. Vandenbussche S, Kochenova S, Vandaele AC, Kumps N, De Mazière M. Retrieval of desert dust aerosol vertical profiles from IASI measurements in the TIR atmospheric window. Atmospheric Measurement Techniques. 2013;6: 2577–2591.
- Cuesta J, Flamant C, Eremenko M, Dufour G, Laurent B, Bergametti G, et al. Three-dimensional distribution of a major Saharan dust outbreak in June 2011 derived from IASI satellite observations. 4th IASI International Conference. 2016. Available: https://hal.univ-lille.fr/LATMOS/insu-01306503

- 28. Cuesta J, Flamant C, Gaetani M, Knippertz P, Fink AH, Chazette P, et al. Three-dimensional pathways of dust over the Sahara during summer 2011 as revealed by new Infrared Atmospheric Sounding Interferometer observations. Quarterly Journal of the Royal Meteorological Society. 2020. pp. 2731–2755. doi:10.1002/qj.3814
- 29. Guermazi H, Sellitto P, Cuesta J, Eremenko M, Lachatre M, Mailler S, et al. Quantitative Retrieval of Volcanic Sulphate Aerosols from IASI Observations. Remote Sensing. 2021;13: 1808.
- 30. Boer MM, Resco de Dios V, Bradstock RA. Unprecedented burn area of Australian mega forest fires. Nat Clim Chang. 2020;10: 171–172.
- 31. Filkov AI, Ngo T, Matthews S, Telfer S, Penman TD. Impact of Australia's catastrophic 2019/20 bushfire season on communities and environment. Retrospective analysis and current trends. Journal of Safety Science and Resilience. 2020;1: 44–56.
- 32. Steck T. Methods for determining regularization for atmospheric retrieval problems. Appl Opt. 2002;41: 1788–1797.
- 33. Liu X, Bhartia PK, Chance K, Spurr RJD, Kurosu TP. Ozone profile retrievals from the Ozone Monitoring Instrument. Atmospheric Chemistry and Physics. 2010. pp. 2521–2537. doi:10.5194/acp-10-2521-2010
- 34. Cuesta J, Eremenko M, Liu X, Dufour G, Cai Z, Höpfner M, et al. Satellite observation of lowermost tropospheric ozone by multispectral synergism of IASI thermal infrared and GOME-2 ultraviolet measurements over Europe. Atmos Chem Phys. 2013;13: 9675–9693.
- 35. Rodgers CD. Inverse Methods For Atmospheric Sounding: Theory And Practice. World Scientific; 2000.
- 36. Ludewig A, Kleipool Q, Bartstra R, Landzaat R, Leloux J, Loots E, et al. In-flight calibration results of the TROPOMI payload on board the Sentinel-5 Precursor satellite. Atmospheric Measurement Techniques. 2020;13: 3561–3580.
- 37. Tropomi monitoring portal. [cited 31 Mar 2022]. Available: http://mps.tropomi.eu/dashboard
- 38. Torres O, Tanskanen A, Veihelmann B, Ahn C, Braak R, Bhartia PK, et al. Aerosols and surface UV products from Ozone Monitoring Instrument observations: An overview. J Geophys Res. 2007;112. doi:10.1029/2007jd008809
- 39. https://s5phub.copernicus.eu. [cited 6 Oct 2021]. Available: https://s5phub.copernicus.eu/
- 40. Platnick S, Meyer K, Wind G, Holz RE, Amarasinghe N. The NASA MODIS-VIIRS continuity cloud optical properties products. Remote sensing. 2021. Available: https://www.mdpi.com/932614
- 41. Earthdata Search. [cited 31 Mar 2022]. Available: https://search.earthdata.nasa.gov/
- 42. Lutz R, Loyola D, García SG, Romahn F. OCRA radiometric cloud fractions for GOME-2 on MetOp-A/B. Atmospheric Measurement Techniques. 2016. pp. 2357–2379. doi:10.5194/amt-9-2357-2016
- 43. Loyola DG, Gimeno García S, Lutz R, Argyrouli A, Romahn F, Spurr RJD, et al. The operational cloud retrieval algorithms from TROPOMI on board Sentinel-5 Precursor. Atmospheric Measurement Techniques. 2018;11: 409–427.
- 44. Dee DP. The ERA-Interim Reanalysis: Configuration and Performance of the Data Assimilation System. European Centre for Medium Range Weather Forecasts; 2011.
- 45. Rothman LS, Gordon IE, Babikov Y, Barbe A, Chris Benner D, Bernath PF, et al. The HITRAN2012 molecular spectroscopic database. J Quant Spectrosc Radiat Transf. 2013;130: 4–50.
- 46. Chance K, Kurucz RL. An improved high-resolution solar reference spectrum for earth's atmosphere measurements in the ultraviolet, visible, and near infrared. Journal of Quantitative Spectroscopy and Radiative Transfer. 2010. pp. 1289–1295. doi:10.1016/j.jqsrt.2010.01.036
- 47. Sentinel-5p+ innovation. [cited 31 Mar 2022]. Available: https://www.grasp-sas.com/projects/aod-brdf_sentinel-5p-innovation/
- 48. Sarpong E, Smith D, Pokhrel R, Fiddler MN, Bililign S. Refractive Indices of Biomass Burning Aerosols Obtained from African Biomass Fuels Using RDG Approximation. Atmosphere. 2020. p. 62. doi:10.3390/atmos11010062
- 49. Tran H, Boulet C, Hartmann J-M. Line mixing and collision-induced absorption by oxygen in the A band: Laboratory measurements, model, and tools for atmospheric spectra computations. J Geophys Res. 2006;111. doi:10.1029/2005jd006869
- 50. ISRF dataset. In: Tropomi [Internet]. [cited 31 Mar 2022]. Available: http://www.tropomi.eu/data-products/isrf-dataset
- 51. Spurr RJD. VLIDORT: A linearized pseudo-spherical vector discrete ordinate radiative transfer code for forward model and retrieval studies in multiple scattering media. J Quant Spectrosc Radiat Transf. 2006;102: 316–342.
- 52. Spurr R, Christi M. On the generation of atmospheric property Jacobians from the (V)LIDORT linearized radiative transfer models. J Quant Spectrosc Radiat Transf. 2014;142: 109–115.
- 53. Zoogman P, Liu X, Chance K, Sun Q, Schaaf C, Mahr T, et al. A climatology of visible surface reflectance spectra. J Quant Spectrosc Radiat Transf. 2016;180: 39–46.
- 54. Clark RN, Swayze GA, Wise RA, Livo KE, Hoefen TM, Kokaly RF, et al. USGS digital spectral library splib06a. US Geological Survey; 2007. Available: https://pubs.er.usgs.gov/publication/ds231
- 55. Tikhonov AN. On the solution of ill-posed problems and the method of regularization. Doklady Akademii Nauk. Russian Academy of Sciences; 1963. pp. 501–504.
- 56. Eremenko M, Dufour G, Foret G, Keim C, Orphal J, Beekmann M, et al. Tropospheric ozone distributions over Europe during the heat wave in July 2007 observed from infrared nadir spectra recorded by IASI. Geophys Res Lett. 2008;35. doi:10.1029/2008gl034803

- 57. Giles DM, Sinyuk A, Sorokin MG, Schafer JS, Smirnov A, Slutsker I, et al. Advancements in the Aerosol Robotic Network (AER-ONET) Version 3 database automated near-real-time quality control algorithm with improved cloud screening for Sun photometer aerosol optical depth (AOD) measurements. Atmospheric Measurement Techniques. 2019. pp. 169–209. doi:10.5194/amt-12-169-2019
- 58. Aerosol Robotic Network (AERONET) Homepage. [cited 6 Oct 2021]. Available: https://aeronet.gsfc.nasa.gov/
- 59. Bilal M, Qiu Z, Campbell JR, Spak SN, Shen X, Nazeer M. A New MODIS C6 Dark Target and Deep Blue Merged Aerosol Product on a 3 km Spatial Grid. Remote Sensing. 2018;10: 463.
- 60. Fernald FG. Analysis of atmospheric lidar observations: some comments. Appl Opt. 1984;23: 652.
- 61. Fernald FG, Herman BM, Reagan JA. Determination of Aerosol Height Distributions by Lidar. J Appl Meteorol Climatol. 1972;11: 482–489.
- 62. Landgraf J, de Brugh JA, Scheepmaker R, Borsdorff T, Hu H, Houweling S, et al. Carbon monoxide total column retrievals from TROPOMI shortwave infrared measurements. Atmospheric Measurement Techniques. 2016. pp. 4955–4975. doi:10.5194/amt-9-4955-2016
- 63. Borsdorff T, Hu H, Hasekamp O, Sussmann R, Rettinger M, Hase F, et al. Mapping carbon monoxide pollution from space down to city scales with daily global coverage. Atmospheric Measurement Techniques. 2018;11: 5507–5518.
- 64. Giglio L, Schroeder W, Justice CO. The collection 6 MODIS active fire detection algorithm and fire products. Remote Sens Environ. 2016;178: 31–41.
- 65. Hersbach H, Bell B, Berrisford P, Hirahara S, Horányi A, Muñoz-Sabater J, et al. The ERA5 global reanalysis. Quart J Roy Meteor Soc. 2020;146: 1999–2049.
- 66. Stull RB. An Introduction to Boundary Layer Meteorology. Springer Science & Business Media; 1988.
- 67. Nanda S, de Graaf M, Veefkind JP, Sneep M, ter Linden M, Sun J, et al. A first comparison of TROPOMI aerosol layer height (ALH) to CALIOP data. Atmospheric Measurement Techniques. 2020;13: 3043–3059.
- 68. Kloss C, Sellitto P, Von Hobe M, Berthet G, Smale D, Krysztofiak G, et al. Australian Fires 2019--2020: Tropospheric and Stratospheric Pollution Throughout the Whole Fire Season, Front. Environ Sci. 2021;9: 10–3389.
- 69. Beringer J, Hutley LB, Abramson D, Arndt SK, Briggs P, Bristow M, et al. Fire in Australian savannas: from leaf to landscape. Glob Chang Biol. 2015;21: 62–81.
- 70. Williams RJ, Gill AM, Moore PHR. Seasonal Changes in Fire Behaviour in a Tropical Savanna in Northern Australia. Int J Wildland Fire. 1998;8: 227–239.
- 71. Chaivaranont W, Evans JP, Liu YY, Sharples JJ. Estimating grassland curing with remotely sensed data. Nat Hazards Earth Syst Sci. 2018;18: 1535–1554.

Supplementary Materials: The following are supporting information:

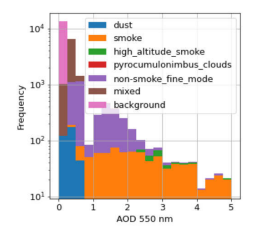


Figure S1: VIIRS AOD 550 nm per aerosol type from 20 to 24 December 2019;

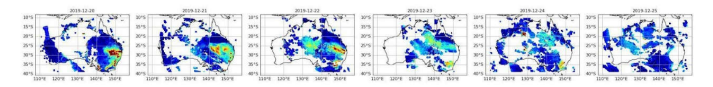


Figure S2: AEROS5P AOD 550 nm from 20 to 24 December 2019.

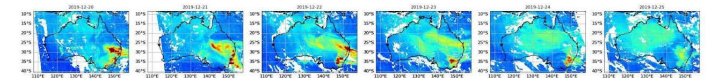


Figure S3: TROPOMI Carbon monoxide total column in 10²⁴ molec.m⁻² from 20 to 24 December 2019.

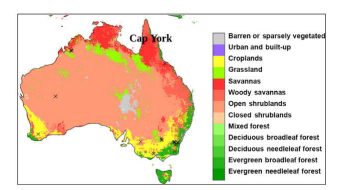


Figure S4: MCD12C1 land cover type map for Australia. Estimating grassland curing with remotely sensed data; (Chaivaranont et al. 2018).