We would like to thank the referees for their reviews. The manuscript has been revised according to their comments.

When rereading the manuscript we noticed that the altitude of the Maïdo observatory has been consistently reported as 2250 a.s.l. This is an error, the real altitude of the observatory is 2160 m a.s.l. This has been corrected everywhere in the manuscript and does not affect the analysis or the discussion presented in the current work.

# **Anonymous Referee #1**

Received and published: 26 August 2020

This is a well-written sound paper describing a campaign in Maïdo for testing the effect of biomass burning on mixing ratios of VOCs and on the model performance if a chemical transport model to calculate levels of O3 and NO2 at the site. By comparing the ratio of measured VOCs against CO and emission ratios, assessments about losses and production during transport are made. In general, the conclusions are a little bit weak but I am not aware of any similar data from this region of the world and therefore, I suggest publishing the manuscript in ACP, taking into account the suggestions from below.

# Major issue:

L9: For C6H6 and CH3OH, the EnR is lower than the ER, indicating a significant net sinks of these compounds. This is overselling as you assess it in line 274ff. as completely in line with the life times and the transport time. I was completely mislead when I read it in the abstract only. I thought you found an exceptional sink but I was then disappointed when I came to line 274. We have nuanced this in the abstract and explicitly mention now that this is in line with the expected atmospheric lifetime. Thank you for this reaction, we did not want to mislead any readers.

# Minor issues:

Abstract: This is obviously a paper focusing on VOC. Thus, mention the VOCs first. The order has been changed.

L35: there is also incomplete understanding about direct emissions. The text has been completed.

# L37: Isn't CO also controlling the O3 levels in remote atmospheres?

We do not expect CO to control the ozone levels in the remote atmosphere as you need NOx for ozone production during the OH initiated CO oxidation. However, the mixing ratios of CO, as well as CH4, have a major impact on the oxidative capacity of the remote marine atmosphere as they account for over 50% of the OH reactivity [Travis et al, 2020]. We refer here specifically to the role of OVOCs compared to all other NMVOCs. This specification has been added to the article.

L65ff: only OVOCs and certain VOCs with a conjugated double bond or heteroatoms can be measured by PTR-MS. Mention somewhere that other VOCs could also be present in BB plumes but were not in the focus of this study.

We have added that other compounds could have been present in biomass burning plumes but that either they were not the focus of this study or local sources interfered too much in order for us to reliably quantify the enhancement ratios (e.g. MEK).

# L249: the primary sink of what?

Of HCOOH when the wet deposition sink is negligible in the free troposphere during the dry searson. The information relating to a specific compound was taken out during a rewrite to generalise the discussion in this section. Thank you for noticing this. We have put the focus on the reduction of the wet deposition sink which is a statement that holds up for all compounds.

L254: . . . CH3OH and are shown. . .

We have restructured the sentence to be grammatically correct.

L296ff: a part of the difference could be that you did not measure the NMHCs, which also have BB sources and could contribute to the O3 formation. This could also be mentioned again in the conclusion, e.g. L 367.

The corresponding line reference is discussing HCOOH formation during transport. Measuring a complete array of NMHCs could indeed help identify precursors present in the plume. However, note that when production of HCOOH is a fast process, the precursor might have been mostly/fully lost during transport.

Instead of focussing on measurements, we searched for known HCOOH precursors in the literature and cross-referenced these with species emitted by biomass burning sources using the emission factors compiled by Adreae (2019). This is the same approach we used in looking for secondary production of CH3COCH3 during transport. We have added this clarification to the manuscript.

*Table 4*: As in the legend for Table 5, mention the values in the brackets. It has been adjusted.

Figure 2: explain MF Ok.

Figure 5: Please indicate Maïdo on the map for those being weak in geography of the South West Indian Ocean

The location of the Island has been added to the maps.

# **Anonymous Referee #2**

Received and published: 22 September 2020

The manuscript is well written with sound scientific results and an abundance of figures and tables to support the text. I suggest it is published to ACP, after addressing these minor technical details.

# Technical comments:

Figure 1: The uncertainty is barely visible in the bottom 6 plots (c,d,e,f,g,h). In most cases it looks as part of the pixelation (maybe increase resolution?)

We have regenerated the figures with a smaller linewidth and in pdf format to increase resolution. For consistency we have recreated all figures in pdf format, increasing the resolution for each.

Table 5 and Table 6 caption: You mention brackets for the standard deviation, when in the table you use parentheses.

Thank you for noticing, it has been corrected.

# References:

\* Travis, K. R., Heald, C. L., Allen, H. M., Apel, E. C., Arnold, S. R., Blake, D. R., Brune, W. H., Chen, X., Commane, R., Crounse, J. D., Daube, B. C., Diskin, G. S., Elkins, J. W., Evans, M. J., Hall, S. R., Hintsa, E. J., Hornbrook, R. S., Kasibhatla, P. S., Kim, M. J., Luo, G., McKain, K., Millet, D. B., Moore, F. L., Peischl, J., Ryerson, T. B., Sherwen, T., Thames, A. B., Ullmann, K., Wang, X., Wennberg, P. O., Wolfe, G. M., and Yu, F.: Constraining remote oxidation capacity with ATom observations, Atmos. Chem. Phys., 20, 7753–7781, https://doi.org/10.5194/acp-20-7753-2020, 2020.

\* Andreae, M. O.: Emission of trace gases and aerosols from biomass burning – an updated as sessment, Atmos. Chem. Phys., 19, 8523–8546, https://doi.org/10.5194/acp-19-8523-2019, 2019.

# Characterisation of African biomass burning plumes and impacts on the atmospheric composition over the South-West Indian Ocean

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Abstract. We present an investigation of biomass burning (BB) plumes originating from Africa and Madagascar based on measurements of a suite of volatile organic compounds (VOCs), carbon monoxide (CO), ozone (O<sub>3</sub>), and nitrogen dioxide (NO<sub>2</sub>) and a suite of volatile organic compounds (VOCs) obtained during the dry season of 2018 and 2019 at the high altitude Maïdo observatory (21.1° S, 55.4° E, 2250-2160 m above sea level), located on the remote island of La Réunion in the South-West Indian Ocean (SWIO). Biomass burning plume episodes were identified from increased acetonitrile (CH<sub>3</sub>CN) mixing ratios. Enhancement ratios (EnRs) — relative to CO — were calculated from in situ measurements for CH<sub>3</sub>CN, acetone (CH<sub>3</sub>COCH<sub>3</sub>), formic acid (HCOOH), acetic acid (CH<sub>3</sub>COOH), benzene (C<sub>6</sub>H<sub>6</sub>), methanol (CH<sub>3</sub>OH) and O<sub>3</sub>. We compared the EnRs to emission ratios (ERs) — relative to CO — reported in literature in order to estimate loss/production of these compounds during transport. For CH<sub>3</sub>CN and CH<sub>3</sub>COOH, the calculated EnRs are similar to the ERs. For C<sub>6</sub>H<sub>6</sub> and CH<sub>3</sub>OH, the EnR is lower than the ER, indicating a significant net sinks net sink of these compounds which was found to be in line with the expected atmospheric lifetime. For CH<sub>3</sub>COCH<sub>3</sub> and HCOOH, the calculated EnRs are larger than the ERs. The discrepancy reaches an order of magnitude for HCOOH  $(18-34 \text{ pptv ppbv}^{-1} \text{ compared to } 1.8-4.5 \text{ pptv ppbv}^{-1})$ . This points to significant secondary production of HCOOH during transport. The Copernicus Atmospheric Monitoring Service (CAMS) global model simulations reproduces well the temporal variation of CO mixing ratios at the observatory but underestimates O<sub>3</sub> and NO<sub>2</sub> mixing ratios in the plumes on average by 16 ppbv and 60 pptv respectively. This discrepancy between modelled and measured O<sub>3</sub> mixing ratios was attributed to i) large uncertainties in VOC and NO<sub>x</sub> (NO+NO<sub>2</sub>) emissions due to BB in CAMS and ii) misrepresentation of NO<sub>x</sub> recycling in the model during transport. Finally, transport of pyrogenically emitted CO is calculated with FLEXPART in order to i) determine the mean plume age during the intrusions at the observatory and ii) estimate the impact of BB on the pristine marine boundary layer (MBL). By multiplying the excess CO in the MBL with inferred EnRs at the observatory, we calculated the expected impact of BB on CH<sub>3</sub>CN, CH<sub>3</sub>COCH<sub>3</sub>, CH<sub>3</sub>OH and C<sub>6</sub>H<sub>6</sub> concentrations in

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the MBL. These excesses constitute increases of  $\sim 20\%$  – 150% compared to background measurements in the SWIO MBL reported in literature.

#### 1 Introduction

Non-methane volatile organic compounds (NMVOCs) are key tropospheric constituents. Many of them are highly reactive with the major atmospheric oxidants, especially with the OH radical, and therefore they strongly affect the oxidation capacity of the troposphere (Atkinson, 2000). By being a strong sink for OH, they also exert control on the lifetime of methane (Zhao et al., 2019) and thus on climate. Moreover, OH-initiated NMVOC oxidation modulates tropospheric  $O_3$  concentrations and is the major source of this secondary pollutant in high  $NO_x$  (NO+NO<sub>2</sub>) environments (Monks et al., 2015). Less volatile NMVOC oxidation products contribute to the formation and growth of secondary organic aerosol which deteriorates air quality and affects radiative forcing, and hence climate, both in a direct (by interacting with solar radiation) and indirect way (by acting as cloud condensation nuclei) (IPCC, 2013).

Whereas atmospheric oxidation of precursor VOC species is the dominant source of many oxygenated VOCs (OVOCs), primary anthropogenic emissions and (bidirectional) exchange with the biosphere and the ocean and biomass and biofuel burning also contribute to the atmospheric OVOC burden (Mellouki et al., 2015). Photochemical degradation and dry and wet deposition are the major sink processes. Global OVOC budgets are still prone to large uncertainties due to an incomplete understanding of direct emissions, photochemical production and loss processes and ocean–atmosphere exchange (Millet et al., 2010; Fischer et al., 2012; Read et al., 2012; Wang et al., 2019), and a paucity of (O)VOC data, especially at remote marine areas where the oxidative capacity of the atmosphere is mainly controlled by OVOCs (Lewis et al., 2005; Carpenter and Nightingale, 2015) in comparison to all other NMVOCs (Lewis et al., 2005; Carpenter and Nightingale, 2015; Travis et al., 2020).

The South-West Indian Ocean (SWIO) is one of the few pristine regions on Earth. It is largely decoupled from emissions originating from large bodies of land and is well suited to characterise remote marine air composition and ocean emissions (Colomb et al., 2009; Mallet et al., 2018). Located in the SWIO is the French overseas department La Réunion, a small volcanic island, home to the high altitude Maïdo atmospheric observatory (21.1° S, 55.4° E, 2250-2160 m above sea level) (Baray et al., 2013), hereafter referred to as RUN. From October 2017 to November 2019, a high-sensitivity quadrupole-based Proton Transfer Reaction Mass Spectrometry VOC analyser (hs-PTR-MS) was deployed at RUN in the framework of the OCTAVE (Oxygenated Compounds in the Tropical Atmosphere: Variability and Exchanges) project (http://octave.aeronomie.be). In combination with other ground-based and satellite data, the resulting near-continuous high time-resolution two-year data set will serve to better constrain VOC emissions in the remote tropical marine atmosphere and to identify missing sources. Part of this dataset has already been used in a source apportionment study of formaldehyde (HCHO) (Rocco et al., 2020).

The present paper contributes to the disentanglement of the different sources contributing to the (O)VOC composition at RUN by focusing on the role of biomass burning (BB). It is established from ground-based remote-sensing Fourier Transform Infrared (FTIR) observations that BB impacts the atmosphere over La Réunion. The BB events affecting the region occur most frequently in southern Africa and Madagascar but impacts from burning in South America and Malaysia has also been iden-

tified (Duflot et al., 2010; Vigouroux et al., 2012). Seasonality of in situ CO concentrations at RUN indicates that BB plumes also impact the atmospheric composition at the surface (Zhou et al., 2018). This was confirmed by the hs-PTR-MS dataset generated at RUN for the OCTAVE project (Fig. A1). Biomass burning represents the second largest global source of NMVOC emissions (Yokelson et al., 2008; Akagi et al., 2011). Pyrogenic emissions are reasonably well constrained by numerous laboratory studies (e.g. Holzinger et al. (1999); Christian et al. (2003); Yokelson et al. (2008)) and observations of BB plumes in the atmosphere (e.g. Lefer et al. (1994); Yokelson et al. (1999, 2003); Jost et al. (2003); de Gouw et al. (2006); Vigouroux et al. (2012); Akagi et al. (2014)). Emission factors for numerous compounds have been compiled recently by Andreae (2019). The investigation of compositional changes during BB plume transport may provide valuable clues for identifying missing sources of reactive trace species (e.g. Jost et al. (2003); de Gouw et al. (2006); Chaliyakunnel et al. (2016)). This is of particular interest for the carboxylic acids, as current models underestimate their observed abundances, possibly in part due to a misrepresentation of the contribution of biomass burning (Paulot et al., 2011; Chaliyakunnel et al., 2016).

In this work, we focused on the first BB plume intrusions from the fire season in 2018 and 2019. Enhancement ratios (EnRs) — relative to excess CO — of excess acetonitrile (CH<sub>3</sub>CN), formic acid (HCOOH), acetone (CH<sub>3</sub>COCH<sub>3</sub>), acetic acid (CH<sub>3</sub>COOH), benzene (C<sub>6</sub>H<sub>6</sub>), methanol (CH<sub>3</sub>OH) and O<sub>3</sub> were calculated from in situ measurements at RUN for the first time. Other VOCs could also be present in the BB plumes but either were not the focus of this study or local emissions from the island interfered too much with the signal to reliably quantify the EnRs (e.g. methyl ethyl ketone, MEK). In addition, observations of BB plumes at RUN were used to evaluate the global near-real time (NRT) CO, O<sub>3</sub> and NO<sub>2</sub> modelled concentrations at RUN from the Copernicus Atmospheric Monitoring Service (CAMS). Finally, we propose a way to use in situ VOC measurements at RUN to estimate the impact of BB plumes on the pristine marine boundary layer (MBL) over the SWIO. This is done for CH<sub>3</sub>CN, CH<sub>3</sub>COCH<sub>3</sub>, C<sub>6</sub>H<sub>6</sub> and CH<sub>3</sub>OH.

In section 2 the instruments, methods and models used in this study are presented. The results are shown in section 3 and discussed in section 4.

#### 2 Observations and Methods

# 2.1 Observations

#### 2.1.1 In situ air mass characterisation

RUN houses routine instruments characterising in situ air constituents in the context of global networks such as GAW (Global Atmospheric Watch), ICOS (Integrated Carbon Observation System) and ACTRIS (European Research Infrastructure for the observation of Aerosol, Clouds and Trace Gases). A summary of routine measurements from the observatory used in this study is shown in Table 1. A detailed description of these and other operational routine instruments at the observatory can be found in Duflot et al. (2019); Baray et al. (2013) and Zhou et al. (2018).

In the frame of the OCTAVE project, a hs-PTR-MS instrument (Ionicon Analytik GmbH, Austria) was deployed at RUN from October 2017 to November 2019. This resulted in a near-continuous high time-resolution two-year data set of volatile

organic compounds (VOCs). The instrument was run in the multiple ion detection mode using H<sub>3</sub>O<sup>+</sup> precursor ions with a total cycle time of  $\sim 2.7$  min. Regular calibrations of the hs-PTR-MS were performed by diluting a gravimetrically prepared VOC/N<sub>2</sub> mixture (Apel-Riemer Environmental Inc., Miami, FL, USA; stated accuracy of 5% for all VOCs) with zero-VOC air obtained by sending ambient air through a catalytic converter (Parker, type HPZA-3500, Haverhill, MA, USA). This resulted in VOC concentrations in the lower ppby range. Calibrations as a function of relative humidity were performed bimonthly by controlling the humidity of the zero air with a dew point generator (LI-COR LI610, Lincoln, Nebraska, USA). The calibration factor (CF) for acetic acid (CH<sub>3</sub>COOH) was estimated from the experimentally determined CF for CH<sub>3</sub>COCH<sub>3</sub>. This is done by considering the calculated collision rate constants of H<sub>3</sub>O<sup>+</sup> with CH<sub>3</sub>COOH and CH<sub>3</sub>COCH<sub>3</sub> (Su, 1994), the contributions of the protonated molecules to the respective product ion distributions (Schwarz et al., 2009; Inomata and Tanimoto, 2010), and by assuming the same hs-PTR-MS transmission efficiency for ions with a mass difference of 2 u. Similarly, the CF of HCOOH was determined from the measured one of acetaldehyde. The humidity dependence of formic and acetic acid CFs obtained at similar hs-PTR-MS operating conditions has been reported in literature (Baasandorj et al., 2015) and has been taken into account for quantification. By considering the uncertainties on the different parameters involved in the carboxylic acid quantification in the present study, the total uncertainty on their mixing ratio is estimated at 50%. Measurements were averaged over 1 hour to lower the limit of detection (LoD) and the random fluctuations of the measurements. A list of masses, and their associated compound(s), recorded by the hs-PTR-MS together with the LoD, dwell time and whether the compounds are directly calibrated is shown in Table 2.

#### 2.1.2 Ground-based remote sensing

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The University of Colorado Multi-AXis Differential Optical Absorption Spectroscopy (CU MAX-DOAS) instrument consists of a scanning (horizon – zenith – horizon) telescope coupled to two ultraviolet-visible grating spectrometers (Coburn et al., 2011). Scattered-light solar spectra are collected along lines of sight at different elevation angles above the horizon (Hönninger et al., 2004), and analyzed using DOAS least-square fitting (Platt and Stutz, 2008) to retrieve trace gas slant column densities (SCD) by the QDOAS software package (Danckaert et al., (accessed June 10, 2019). For this analysis, NO<sub>2</sub> (Vandaele et al., 1998) and O<sub>2</sub>-O<sub>2</sub> (Thalman and Volkamer, 2013) were retrieved in a fitting window from 425 – 490 nm, using the further fit settings as described in Kreher et al. (2020). Near-surface volume mixing ratios of NO<sub>2</sub> were retrieved from limb (0° elevation angle) spectra following Dix et al. (2016). This approach takes advantage of the fact that the limb viewing geometry is highly sensitive to absorbers near instrument altitude, O<sub>2</sub>–O<sub>2</sub> is used to parameterise aerosol extinction near instrument altitude, avoiding the need for complex aerosol profile information (Sinreich et al., 2013; Dix et al., 2016). The NO<sub>2</sub> profile shape was constructed using a typical tropical background with BB enhancements collocated to excess CO from FLEXPART (see section 2.3.2). Variations on the retrieval settings and profile assumptions indicate that  $\sim 10$  pptv NO<sub>2</sub> can be quantified with an uncertainty of 5 pptv using this approach. Further tests using NO<sub>2</sub> and O<sub>4</sub> fits at shorter wavelengths (Kreher et al., 2020) determined that the retrieved NO2 volume mixing ratios generally agree within the reported uncertainty, despite the different spectral ranges average NO<sub>2</sub> over different horizontal spatial scales. This indicates that the NO<sub>2</sub> mixing ratio is representative of the regional lower troposphere predicted by the CAMS model.

#### 120 2.2 Enhancement ratios

The impact of BB events on an atmospheric species X is often quantified by an emission factor (EF<sub>X</sub>) or an enhancement ratio relative to a compound Y (EnR<sub>X/Y</sub>). The first is defined as the mass of compound X that is released by burning 1 kg of dry fuel, whereas the second is defined as the excess mixing ratio — due to BB — of compound X ( $\Delta$ X), with respect to that of a reference species Y ( $\Delta$ Y). If the EnR is measured close to the source and/or if both X and Y were minimally affected by physico-chemical interactions, it is also referred to as the emission ratio of compound X normalised to Y (ER<sub>X/Y</sub>). The ER can be computed from the EF by taking the molecular weights (MW) of both species into account:

$$ER_{X/Y} = \frac{EF_X}{EF_Y} \frac{MW_Y}{MW_X}.$$
 (1)

A list of EFs with the associated fuel type has been compiled most recently by Andreae (2019). When comparing the EnR

values derived from our observations to ERs from literature, production/loss of plume constituents during transport should be taken into consideration. Enhancement and emissions ratios are often used with CO as the reference species Y. Hereafter, enhancement ratios are always considered with respect to CO unless specifically stated otherwise.

Excess mixing ratios are determined above the background — unaffected by BB — diel profiles which were approximated by the seasonal median diel profiles (appendix A2). During the day, mesoscale transport at La Réunion results in the observatory being located in the planetary boundary layer (PBL). The chemical composition of the PBL is determined by marine, biogenic and anthropogenic sources and sinks interacting in physicochemical atmospheric processes. At night, air masses arriving at RUN originate primarily from the free troposphere (FT). This mesoscale transport results in a natural diel variation of com-

pound mixing ratios which needs to be taken into account when calculating EnR.

## 2.3 Modelling

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Below we discuss the model simulations used in this study. Each model is used with a specific goal in mind. First, we evaluate the CAMS NRT atmospheric composition service using in situ measurements. It is important that CAMS correctly reproduces CO concentrations at RUN as pyrogenic emissions used in this service will be used to calculate transport of excess CO (ΔCO) over the SWIO with the Lagrangian FLEXible PARTicle dispersion model, FLEXPART (Stohl et al., 1998; Stohl and Thomson, 1999; Stohl et al., 2005; Pisso et al., 2019). We use FLEXPART to calculate the mean plume ages during the BB episodes at RUN but also to calculate the impact of pyrogenic emissions on the pristine MBL over the SWIO. Finally FLEXPART-AROME (Verreyken et al., 2019) is used to simulate mesoscale transport in complex terrain towards the observatory. This last simulation is performed in an effort to quantify the PBL–FT mixing during BB intrusions and identify the main transport mode of the plumes.

#### 2.3.1 CAMS NRT

The CAMS NRT service was developed based on a series of Monitoring Atmospheric Composition and Climate (MACC) research projects. It provides daily forecasts of reactive trace gases, greenhouse gases and aerosol concentrations. The data are generated by the Integrated Forecast System (IFS) at the European Centre for Medium-Range Weather Forecasts (ECMWF). The chemical mechanism used is an extended version of the Carbon Bond 2005 lumped chemistry scheme (Flemming et al., 2015). BB emissions implemented in the NRT service rely on the Global Fire Assimilation System v1.2 (GFAS v1.2) inventory. The GFAS assimilates fire radiative power observations from the NASA MODIS satellites to quantify BB emissions (Giuseppe et al., 2018; Rémy et al., 2017; Kaiser et al., 2012). On 9 July 2019, the model was updated to use the CAMS emission inventories, CAMS\_GLOB\_ANT v2.1 and CAMS\_GLOB\_BIO v1.1 (Granier et al., 2019), instead of the previous MACCity (Lamarque et al., 2010) and the MEGAN\_MACC (Sindelarova et al., 2014) inventories. BB plume injection heights were also introduced in this update. A full description and validation of the update was reported by Basart et al. (2019).

We used modelled mass mixing ratios at the location of RUN calculated on different pressure levels (1000, 950, 925, 900, 850, 800, 700 and 600 mbar levels) every three hours (0, 3, 6, 9, 12, 15, 18, 21 UT) from the midnight forecast at  $0.5^{\circ} \times 0.5^{\circ}$  resolution  $^{1}$ . The CO,  $O_{3}$  and  $NO_{2}$  mass mixing ratios are transformed to volume mixing ratios and compared to the in situ measurements.

#### 2.3.2 FLEXPART

FLEXPART, driven by ECMWF IFS meteorology at 0.5° × 0.5° horizontal resolution was used to calculate the transport of ΔCO due to BB during 15 June – 31 August 2018 and 17 June – 22 August 2019. The CO emissions are provided by the GFAS v1.2 inventory. Three-hourly mean mixing ratios of CO were generated on vertical layers of 500 m depth between 0 and 3500 m above ground level (a.g.l.). The output was given on a 0.5° × 0.5° grid. Due to the low horizontal resolution, the orographic profile of La Réunion is not well resolved. For example, the ground level of RUN is only 284 m above sea level (a.s.l.) in the model, much below the true altitude of 2250-2160 m a.s.l.

Age classes (AC) are used to estimate the mean plume age (T) for the different intrusions. The CO plumes are categorised by age with 2 day resolution ( $T_{AC} = 1 \pm 1, 3 \pm 1, ..., 23 \pm 1$  days). BB plume excesses are traced for 24 days, after which the plume is assumed to be diluted to negligible background levels. The mean BB plume age is obtained from the FLEXPART output by:

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$$T = \frac{\sum_{j=0}^{11} \Delta \text{CO}_j \times T_j}{\sum_{j=0}^{11} \Delta \text{CO}_j},$$
 (2)

<sup>&</sup>lt;sup>1</sup> available at https://apps.ecmwf.int/datasets/cams-nrealtime/levtype=pl/

where  $\Delta CO_i$  is the mean mixing ratio calculated by FLEXPART with AC=j.

To estimate the impact of BB on the MBL for compound X, we use:

$$\Delta X_{\text{estimate}} = \Delta CO \times EnR_X,$$
 (3)

where  $\Delta CO$  is calculated by FLEXPART and EnR<sub>X</sub> is inferred from data. In this approach, the role of an ocean sink is neglected.

#### 2.3.3 FLEXPART-AROME

FLEXPART-AROME 24-hour backtrajectory simulations are used to estimate the respective contribution of the PBL and the free troposphere to the in situ measurements at RUN. Lesouëf et al. (2011) characterised the PBL impact on the Maïdo mountain region by using a passive boundary layer tracer initialised in an approximation of the minimal boundary layer. This PBL proxy is defined as 500 m a.g.l., capped at 1000 m a.s.l. Here, the inverse approach is used by calculating the fraction of time air parcels have spent in the PBL-proxy during the 24-hour backtrajectory simulation. This fraction measures the potential impact of surface emissions on the in situ measurements. We will split this fraction up according to surface type (land/ocean) and call the separate components the mixing fraction (MF). Given the lack of a high-resolution anthropogenic emission inventory over La Réunion, we are not able to use the model to quantify mixing ratios unperturbed by BB plumes and instead use the median diel profile as stated in section 2.2.

## 3 Results

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# 3.1 Data analysis

Six episodes of enhanced CH<sub>3</sub>CN, which is a typical BB compound, were identified in August 2018 and August 2019 (Fig. 1). The correlation (*r*) between the excess mixing ratio of the monitored trace gases and ΔCH<sub>3</sub>CN, during the identified intrusions, is shown in Table 3. As dimethyl sulphide (DMS) is only marginally present in pyrogenic emissions (0.0022 – 0.05 g emitted per kg dry matter burned from tropical forest and agricultural residue burning respectively (Andreae, 2019)) and has a short atmospheric lifetime (less than 1 day (Blake et al., 1999)), the correlation between ΔDMS and ΔCH<sub>3</sub>CN is not expected to be directly related to the BB emissions. For this reason, compounds that correlated less well with ΔCH<sub>3</sub>CN than ΔDMS were not considered as plume constituents. Plume constituents in this analysis are thus limited to CH<sub>3</sub>CN, HCOOH, CH<sub>3</sub>COCH<sub>3</sub>, CH<sub>3</sub>COOH, O<sub>3</sub>, C<sub>6</sub>H<sub>6</sub> and CH<sub>3</sub>OH.

Mean background (i.e. outside BB episodes) concentrations of plume constituents in austral winter together with the mean excesses during the different BB intrusions (in %) are shown in Table 4. Correlation with CH<sub>3</sub>CN is especially strong for compounds showing large excesses compared to the diel background pattern (illustrated in appendix A2). We note that trace species such as HCHO, MEK and acetaldehyde (CH<sub>3</sub>CHO) and methyl ethyl ketone (MEK) show elevated concentrations during the night in BB episodes, which suggests that they are related to BB. However, as the diel patterns for these compounds are subject to strong variability, excesses are poorly characterised during the day and not analysed further here.

For each of the intrusions, the EnR is computed for CH<sub>3</sub>CN, CH<sub>3</sub>OH, CH<sub>3</sub>COCH<sub>3</sub>, C<sub>6</sub>H<sub>6</sub>, HCOOH, CH<sub>3</sub>COOH and O<sub>3</sub>. Figure 1 shows the scatter plots correlating the excess of the trace species monitored by the hs-PTR-MS instrument and  $\Delta$ CO. The calculated EnRs are found in Table 5.

# 3.2 Comparison with model

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## 3.2.1 FLEXPART-AROME

Figure 2 shows the fraction of time spent in the PBL-proxy from Lesouëf et al. (2011) over sea (blue) and land (brown), during the 24-hour backtrajectory calculations with FLEXPART-AROME, together with the relative humidity (RH) at the observatory.

Biomass burning intrusions have lower than average RH values. The humidity peaks during the BB episodes are coincident with peak impacts of the MBL. It is also shown that the impact of mesoscale PBL emissions on the VOC concentrations is lower during the BB intrusions in August 2018 than in August 2019.

## 3.2.2 CAMS near-real-time model simulations

The modelled mixing ratios at RUN calculated by the CAMS NRT service are compared to data recorded at the observatory for CO, O<sub>3</sub> and NO<sub>2</sub> (Fig. 3). The model bias for CO, during the BB intrusions, is lowest on the 800 mbar pressure level (bias of 9.7 ppbv), which is closest to the mean pressure measured at the observatory during the same period (792.8 mbar). Note that CAMS reflects well the CO mixing ratios at Maïdo both during and outside (5.1 ppbv bias) BB episodes. As CO is a chemically stable compound in the atmosphere, the agreement between model and measurements indicates that synoptic scale transport and mesoscale mixing with the BB plumes at the location of RUN is sufficiently reproduced by the CAMS NRT model.

The O<sub>3</sub> model bias is 16 ppbv during the BB episodes with a maximum bias of 39 ppbv (67% above the calculated value). Outside the BB episodes, the CAMS O<sub>3</sub> concentrations show only a small bias (0.8 ppbv), within the uncertainty of measurements. This good agreement outside of the BB events suggests that mesoscale O<sub>3</sub> sources and sinks either have a limited impact or are correctly calculated by the model at the location of RUN.

The NO<sub>2</sub> bias reaches 60 pptv during BB episodes, while it is only 9 pptv (within 10 pptv DOAS accuracy error) in other periods. Note that the NO<sub>2</sub> measurements are from the ground-based remote sensing CU MAX-DOAS instrument and reflect the NO<sub>2</sub> mixing ratio in the lower free troposphere. The large discrepancy in modelled and measured NO<sub>2</sub> on 3 August 2019 may be due to a weak BB plume passing near RUN (appendix B).

#### 3.2.3 FLEXPART forward simulation

A comparison between ΔCO obtained from measurements and the calculated ΔCO from transport of GFAS v1.2 emission inventory, simulated by FLEXPART, is shown in Fig. 4. Due the misrepresentation of the orographic profile of La Réunion, ground level at the location of the observatory is only 284 m above sea level in the model. The real altitude of RUN (2250 2160 m a.s.l.) is situated near the boundary between layers 1500 – 2000 m a.g.l. and 2000 – 2500 m a.g.l. in the FLEXPART

output. In reality, mesoscale transport, not resolved in FLEXPART, mixes the different vertical layers and data recorded at Maïdo correspond to a mixture between different output levels. In what follows, we consider RUN to be located in the layer between 2000 and 2500 m a.g.l.

The model overestimates  $\Delta$ CO mixing ratios at RUN by 37 ppbv and 17 ppbv on average during the BB episodes in 2018 and 2019 respectively. Peak differences between modelled and observed mixing ratios are 340 ppbv during the BB episodes in 2018 and 162 ppbv during those in 2019. The model bias outside BB episodes, reduces to 3 ppbv for both 2018 and 2019.

As the timing of BB intrusions is well represented in the model, as can be visually confirmed from Fig. 4, calculated mean plume ages during the different episodes are expected to be accurate. The calculated plume ages are, in chronological order of arrival at RUN, 7.5, 10.6 and 11.3 days in 2018 and 7.4, 9.3 and 13.7 days in 2019.

## 4 Discussion

## 4.1 Transport and dominant sink

The relative humidity during the BB intrusions was generally low (see Fig. 2). Peak RH values correspond to large impact of the MBL and often lower ΔCH<sub>3</sub>CN concentration (e.g. 7 and 17 August 2019, Fig. 2). From this, we expect the plume to be primarily located in the free troposphere, which is drier than the PBL. This is consistent with results from FLEXPART (Fig. 4), where ΔCO is especially significant in layers above 1500 m a.g.l. The primary sink in the FT during austral winter (dry season) is due to photochemical interaction. This is also evident same is also found from the CAMS NRT model where elevated CO mixing ratios are calculated between the 850 mbar and 700 mbar pressure levels (~ 1500 – 3000 m a.s.l.). As the BB plume is primarily located in the FT during austral winter (dry season) we expect that wet deposition is negligible for all compounds.

# 4.2 Plume characterisation

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The emission ratios are computed based on emission factors from (Andreae, 2019) for of  $CH_3CN$ , HCOOH,  $CH_3COOH$ ,  $CH_3COCH_3$ ,  $C_6H_6$  and  $CH_3OH$  are shown in Table 6. Possible fuel types for BB plumes arriving at RUN are: savanna and grassland, tropical forest or agricultural residue. Enhancement ratios are compared to the emission ratios to check for consistency with accepted knowledge regarding sources/sinks during transport.

#### 4.2.1 Acetonitrile, acetone, methanol and benzene

During the synoptic scale transport in the free troposphere, the photochemical sink is expected to be dominant over wet scavenging. As the lifetime with regards to this sink is larger than the maximum plume age (13.7 days) for both CH<sub>3</sub>CN ( $\tau_{\text{CH}_3\text{CN}}$ = 1.4 years (de Gouw et al., 2003)) and CH<sub>3</sub>COCH<sub>3</sub> ( $\tau_{\text{CH}_3\text{COCH}_3}$ = 36 – 39 days (Arnold et al., 2005; Fischer et al., 2012)), the EnRs are expected to correspond well with the ERs from literature. This is the case for CH<sub>3</sub>CN (Table 6). In contrast, the EnR

of acetone (~ 8 pptv ppbv<sup>-1</sup>) is at least a factor of ~ 2 larger than the ER from the literature (Table 6), a likely indication of secondary CH<sub>3</sub>COCH<sub>3</sub> formation in the BB plume. Acetone production has been recorded in BB plumes over the Eastern 270 Mediterranean (Holzinger et al., 2005) and over Namibia (Jost et al., 2003). In contrast, aged BB plumes over Eastern Canada and Alaska did not show evidence of acetone production (de Gouw et al., 2006). Known pyrogenic CH<sub>3</sub>COCH<sub>3</sub> precursors are propane, i-butane and i-butene (Singh et al., 1994). Using the EFs from Andreae (2019), we find ER<sub>propane</sub>= 1.2 – 3.2 pptv ppbv<sup>-1</sup>, ER<sub>i-butane</sub>= 0.05 – 0.1 pptv ppbv<sup>-1</sup> and ER<sub>i-butene</sub>= 0.30 – 0.52 pptv ppbv<sup>-1</sup>. Taking these known precursors of secondary CH<sub>3</sub>COCH<sub>3</sub> into account, as well as acetone formation yields at high NO<sub>x</sub> estimated based on the Master Chemical Mechanism MCMv3 (http://mcm.leeds.ac.uk/MCM/) (Saunders et al., 2003), the secondary production of acetone can be estimated. It is found to enhance the acetone EnR by 1.16 – 2.80 pptv ppbv<sup>-1</sup>, therefore explaining the major part of the discrepancy. This is at odds with results from Jost et al. (2003) where fast CH<sub>3</sub>COCH<sub>3</sub> production is observed and propane could not be considered as a precursor since this conversion is a slow process.

Both methanol and benzene have shorter expected lifetimes compared to the age of the BB plume arriving at RUN ( $\tau_{CH_3OH}$ = 7 days (Jacob et al., 2005),  $\tau_{C_6H_6}$ = 9 days (Monod et al., 2001)). This is consistent with the reduced EnRs inferred from data at RUN compared to the reported average emission ratios from literature (Table 6).

# 4.2.2 Carboxylic acids

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Due to the relatively short global average atmospheric lifetime of HCOOH ( $\tau_{HCOOH}$ = 2 – 4 days (Stavrakou et al., 2012)) and CH<sub>3</sub>COOH ( $\tau_{CH_3COOH} \approx 2$  days (Khan et al., 2018)), EnRs in aged BB plumes should not be compared to emissions ratios from literature (Paulot et al., 2011). However, as wet- and dry deposition are dominant sinks for both CH<sub>3</sub>COOH and HCOOH, their effective lifetime during transport in the FT is expected to be much longer ( $\tau_{HCOOH} \approx 25$  days from photochemical oxidation (Millet et al., 2015)).

The much higher HCOOH enhancement ratio estimated from RUN data (20 – 30 pptv ppbv<sup>-1</sup>) compared to reported emission ratios (2 – 4 pptv ppbv<sup>-1</sup>) points to significant secondary production during transport to RUN. Similar to secondary production of CH<sub>3</sub>COCH<sub>3</sub>, we identified HCOOH precursor species from literature and cross-referenced these with pyrogenic EFs (Andreae, 2019). A potential precursor to HCOOH strongly emitted by agricultural residue burning is glycolaldehyde (ER = 19 ± 12 pptv ppbv<sup>-1</sup> (Andreae, 2019)). The yield of HCOOH from glycolaldehyde oxidation has been measured to be 18% at 296 K and 52% at 233 K (Butkovskaya et al., 2006). This may account for part of the HCOOH production during transport. However, recent studies indicate that this production is effective only in high NO<sub>x</sub> conditions that are not realistic in a natural environment (Orlando et al., 2012; Orlando and Tyndall, 2020). Production of HCOOH from glycolaldehyde is thus most likely only a minor source. No other known precursors were identified of HCOOH were identified in pyrogenic emissions to account for the high HCOOH production during transport to RUN suggesting a missing source in current knowledge.

Secondary production of HCOOH was also found in BB plumes over Canada (Lefer et al., 1994) but was not observed in previous ground-based FTIR studies at La Réunion (Vigouroux et al., 2012). Enhancement ratios of HCOOH calculated from the Tropospheric Emission Spectrometer instrument aboard the NASA's aura spacecraft over Africa ranged from 26 to 28 pptv

ppbv<sup>-1</sup> (Chaliyakunnel et al., 2016), consistent with our results. This secondary HCOOH production in BB plumes could account for part of the discrepancy in global HCOOH budget between models and observations (Chaliyakunnel et al., 2016). As these EnRs are inferred from data over biomass burning hotspots in Africa, HCOOH is probably formed primarily close to the source and conserved during synoptic scale transport towards RUN.

For  $CH_3COOH$  the enhancement ratio ( $EnR_{CH_3COOH} \approx 14 \text{ pptv ppbv}^{-1}$ ) is of the same order of magnitude as the emission ratios from literature (Table 6). Therefore, in contrast with the case of HCOOH, no significant secondary production of acetic acid in BB plumes is identified.

# 310 **4.2.3** Ozone and NO<sub>2</sub>

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It is generally accepted that  $O_3$  is produced in BB plumes during transport (Taupin et al., 2002; Jaffe and Wigder, 2012; Parrington et al., 2013; Arnold et al., 2015; Brocchi et al., 2018). The EnRs obtained in this study (410 – 640 pptv ppbv<sup>-1</sup>) are in agreement with the range of EnRs obtained in tropical BB plumes older than 5 days, compiled by Jaffe and Wigder (2012), 410 - 750 pptv ppbv<sup>-1</sup>.

Figure 3 shows that the CAMS model reproduces correctly the  $O_3$  concentrations at RUN outside the BB episodes but underestimates  $O_3$  during these episodes. The large underestimation of  $O_3$  during these episodes indicates a misrepresentation of the BB emissions at the source and/or missing  $O_3$  production during transport in the chemically complex plumes.

The  $O_3$  production in the troposphere is highly dependent on the ratio between VOCs and  $NO_x$ . The CAMS NRT service is known to overestimate  $NO_2$  over southern Africa in austral winter/spring (Flemming et al., 2015; Basart et al., 2020). However, this overestimation was reduced since the upgrade in 2017 (Basart et al., 2020). Total BB VOC emissions in the IFS of ECMWF was  $\sim 40$  Tg in the year 2008 (Flemming et al., 2015). This is too low in comparison with the top-down estimate by Stavrakou et al. (2015) where the global pyrogenic VOC emissions are estimated to be 67 - 75 Tg yr<sup>-1</sup>.

Ozone production in BB plumes tends to be  $NO_x$ -limited (Jaffe and Wigder, 2012). The measured  $NO_2$  mixing ratio during BB episodes is significantly higher than those calculated by CAMS (Fig. 3). The largest and smallest difference between model and measurements for both  $NO_2$  and  $O_3$  were recorded during the first and last BB intrusion in 2019 respectively. This mismatch for  $NO_2$  may be caused by an underestimation of  $NO_x$  emissions by fires or by a misrepresentation of  $NO_x$  recycling (e.g. through peroxyacetyl nitrate or PAN). BB plumes reaching RUN are located at relatively low altitudes where warmer temperatures make thermal decomposition of PAN a likely source of  $NO_x$ . This could be a decisive factor in harmonising modelled and recorded  $O_3$  mixing ratios as an increase in VOC emissions related to BB is unlikely to lead to  $O_3$  production in the absence of  $NO_x$ .

Uncertainties on VOC and  $NO_x$  emissions by BB and misrepresentations of  $NO_x$  recycling during transport are both likely contributors to the misrepresentation of  $O_3$  mixing ratios at the location of RUN.

# 4.3 Plume dispersion over the SWIO

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335 Transport of BB plumes recorded by the hs-PTR-MS at RUN takes place primarily in the lower FT. This implies that dispersion of the plume into the MBL is possible through turbulent mixing in shallow cumulus clouds and development of the MBL. Figure 5 shows  $\Delta$ CO due to pyrogenic emissions from plumes between 4 and 16 days old (corresponding to the extremes of plume ages observed at Maïdo) as calculated with FLEXPART on the model output layer 0 – 500 m a.g.l. By using equation 3, estimates of  $\Delta CH_3CN$ ,  $\Delta CH_3COCH_3$ ,  $\Delta CH_3OH$  and  $\Delta C_6H_6$  in the pristine marine boundary layer environment were made 340 (Fig. 5). To illustrate the importance of these BB plumes on the MBL composition, these expected excesses are compared with background VOC measurements performed in the SWIO during the MANCHOT campaign that took place December 2004 (Colomb et al., 2009). Shipborne measurements of VOC concentrations were performed South of La Réunion to characterise the impact of oceanic fronts on MBL composition (Colomb et al., 2009). We use background measurements North (zone I, 24.2° - 30.2° S) and South (zone III, 45.9° - 49.2° S) of the different oceanic fronts that were under consideration (Colomb et al., 2009). Due to the higher concentrations of anthropogenic tracers in zone I of the campaign, it was suggested that there may have been an impact of African outflow on these backgrounds (Colomb et al., 2009). Note that MANCHOT took place in December 2004, which is typically the end of the BB season over the SWIO. Due to the long lifetime of CH<sub>3</sub>CN and to a lesser extent CH<sub>3</sub>COCH<sub>3</sub>, part of these concentrations in zone I may be originating from accumulation of BB plumes in the troposphere.

The low variability in EnRs, between different BB intrusions at RUN, for both CH<sub>3</sub>CN and CH<sub>3</sub>COCH<sub>3</sub> allows for characterisation of mixing ratios in the marine boundary layer with small relative uncertainties (8.3% and 13.5% respectively). The local impact of  $\Delta$ CH<sub>3</sub>CN in the SWIO MBL during the August BB episodes ( $\sim$  50 pptv) constitutes an increase of  $\sim$  60 – 150% over the SWIO as measured during the MANCHOT campaign (zone I:  $80\pm20$  pptv, zone III:  $20\pm10$  pptv). Acetone excesses are based on the assumption that acetone production in the BB plume is similar in the free troposphere and in the marine boundary layer. The excesses over the SWIO can reach up to 300 pptv,  $\sim$  30 – 75% above the backgrounds recorded during MANCHOT.

The relatively short lifetimes of  $CH_3OH$  and  $C_6H_6$  result in a larger variability of the enhancement ratios between different BB intrusions. This is reflected in the larger relative uncertainty in the calculated excesses over the SWIO (21.7% and 32.6% for  $CH_3OH$  and  $C_6H_6$  respectively). Calculated  $\Delta CH_3OH$  over the SWIO are  $\sim 0.5$  ppbv, corresponding to an increase of 25% (zone I) to at least 100% (zone III) compared to the values recorded during MANCHOT (Colomb et al., 2009). The expected  $\Delta C_6H_6$  over the SWIO is 30 pptv. This is only a minor increase compared to zone I of the MANCHOT campaign (160 $\pm$ 40 pptv) but constitutes a significant increase (150%) in zone III, further south over the SWIO.

Due to the short lifetime of carboxylic acids in the humid marine boundary layer, the method used above to estimate the BB impact on the SWIO is not valid for HCOOH CH<sub>3</sub>COOH.

## 5 Conclusions

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We have shown that BB plumes were recorded with the hs-PTR-MS instrument deployed at the high-altitude Maïdo observatory located in the South-West Indian Ocean. Six different episodes of biomass burning plumes have been identified and studied in August 2018 and August 2019. Enhancement ratios relative to CO have been calculated for CH<sub>3</sub>CN (1.61 – 2.06 pptv ppbv<sup>-1</sup>), 4700 HCOOH (17.5 – 33.8 pptv ppbv<sup>-1</sup>), CH<sub>3</sub>COCH<sub>3</sub> (6.84 – 10.0 pptv ppbv<sup>-1</sup>), CH<sub>3</sub>COOH (9.8 – 18.0 pptv ppbv<sup>-1</sup>), C<sub>6</sub>H<sub>6</sub> (0.27 – 0.83 pptv ppbv<sup>-1</sup>), CH<sub>3</sub>OH (8.7 – 18.8 pptv ppbv<sup>-1</sup>) and O<sub>3</sub> (410 – 640 pptv ppbv<sup>-1</sup>). Comparison between these EnRs and the ERs calculated from literature showed production of CH<sub>3</sub>COCH<sub>3</sub> and HCOOH. Secondary production of CH<sub>3</sub>COCH<sub>3</sub> was accounted for by pyrogenic emission of precursor species propane and to a lesser extent i-butane and i-butene. Production was especially significant for HCOOH with EnRs about 10 times larger than the ERs. This HCOOH production can not be accounted for by known precursor species.

The CAMS NRT atmospheric composition service was shown to reproduce well the CO concentrations at RUN both during and outside BB episodes. In contrast, O<sub>3</sub> concentrations were only correctly reproduced outside the BB episodes. The large underestimation of O<sub>3</sub> concentrations during the BB episodes were linked to i) large uncertainties in VOC and NO<sub>x</sub> emissions and ii) misrepresentation of NO<sub>x</sub> recycling during transport of the BB plume in the CAMS NRT service. FLEXPART-AROME mesoscale backtrajectory simulations showed that biomass burning plumes were diluted at the observatory when the impact of PBL air increased. This implies that the BB plume recorded at the observatory is primarily transported through the FT. Large scale transport of  $\Delta$ CO originating from pyrogenic emissions, simulated with FLEXPART supported this by showing larger  $\Delta$ CO concentrations at higher altitudes. Finally, the horizontal distribution of  $\Delta$ CO in the SWIO MBL — calculated with FLEXPART — is multiplied with the EnR values inferred from data at the Maïdo observatory. This provided estimates for the impact of BB on air mass composition in the MBL over the SWIO. We compared the calculated estimates with background VOC measurements in the region reported in literature. Expected excesses for CH<sub>3</sub>CN, CH<sub>3</sub>COCH<sub>3</sub>, H<sub>6</sub>H<sub>6</sub> and CH<sub>3</sub>OH represent an increase of background concentrations by 60 - 150%, 30 - 75%, 15 - 150% and 25 - >100%, respectively. In the future, synchronous VOC measurements at RUN and marine campaigns should be conducted in order to i) better quantify the Ocean-Atmosphere interaction in regions with locally enhanced atmospheric concentrations of these species from BB and ii) identify the different ageing mechanisms during transport in the MBL compared to transport in the FT. This would be especially valuable for CH<sub>3</sub>COCH<sub>3</sub> and CH<sub>3</sub>OH, for which the role of the ocean on the total atmospheric budget remains uncertain.

Data availability. The core hs-PTR-MS dataset is available at https://octave.aeronomie.be/index.php/datasets (last access August 2020)can be downloaded from https://doi.org/10.18758/71021059. Other data is available upon request.

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tion and management, R.V., T.K.K. and C.F.L.; FLEXPART and FLEXPART-AROME simulations, B.V. and J.B.; original draft preparation, B.V.; review and editing, C.A., J.-F.M., T.S., J.B., R.V., N.S. and A.C.

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#### Appendix A: In situ measurement data visualisation

#### A1 Seasonal biomass burning profile

Hourly averages of CO and CH<sub>3</sub>CN are shown in Fig. A1. Both CO and CH<sub>3</sub>CN have large peak values from August to November. This corresponds to the biomass burning season as determined from ground-based remote-sensing data studies performed at La Réunion (Duflot et al., 2010; Vigouroux et al., 2012). The analysis presented in this study focuses on the first biomass burning intrusions measured for each season. The motivation for this choice is that the variability in diel profiles between different days is less pronounced during this period and backgrounds do not suffer from accumulated BB tracers for compounds with long atmospheric lifetimes.

# A2 Austral winter variation of in situ measurements at RUN

The temporal evolution of biomass burning plume constituents during austral winter 2018 and austral winter 2019 are shown together with the diel distribution of hourly averaged mixing ratio from Fig. A2 to Fig. A10. The median diel profile is used as an estimate of background variation above which the biomass burning excesses are determined. This works especially well for compounds with relatively small variability between different days compared to the excesses due to biomass burning (e.g. CO, CH<sub>3</sub>CN, HCOOH, CH<sub>3</sub>COCH<sub>3</sub> and CH<sub>3</sub>COOH) but may introduce errors for other compounds (e.g. C<sub>6</sub>H<sub>6</sub>, CH<sub>3</sub>OH and O<sub>3</sub>).

When this difference becomes negligible, the analysis no longer works and these compounds are not considered (e.g. HCHO

and CH<sub>3</sub>CHO).

# Appendix B: NO<sub>2</sub> coincidence with FLEXPART simulations

NO $_2$  volume mixing ratios from the CU MAX-DOAS instrument are generally lower than 100 pptv outside of the BB episodes. A notable exception to this is 3 August 2019 when it reaches  $\sim$  280 pptv. This coincides with slightly elevated  $\Delta$ CO signals simulated by FLEXPART at RUN (Fig. A11). At the visible wavelengths, the horizontal spatial scale probed is about 40 km and the overlap with the PBL is only a few km. As a results, measurements from the CU MAX-DOAS instrument are expected to compare well to the FLEXPART and CAMS models which have a low spatial resolution. Remark that when the NO $_2$  mixing ratio from the CU MAX-DOAS instrument is above 100 pptv, FLEXPART  $\Delta$ CO is generally enhanced between 1000 – 1500 m a.g.l.

As the plume on 3 August is not clearly observed in the in situ measurements we assume that it is not well mixed with boundary layer air at RUN and do not investigate it further here.

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**Table 1.** In situ routine observations at RUN and their respective limits of detection (LoD) and statistical uncertainties ( $\sigma$ ). (Zhou et al., 2018; Duflot et al., 2019).

Compound	Instrument	Technique	LoD [ppbv]	σ [ppbv]
СО	Picarro G2401	Cavity ring down spectroscopy	1	1.5
$O_3$	Thermo Scientific model 49i	UV photometric analyser	0.05	1

**Table 2.** A list of mass-to-charge ratios (m/z) observed in multiple ion detection mode by the hs-PTR-MS at RUN with the associated chemical compounds, dwell times and information about the instrument calibration limit of detection (LoD) per compound (Y: yes, N: no, N/A: not applicable). Dwell time is shown for 1 cycle. The LoD is computed for the hourly averages used here. The corresponding accumulated dwell times are about 22 time the stated dwell times of an individual cycle.

m/z	Compound	Dwell time [s]	Calibrated	LoD [pptv]
21	$H_3^{18}O^+$	2	N/A	N/A
31	formaldehyde (HCHO)	10	Y	100
32	$\mathrm{O}_2^+$	0.1	N/A	N/A
33	methanol (CH <sub>3</sub> OH)	10	Y	40
37	$\mathrm{H_{3}O^{+}.H_{2}O}$	0.1	N/A	N/A
42	acetonitrile (CH <sub>3</sub> CN)	10	Y	1
45	acetaldehyde (CH <sub>3</sub> CHO)	10	Y	18
47	formic acid (HCOOH)	10	N	50
59	acetone (CH <sub>3</sub> COCH <sub>3</sub> )	10	Y	4
61	acetic acid (CH <sub>3</sub> COOH)	10	N	7
63	dimethyl sulphide sulfide (DMS)	10	Y	6
69	isoprene $(C_5H_8)$	10	Y	5
71	methyl vinyl ketone (MVK)/ methacrolein (MACR)/	10	Y	2
	hydroxy hydroperoxides from isoprene (ISOPOOH)			
73	methyl ethyl ketone (MEK)	10	Y	3
79	benzene (C <sub>6</sub> H <sub>6</sub> )	10	Y	2
81	sum of monoterpenes <sup><math>\alpha</math></sup> ( $C_{10}H_{16}$ )	10	Y	5
93	toluene $(C_7H_8)$	10	Y	7
107	xylenes $^{\alpha}$ (C <sub>8</sub> H <sub>10</sub> )	10	Y	7
137	sum of monoterpenes $^{\alpha}$ (C <sub>10</sub> H <sub>16</sub> )	10	Y	8

 $<sup>^{\</sup>alpha}$ o-xylene and limonene were used for calibration.

**Table 3.** Pearson correlation coefficients (r) between the excess of chemical compound X ( $\Delta X$ ) and the excess of the typical BB marker  $CH_3CN$  during the BB episodes.

X	r	X	r
СО	0.98	DMS	0.60
НСООН	0.89	НСНО	0.55
$CH_3COCH_3$	0.88	MEK	0.39
$CH_3COOH$	0.87	CH <sub>3</sub> CHO	0.12
$O_3$	0.83	$C_5H_8$	-0.08
$C_6H_6$	0.81	MVK/MACR/ISOPOOH	-0.22
$CH_3OH$	0.71		

**Table 4.** Mean background mixing ratio ppbvduring the daytime (10 a.m. – 4 p.m., reflecting planetary boundary layer air composition) and nighttime (10 p.m. – 4 a.m., measuring free tropospheric air masses),  $\mu_{PBL}$  [ppby] and  $\mu_{FT}$  [ppby] respectively, and mean excesses [%] during the Aug 2018 and Aug 2019 BB intrusions. The reported background values are mean mixing ratios recorded during austral winter (June, July, August), excluding BB incidents.

	$\mu_{\mathrm{PBL}}  [\mathrm{ppbv}]$	$\mu_{\rm FT}  [{\rm ppbv}]$	03-05 Aug 2018	08-14 Aug 2018	17-19 Aug 2018	$\mu_{\mathrm{PBL}} \; [\mathrm{ppbv}] \qquad \mu_{\mathrm{FT}} \; [\mathrm{ppbv}]  03-05 \; \mathrm{Aug} \; 2018  08-14 \; \mathrm{Aug} \; 2018  17-19 \; \mathrm{Aug} \; 2018  06-08 \; \mathrm{Aug} \; 2019  10-11 \; \mathrm{Aug} \; 2019  15-18 \; \mathrm{Aug} \; 2019  10-11 \; \mathrm{Aug} \; 2019  10-11 \; \mathrm{Aug} \; 2019  10-11 \; \mathrm{Aug} \; 2010  10-$	10-11 Aug 2019	15-18 Aug 2019
$CH_3CN$	0.091 (0.035) 0.092 (0.042)	0.092 (0.042)	%96	164%	%66	182%	128%	103%
00	67 (15)	61 (17)	81%	129%	78%	140%	110%	%89
НСООН	0.87 (0.62)	0.40 (0.62)	466%	9089	285%	942%	515%	379%
$\mathrm{CH_{3}COCH_{3}}$	0.40 (0.17)	0.30 (0.19)	153%	221%	118%	227%	157%	147%
$\mathrm{CH_{3}COOH}$	0.40 (0.30)	0.17 (0.31)	%602	863%	398%	1155%	754%	450%
${\sf C_6H_6}$	0.025 (0.014)	0.025 (0.014) 0.012 (0.012)	303%	314%	123%	360%	274%	156%
$CH_3OH$	1.2 (0.42)	0.63 (0.36)	131%	163%	63%	188%	132%	101%

Table 5. Calculated EnRs [pptv ppbv<sup>-1</sup>] — relative to CO — for the identified BB intrusions. Between brackets parentheses are the standard deviations of the EnR [pptv ppbv $^{-1}$ ] obtained from the linear fit.

	3-5 Aug 2018	8-14 Aug 2018	3-5 Aug 2018 8-14 Aug 2018 17-19 Aug 2018 6-8 Aug 2019 10-11 Aug 2019 15-18 Aug 2019	6-8 Aug 2019	10-11 Aug 2019	15-18 Aug 2019
$CH_3CN$	1.61 (0.02)	1.71 (0.01)	1.69 (0.04)	1.79 (0.01)	1.69 (0.04)	2.06 (0.03)
НСООН	29.3 (0.5)	23.5 (0.3)	17.5 (0.5)	33.8 (0.5)	24.6 (0.9)	31.2 (0.6)
$\mathrm{CH_{3}COCH_{3}}$	8.85 (0.22)	7.76 (0.13)	6.84 (0.14)	7.85 (0.08)	7.06 (0.26)	10.0 (0.3)
$CH_3COOH$	18.0 (0.3)	12.9 (0.2)	9.8 (0.3)	16.3 (0.2)	13.2 (0.5)	13.7 (0.2)
03	640 (19)	438 (9)	635 (27)	461 (9)	410 (20)	422 (16)
$\mathrm{C_6H_6}$	0.83 (0.04)	0.46 (0.01)	0.27 (0.01)	0.50(0.01)	0.50 (0.02)	0.46 (0.02)
$CH_3OH$	18.8 (0.6)	13.5 (0.4)	8.7 (0.5)	15.9 (0.3)	14.2 (0.8)	16.8 (0.7)

Table 6. ER [pptv ppbv<sup>-1</sup>] — relative to CO — for the 3 most probable fuel types (Savanna and grassland, tropical forest and agricultural residue) of the BB plume sampled at RUN. Between brackets parentheses is the uncertainty obtained by combining the standard deviations of the EFs recorded by Andreae (2019).

	CH <sub>3</sub> CN	НСООН	СН3СООН	CH <sub>3</sub> COCH <sub>3</sub>	$C_6H_6$	CH <sub>3</sub> OH
Savanna and grassland	1.68 (1.62)	1.68 (1.62) 1.85 (1.59)	15.6 (10.3)	3.28 (2.24)	1.72 (0.87)	17.1 (21.1)
Tropical forest	3.21 (3.10)	2.87 (2.47)	14.8 (9.76)	2.92 (-)	1.31 (0.664)	23.5 (29.1)
Agricultural residue	2.24 (2.17)	4.48 (3.86)	37.4 (24.7)	4.5 (3.07)	1.27 (0.646)	38.0 (46.9)
Vigouroux et al. $(2012)^{\alpha}$		4.6 (0.3)				11.6 (0.6)
de Gouw et al. $(2006)^{\beta}$	1.18 (0.14) - 3.24 (0.09)		0.9 (0.3) - 12.9 (0.5)	0.9 (0.3) - 12.9 (0.5) $2.6 (0.3) - 22.8 (1.0)$	0.8(0.2) - 1.41(0.04)	2(2) - 21(2)
Lefer et al. $(1994)^{\gamma}$		8.2 - 62	6 – 34			
This work $^\delta$	1.76 (0.146) 26.7 (5.44)	26.7 (5.44)	14.0 (2.61)	8.06 (1.09)	0.50 (0.16)	14.6 (3.18)
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 $<sup>^{\</sup>alpha}$  EnRs derived from FTIR measurements at RUN.

 $<sup>^{\</sup>beta}$  EnR ranges from 11 forest fire plumes Sampled during NEAQS-ITCT 2k4.

 $<sup>^{\</sup>gamma}$  EnR ranges from 10 subarctic forest fire plumes sampled during ABLE 3B.

 $<sup>^{\</sup>delta}$  This work, mean EnR and the standard deviation.

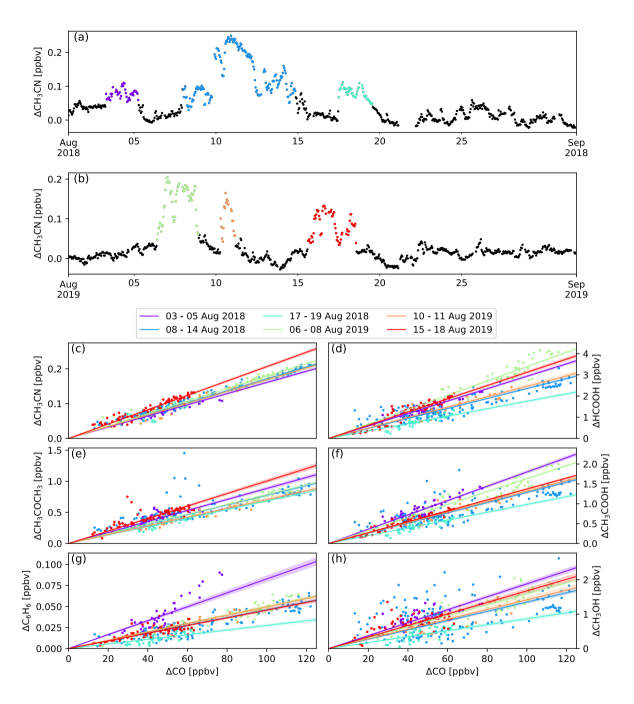
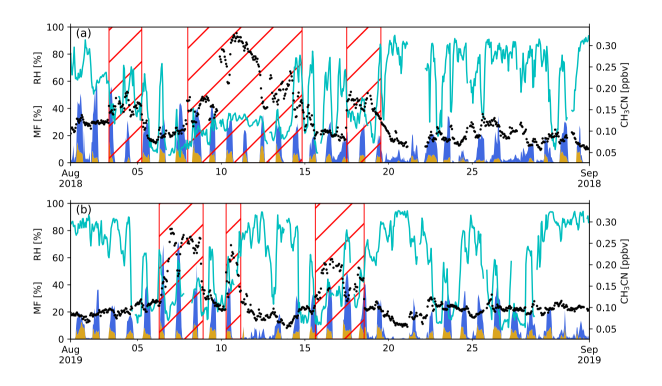


Figure 1. The top two panels (a and b) show the six BB intrusions identified using  $\Delta$ CH<sub>3</sub>CN (purple: 03 – 05 Aug 2018, blue: 8 – 14 Aug 2018, cyan: 17 – 19 Aug 2018, green: 6 – 8 Aug 2019, orange: 10 – 11 Aug 2019, red: 15 – 18 Aug 2019). The bottom six panels show the EnR fits for CH<sub>3</sub>CN (c), HCOOH (d), CH<sub>3</sub>COOH (e), CH<sub>3</sub>COCH<sub>3</sub> (f), C<sub>6</sub>H<sub>6</sub> (g) and CH<sub>3</sub>OH (h). EnRs are normalised to the excess mixing ratio of CO for the six intrusions in the colors used in the top two panels. Uncertainty on the linear regression is shown as a coloured band around the curves.



**Figure 2.** Measured relative humidity (RH, cyan curve), CH<sub>3</sub>CN mixing ratio [ppbv] (black points) and modelled mesoscale MF (fraction of time air parcels are situated over a certain source in the PBL proxy from Lesouëf et al. (2011)) [%] from 24 hour backtrajectories using FLEXPART-AROME in August 2018 (a) and August 2019 (b). Blue denotes the marine boundary layer MF, brown represents the island surface PBL MF. The hatched red area represents the different BB intrusions.

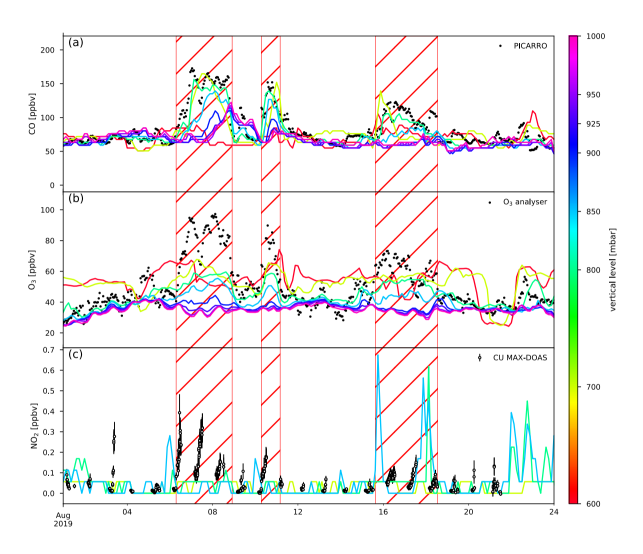
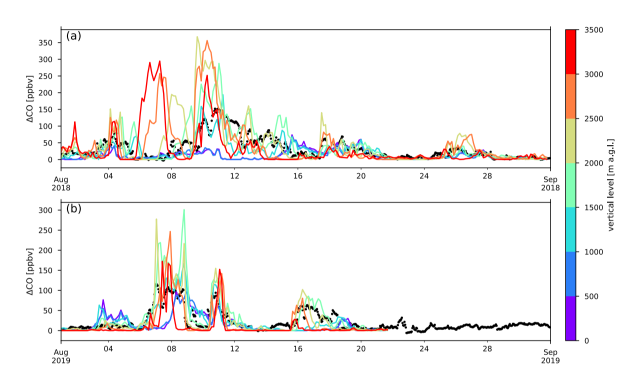


Figure 3. Comparison between measured (black dots) and calculated mixing ratios (coloured lines) of CO (a),  $O_3$  (b) and  $O_2$  (c) from CAMS. The coloured lines indicate the lowest eight pressure levels in the model (1000, 950, 925, 900, 850, 800, 700 and 600 mbar). The hatched red area represents the different BB intrusions.



**Figure 4.** Comparison between measured excess CO (black) at RUN to that modelled on different vertical levels of the FLEXPART model during August 2018 (a) and August 2019 (b). FLEXPART output levels are defined in meters above ground level (m a.g.l.).

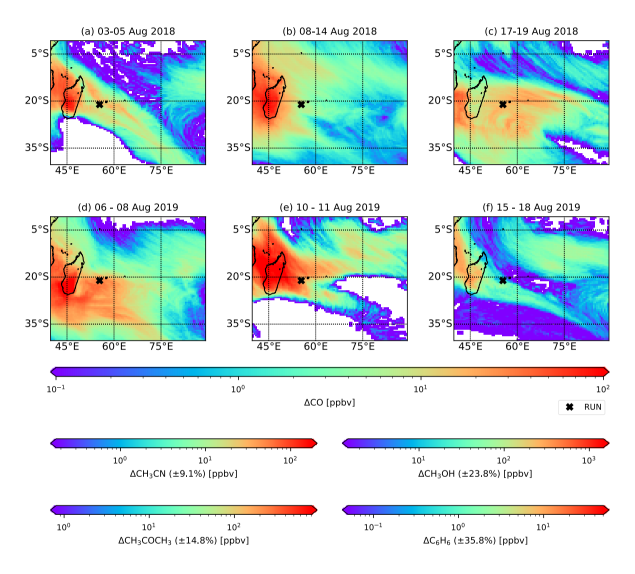


Figure 5. Excess CO over the South West Indian Ocean between 0 and 500 m a.g.l. from BB emissions as simulated by FLEXPART. Additional color scales quantify the projected  $CH_3CN$ ,  $CH_3COCH_3$ ,  $C_6H_6$  and  $CH_3OH$  excesses.

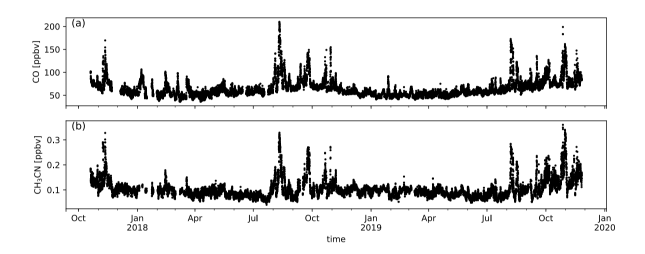
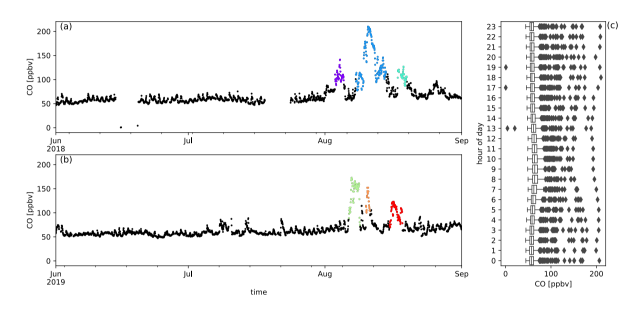
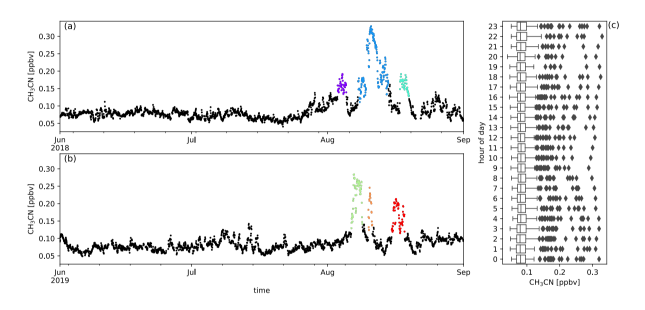


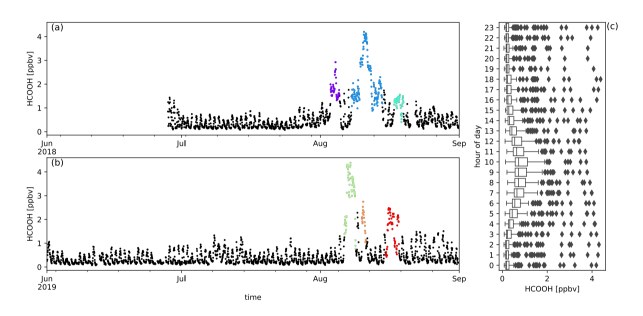
Figure A1. Measured CO (a) and CH<sub>3</sub>CN (b) mixing ratios [ppbv] during the deployment of the hs-PTR-MS for the OCTAVE project.



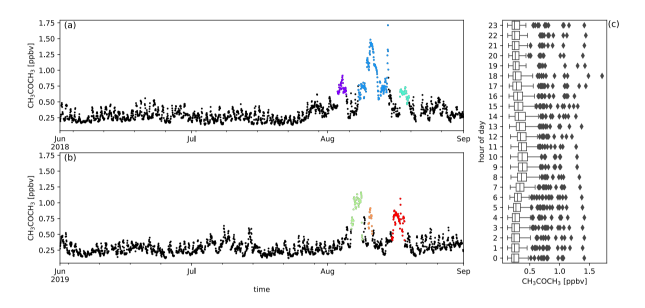
**Figure A2.** Measured CO mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.



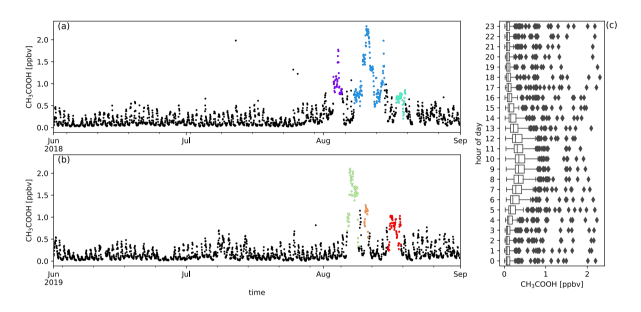
**Figure A3.** Measured CH<sub>3</sub>CN mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.



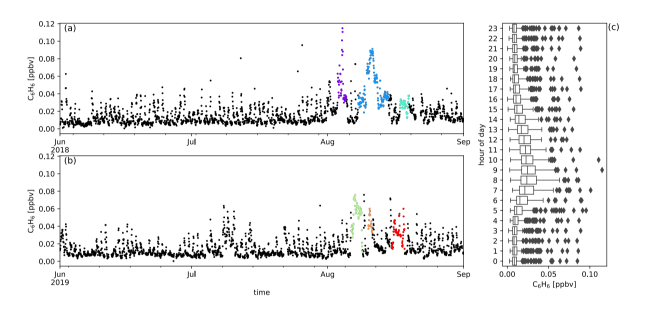
**Figure A4.** Measured HCOOH mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.



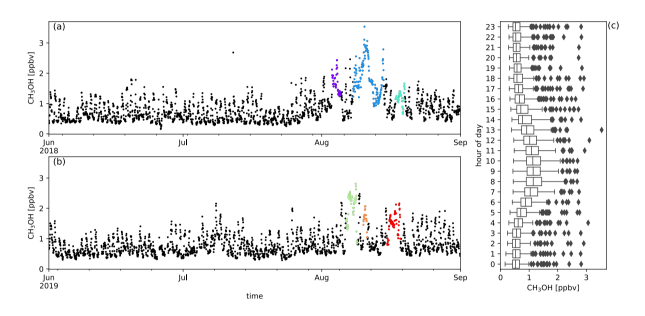
**Figure A5.** Measured CH<sub>3</sub>COCH<sub>3</sub> mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.



**Figure A6.** Measured CH<sub>3</sub>COOH mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.



**Figure A7.** Measured  $C_6H_6$  mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.



**Figure A8.** Measured CH<sub>3</sub>OH mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.

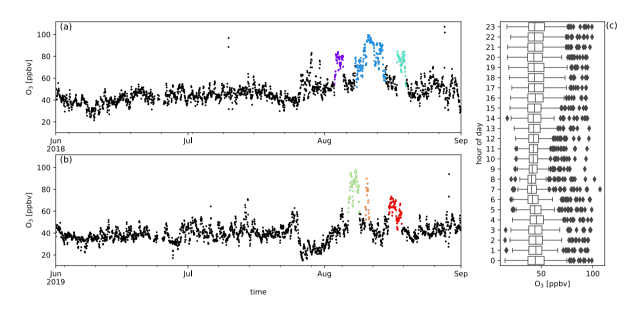
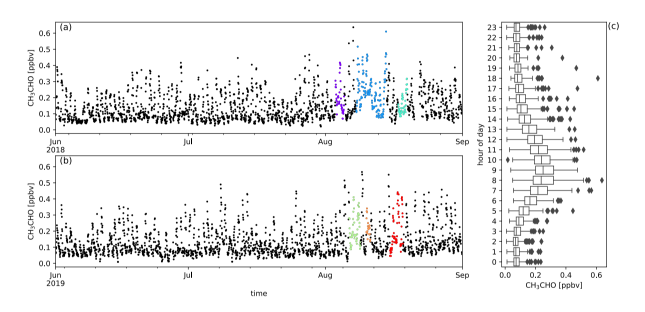


Figure A9. Measured  $O_3$  mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.



**Figure A10.** Measured CH<sub>3</sub>CHO mixing ratios [ppbv] during austral winter 2018 (a) and austral winter 2019 (b) together with the diel distribution of hourly averages (c). Biomass burning plumes under investigation are highlighted in colors.

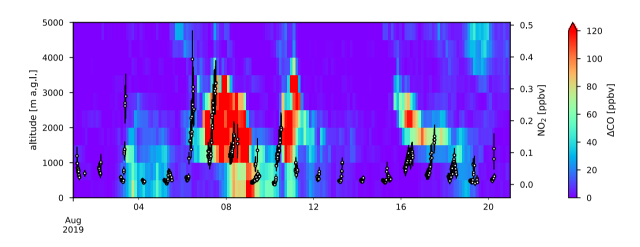


Figure A11. NO<sub>2</sub> [ppbv] from CU MAX-DOAS in comparison with  $\Delta$ CO [ppbv] from FLEXPART simulations. Altitude [m a.g.l.] corresponds with vertical layers in the FLEXPART output at the location of RUN. The ground level of RUN in the model is 284 m a.s.l.