Investigation of 3D radiative transfer effects for UV / vis satellite and ground based observations of volcanic plumes

Abstract. We investigate effects of the 3-dimensional (3D) structure of volcanic plumes on the retrieval results of satellite

5 Thomas Wagner¹, Simon Warnach^{1,2}, Steffen Beirle¹, Nicole Bobrowski^{2,3}, Adrian Jost¹, Janis Puķīte¹, and Nicolas Theys⁴

Correspondence to: Thomas Wagner (thomas.wagner@mpic.de)

and ground based UV-vis observations. For the analysis of such measurements usually 1D scenarios are assumed (the 15 atmospheric properties only depend on altitude). While 1D assumptions are well suited for the analysis of many atmospheric phenomena, they are usually less appropriate for narrow trace gas plumes. For UV/vis satellite instruments with large ground pixel sizes like GOME-2, SCIAMACHY, or OMI, 3D effects are of minor importance, but usually these observations are not sensitive to small volcanic plumes. In contrast, observations of TROPOMI aboard Sentinel-5P have a much smaller ground pixel size (3.5 x 5.5 km²). Thus on the one hand, TROPOMI can detect much smaller plumes than previous instruments. On the other hand 3D effects become more important, because the TROPOMI ground pixel size is smaller than the height of the troposphere and also smaller than horizontal atmospheric photon path lengths in the UV/vis spectral range. In this study we investigate the following 3D-effects using Monte-Carlo radiative transfer simulations: 1. the light mixing effect caused by horizontal photon paths, 2. the saturation effect for strong SO₂ absorption, 3. geometric effects related to slant illumination and viewing angles, and 4. Plume side effects related to slant illumination angles and photons reaching the 25 sensor from the sides of volcanic plumes. Especially the first two effects can lead to a strong and systematic underestimation of the true trace gas content if 1D retrievals are applied (more than 50% for the light mixing effect, and up to 100% for the saturation effect). Besides the atmospheric radiative transfer, the saturation effect also affects the spectral retrievals. Geometric effects have a weaker influence on the quantitative analyses, but can lead to a spatial smearing of elevated plumes or even to virtual double plumes. Plume side effects are small for short wavelengths, but can become large for longer wavelengths (up to 100% for slant viewing and illumination angles). For ground based observations, most of the above mentioned 3D effects are not important, because of the narrow FOV and the closer distance between the instrument and the volcanic plume. However, the light mixing effect shows a similar strong dependence on the horizontal plume extension as for satellite observations and should be taken into account for the analysis of ground based observations.

35 1 Introduction

SO₂ emitted from volcanoes has been observed from satellites since about 40 years. After successful first detections for large eruptions by the TOMS instrument (Krueger, 1983), subsequent UV/vis satellite instruments with continuous spectral coverage allowed also the observation of smaller plume amounts (Eisinger and Burrows, 1998; Afe et al., 2004, Khokhar et al., 2005, Krotkov et al., 2006; Yang et al., 2007, 2010; Nowlan et al., 2011; Rix et al., 2012; Hörmann et al., 2013; Li et al., 2013; Penning de Vries et al., 2014; Theys et al., 2015, Fioletov et al., 2016; Zhang et al., 2017; Theys et al., 2017, 2019, 2021a). Furthermore, besides SO₂, also other trace gases like BrO, OCIO and IO in volcanic plumes could be analysed from these observations (Theys et al., 2009; Heue et al., 2011; Rix et al., 2012; Hörmann et al., 2013; Theys et al., 2014; Schönhardt et al., 2017; Suleiman et al., 2019).

¹Satellite Remote Sensing Group, Max Planck Institute for Chemistry, Mainz, Germany

²Institute of Environmental Physics, University of Heidelberg, Heidelberg, Germany

³Istituto Nazionale Geofisica e Vulcanologia Catania, Italy

^{10 &}lt;sup>4</sup>Royal Belgian Institute for Space Aeronomy (BIRA-IASB), Brussels, Belgium

Since the launch of the GOME-1 instrument aboard ERS-2 in 1995 (Burrows et al., 1999), the ground pixel size of UV/vis sensors has strongly decreased from 40 x 320 km² (GOME-1) down to 3.5 x 5.5 km² (TROPOMI), see Table 1. The ground pixel size of TROPOMI is thus only slightly larger or even similar to the extension of plumes emitted from point sources, in particular to those from small/medium volcanic eruptions or passive degassing. Thus with TROPOMI, many small/medium volcanic plumes have become detectable, which could not be detected with the former instruments (also the high signal to 50 noise ratio of TROPOMI contributes to the increased sensitivity to small volcanic plumes). An example of TROPOMI observations of SO₂ and BrO for a narrow volcanic plume is shown in Fig. 1 (Warnach, 2022). Shown are the slant column densities (SCD, commonly interpreted as the integrated concentrations along the light path) of both trace gases.

To illustrate the effects of this improved horizontal resolution, we compare the expected trace gas absorptions of small/medium volcanic plumes for different satellite instruments. As a reference, we take the GOME-2 instrument, because from GOME-2 observations it was possible for the first time to observe enhanced BrO amounts for several volcanic plumes (Hörmann et al., 2013). A typical detection limit for the SO₂ SCD in the SO₂ standard fit range (about 312 to 324 nm, Hörmann et al., 2013; Theys et al., 2017, 2021b) is about 1×10^{16} molec/cm². The detection limit is similar for the different instruments, but depends slightly on the signal to noise ratio (see Theys et al., 2019). A SCD of 1×10^{16} molec/cm² corresponds to a differential SO₂ optical depth of about 0.001 when measured with a FWHM of about 0.5 nm, as for the 60 TROPOMI instrument. We will use this optical depth as detection limit for SO₂ in this study. This choice is somewhat arbitrary, but it will serve as a realistic reference point. The detection limit will probably improve in the future using advanced analyses techniques, see e.g. the recent study by Theys et al. (2021a).

55

65

70

75

If we use the above mentioned SO_2 SCD of 1×10^{16} molec/cm² for a GOME-2 observation (as a reference case) and assume that the horizontal extension of an observed volcanic plume is smaller than or equal to the size of a TROPOMI ground pixel (e.g. 1 x 1 km² compared to 3.5 x 5.5 km²), we can estimate the SO₂ SCDs observed for instruments with different ground pixel sizes assuming that the measured SCD scales according to the geometric coverage of the volcanic plume. We will see later (light mixing effect) that assuming simple geometry does not perfectly describe these observations, but nevertheless provides a good estimate for the overall effect. The resulting SO₂ SCDs for the different satellite ground pixels are shown in Table 1. The SO₂ SCD for TROPOMI can be higher than the SO₂ SCD for GOME-2 observations by more than 2 orders of magnitude. Even compared to OMI the increase is larger than a factor of 10.

The results of these simple calculations indicate the great potential of TROPOMI observations for the detection of small/medium volcanic plumes; many weak plumes on the scale of TROPOMI ground pixels were invisible for previous sensors. The corresponding increase in the frequency of detection is difficult to quantify, because the exact frequency distribution of plume sizes and amounts of SO₂ and aerosols is not known. In addition, also the self-shielding of plumes by aerosols or clouds formed from the plume probably depends on the plume sizes themselves. Nevertheless, we can quantify the increase in the detection frequency from the satellite observations themselves. For example, Hörmann et al. (2013) found 220 volcanic SO₂ plumes per year in GOME-2 data, whereas Warnach (2022) found 870 SO₂ plumes per year in TROPOMI data (both studies cover different time periods). In view of the SCD ratios discussed above (see Table 1), this increase by a factor of about 4 in the detection frequency seems rather small (see also Theys et al., 2019). The main and simple reason for the small increase of the number of detections is that many volcanic plumes are larger than the size of a TROPOMI pixel. For such plumes, the SO₂ SCD detected by TROPOMI will be similar to those observed by instruments with larger pixels, up to the size of the plume itself.. However, another reason for the rather low increase of detections is related to 3D effects which are the topic of this paper. As shown below, for TROPOMI observations, 3D effects can cause an underestimation of the true trace gas content of the plume by more than 50 % (light mixing effect). For plumes with strong SO₂ absorptions, the underestimation can become even larger.

3D effects are fundamental effects and do not only affect UV/vis satellite observations with small ground pixel sizes like TROPOMI. However, for instruments with large ground pixel sizes, 3D effects are typically much smaller than for TROPOMI, and the related errors are typically ignored, because they are smaller than other measurement uncertainties (e.g. related to aerosols or the layer height). For small ground pixel sizes (like for TROPOMI), however, the importance of 3D effects increases because of the following reasons:

- 1) the horizontal pixel dimensions are typically smaller (except for situations with low visibility) than atmospheric photon path lengths (see table 1). Thus light scattered horizontally across the borders of the satellite ground pixels becomes important.
- 2) the horizontal pixel dimensions are smaller than the vertical extent of the troposphere. Thus geometric effects related to slant illumination and viewing geometry become important.
 - 3) the horizontal pixel dimensions are similar to those of volcanic plumes. Thus the relative contribution of photons from the sides of the plume increases.
 - 4) one TROPOMI ground pixel might cover the very early part of a volcanic plume, for which the SO_2 concentration can be extremely high. Due to the strong SO_2 absorption, photons will then not penetrate the full plume.
- Based on these general aspects, different specific 3D effects can be deduced. In this study we investigate and quantify four specific 3D effects. Two of them (light mixing effect and geometrical effect) were already investigated by Schwärzel et al. (2020, 2021) for ground based and aircraft measurements of NO₂. In this study, we extend these investigations to satellite observations (with different ground pixel sizes). We also investigate two additional 3D effects (saturation effects and plume side effects). The four specific 3D effects are described in more detail below (see also Fig. 2):
- a) Light mixing effect: part of the detected photons originates from air masses outside the observed ground pixel (and also from outside the trace gas plume). This leads to a reduction of the trace gas absorption compared to the scenario of a horizontally extended plume (Fig. 2a). Here it should be noted that in case of spatially varying surface albedo or aerosol distributions, the light mixing effect could be further enhanced (over a dark surface surrounded by a bright surface) or decreased (over a bright surface surrounded by a dark surface), which is especially important for aerosol retrievals from satellite (Richter, 1990; Lyapustin and Kaufman, 2001). Also the contributions from different parts within the ground pixel depend on the brightness distribution within the pixel. But here we focus on scenarios with constant surface albedo.
- b) Saturation effect (Fig. 2b): for strong trace gas absorptions (especially for SO₂ in volcanic plumes) the exact spatial extent of the plume (depending e.g. on the mixing with air from outside) becomes important. If e.g. a trace gas amount is confined in a small part of the satellite ground pixel, the absorption in that part can become very strong. In extreme cases the backscattered light intensity can even approach zero. Then all light measured by the satellite will originate from the remaining part of the satellite ground pixel (outside the narrow plume) where no SO₂ absorption takes place. In such extreme cases, no trace gas absorption will effectively be observed by the satellite. If instead the same amount of molecules is distributed over a larger volume (and thus a larger fraction of the satellite ground pixel), an increased (but typically still too small) SO₂ aborption will be seen. Thus the effective trace gas absorption systematically depends on the horizontal extent of a plume and thus on the temporal evolution of the plume and its mixing with air from outside.
 - c) Geometric effects: for TROPOMI, the ground pixel size is smaller than the vertical extent of the troposphere. Thus, especially for elevated plumes, geometric effects caused by slant viewing directions and/or slant solar illumination angles can cause a smearing-out and apparent displacement of the volcanic plume compared to its true location. In extreme cases, even double plumes might result (Fig. 2c).
- d) Plume side effects: especially if aerosols are present within the plume, edge effects can become important. Their importance increases with decreasing plume size, because a larger fraction of the measured light then reaches the detector from the sides of the plume compared to the top (Fig. 2d). The plume side effect includes the light mixing effect, but adds the dependence on illumination and viewing angle.

Usually all four effects occur for satellite observations at the same time. But in this study we investigate them separately in order to assess their importance for specific measurement scenarios. For these investigations we use the 3D Monte Carlo model TRACY-2 (Wagner et al., 2007; Kern et al., 2010).

It should be noted that for ground based measurements, a fifth 3D effect becomes important, the so-called light dilution effect (Kern et al., 2010). It has contributions from the light mixing effect, but is mainly caused by light scattered into the line of sight of the instrument between the plume and the instrument without having crossed the (localised) trace gas plume.

Since for satellite observations, the observed light typically has not traversed the whole atmosphere, but part of it is scattered above the trace gas layer (leading to air mass factors (AMFs) below unity; for the definition of the AMF, see section 2), the light dilution effect is already implicitly considered in radiative transfer simulations (even in calculations of 1D AMFs). In this study the light dilution effect is thus explored as a separate effect only for ground based observations.

The paper is organised as follows. In section 2 the 3D Monte-Carlo radiative transfer model TRACY-2 is introduced.

Sections 3 to 6 present simulations of the above mentioned four 3D-effects. Section 7 discusses 3D-effects for ground based observations, and section 8 presents a summary and outlook.

2 The Monte Carlo radiative transfer model TRACY-2

155

165

170

145 The Monte Carlo radiative transfer model TRACY-2 was developed by Tim Deutschmann at University of Heidelberg, Germany. TRACY-2 allows the simulation of individual photon paths between the Sun and an observing instrument with flexible boundary conditions. The location and properties of the detector (e.g. width and shape of the field of view, FOV) can be freely chosen. The atmospheric properties, especially the trace gas and aerosol distributions, can be varied in all three dimensions. TRACY-2 is based on the backward Monte Carlo method: a photon emerges from a detector and undergoes various interactions in the atmosphere until the photon leaves the top of the atmosphere (where it is 'forced' into the sun with appropriate weighting factors). A large number of random photon paths is generated reproducing the light contributing to the simulated measurement. Depending on the atmospheric conditions and the FOV, the number of simulated photons in this study ranges from 10 thousand to 1 million. Air mass factors (AMFs) are derived from the modelled radiances with (I) and without (I₀) the absorber of interest:

$$AMF = \frac{\ln\left(\frac{I_0}{I}\right)}{VCD \cdot \sigma} \tag{1}$$

with σ being the absorption cross section of the considered trace gas and VCD the vertical column density (the vertically integrated trace gas concentration). Good agreement between TRACY-2 and other radiative transfer models was found in an extensive comparison exercise (Wagner et al., 2007). In Fig. 3 an exemplary simulation with TRACY-2 is shown for a TROPOMI observation of an idealised narrow plume.

For the simulations, an atmospheric domain extending $\pm 20^{\circ}$ in latitude and longitude around the plume location (centered at 0° longitude and 0° latitude) and from the surface to 1000 km altitude is chosen. The horizontal resolution is high close to the center (500 m from the center until 10 km) and coarser outside (11.1 km until 111 km; 55.5 km until 555 km, and one grid cell from 555 km to 2222 km). For some simulations (see Figs. 5 and 20) even a finer horizontal grid was used to investigate the effect of very narrow plumes (100 m until 22.5 km, 55.5 km until 555 km). The vertical resolution was set to 200 m below 6 km, 1 km until 25 km, 2 km until 45 km, and 5 km until 100 km altitude, and one layer from 100 km to 1000 km. The surface albedo was set to 5%. This value was chosen, because typical albedo values in the considered wavelength ranges over volcanic areas are close to this value (see Fig. A1.1 in appendix A1). The exact choice of the surface albedo is, nevertheless, not critical, because the ratio of AMFs for narrow plumes and those for horizontally extended plumes hardly

depends on the surface albedo (see Fig. A1.2 in appendix A1). Vertical profiles of temperature, pressure and O_3 concentration are taken from the US standard atmosphere (United States Committee on Extension to the Standard Atmosphere, 1976).

The satellite instrument is placed at 824 km above sea level representing the altitude of the TROPOMI instrument. For the investigation of the effects of very narrow plumes (Figs. 5 and 20) as well as for the plume scans (like in Fig. 6) a narrow FOV of 0.014° (corresponding to a ground pixel diameter of 200 m) is used. For the simulations of real satellite measurements, rectangular FOVs corresponding to the nominal ground pixel sizes of the different satellite instruments (see table 1) are used.

Volcanic plumes are defined for SO₂, BrO, and IO based on ground based and satellite observations (for details see section 2.1). Simulations are performed for selected wavelengths relevant for the considered trace gases: for SO₂, four wavelengths (313.1, 324.15, 332.0, 370.3 nm) are chosen for the dominant absorption bands of the different fit ranges (Theys et al., 2017). Simulations for BrO are performed at 340 nm, and for IO (and NO₂, and H₂O) at 440 nm. Note that substantial amounts of NO₂ are not expected in volcanic plumes. Also, the increase of the atmospheric waper vapor absorption due to water vapor inside the volcanic plume is expected to be very small. Nevertheless, because NO₂ and H₂O are often analysed in the same spectral range as IO, the IO results are also representative for H₂O and NO₂ observations if the ratios of the respective differential cross sections in the spectral range around 440 nm are applied (see section 2.1.3 and appendix A2). Thus the IO results could also be used for the potential detection of enhanced NO₂ or H₂O in volcanic (or other emission) plumes.

AMFs are calculated according to equation 1 for single wavelengths ('monochromatic AMFs'). Thus the wavelength dependence of the AMF (Marquardt et al. 2000; Pukīte et al., 2010, 2016) is not explicitly taken into account, which for strong absorbers usually causes an overestimation of the monochromatic AMFs compared to the true AMFs. In a case study (section 4.3) the effect of the wavelength dependence of the AMF was investigated. It was found that the wavelength dependence of the AMF is negligible for most scenarios (with weak absorptions) considered here. Only for scenarios with strong SO₂ absorptions (with optical depth larger than 0.05 to 0.1) the wavelength dependence of the AMF becomes important and further amplifies the underestimation of the true AMF. But such scenarios can easily be identified and excluded from the data analysis, and for such scenarios wavelengths with weaker SO₂ absorptions should be selected. Thus, the effect of the wavelength dependence of the AMF is ignored in this study, and the main findings can be derived from monochromatic AMFs (equation 1). While this strongly reduces the computational effort, the computation time for the 3D simulations still remains rather high. The calculation of a single monochromatic 3D AMF takes about 5 min on a state of the art notebook.

2.1 Selected trace gas scenarios

180

185

190

195

200

205

210

The construction of realistic trace gas scenarios is a challenging task. The assumed trace gas plumes should represent realistic distributions, which are based on ground based observations. They should also fit to satellite observations of narrow plumes, which, however, are often close to the detection limit, because the plumes only cover part of the satellite ground pixel. From the satellite observations no information on the sub-ground pixel scale can be obtained.

For this study, trace gas scenarios were set up for SO₂, BrO, and IO. BrO and IO are typically weak absorbers with absorption optical depths clearly below 0.1. Thus they do not significantly affect the atmospheric light path distribution, and only the relative spatial distribution, but not the absolute trace gas amount is important for the calculation of the AMFs. The scenarios for SO₂, BrO, and IO were chosen, because these trace gases were already observed in volcanic plumes. Moreover, the corresponding wavelengths cover the spectral range from about 310 to 440 nm, over which the probability of Rayleigh scattering strongly changes (by a factor of 4). Thus the light mixing effect (and also the geometric effect and the side

scattering effect) are expected to differ substantially for the chosen wavelengths. A second set of wavelengths is selected for the SO_2 scenarios (4 wavelengths covering the spectral range from about 313 to 370 nm, for details see section 2.14). But here we are not primarily interested in the effect of the wavelength dependence of Rayleigh scattering, but rather on the strong wavelength dependence of the SO_2 absorption cross section. This second set of wavelengths is used to study the saturation effect for SO_2 .

The trace gas scenarios defined below will be used to decide whether the trace gas absorption of a given measurement scenario will be above the detection limit or not. For that purpose we also chose the following detection limits: optical depths of 0.0005 for BrO and IO, and 0.001 for SO₂. Of course these choices are somewhat arbitrary, but represent realistic reference points for single observations. Here it should be noted that in some cases (e.g. for continuous degassing volcanoes and stable wind fields) many observations might be averaged. In such cases the detection limits will decrease for the statistical mean. Also, for possibly improved future algorithms and instruments the detection limit might be reduced. Then also observations with trace gas absorptions currently below the detection limit will become significant leading to a larger total number of significant observations. Thus the results obtained in this study should be seen as a general orientation. The differential cross sections (high-frequency amplitudes) used to convert the SCDs of the different trace gases into optical depths are given in Table A2.1 in the appendix A2. Here it should be noted that in our radiative transfer simulations the absorption cross sections were used.

230

235

240

2.1.1 Size of the volcanic plumes

Plume diameters close to volcanic vents are typically ≤ 1 km (see e.g. Bobrowski et al., 2003; Bobrowski and Platt, 2007; Dinger et al., 2021). Thus we assume as a standard scenario a plume with an extension of 1 x 1 x 1 km³. Of course this is a simplification, since realistic plumes close to the volcanic vent will be more complex (including e.g. a vertical updraft above the vent, and a horizontal outflow part). But here we are interested in the basic effects. In addition to the standard scenario, we also assume plumes with larger horizontal extensions, but with the same vertical extent and the same total amount of molecules. This allows us to study the light mixing effect and saturation effect depending on the turbulent dilution state of the plume. For simplicity we assume that the dilution will only occur in the horizontal dimensions. For most simulations (light mixing effect and saturation effect) we investigate plumes in four altitude ranges: 0-1 km, 5-6 km, 10-11 km, and 15-16 km. For the study of the geometric effects and the side scattering effects only plumes at specific heights are chosen to illustrate the general dependencies.

2.1.2 BrO

245

250

255

2.1.3 IO

- Also IO is a weak absorber, but because of the sparseness of IO observations in volcanic plumes (e.g. Schönhardt et al., 2017), we simply use the same VCD (5 · 10¹⁵ molec/cm²) as for BrO for the standard plume (1 x 1 x 1 km³) corresponding to a VCD of 2.6 · 10¹⁴ molec/cm² for a TROPOMI observation. This value probably overestimates the true IO amounts in volcanic plumes. But since IO is a weak absorber, the results are representative also for plumes with smaller IO amounts. The corresponding VCDs for the different horizontal plume extensions are shown in Table 3.
- The results of the IO scenarios could be also transferred to measurements of NO_2 and H_2O , if they were analysed in the same spectral range (around 440 nm). To convert the IO results to the corresponding NO_2 and H_2O results, just the ratios of the cross sections have to be applied. With a (differential) absorption cross section of $3 \cdot 10^{19}$ cm² for NO_2 , and $3 \cdot 10^{26}$ cm² for H_2O (see table A1.1), all IO VCDs and SCDs would have to be multiplied by 83 and $8.3 \cdot 10^8$ for NO_2 and H_2O , respectively. The IO VCD for the standard plume ($5 \cdot 10^{15}$ molec/cm²) corresponds to a NO_2 VCD of $4.15 \cdot 10^{17}$ molec/cm² and a H_2O VCD of $4.15 \cdot 10^{24}$ molec/cm².

2.1.4 SO₂

- SO_2 can act as weak, but also as a very strong absorber, depending on the one hand on the amount of molecules in a plume, which can vary over many orders of magnitude. On the other hand it strongly depends on the selected wavelength range. At short wavelengths, the SO_2 absorption cross section is much stronger than at longer wavelengths. Around 310 nm, the absorption cross section is about $3.5 \cdot 10^{-19}$ cm², around 320 nm about $5 \cdot 10^{-20}$ cm², around 330 nm about $2 \cdot 10^{-21}$ cm², and around 370 nm $6 \cdot 10^{-22}$ cm² (see Fig. 4).
- The standard wavelength range for the TROPOMI SO₂ analysis is 312 to 326 nm (Theys et al., 2021b). This wavelength range is especially well suited for plumes with small SO₂ amounts. For the scenario with SO₂ as a weak absorber we chose a VCD of $1 \cdot 10^{18}$ molec/cm² for the standard plume (1 x 1 x 1 km³). For a TROPOMI observation this corresponds to a VCD of $5.2 \cdot 10^{16}$ molec/cm² and a differential optical depth of about 0.002 around 313 nm assuming an AMF of 0.4 and a differential optical depth of about $1 \cdot 10^{-19}$ cm², see table A1.1 (the differential optical depth would be by about 20% larger without the adaptation of the SO₂ absorption cross section to the TROPOMI spectral resolution). Here it should be noted that the (absolute) optical depth of the standard plume itself (1 x 1 x 1 km³) is much larger (about 0.25). Thus for very narrow plumes, the weak SO₂ scenario is already at the edge between a weak and strong absorber. But for larger plumes, the corresponding SO₂ absorption quickly decreases (for a 2 x 2 x 1 km³ plume it is about 0.06).
- In addition to this scenario, we also consider 4 scenarios with SO₂ as a strong absorber (in the standard TROPOMI fit range). The maximum SO₂ VCD is chosen according to Kern et al. (2020), who observed a very high SO₂ VCD (about 2 to 3 · 10²⁰ molec/cm² over an about 2 km wide plume from the Kilauea volcano about 4 km downwind of the vent. Thus we use a VCD of 4 · 10²⁰ molec/cm² as maximum VCD for the standard plume (1 x 1 x 1 km³). A very high SO₂ SCD of 3.2· 10¹⁹ molec/cm² was also observed at Mount Etna by Bobrowski et al. (2010). Very high SO₂ VCDs up to 5 · 10¹⁹ molec/cm² were also observed by TROPOMI for the La Palma eruption in September to December 2021 (e.g. Warnach, 2022). Here the plume usually covered several TROPOMI ground pixels, but with maximum SO₂ VCDs for individual TROPOMI pixels indicating a characteristic horizontal plume extension similar to a TROPOMI pixel. To cover a large range of strong SO₂ absorptions, we chose the following VCDs (for a 1 x 1 x 1 km³ plume): 1 · 10¹⁹ molec/cm², 2.5 · 10¹⁹ molec/cm², 1 · 10²⁰ molec/cm², and 4 · 10²⁰ molec/cm². The corresponding VCDs for the different SO₂ scenarios and horizontal plume extensions are given in Table 3. The standard fit range is represented by simulations at 313.1 nm (where the strongest SO₂

absorption peak is located). In addition, also simulations at 324.15 nm, 332.0 nm, 370.3 nm are performed. The corresponding optical depths for the chosen wavelengths are summarised in Table 4.

2.2 Aerosol input data

300

325

335

340

For the investigation of aerosol effects two scenarios are considered. For both scenarios an aerosol optical depth (AOD) of 10 (for the 1 x 1 x 1 km³ plume) is assumed, either with purely scattering aerosols (SSA = 1) or strongly absorbing aerosols (SSA=0.8). The first case represents plumes with sulfuric acid aerosols, the second case plumes with ash particles. The phase function is represented by the Henyey-Greenstein model with an asymmetry parameter of 0.68. These simplified scenarios were chosen to represent the basic characteristics of aerosol containing volcanic plumes. It should be noted that there is a lack of observations of the total amount of aerosols in fresh plumes, mainly because the high AOD in such situations itself prevents accurate measurements of the AOD. Like for the trace gases, also the aerosol amount is kept constant for the different horizontal plume extension leading to different AODs for different horizontal plume extensions (see table 3).

3 Light mixing effect

- The effect of horizontal light paths on satellite observations has first been investigated for aerosol retrievals from satellites (Richter, 1990; Lyapustin and Kaufman, 2001). Such retrievals are based on radiance measurements, and the effect of horizontal light paths can strongly affect the measurements in the presence of strong spatial radiance contrasts, e.g. caused by sea-land boundaries or cloud edges. In such cases, horizontal light paths cause an increase (decrease) of the radiance above the dark (bright) scene and thus systematically affect the aerosol retrieval. This effect (for absolute radiance measurements) was referred to as adjacency effect (Richter, 1990).
 - Trace gas measurements are usually not based on absolute radiance measurements but on narrow-band relative (differential) absorptions. For such measurements, horizontal light paths can also play an important role. But in contrast to aerosol measurements (based on absolute radiances), horizontal light paths affect the measurements even for scenes without spatial radiance contrasts (if spatial gradients of the trace gas of interest are present). In order to distinguish the effect of horizontal light paths on trace gas measurements from those on absolute radiance measurements (adjacency effect), we refer to it in this study as light mixing effect.
 - Light mixing effects for trace gas measurements have first been investigated by Schwärzel et al. (2020; 2021) for ground based and aircraft measurements. They found that like for the aerosol measurements, the light mixing effect for trace gas measurements causes a spatial smoothing of the trace gas signals.
- Here we extend these investigations to satellite observations. First we investigate the fundamental effects of horizontal photon paths (for scenarios without aerosols). Then we quantify the light mixing effect for ground pixel sizes of different satellite sensors and also scenarios with aerosols taking into account also realistic detection limits.

3.1 General findings and dependencies of the light mixing effect for trace gas observations

Figure 5 presents trace gas AMFs for different plume altitudes and wavelengths as function of the horizontal plume extension. The vertical plume thickness is 1 km. The wavelengths were chosen to represent typical trace gas analyses relevant for volcanic studies (SO₂ around 313 nm, BrO around 340 nm, IO around 440 nm, see section 2.1). The AMFs strongly depend on the horizontal plume extension. This dependency gets stronger towards shorter wavelengths, because of the higher probability of Rayleigh scattering and thus a larger contribution of photons originating from outside the satellite's FOV. Interestingly, the normalised AMFs (divided by the corresponding 1D AMFs) for different altitudes show a similar

dependence on the horizontal plume extension indicating compensating effects of the increasing free path lengths and the decreasing scattering probability with increasing altitude (because of the decreasing air density). The results in Fig. 5 show that the light mixing effect can cause a strong underestimation of the trace gas amounts of a volcanic plume if a 1D AMF is used in the data analysis. The underestimation is largest for small plumes and short wavelengths (up to > 70%).

345

350

355

360

365

370

375

In Fig. 6 so-called plume scans for plumes with different horizontal extensions are presented for a plume altitude of 5 - 6 km (results for other plume altitudes are shown in Fig. A3.1 the appendix A3). The highest AMFs occur directly above the plume center, but even these AMFs are systematically smaller compared to the corresponding 1D AMFs (orange horizontal lines). The deviation is strongest for short wavelengths and narrow plumes. The AMFs slightly decrease from the plume center towards the edges. Outside the plume they decrease rapidly, but stay clearly above zero.

This ,smoothing' of the AMF (compared to the horizontal extent of the plume) reflects the effect of horizontal photons paths and is directly connected to the atmospheric visibility. Towards short wavelengths, the free photon paths get shorter and the scattering probability gets higher. Thus the smoothing effect is strongest at short wavelengths. These general dependencies are also found for other plume altitudes (see Fig. A3.1 in appendix A3), but the absolute values of the AMFs strongly increase with increasing plume height both for narrow plumes and horizontally extended plumes (while the ratios of the AMFs for narrow plumes to those of horizontally extended plumes are similar for the different plume altitudes).

One interesting question is whether the decrease of the AMF above the plume (compared to the 1D AMF) and the increase outside the plume (compared to a zero AMF) exactly compensates each other. Indications for such a compensation are illustrated in Fig. 7 (note that for a quantitative interpretation, two horizontal dimensions have to be considered). To answer this question, simulations for satellite observations with a very large FOV corresponding to a ground pixel size of 200 x 200 km² are performed. Such large ground pixels are larger than typical horizontal photon paths (see Table 2). Thus almost all photons which have ,seen' the trace gas plume will stay inside the FOV.

The simulation results (together with similar results for a TROPOMI FOV) for a wavelength of 313 nm and a plume height of 5-6 km are shown in Fig. 8 (results for other combinations of wavelengths and plume altitudes are shown in Fig. A3.2 in appendix A3). The top panels present the retrieved fraction of the plume content as function of the horizontal plume extension. This fraction is defined as the ratio of the retrieved number of trace gas molecules using a 1D AMF and the true number of trace gas molecules of the plume (as used as input in the radiative transfer simulations). The retrieved fraction of the plume content is calculated as follows: First the simulated SCD for 3D plumes is divided by the corresponding 1D AMF yielding the average VCD across the satellite ground pixel. This VCD is then multiplied by the area of the ground pixel size yielding the total number of molecules. Finally, this value is divided by the number of trace gas molecules used as input for the simulations. Note that if the satellite ground pixel is smaller than the plume, only the input number of plume molecules covered by the satellite ground pixels is used to calculate the retrieved fraction of the plume content.

For TROPOMI, the retrieved fractions of the plume amount largely underestimate (between about 15 and 60%, depending on altitude and wavelength) the input trace gas amounts for small plume sizes, in agreement with the results shown in Figs. 5 and 6. The largest underestimation occurs for short wavelengths and a plume size similar to the ground pixel size. This can be understood by the results of the plume scans shown in Fig. 6, where the AMFs at the plume edges are the smallest within the plume. For plumes larger than the TROPOMI ground pixel, the retrieved fraction of the plume content (within the ground pixel size) then increases with plume size, because these scenarios converge to the 1D scenario (horizontally homogeneous trace gas layer).

Interestingly, for the large FOV (corresponding to a satellite ground pixel of 200 x 200 km²) the retrieved fractions of the plume amount are close to unity independent of the plume size. This confirms the above expectation that the decrease of the AMF in the area of the plume (compared to the 1D AMF) and the increase of the AMF outside the plume (compared to zero) compensate each other. In conclusion we find that the correct plume amount can be retrieved using 1D AMFs only for two extreme cases:

- a) The ground pixel size is much larger than the free photon path length and the plume size (Fig. 8 right). However, in such cases the trace gas absorption is very weak and usually below the detection limit (see Fig. 8 bottom right). Thus small/medium volcanic plumes can usually not be detected by instruments with a large ground pixel size.
 - b) The plume is homogenous and much larger than the free photon path length (and the ground pixel size). While the first condition is usually not well fulfilled, the second condition is typically fulfilled for strong eruptions. But such plumes are large and can usually be detected by satellite instruments with low spatial resolution (no observations with high spatial resolution are needed). For such large ground pixel sizes, also the effect of horizontal gradients will be rather small.

Thus small plumes (like for small/medium volcanic eruptions or degassing events) can only be observed by sensors with a small ground pixel size (like TROPOMI). But for such observations the retrieved trace gas amount will be systematically and strongly underestimated if 1D AMFs are applied in the data analysis.

395

415

390

3.2 Quantitative analysis for different ground pixel sizes taking into account the detection limit of the spectral retrieval

- As shown in the previous section, all molecules of a volcanic plume could in principle be retrieved with a 1D AMF if the satellite ground pixel would be large (> about 100 x 100 km²). Thus, even for satellite measurements with small ground pixels it should be possible to retrieve all molecules of a volcanic plume if the results of many neighboring ground pixels (surrounding the volcanic plume) were summed up. However, since the AMFs for pixels outside the volcanic plume are typically rather small (Fig. 6), for most of the observations of neighboring ground pixels the trace gas absorption will be below the detection limit. This finding is confirmed in Fig. 9, which shows simulation results for the trace gas scenarios for the weak absorbers introduced in section 2.1. Here again it should be noted that these results are valid only for the assumed trace gas scenarios and detection limits. For higher plume amounts and/or lower detection limits, lower SCDs, e.g. those for the neighboring pixels might be above the detection limit. Moreover, for scenarios with constant emissions (degassing) and stable wind fields, several measurements could be averaged, which would also lower the detection limit which is usually determined by photon noise. Nevertheless, from these simulations we derive the following general conclusions:
- 410 a) The trace gas SCDs from neighboring pixels are usually rather small. For TROPOMI observations the ratios of the maximum SCDs from the neighboring pixels relative to the SCDs of the center pixel range from 2% for 440 nm to 7% for 313 nm. Thus these measurements will often be below the detection limit.
 - b) even if measurements of the neighboring pixels are above the detection limit, their contribution to the detected total plume amount is small. For the two closest neighboring pixels (because of symmetry in total 4) their additional contribution ranges from about 4% (440 nm) to about 11% (313 nm) (relative contributions with respect to the true plume content).
 - c) for OMI, SCIAMACHY, or GOME-2, the SCDs of the neighboring pixels are always below the detection limit for the scenarios (with narrow volcanic plumes) considered in Table 3.
- Fig. 10 shows the retrieved fraction of the plume molecules (for a 1 x 1 x 1 km³ plume at different altitudes) as function of the ground pixel size for the three selected wavelengths. As discussed before, the strongest underestimation is found for TROPOMI and for short wavelengths. For SCIAMACHY and GOME-2 observations, the underestimation is only between a few and 15 percent. However, for these sensors the retrieved trace gas SCDs are below the detection limit for the narrow plumes considered here. Interestingly, very similar results are found for 313 and 340 nm, probably because of some compensating effects of the increase of the light mixing effect, but also an increase of the probability for multiple scattering towards shorter wavelengths. For plumes close to the surface (0 1 km altitude) very similar fractions are found as for the plume between 5 and 6 km, but because of the decreasing sensitivity of the satellite observations towards the ground, the corresponding trace gas absorptions are smaller and for the weak SO₂ scenario even below the detection limit. For plumes at

higher altitudes, the underestimation becomes smaller because the probability of multiple scattering due to Rayleigh scattering decreases and thus the differences of the AMFs for narrow and horizontally extended plumes become smaller.

430 3.3 Influence of aerosols

435

450

455

460

465

Fig. 11 presents AMFs (similar to Fig. 6) and normalised radiances (radiance/irradiance) for plume scans in near-nadir viewing geometry (solar zenith angle, SZA=0°) for different wavelengths (for SO₂ at 313 nm, BrO at 340 nm, and IO at 440 nm) and aerosol scenarios. The plumes are located between 5 and 6 km and contain the trace gas together with the aerosols (results for other plume heights are shown in Fig. A3.3 in appendix A3). Again the amount of molecules in the plume is assumed to be constant (and homogenously distributed), thus the trace gas VCDs and AOD vary with the horizontal plume extent (see Table 3). The aerosols are assumed either purely scattering (SSA=1) or strongly absorbing (SSA=0.8). The different colours in Fig. 11 and Fig. A3.3 represent AMFs and normalised radiances for plumes with different horizontal extensions (from 1 x 1 km² to 40 x 40 km²).

440 For small plume extensions (and thus high AOD in the plume) a strong effect of aerosols is seen (except for short wavelengths and plumes close to the surface): for the scenarios with scattering aerosols, both the AMF and the normalised radiance increase compared to the scenarios without aerosols (Fig. 6). For the scenarios with absorbing aerosols the opposite is found. For the interpretation of the simulated radiances it is important to take into account that especially for the low altitude plumes a substantial fraction of the atmospheric molecules is still located above the volcanic plume, which scatter the sun light towards the instruments without having 'seen' the plume. Thus, even for narrow plumes with absorbing aerosols the reduction of the observed radiance above the plume is relatively small. The low radiances at 313 nm are caused by the stratospheric ozone absorption, which is negligible at longer wavelengths.

For observations with extended FOV, the effective AMF results from the spatial averaging of the AMF weighted by the radiance. In that way the increased (decreased) radiance above scattering (absorbing) aerosols further increases (decreases) the AMFs for scattering (absorbing) aerosols. As a consequence, aerosols can have a rather strong impact on the effective AMF compared to the scenarios without aerosols.

In Fig. 12 the retrieved ratios of the plume content (ratio of the retrieved number of trace gas molecules assuming a 1D AMF and the true number of trace gas molecules in the plume) for TROPOMI observations are shown for aerosol scenarios (for a 1 x 1 x 1 km³ plume). Compared to the results for the aerosol-free cases the results are higher (for scattering aerosols) or lower (for absorbing aerosols). The strongest differences are found for IO at 440 nm, because the relative effect of the aerosols compared to Rayleigh scattering increases towards longer wavelengths. For scattering aerosols, the true plume content can even be overestimated due to the light path enhancement from increased multiple scattering in the plume as well as more backscattered light from the plume (higher reflectance above the plume). Very similar to the scenarios without aerosols, the ratios of the maximum SCDs of the neighboring pixels to the SCDs of the center pixels range from 2 % to 9%.

4 Saturation effect

For strong atmospheric absorbers like SO_2 , the exact plume extent becomes especially important. If the same amount of molecules is confined in a small or large volume (depending on the dilution state of the plume), the corresponding VCDs and vertical optical depths vary. For narrow horizontal plume extensions (e.g. in fresh plumes before effective mixing with ambient air) the optical depth is higher than for more diluted plumes (Fig. 2b). This effect is illustrated in Fig. 13 for SO_2 observations at a wavelength of 313 nm (representative for the standard SO_2 fit range, see Fig. 4) and a plume height of 5-6 km. Results for other plume heights are shown in Fig. A3.4 in appendix A3. In the left part of the figure the spatial distributions of the SO_2 VCDs are shown for the scenarios with SO_2 as weak absorber (top), ,strong,1' (middle), and

, strong, 4' (bottom), see table 3. The different colours indicate the different horizontal extensions of the plume (with the total amount of molecules kept constant). In the center and right part of the figure, plume scans of the AMF and the backscattered normalised radiance are shown. As expected, for the scenario with SO2 as weak absorber even for the smallest horizontal plume extension (1 x 1 km²) the radiance above the plume is the same as outside the plume (top, right). The decrease of the AMF compared to the 1D-AMF is thus simply a result of the light mixing effect (section 3). For higher SO₂ concentrations, 475 however (middle and bottom panel), the radiance is systematically reduced above the plume (like for the scenario with strong aerosol absorption, see Fig. 11 bottom), because a substantial fraction of the photons is absorbed by SO₂. Here it is interesting to note that even for the highest SO₂ concentrations the radiance does not approach zero indicating that part of the light is backscattered from molecules above the volcanic plume (while almost all photons reaching the volcanic plume are probably absorbed by SO₂). Accordingly, with increasing plume height a stronger reduction of the observed radiance for 480 plumes with high SO₂ amounts is found (see Fig. A3.4 in the appendix A3). For the scenarios with high SO₂ amounts, also the AMFs above the plume are systematically reduced compared to the scenarios with weak SO₂ absorption, because the SO₂ absorption is so strong that the observed photons have only ,seen' part of the vertical plume extension (similar to the scenario with strong aerosol absorption, see Fig. 11 bottom). The combined effects of reduced backscattered radiance and reduced AMFs strongly reduce the effective AMFs, when averaged over a satellite ground pixel. The reduction of the AMF 485 is largest for the narrow plumes.

4.1 SO₂ results for TROPOMI observations for different plume contents and wavelengths

Fig. 14 shows the fraction of the retrieved plume content for a plume at 5 – 6 km altitude as function of the plume size derived from TROPOMI observations if a 1D-AMF is assumed in the analysis. In addition, the retrieved SO₂ SCDs are shown. Similar results for other plume altitudes are shown in Fig. A.3.5 in appendix A3. The simulations are performed for the 4 scenarios with strong SO₂ absorption (see table 3) and 4 wavelengths (313.1 nm, 324.15 nm, 332.0 nm, 370.3 nm, see table 4 and Fig. 4). From the results in Fig. 14 and Fig. A3.5 several conclusions can be drawn:

a) a systematic and strong underestimation for short wavelengths is found (in addition to the light mixing effect). This additional underestimation is caused by the strong SO₂ absorption for narrow plumes and is referred to in this study as saturation effect. The underestimation caused by the saturation effect is similar for the different plume heights.

b) in spite of the strong SO_2 absorption, in some cases, the retrieved SO_2 SCDs are below the threshold value (red horizontal line), above which the operational SO_2 retrieval switches to the alternative fit windows (at longer wavelengths). The threshold values are $4 \cdot 10^{17}$ molec/cm² for the switch from fit range 1 to 2 and $6.7 \cdot 10^{18}$ molec/cm² for the switch from fit range 2 to 3, see Theys et al. (2021b). This is an important finding, because in such cases no analyses in the alternative fit windows will be performed and such high SO_2 amounts stay undetected in the current TROPOMI SO_2 retrieval. In contrast to the underestimation caused by the saturation effect the retrieved trace gas SCDs depend strongly on the plume altitude.

c) To retrieve the correct SO₂ SCD (and to avoid the saturation effect), it is recommended that simultaneous analyses in different wavelength ranges are performed. If the SCDs at shorter wavelengths are found to be systematically smaller than those at longer wavelengths, the results of the retrievals at the longer wavelengths should be considered. Only if in the standard fit window similar SCDs are obtained as at longer wavelengths, the results from the standard fit window should be used. This procedure would avoid that high SO₂ amounts stay undetected. Specific thresholds for the differences of the results in the different fit windows should be defined in a dedicated and more detailed study taking also into account the increasing detection limits towards longer wavelengths.

4.2 SO₂ results for observations from different satellites

490

495

500

505

Fig. 15 presents an overview on the fraction of the retrieved SO₂ content of the plume by different satellite instruments for scenarios with strong absorptions (see table 3) for a 1 x 1 x 1 km³ plume at 5 – 6 km altitude. Similar results for other plume altitudes are shown in Fig. A3.6 in appendix A3. For all SO₂ scenarios a strong dependence of the retrieved SO₂ SCD (and the retrieved fraction of the plume content) is found. The wavelength above which the underestimation becomes negligible depends on the SO₂ scenario (see table 3). For high SO₂ amounts, only analyses at long wavelengths yield reasonable results (but are of course still affected by the light mixing effect). Interestingly, the saturation effect is similar for the different instruments (with different ground pixels sizes), indicating that the saturation effect is mainly determined by the plume properties. Overall, similar results for the different plume heights are found, but for plumes at high altitudes a difference is found for the longer wavelengths compared to plumes at lower altitudes: there the retrieved fraction of the plume content increases especially for the instruments with small pixel sizes (TROPOMI and OMI), because the light mixing effect becomes less efficient with increasing altitude due to the reduced probability of multiple Rayleigh scattering.

4.3 Effect of the wavelength dependence of the AMF

So far the simulations were performed at single wavelengths in order to save computation time. These wavelengths were chosen to represent the dominant differential absorption cross sections for the different SO_2 fit windows (see Fig. 4 and table A2.1). While these monochromatic simulations can give a good first indication of the saturation effect, they neglect the effect of the wavelength dependence of the AMF (Marquard et al., 2000; Puķīte et al., 2010, 2016). This wavelength dependence is mainly caused by the wavelength dependence of the SO_2 absorption cross section, because for increasing absorption cross sections the penetration depth of the backscattered photons (and hence the AMF) decreases: the AMF is smallest for the highest values of the cross sections, and the wavelength-dependence of the AMF leads to a further decrease of the differential optical depth:

$$\tau(\lambda) = VCD \cdot AMF(\lambda, \sigma(\lambda)) \cdot \sigma(\lambda) \tag{2}$$

Note that an additional wavelength dependency of the AMF due to Rayleigh scattering is less important for the rather small fit windows considered here. We quantify the effect of the wavelength dependence of the AMF as described in appendix A4. It is found that the effect of the wavelength dependence of the AMF is small for cases with small optical depths. For cases with high optical depths and thus strong saturation effects, the true underestimation becomes larger than for the monochromatic AMFs. But such cases can be easily identified by the comparison of the retrieved SO₂ SCDs from different spectral ranges. For the SO₂ analysis, then an appropriate spectral range (with negligible saturation effect) has to be selected. For such analyses, also the effect of the wavelength dependence of the AMF becomes negligible.

545

550

540

515

520

525

530

535

5 Geometric effects for elevated plumes

The simulations presented so far were carried out for scenarios with overhead sun (SZA = 0) and nadir-looking instruments (viewing zenith angle, VZA = 0). These scenarios were chosen to investigate the light mixing and saturation effects in a fundamental way. In typical measurement situations, however, the illumination and viewing directions deviate from the vertical axes. Especially for elevated plumes, the trace gas absorption will then not be seen for the ground pixel exactly below the plume (Fig. 2c). In extreme cases, even double plumes might be observed. These effects of slant illumination and viewing angles are referred to as geometric effects in this study and are investigated in this section. Geometric effects lead to an increase of the horizontal extension over which the plume signal is seen compared to the observation geometry with SZA = VZA = 0. But at the same time, also the magnitude of the absorption is decreased, and might effectively fall below the

detection limit. This reduction of the absorption is simply caused by the fact that for such cases the atmospheric light paths cross the plume only once (or less) for a given viewing angle.

In Fig. 16 the AMFs for plume scans for observations of elevated plumes with various combinations of SZA and VZA are shown. The blue lines show the AMFs for observations with a narrow FOV (right y axis); the bright blue horizontal bars represent simulations for a TROPOMI FOV (3.5 km in across track dimension, left axis). Note that we kept the ground pixel size constant (3.5 x 5.5. km²) for all viewing angles in order to study the basic effects in a systematic way. For real TROPOMI observations, the ground pixel sizes increase for slant viewing angles. The TROPOMI AMFs are systematically lower than the AMFs for the narrow FOV, because the plume covers only a small fraction of the TROPOMI ground pixel (for nadir view ~5%).

565 The simulations shown in Fig. 16 are performed for 440 nm (IO). Similar results, but with less clear peaks, are obtained for the other wavelengths (for SO₂ at 313 nm and BrO at 340 nm see Fig. A5.1 and A5.2 in appendix A5). In addition to the AMFs, also sketches of the corresponding observation geometries are shown above the sub figures. As expected, the plume signals seen by the satellite are usually not found for those ground pixels covering the plume. Instead, maxima of the AMF are found for the downward and upward light paths crossing the plume, but slightly enhanced values are also found in 570 between these maxima. They are caused by sunlight which has crossed the plume and is then scattered towards the instrument before it reached the ground. As a general finding, the apparent horizontal extent of the plume increases for observations with slant illumination and/or slant viewing angles compared to observations with SZA = VZA = 0. At the same time the magnitude of the AMF decreases for observations with slant illumination and/or slant viewing angles. Both aspects have a direct influence on the retrieved fraction of the plume content: on the one hand, the ,smearing' of the plume 575 signal can lead to a larger number of ground pixels with enhanced absorptions and thus to a larger total covered area. On the other hand, ground pixels with weak absorptions (e.g. between the two AMF maxima) might fall below the detection limit (especially for scenarios with low trace gas VCDs). Both dependencies are seen in Fig. 17 where the number of pixels above the detection limit and the retrieved fraction of the plume content for TROPOMI observations with various viewing and illumination angles are shown for different scenarios of weak absorbers (see table 3). For the different trace gases, different 580 numbers of pixels with SCDs above the detection limit are found. The largest numbers are found for IO at 440 nm (weakest probability of Rayleigh scattering) and slant viewing and illumination angles (with opposite direction, relative azimuth angle, RAA = 0°), for which the apparent spatial extent of the plume gets largest. Interestingly, the retrieved fraction of the plume content is similar for all viewing geometries.

6 Plume side effects for low plumes

585

590

For slant illumination and viewing angles, interactions of photons with the sides of the plumes can become important, especially for plumes containing aerosols. However, even for plumes free of aerosols, photons entering or leaving the plumes from the sides have ,seen' different trace gas absorptions compared to the 1D scenarios. In this section plume side effects are investigated for TROPOMI observations and for a narrow plume (1 x 1 x 1 km³) located directly above the surface. We chose this low plume altitude, because for such plumes the side scattering effect can be investigated without interference with the geometric effects (section 5). Of course, the side scattering effect also affects plumes at higher altitudes (in addition to geometric effects). The aerosol scenarios are similar as in section 2.2 with an AOD of 10 and a single scattering albedo of 1.0 (purely scattering aerosols) or 0.8 (strongly absorbing aerosols).

In the middle panels of Fig. 18a,b the AMFs and normalised radiances for plume scans with a narrow FOV (0.014°) for (a) IO at 440 nm and (b) SO₂ 313 nm are shown (blue symbols). The simulations are performed for a slant viewing angle (VZA = 60°) and selected SZAs (0° and 50° in forward and backward direction). The plumes of 1 x 1 x 1 km³ are located at the surface (red boxes) and contain the trace gas and scattering aerosols with an AOD of 10. In addition to the AMFs for the

narrow FOV, also AMFs for a TROPOMI FOV (3.5 km in across track direction) are shown (horizontal magenta bars). 600 Especially for IO at 440 nm the normalised radiances and AMFs for the narrow FOV show complex dependencies on the viewing angle. Moreover, while the enhanced values of the radiances and the AMFs are found at similar viewing angles, there are also systematic differences in the details of their viewing angle dependencies. These are caused by the different sensitivities of both quantities on the atmospheric light paths. While the radiance mainly depends on the probability of scattering (on molecules and aerosols), the AMF also strongly depends on the length of the light path through the trace gas 605 plume. These dependencies can be e.g. seen in the upper left part of Fig. 18a, where the maximum of the radiance is found at about 1.5 km, while the maximum AMF is found at a distance of about 0.8 km from the plume center. Also clear differences for forward and backward illumination are found. The stronger and more structured signals for IO at 440 nm are caused by the fact that the direct sun light can penetrate deeper into the atmosphere than at shorter wavelengths and thus geometric effects (like shadowing for a VZA of 60° and a SZA of -50°) are more clearly visible. Also the AMFs for the TROPOMI 610 FOV differ systematically for different illumination directions. The TROPOMI AMFs (for ground pixels covering the plume) as function of the SZA and for different aerosol scenarios) are shown in the bottom panels of Fig. 18 (also for a VZA of 60°). Besides the 3D-AMFs also the corresponding 1D-AMF (without aerosols) are presented. A clear dependence on the SZA is found for all AMFs, which is slightly asymmetric (for forward and backward illumination) for the non-nadir viewing direction (VZA = 60°). These dependencies are caused by two effects: first by the angular dependence of Rayleigh scattering 615 which is relevant for both the 1D and 3D simulations; second by the angular dependence of Rayleigh scattering and aerosol scattering (if aerosols are present) for the 3D plumes).

620

625

630

635

640

In contrast to 313 nm and 340 nm, for IO at 440 nm systematic dependencies of the retrieved fraction of the plume content on the SZA are found, indicating the increased penetration depth of the direct sun light. For plumes without aerosols, almost constant values are found for SZA $< 70^{\circ}$, while for SZA $\ge 70^{\circ}$ the values strongly decrease. For observations of 3D plumes under such slant illumination angles part of the direct sun light has not crossed the plume (like for the 1D scenario) before it is reflected / scattered towards the instrument. For the scenario with scattering aerosols, the retrieved fraction of the plume content is always > 1 indicating the effect of multiple scattering (together with the increased radiance from the plume providing more weight for the trace gas signal from the plume). The values systematically increase with increasing SZA (except for the highest SZA). This increase is related to the higher contribution of photons entering the plume from the sides (pronounced forward scattering). For the non-nadir observations (VZA $\neq 0$) also a slight asymmetric dependence on the SZA is found.

For the scenario with absorbing aerosols, low values and almost no SZA-dependence are found for all wavelengths indicating that the aerosol absorption prevents the penetration of the solar photons deep into the volcanic plume (together with the reduced radiance from the plume). In summary, as already shown in section 3.3, the 3D-AMFs (and thus the retrieved fraction of the plume content) for plumes with scattering aerosols can be strongly increased compared to aerosol-free plumes. This overestimation is strongest for long wavelengths. In such cases the satellite measurements strongly overestimate the true plume content if a 1D AMF is applied up to >100% towards high SZA and high VZA (note that the largest VZAs for such observations are about 66° as for example for TROPOMI). Interestingly, the retrieved fraction of the plume content at 313 nm is close to unity regardless of the geometry.

For observations of plumes without aerosols or with absorbing aerosols the satellite measurements strongly underestimate the number of plume molecules if a 1D AMF (without aerosols) is applied in the data analysis (in agreement with the results shown in section 3.3).

645 7 Ground based measurements

650

675

680

3D effects are not only important for satellite observations of volcanic plumes, but also for ground based observations. In this section we briefly deal with this topic but only discuss the light mixing effect for ground based observations, because it is the most fundamental 3D radiative transfer effect. Kern et al. (2010) investigated 3D effects for ground based observations of volcanic plumes. Their focus was on the effect of sun light scattered into the line of sight of the instrument without having crossed the volcanic plume before, is referred to as light dilution effect (see also Millán, 1980). The light dilution effect is usually the dominant effect for such observations. While the exchange of light between the plume and outside the plume by multiple scattered photon was also included in their simulations, its contribution to the 3D radiative effects was not explicitly discussed.

- 655 In our simulations for ground based observations we therefore consider two scenarios (see Fig. 20):
 - a) ground based observations for zenith direction and very small SZA (here 0.3°): In this scenario the single-scattered light must have crossed the narrow plume before it is scattered into the instrument (like for the satellite observations with SZA = VZA = 0, see section 3). Thus no direct sun light is scattered into the line of sight of the instrument without having crossed the plume before. These results can be directly compared to the results for satellite observations.
- b) ground based observations for zenith direction and slightly larger SZA (here 10°): In this scenario the single-scattered light has not crossed the narrow plume before it is scattered into the instrument (like for scenarios in Kern et al., 2010).
 Note that for both scenarios single scattered light can also be scattered above the plume into the line of sight of the
 - Note that for both scenarios single scattered light can also be scattered above the plume into the line of sight of the instrument, but these light paths do not fall under the definition of the light dilution effect.

 The results for both scenarios are shown in Fig. 21. For the first scenario, like for the satellite observations (Fig. 5), the
- strongest dependence is found for 313 nm, because the probability of Rayleigh scattering is largest for the short wavelengths. But the altitude dependence is opposite to that for the satellite observations. Here it is interesting to note that for the AMF ratio between narrow and horizontally extended plumes, no monotonous altitude dependence is found. This is caused by the non-monotonous altitude dependence of the 1D AMFs for horizontally extended plumes. Interestingly, the highest 1D AMFs are found for plumes at medium altitudes (here altitudes of 5 and 10 km). For plumes at lower altitudes, the effect of multiply scattered photons is reduced because of the low surface albedo. For higher altitudes, it is reduced because of the low probability of Rayleigh scattering with decreasing air density.

The results for the second scenario show even lower AMFs for the narrow plumes than those for the first scenario. This is caused by the direct sun light scattered into the line of sight without having crossed the plume before (similar as for the scenarios in Kern et al., 2010). However, a direct comparison of the results of Kern et al. (2010) to the results of this study is complicated, because of two reasons: first, Kern et al. (2010) assumed a plume with infinite extension in one horizontal direction, while we assumed confined plumes in all dimensions. Second, they compared the AMFs of their 3D radiative transfer simulations with the assumption of a geometric light path through the plume, while in our study we compare the 3D AMFs with the corresponding 1D AMFs for horizontally extended plumes. Despite these differences, the general dependencies of the results of Kern et al. (2010) and our results are very similar: the AMF can be strongly reduced by the 3D effects. While for plumes in close proximity to the ground, the AMF is similar to the geometric AMF (but smaller than the corresponding 1D-AMF), it strongly decreases with increasing plume altitude (or horizontal distance between instrument and plume). These results indicate that usually for ground based observations not only the dilution effect (as described in Kern et al. (2010) but also the light mixing effect is important.

The results of this study also indicate that the correction factors presented by Kern et al. (2010) are only valid for the chosen plume size (vertical extent of 500 m, horizontal extent in one dimension: 600 m, and infinite in the other dimension). An assumed change of the plume extent from 200 m to 4 km changes the AMFs by about 5 % (for 440 nm) and 30 % (for 313 nm). The dependence of the AMFs on the horizontal plume extent is probably smaller for more realistic plumes (like those in Kern et al., 2010). Nevertheless, for future analyses of ground based observations, the plume size should be also taken into account. It should be noted that our simulations cover only a limited set of scenarios (zenith view, small SZA, no aerosols, no strong SO₂ absorptions). While our results are useful to illustrate the general influence of the light mixing effect and dilution effect on ground based observations, they are not representative for all measurement situations. Simulations for a more comprehensive set of scenarios (including more realistic plumes) and covering also the other 3D effects should be performed in future studies.

695

700

710

725

8 Conclusions

In this study different 3D effects were investigated with a focus on satellite observations. The new TROPOMI instrument aboard Sentinel-5P has a much smaller ground pixel size (down to 3.5 x 5.5 km²) than previous sensors. Thus TROPOMI can detect small volcanic plumes, which were invisible for previous instruments. However, for observations with small ground pixels, 3D effects become important, which were of minor importance for the observations with larger ground pixels. We investigated the following four 3D effects: the light mixing effect caused by horizontal photon paths, the saturation effect caused by strong SO₂ absorption, geometric effects related to slant illumination and viewing angles, and side-scattering effects related to photons originating from the sides of volcanic plumes.

Especially the first two effects can lead to a strong and systematic underestimation of the true trace gas content of the plume if 1D assumptions are used in the data analysis (up to more than 50% for the light mixing effect, and up to 100% for the saturation effect).

Thus for TROPOMI analyses of small volcanic plumes the light mixing effect has to be considered if the volcanic plume covers only 1 satellite ground pixel or a small number of ground pixels with a clear maximum for a single ground pixel (or in general if strong gradients on spatial scales of the TROPOMI ground pixel size are present). In such cases, wavelength-dependent correction factors (according to the results in Fig. 10) have to be applied to the results if a 1D AMF is used. The strongest underestimation (>50%) caused by the light mixing effect occurs for observations at short wavelengths and plumes below 10 km.

The saturation effect can lead to a further strong underestimation of the analysis results for cases with strong SO₂ absorptions. The strongest saturation effect occurs for observations at short wavelengths and for narrow plumes with high SO₂ VCDs (e.g. about > 95% underestimation for observations at 313 nm of a 1 x 1 x 1 km³ plume with SO₂ VCDs > 2.5 · 10¹⁹ molec/cm²). In extreme cases, in spite of the strong SO₂ absorption, the SO₂ SCDs retrieved in the standard fit window might even stay below the threshold for the switch of fit windows, and such plumes might remain undetected. For future analyses (not only for TROPOMI, but also for other sensors) we recommend to always retrieve SO₂ SCDs simultaneously in different fit windows. If the SCDs retrieved at short wavelengths (standard analysis) are found to be systematically smaller than at the other wavelengths, the SCDs from the alternative fit windows should be used.

Geometric effects typically have a weaker influence on the quantitative results, but they can affect the spatial pattern under which elevated plumes are ,seen' by the satellite instrument. Usually, localised elevated plumes are widened and even double peaks can occur. These effects should be kept in mind if complex plume patterns are observed. They might be simply caused by slant viewing and/or illumination angles.

Plume side effects are usually small for short wavelengths, but become larger for longer wavelengths. Especially if scattering aerosols are present in volcanic plumes they can lead to a strong overestimation of the true plume content if 1D AMFs (without aerosols) are used. While this overestimation is rather small for nadir observations and overhead sun (between about 5% and 50% depending on wavelength), it can reach much higher values (up to >100%) for slant viewing and illumination angles. Here it is important to note that such effects will probably play a rather important role for future sensors like the upcoming geostationary Sentinel-4 mission (Bazalgette Courrèges-Lacoste et al., 2017) which measures under rather slant viewing and illumination angles at northern latitudes.

Also for ground based observations, 3D effects can become important. In addition to the dilution effect (see Kern et al. 2010), which mostly depends on the distance between the ground based instrument and the volcanic plume, also the light mixing effect plays an important role, and a similarly strong dependence on the horizontal plume extension as for satellite observations is found. Thus the horizontal plume extension should also be taken into account for the analysis of ground based observations.

In this study the most fundamental 3D effects were investigated, but some important questions remain open:

- a) How representative are the assumed aerosol properties for real volcanic plumes? There is a huge knowledge gap about aerosol properties in volcanic plumes, mainly because especially for absorbing aerosols, the dense volcanic plumes can not be penetrated by optical remote sensing techniques. One option to overcome this difficulty could be to perform in situ and remote sensing measurements from drones flying into volcanic plumes. At least close to the edge of the plume, the aerosol extinction might be retrieved from direct sun measurements.
- b) What is the consequence of 3D effects for the quantification of emission fluxes from real volcanoes? The simulations of this study were made for idealised volcanic plumes. In reality, more complex plume configurations will occur. Thus in particular two scenarios should be investigated: 1) confined or vertical plumes: in many cases (for small plumes) enhanced trace gas absorptions are found only directly above the volcano or close to it. If the horizontal wind speeds are low, also a vertically oriented plume might contribute to the observed trace gas absorption. 2) horizontal plumes: for larger eruptions often elongated plumes are found, which are dominated by horizontal transport. Both scenarios should be simulated and eventually a direct connection between the observed trace gas SCDs and the emission flux from the volcano should be established. Such simulations will be carried out in a follow-up study.
 - c) How important are 3D effects for other confined trace gas plumes? Confined trace gas plumes also occur for other localised emission sources, in particular power plants. For measurements of the SO₂ emissions from power plants (in the UV spectral range), a similar strong underestimation will occur (up to more than 50%) as for the volcanic plumes. As shown in this study, for NO₂ observations in the blue spectral range the light mixing effect is much smaller than for the UV spectral range. But these results were obtained for a pure Rayleigh atmosphere outside the plume. Since power plant emissions usually occur in polluted regions with high aerosol concentrations, aerosol scattering will enhance the diffuse atmospheric radiation compared to a pure Rayleigh atmosphere, which will possibly increase the light mixing. Also 3D effects for realistic power plant plumes will be addressed in a forthcoming study.

Author contributions

TW performed the simulations and wrote the manuscript with contributions from all co-authors.

765 Acknowledgements

730

735

755

760

The study received funding from BMBF/DLR Bonn (contract no. 50EE1811B). We thank Tjarda Roberts (Université d'Orléans) and Luke Surl (University of Aberdeen) for helpful discussions about volcanic plumes.

770 Competing interests

Thomas Wagner is executive editor of AMT. Steffen Beirle is AMT associate editor.

References

- Afe, O. T., Richter, A., Sierk, B., Wittrock, F., and Burrows, J. P.: BrO emissions from volcanoes: a survey using GOME and SCIAMACHY measurements, Geophys. Res. Lett., 31, L24113, https://doi.org/10.1029/2004GL020994, 2004.
 - Bazalgette Courrèges-Lacoste, G., M. Sallusti, G. Bulsa, G. Bagnasco, Ben Veihelmann, S. Riedl, D. J. Smith, R. Maurer, "The Copernicus Sentinel 4 mission: a geostationary imaging UVN spectrometer for air quality monitoring," Proc. SPIE
- 780 10423, Sensors, Systems, and Next-Generation Satellites XXI, 1042307 (29 September 2017); https://doi.org/10.1117/12.2282158
 - Bobrowski, N., Hönninger, G., Galle, B., Platt, U., Detection of bromine monoxide in a volcanic plume. Nature 423, 273–276, 2003.
 - Bobrowski, N., U. Platt, SO₂/BrO ratios studied in five volcanic plumes, Journal of Volcanology and Geothermal Research,
- 785 Volume 166, Issues 3–4, Pages 147-160, 2007.
 - Bobrowski, N., Kern, C., Platt, U., Hörmann, C., and Wagner, T.: Novel SO₂ spectral evaluation scheme using the 360–390 nm wavelength range, Atmos. Meas. Tech., 3, 879-891, doi:10.5194/amt-3-879-2010, 2010.
 - Bobrowski, N. and Giuffrida, G.: Bromine monoxide / sulphur dioxide ratios in relation to volcanological observations at Mt. Etna 2006–2009, Solid Earth, 3, 433–445, https://doi.org/10.5194/se-3-433-2012, 2012.
- Bogumil, K., J. Orphal, J. P. Burrows, Temperature dependent absorption cross sections of O3, NO2, and other atmospheric trace gases measured with the SCIAMACHY spectrometer, Proc. ERS Envisat Symposium. Looking down at our Earth in the New Millennium, Gothenburg 16 20 October 2000.
 - Bovensmann, H., J. P. Burrows, M. Buchwitz, J. Frerik, S. Noe"l, V. V. Rozanov, K. V. Chance, and A. Goede (1999), SCIAMACHY—Mission objectives and measurement modes, J. Atmos. Sci., 56, 127–150.
- Burrows, J. P., et al. (1999), The Global Ozone Monitoring Experiment (GOME): Mission concept and first scientific results, J. Atmos. Sci., 56, 151–175.
 - Callies, J., Corpaccioli, E., Eisinger, M., Hahne, A., and Lefebvre, A.: GOME-2 MetOp's second-generation sensor for operational ozone monitoring, ESA Bull., 102, 28–36, 2000.
- Dinger, F., Kleinbek, T., Dörner, S., Bobrowski, N., Platt, U., Wagner, T., Ibarra, M., and Espinoza, E.: SO2 and BrO emissions of Masaya volcano from 2014 to 2020, Atmos. Chem. Phys., 21, 9367–9404, https://doi.org/10.5194/acp-21-9367-2021, 2021.
 - Eisinger, M. and Burrows, J. P.: Tropospheric sulfur dioxide observed by the ERS-2 GOME instrument, Geophys. Res. Lett., 25, 4177–4180, 1998.
- Fioletov, V. E., McLinden, C. A., Krotkov, N., Li, C., Joiner, J., Theys, N., Carn, S., and Moran, M. D.: A global catalogue of large SO₂ sources and emissions derived from the Ozone Monitoring Instrument, Atmos. Chem. Phys., 16, 11497–11519, https://doi.org/10.5194/acp-16-11497-2016, 2016.
 - Flynn, L. E., Seftor, C. J., Larsen, J. C., and Xu, P.: The Ozone Mapping and Profiler Suite, in: Earth Science Satellite Remote Sensing, edited by: Qu, J. J., Gao, W., Kafatos, M., Murphy, R. E., and Salomonson, V. V., Springer, Berlin, 279–296, doi:10.1007/978-3-540-37293-6, 2006.
- 810 Heue, K.-P., Brenninkmeijer, C. A. M., Baker, A. K., Rauthe-Schöch, A., Walter, D., Wagner, T., Hörmann, C., Sihler, H., Dix, B., Frieß, U., Platt, U., Martinsson, B. G., van Velthoven, P. F. J., Zahn, A., and Ebinghaus, R.: SO₂ and BrO observation in the plume of the Eyjafjallajökull volcano 2010: CARIBIC and GOME-2 retrievals, Atmos. Chem. Phys., 11, 2973–2989, https://doi.org/10.5194/acp-11-2973-2011, 2011.

- Hönninger, G., Referenzspektren reaktiver Halogenverbindungen für DOAS-Messungen, Diploma thesis, Univ. of Heidelberg, Heidelberg, Germany, 1999.
 - Hörmann, C., Sihler, H., Bobrowski, N., Beirle, S., Penning de Vries, M., Platt, U., and Wagner, T.: Systematic investigation of bromine monoxide in volcanic plumes from space by using the GOME-2 instrument, Atmos. Chem. Phys., 13, 4749–4781, https://doi.org/10.5194/acp-13-4749-2013, 2013.
- Kern, C., T. Deutschmann, L. Vogel, M. Wöhrbach, T. Wagner and U. Platt, Radiative transfer corrections for accurate spectroscopic measurements of volcanic gas emissions. Bull Volcanol 72, 233–247 (2010). https://doi.org/10.1007/s00445-
- 009-0313-7.

 Kern, C., A.H. Lerner, T. Elias, P. Nadeau, P.J. Kelly, C.A. Werner, L.E. Chlor, M. Cappos, Quantifying gas emissions
 - Kern, C., A.H. Lerner, T. Elias, P. Nadeau, P.J. Kelly, C.A. Werner, L.E. Chlor, M. Cappos, Quantifying gas emissions associated with the 2018 rift eruption of Kīlauea Volcano using ground-based DOAS measurements. *Bull Volcanol* **82**, 55 (2020).
- Khokhar, M. F., Frankenberg, C., Van Roozendael, M., Beirle, S., Kühl, S., Richter, A., Platt, U., and Wagner, T.: Satellite Observations of Atmospheric SO2 from Volcanic Eruptions during the Time Period of 1996 to 2002, J. Adv. Space Res., 36, 879–887, https://doi.org/10.1016/j.asr.2005.04.114, 2005.
 - Krotkov, N. A., Carn, S. A., Krueger, A. J., Bhartia, P. K., and Yang, K.: Band residual difference algorithm for retrieval of SO₂ from the Aura Ozone Monitoring Instrument (OMI), IEEE T. Geosci. Remote Sens., AURA Special Issue, 44, 1259–
- 830 1266, https://doi.org/10.1109/TGRS.2005.861932, 2006.
 - Krueger, A. J.: Sighting of El Chichon sulfur dioxide clouds with the Nimbus 7 total ozone mapping spectrometer, Science, 220, 1377–1379, 1983.
 - Levelt, P. F., van den Oord, G. H. J., Dobber, M. R., Malkki, A.,
 - Visser, H., de Vries, J., Stammes, P., Lundell, J. O. V., and Saari, H.: The Ozone Monitoring Instrument, IEEE T. Geosci.
- 835 Remote, 44, 1093–1101, https://doi.org/10.1109/TGRS.2006.872333, 2006.
 - Li, C., Joiner, J., Krotkov, N. A., and Bhartia, P. K.: A fast and sensitive new satellite SO2 retrievalalgorithm based on principal component analysis: Application to the ozone monitoring instrument, Geophys. Res. Lett., 40, 6314–6318, https://doi.org/10.1002/2013GL058134, 2013.
- Lyapustin, A. and Kaufman, Y., Role of adjacency effect in the remote sensing of aerosol, J. Geophys. Res. Atmos., 106, 11909–11916, https://doi.org/10.1029/2000JD900647, 2001.
 - Marquard, L.C., T. Wagner, and U. Platt, Improved Air Mass Factor Concepts for Scattered Radiation Differential Optical Absorption Spectroscopy of Atmospheric Species, *J. Geophys. Res.*, 105, 1315-1327, 2000.
 - Millán, M.,M., Remote sensing of air pollutants. A study of some atmospheric scattering effects, Atmospheric Environment, 14, 1241 1253, 1980.
- Nowlan, C. R., Liu, X., Chance, K., Cai, Z., Kurosu, T. P., Lee, C., and Martin, R. V.: Retrievals of sulfur dioxide from the Global Ozone Monitoring Experiment 2 (GOME-2) using an optimal estimation approach: Algorithm and initial validation, J. Geophys. Res., 116, D18301, https://doi.org/10.1029/2011JD015808, 2011.
 - Penning de Vries, M. J. M., Dörner, S., Puķīte, J., Hörmann, C., Fromm, M. D., and Wagner, T.: Characterisation of a stratospheric sulfate plume from the Nabro volcano using a combination of passive satellite measurements in nadir and limb geometry, Atmos. Chem. Phys., 14, 8149–8163, https://doi.org/10.5194/acp-14-8149-2014, 2014.
- geometry, Atmos. Chem. Phys., 14, 8149–8163, https://doi.org/10.5194/acp-14-8149-2014, 2014.

 Puķīte, J., Kühl, S., Deutschmann, T., Platt, U., and Wagner, T.: Extending differential optical absorption spectroscopy for
 - limb measurements in the UV, Atmos. Meas. Tech., 3, 631–653, https://doi.org/10.5194/amt-3-631-2010, 2010.
 - Puķīte, J. and Wagner, T.: Quantification and parametrization of non-linearity effects by higher-order sensitivity terms in scattered light differential optical absorption spectroscopy, Atmos. Meas. Tech., 9, 2147-2177, doi:10.5194/amt-9-2147-
- 855 2016, 2016.

- Richter, R., A fast atmospheric correction algorithm applied to Landsat TM images, Remote Sens., 11, 159–166, https://doi.org/10.1080/01431169008955008, 1990.
- Rix, M., Valks, P., Hao, N., Loyola, D. G., Schlager, H., Huntrieser, H. H., Flemming, J., Koehler, U., Schumann, U., and Inness, A.: Volcanic SO2, BrO and plume height estimations using GOME-2 satellite measurements during the eruption of
- Rothman, L. S., Jacquemart, D., Barbe, A., Benner, D. C., Birk, M., Brown, L. R., Carleer, M. R., Chackerian Jr., C., Chance, K., Coudert, L. H., Dana, V., Devi, V. M., Flaud, J.-M., Gamache, R. R., Goldman, A., Hartmann, J.-M., Jucks, K.

Eyjafjallajökull in May 2010, J. Geophys. Res., 117, D00U19, https://doi.org/10.1029/2011JD016718, 2012.

860

880

W., Maki, A.

- Mandin, G., J.-Y., Massie, S. T., Orphal, J., Perrin, A., Rinsland, C. P., Smith, M. A. H., Tennyson, J., Tolchenov, R. N.,
- Toth, R. A., Vander Auwera, J., Varanasi, P., and Wagner, G.: The HITRAN 2004 molecular spectroscopic database, J. Quant. Spectrosc. Ra., 96, 139–204, 2005.
 - Schönhardt, A., Richter, A., Theys, N., and Burrows, J. P.: Space-based observation of volcanic iodine monoxide, Atmos. Chem. Phys., 17, 4857–4870, https://doi.org/10.5194/acp-17-4857-2017, 2017.
 - Schwaerzel, M., Emde, C., Brunner, D., Morales, R., Wagner, T., Berne, A., Buchmann, B., and Kuhlmann, G., Three-
- dimensional radiative transfer effects on airborne and ground-based trace gas remote sensing, Atmos. Meas. Tech., 13, 4277–4293, https://doi.org/10.5194/amt-13-4277-2020, 2020.
 - Schwaerzel, M., Brunner, D., Jakub, F., Emde, C., Buchmann, B., Berne, A., and Kuhlmann, G.: Impact of 3D radiative transfer on airborne NO2 imaging remote sensing over cities with buildings, Atmos. Meas. Tech., 14, 6469–6482, https://doi.org/10.5194/amt-14-6469-2021, 2021.
- 875 Suleiman, R. M., Chance, K., Liu, X., González Abad, G., Kurosu, T. P., Hendrick, F., and Theys, N.: OMI total bromine monoxide (OMBRO) data product: algorithm, retrieval and measurement comparisons, Atmos. Meas. Tech., 12, 2067–2084, https://doi.org/10.5194/amt-12-2067-2019, 2019.
 - Theys, N., Van Roozendael, M., Dils, B., Hendrick, F., Hao, N., and De Mazière, M. (2009), First satellite detection of volcanic bromine monoxide emission after the Kasatochi eruption, Geophys. Res. Lett., 36, L03809, doi:10.1029/2008GL036552.
 - Theys, N., De Smedt, I., Van Roozendael, M., Froidevaux, L., Clarisse, L., and Hendrick, F., First satellite detection of volcanic OCIO after the eruption of Puyehue-Cordón Caulle, Geophys. Res. Lett., 41, 667–672, doi:10.1002/2013GL058416, 2014.
 - Theys, N., De Smedt, I., van Gent, J., Danckaert, T., Wang, T., Hendrick, F., Stavrakou, T., Bauduin, S., Clarisse, L., Li, C.,
- Krotkov, N. A., Yu, H., Van Roozendael, M.: Sulfur dioxide vertical column DOAS retrievals from the Ozone Monitoring Instrument: Global observations and comparison to ground-based and satellite data, J. Geophys. Res.-Atmos., 120, 2470–2491, https://doi.org/10.1002/2014JD022657, 2015.
 - Theys, N., De Smedt, I., Yu, H., Danckaert, T., van Gent, J., Hörmann, C., Wagner, T., Hedelt, P., Bauer, H., Romahn, F., Pedergnana, M., Loyola, D., and Van Roozendael, M.: Sulfur dioxide retrievals from TROPOMI onboard Sentinel-5
- Precursor: algorithm theoretical basis, Atmos. Meas. Tech., 10, 119–153, https://doi.org/10.5194/amt-10-119-2017, 2017.
 Theys, N., Hedelt, P., Smedt, I. De, Lerot, C., Yu, H., Vlietinck, J., Pedergnana, M., Arellano, S., Galle, B., Fernandez, D., Barrington, C., Taine, B., Loyola, D., and Van Roozendael, M.: Global monitoring of volcanic SO2 degassing from space with unprecedented resolution, Nature Scientific Reports, 9, 2643, https://doi.org/10.1038/s41598-019-39279-y, 2019.
 - Theys, N., Hedelt, P., Smedt, I. De, Lerot, C., Yu, H., Vlietinck, J., Pedergnana, M., Arellano, S., Galle, B., Fernandez, D.,
- Barrington, C., Taine, B., Loyola, D., and Van Roozendael, M.: Global monitoring of volcanic SO₂ degassing from space with unprecedented resolution, Nature Scientific Reports, 9, 2643, https://doi.org/10.1038/s41598-019-39279-y, 2019.
 - Theys, N., Fioletov, V., Li, C., De Smedt, I., Lerot, C., McLinden, C., Krotkov, N., Griffin, D., Clarisse, L., Hedelt, P., Loyola, D., Wagner, T., Kumar, V., Innes, A., Ribas, R., Hendrick, F., Vlietinck, J., Brenot, H., and Van Roozendael, M.: A

- sulfur dioxide Covariance-Based Retrieval Algorithm (COBRA): application to TROPOMI reveals new emission sources,
- 900 Atmos. Chem. Phys., 21, 16727–16744, https://doi.org/10.5194/acp-21-16727-2021, 2021a.
 - Theys, N., I. De Smedt, C. Lerot, H. Yu, M. Van Roozendael, S5P/TROPOMI SO2ATBD, Doc. No.: S5P-BIRA-L2-ATBD-400E, Issue 2.3.0, https://sentinels.copernicus.eu/documents/247904/2476257/Sentinel-5P-ATBD-SO2-TROPOMI, 2021b.
 - United States Committee on Extension to the Standard Atmosphere: U.S. Standard Atmosphere, 1976, National Oceanic and Atmospheric Administration, National Aeronautics and Space Administration, United States Air Force, Washington D.C.,
- 905 1976.
 - Tilstra, L. G., Tuinder, O. N. E., Wang, P., and Stammes, P.: Directionally dependent Lambertian-equivalent reflectivity (DLER) of the Earth's surface measured by the GOME-2 satellite instruments, Atmos. Meas. Tech., 14, 4219–4238, https://doi.org/10.5194/amt-14-4219-2021, 2021.
 - Vandaele, A. C., Hermans, C., Simon, P. C., Carleer, M., Colin, R., Fally, S., Mérienne, M.-F., Jenouvrier, A., and Coquart,
- 910 B.: Measurements of the NO2 Absorption Cross-section from 42000 cm-1 to 10000 cm-1 (238–1000 nm) at 220K and 294 K, J. Quant. Spectrosc. Ra., 59, 171–184, 1997.
 - Veefkind, J., Aben, I., McMullan, K., Förster, H., de Vries, J., Otter, G., Claas, J., Eskes, H., de Haan, J., Kleipool, Q., van Weele, M., Hasekamp, O., Hoogeveen, R., Landgraf, J., Snel, R., Tol, P., Ingmann, P., Voors, R., Kruizinga, B., Vink, R., Visser, H., and Levelt, P.: TROPOMI on the ESA Sentinel-5 Precursor: A GMES mission for global observations of the
- 915 atmospheric composition for climate, air quality and ozone layer applications, Remote Sens. Environ., 120, 70–83, https://doi.org/10.1016/j.rse.2011.09.027, 2012.
 - Wagner, T., J. P. Burrows, T. Deutschmann, B. Dix, C. von Friedeburg, U. Frieß, F. Hendrick, K.-P. Heue, H. Irie, H. Iwabuchi, Y. Kanaya, J. Keller, C. A. McLinden, H. Oetjen, E. Palazzi, A. Petritoli, U. Platt, O. Postylyakov, J. Pukite, A. Richter, M. van Roozendael, A. Rozanov, V. Rozanov, R. Sinreich, S. Sanghavi, F. Wittrock, Comparison of Box-Air-Mass-
- 920 Factors and Radiances for Multiple-Axis Differential Optical Absorption Spectroscopy (MAX-DOAS) Geometries calculated from different UV/visible Radiative Transfer Models, Atmos. Chem. Phys., 7, 1809-1833, 2007.
 - Warnach, S., Bromine monoxide in volcanic plumes A global survey of volcanic plume composition and chemistry derived from Sentinel-5 Precursor/TROPOMI data, PhD thesis, University of Heidelberg, 2022.
- Wilmouth, D. M., Hanisco, T. F., Donahue, N. M., and Anderson, J. G.: Fourier Transform Ultraviolet Spectroscopy of the A 2P3/2–X 2P3/2 Transition of BrO, J. Phys. Chem. A, 103, 8935–8945, doi:10.1021/jp9916510, 1999.
 - Yang, K., Krotkov, N., Krueger, A., Carn, S., Bhartia, P. K., and Levelt, P.: Retrieval of Large Volcanic SO2 columns from the Aura Ozone Monitoring Instrument (OMI): Comparisons and Limitations, J. Geophys. Res., 112, D24S43, https://doi.org/10.1029/2007JD008825, 2007.
- Yang, K., Liu, X., Bhartia, P., Krotkov, N., Carn, S., Hughes, E., Krueger, A., Spurr, R., and Trahan, S.: Direct retrieval of sulfur dioxide amount and altitude from spaceborne hyperspectral UV measurements: Theory and application, J. Geophys. Res., 115, D00L09, https://doi.org/10.1029/2010JD013982, 2010.
 - Zhang, Y., Li, C., Krotkov, N. A., Joiner, J., Fioletov, V., and McLinden, C.: Continuation of long-term global SO2 pollution monitoring from OMI to OMPS, Atmos. Meas. Tech., 10, 1495–1509, https://doi.org/10.5194/amt-10-1495-2017, 2017.

950

960

965

970

975

Tables

Table 1: Ground pixel sizes* and expected SO_2 SCDs for an SO_2 plume with a horizontal extension smaller than or equal to the size of a TROPOMI pixel and a SO_2 SCD of $1 \cdot 10^{16}$ molec/cm² (close to the detection limit) for a GOME-2 observation.

T 4	D. 1 .	A [1 2]	don don	CCD 1: 11	
Instrument	Pixel size	Area [km²]	SO_2 SCD	SCD ratio with	
	[km x km]		[molec/cm ²]	respect to the	
				GOME-2 SCD	
GOME-1	40 x 320	12800	$2.5 \cdot 10^{15}$	0.25	
(Burrows et al., 1999)					
GOME-2	40 x 80	3200	$1 \cdot 10^{16}$	1	
(Callies et al., 2000)					
OMPS	50 x 50	2500	$1.28 \cdot 10^{16}$	1.28	
(Flynn et al., 2006)					
SCIAMACHY	30 x 60	1800	$1.78 \cdot 10^{16}$	1.78	
(Bovensmann et al., 1999)					
OMI	13 x 23	299	$1.07 \cdot 10^{17}$	10.7	
(Levelt et al., 2006)					
TROPOMI	3.5 x 5.5**	19.25	$1.66 \cdot 10^{18}$	166.2	
(Veefkind et al., 2021)					

^{*}for OMI and TROPOMI at nadir

Table 2: Free horizontal light paths (e-folding lengths) for different wavelengths and altitudes considering only Rayleigh scattering.

Altitude	light path length at	light path length at	light path length at			
	313 nm (km)	350 nm (km)	440 nm (km)			
0-1 km	9	16	30			
5-6 km	14	25	47			
10-11 km	35	66	122			

^{**}originally 3.5 x 7 km²

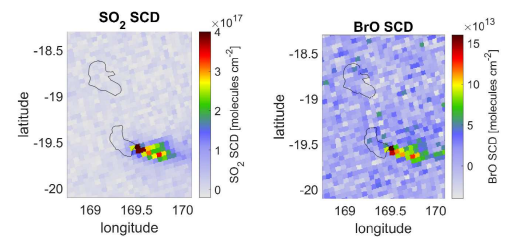
Table 3: Plume extensions and corresponding trace gas VCDs and AODs (lowest row) used in this study. For a given scenario the amount of trace gas is assumed to be constant but is distributed over different plume volumes (depending on the horizontal plume extension). The trace gas VCDs of the scenarios with $1 \times 1 \text{ km}^2$ horizontal extension is used to identify the corresponding scenario in this study.

Scenario	Trace gas VCDs for different horizontal plume extension								
	1 km	2 km	4 km	6 km	10 km	14 km	20 km	30 km	40 km
BrO VCD (1 · 10 ¹³ molec/cm ²)	500	125	31.3	13.9	5	2.55	1.25	0.56	0.31
IO VCD (1 · 10 ¹³ molec/cm²)	500	125	31.3	13.9	5	2.55	1.25	0.56	0.31
SO ₂ VCD _{weak} (1 · 10 ¹⁷ molec/cm ²)	10	2.5	0.625	0.28	0.1	0.051	0.025	0.011	0.006
$SO_2 VCD_{strong, 1}$ (1 · 10 ¹⁷ molec/cm ²)	100	25	6.25	2.78	1	0.51	0.25	0.11	0.06
$SO_2 VCD_{strong, 2}$ $(1 \cdot 10^{17} molec/cm^2)$	250	62.5	15.6	6.9	2.5	1.28	0.63	0.28	0.16
SO ₂ VCD _{strong, 3} (1 · 10 ¹⁷ molec/cm ²)	1000	250	62.5	27.8	10	5.10	2.50	1.11	0.63
$\frac{\text{SO}_2 \text{ VCD}_{\text{strong, 4}}}{(1 \cdot 10^{17} \text{molec/cm}^2)}$	4000	1000	250	111	40	20.4	10	4.44	2.5
Aerosol AOD	10	2.5	0.625	0.278	0.1	0.051	0.025	0.011	0.006

Table 4: SO_2 VCDs (for a 1 x 1 x 1 km³ plume), chosen wavelengths, total absorption cross sections and corresponding vertical optical depths for the strong SO_2 scenarios (see table 3).

corresponding vertical optical depths for the strong 502 sectiatios (see table 5).											
Wavelength /	SO ₂ VCD _{strong, 1}	SO ₂ VCD _{strong, 2}	SO ₂ VCD _{strong, 3}	SO ₂ VCD _{strong, 4}							
Absorption	10	10	20	20							
cross section	1·10 ¹⁹ molec/cm ²	2.5·10 ¹⁹ molec/cm ²	1·10 ²⁰ molec/cm ²	4.10 ²⁰ molec/cm ²							
313.1 nm /	2.45	6.13	24.5	98							
2.45 · 10 ⁻¹⁹ cm ²											
324.15 nm /	0.16	0.39	1.55	6.2							
1.55 · 10 ⁻²⁰ cm ²											
332.0 nm /	0.01	0.025	0.1	0.4							
1 · 10 ⁻²¹ cm ²											
370.3 nm /	0.006	0.015	0.06	0.24							
6 · 10 ⁻²² cm ²											

1010 Figures



 $Fig.\ 1:\ TROPOMI\ observations\ of\ the\ plume\ of\ Mount\ Yasur\ on\ 24.06.2020\ (Warnach,\ 2022).$



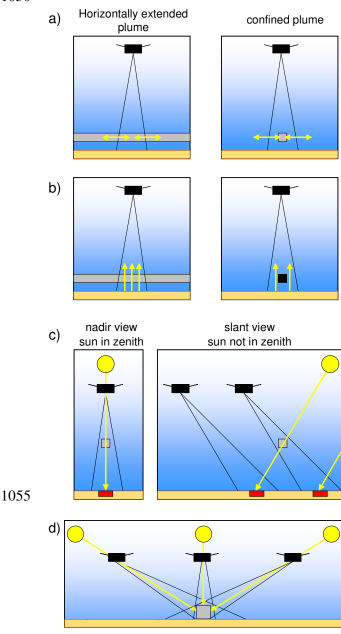


Fig. 2: The four 3D effects investigated in this study. a) Light mixing effect: part of the light seen by the satellite sensor originates from outside the satellite ground pixel. For horizontally confined plumes (right), such light paths contain no trace gas absorption. b) Saturation effect: for strongly absorbing species like SO₂, the incoming sun light might be almost fully absorbed in the plume. This can lead to a strong underestimation for narrow plumes with high trace gas concentration. c) geometric effect: depending on the illumination and viewing geometry, the location of the ground pixel with enhanced trace gas absorption (the projection of the plume along the direction of the incoming sun light, red marks at the surface) can systematically differ from the true plume location (grey square). d) Plume side effect: for narrow volcanic plumes the contributions of photons reaching the sensor from the sides of the plume becomes important.

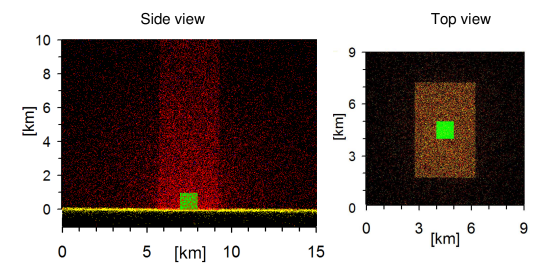


Fig. 3: Exemplary simulation results from the Monte Carlo radiative transfer model TRACY-2. An idealised narrow volcanic plume (1 x 1 x 1 km³) is observed by TROPOMI with a FOV according to a ground pixel size of 3.5 x 5.5 km². Red dots indicate Rayleigh scattering events, blue dots rotational Raman scattering events, yellow dots surface reflections, and green dots aerosol scattering events (the volcanic plume is filled with purely scattering aerosols). Simulations are performed for 340 nm with 25000 photons for overhead sun and a nadir-looking satellite instrument.

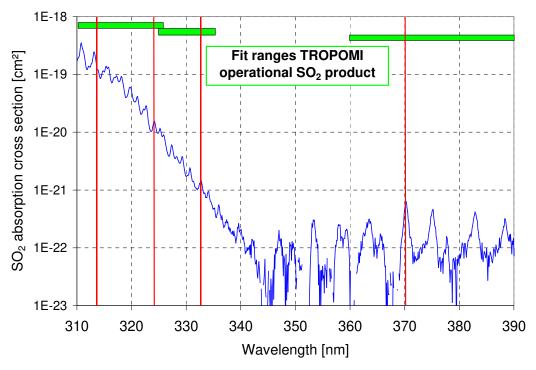
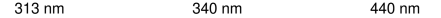


Fig. 4 SO_2 absorption cross section (273K, Bogumil et al., 2000, see also https://www.iup.uni-remen.de/gruppen/molspec/databases/sciamachydata/index.html). The green bars indicate different SO_2 fit ranges used in the operational TROPOMI product. The vertical bars indicate wavelengths used in the 3D RTM simulations.



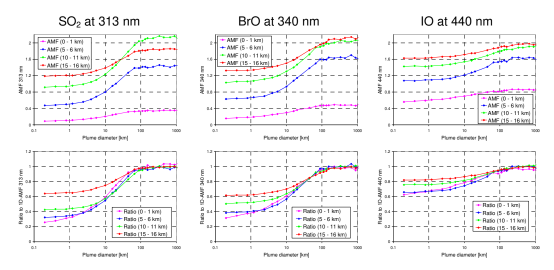


Fig. 5: Dependence of the AMF on the horizontal plume extension. Top: AMFs for different plume altitudes and wavelengths; bottom: AMFs normalised by the corresponding 1D AMFs (assuming a horizontally extended plume. The simulations are performed for SZA: 0° , VZA: 0° , FOV: 0.014° , satellite altitude: 824 km asl, no aerosols.

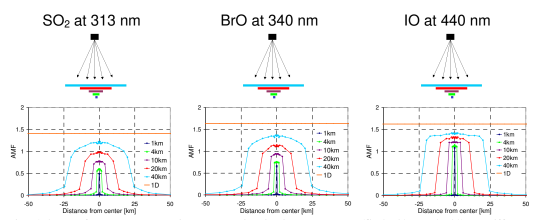


Fig. 6 AMFs for plume scans in near-nadir viewing geometry (SZA=0 $^{\circ}$, VZA=0 $^{\circ}$) at different wavelengths. It is assumed that the satellite scans the plume with a narrow FOV ($\sim 0.014^{\circ}$). The different colours represent AMFs for plumes at 5-6 km altitude and with different horizontal extensions (from 1 x 1 km² to 40 x 40 km²). Results for other plume altitudes are shown in appendix A3.

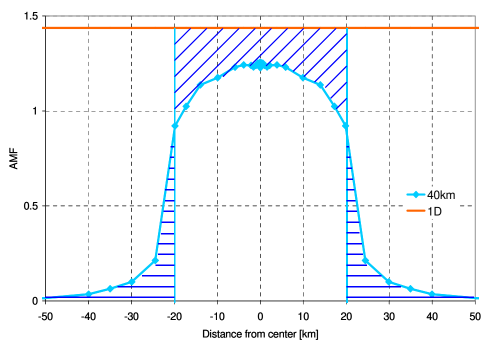


Fig. 7 AMFs for plume scans (as in Fig. 5) in near-nadir viewing geometry (SZA=0°) for SO_2 (weak) at 313 nm and a plume at 5-6 km altitude. In the area of the plume the AMFs are smaller than the 1D-AMF, whereas outside they are larger than zero. The differences are indicated by the blue marked areas for a plume of 40 x 40 km².

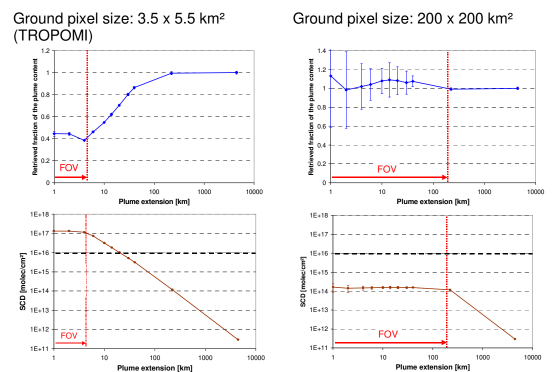
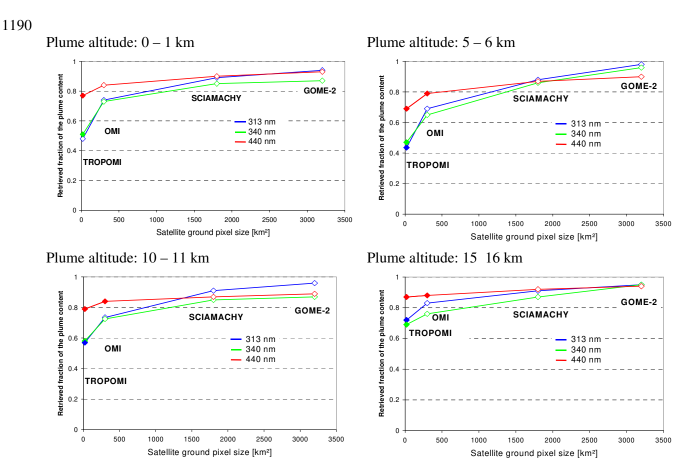


Fig. 8 Top: Retrieved fraction of the plume content as a function of the horizontal plume extension (note the logarithmic scale) for measurements of SO₂ at 313 nm. Bottom: measured SO₂ for a SO₂ VCD (for a 1 x 1 km² pixel) of 1· 10¹⁸ molec/cm². The horizontal dashed line indicates the detection limit (assuming an OD threshold of 0.001). Left: results for a TROPOMI ground pixel; right: results for a large ground pixel of 200 x 200 km². Note that in the case of plume size > ground pixel size, only the fraction of the plume within the ground pixel size is considered. Simulations for a plume altitude from 5 to 6 km, wavelength of 313 nm and for VZA=0, and SZA=0°. The error bars represent the standard deviation calculated from 40 individual simulations. Results for other plume altitudes and molecules (wavelengths) are shown in appendix A3.

SO ₂ at 313 nm					BrO at 340 nm					IO at 440 nm					
0.9%	1.5%	2.1%	1.5%	0.9%	0.6%	1.3%	1.9%	1.3%	0.6%		0.3%	0.6%	0.7%	0.6%	0.3%
<0.001	<0.001	<0.001	<0.001	<0.001	<0.0005	<0.0005	<0.0005		<0.0005		<0.0005	<0.0005		<0.0005	<0.0005
1.3%	3.2%	43.6%	3.2%	1.3%	1.1%	2.8%	47.1%	2.8%	1.1%		0.4%	1.4%	71.3%	1.4%	0.4%
<0.001	<0.001	0.003	<0.001	<0.001	<0.0005	<0.0005	0.004	<0.0005	<0.0005		<0.0005	<0.0005	0.006	<0.0005	<0.0005
0.9%	1.5%	2.1%	1.5%	0.9%	0.6%	1.3%	1.9%	1.3%	0.6%		0.3%	0.6%	0.7%	0.6%	0.3%
<0.001	<0.001	<0.001	<0.001	<0.001	<0.0005	<0.0005	<0.0005	<0.0005	<0.0005		<0.0005	<0.0005	<0.0005	<0.0005	<0.0005

Fig. 9 Horizontal coverage of a 1 x 1 x 1 km³ plume (yellow hatched areas) by TROPOMI observations at different wavelengths. The numbers at the top of the TROPOMI pixels indicate the fractions of the plume molecules retrieved from that ground pixel. The numbers at the bottom indicate the optical depth of the trace gas absorption. Red values represent results above the respective detection limit.



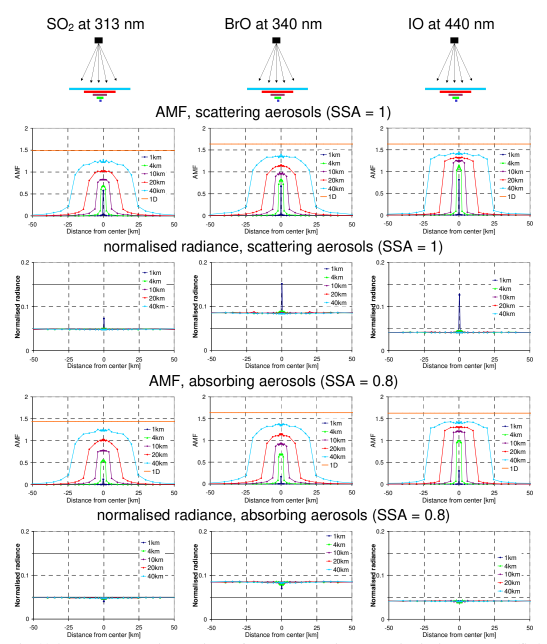
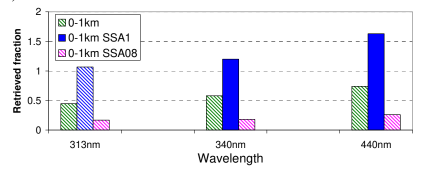
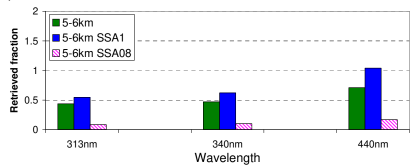


Fig. 11 AMFs and normalised radiances for plume scans in near-nadir viewing geometry (SZA=0°) for different wavelengths and aerosol contents. The satellite scans the plume with a narrow FOV ($\sim 0.014^{\circ}$). The different colours represent AMFs for plumes at 5 – 6 km altitude and with different horizontal extensions (from 1 x 1 km² to 40 x 40 km²). The rather low radiance at 313 nm (in spite of the high probability of Rayleigh scattering) is caused by the stratospheric ozone absorption. Results for other plume altitudes are shown in appendix A3.

A) Plume altitude: 0 – 1 km



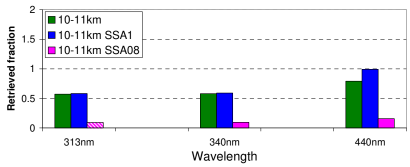
B) Plume altitude: 5 – 6 km



1220

1230

C) Plume altitude: 10 - 11 km



1225 **D) Plume altitude: 15 – 16 km**

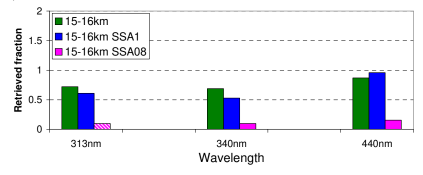


Fig. 12 Retrieved fraction of the plume content at different wavelengths for different plume heights and aerosol scenarios for TROPOMI observations (horizontal plume extension 1 x 1 km²). Hatched bars indicate scenarios, for which the absorption is below the detection limit for the weak trace gas scenarios described in section 2.1 (results for the satellite pixel directly above the plume).

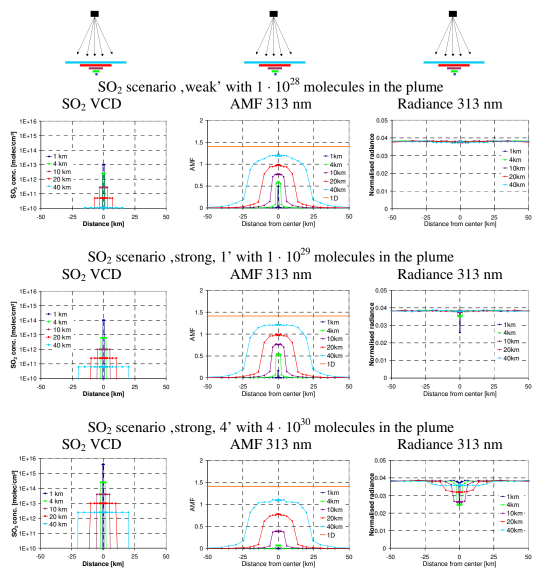


Fig. 13 Plume scans for SO₂ plumes with different amounts of molecules and different horizontal extensions (for the different SO₂ scenarios, see table 3). Left: SO₂ VCDs of the plumes, middle: AMFs at 313 nm; right: normalised radiances at 313 nm. Here only the scenarios 'weak, 'strong,1' and 'strong,4' are considered to illustrate the transition from cases with no saturation to cases with medium or strong saturation. The additional intermediate scenarios 'strong,2' and 'strong,3' are later also used for the quantification of the saturation effect for the different plume extensions (Fig. 14) and satellite instruments (Fig. 15). Results for other plume altitudes are shown in Fig. A3.4 in appendix A3.

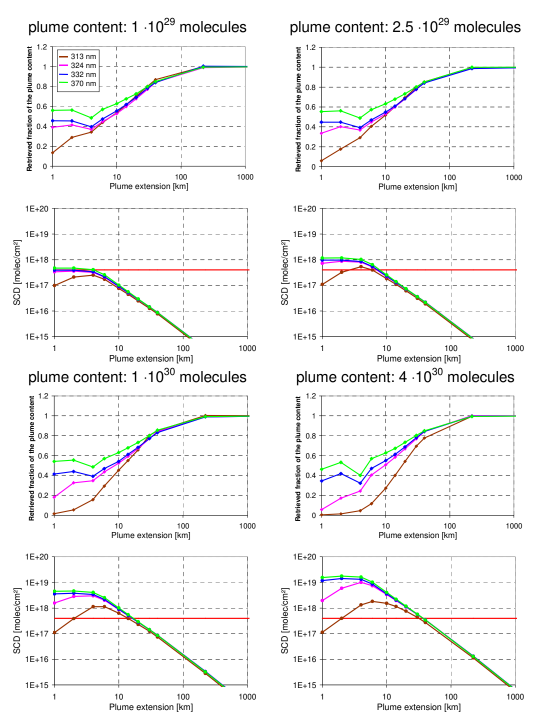


Fig. 14 Retrieved fraction of the plume content (top) and SO₂ SCDs (bottom) from TROPOMI measurements at different wavelengths as function of the plume size and for different amounts of molecules in the plume. The red horizontal line indicates the threshold, above which the operational SO₂ retrieval switches from the standard fit window to the first alternative fit window (at longer wavelengths). Results for other plume altitudes are shown in appendix A3.

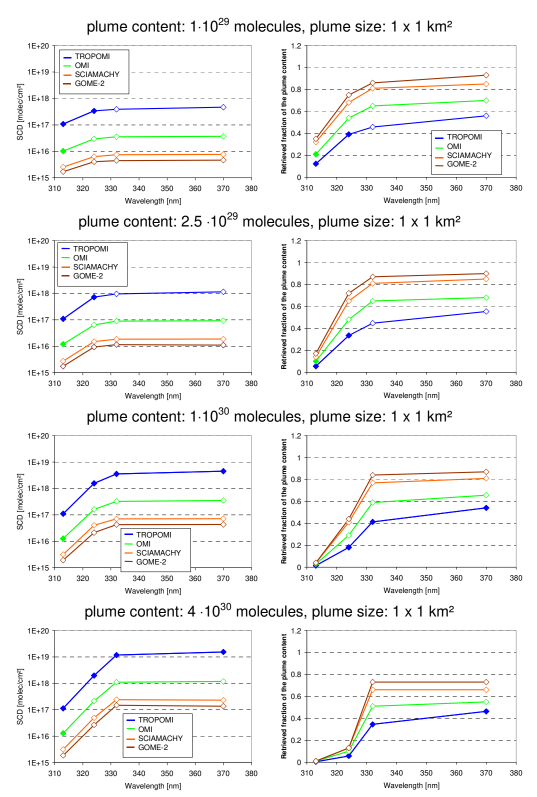


Fig. 15 Retrieved fractions of the plume content and SO_2 SCDs for different satellite instruments and for a plume size of $1 \times 1 \text{ km}^2$ as function of the wavelength. The full symbols represent measurements with SCDs above the detection limit. Note that for the rather large ground pixel sizes of SCIAMACHY and GOME-2 the errors are between 5 and 10%. Results for other plume altitudes are shown in appendix A3.

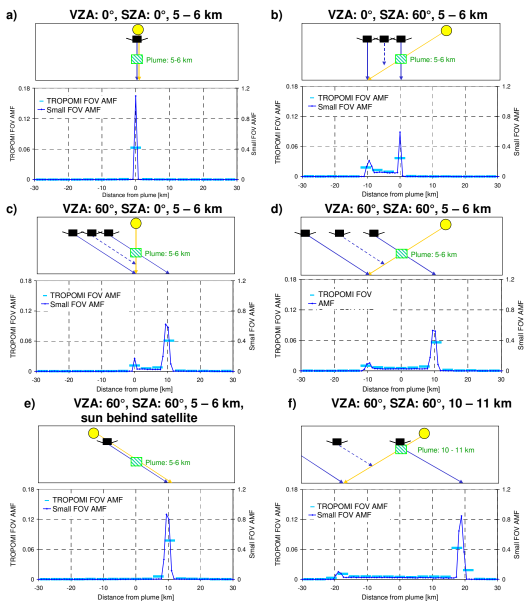
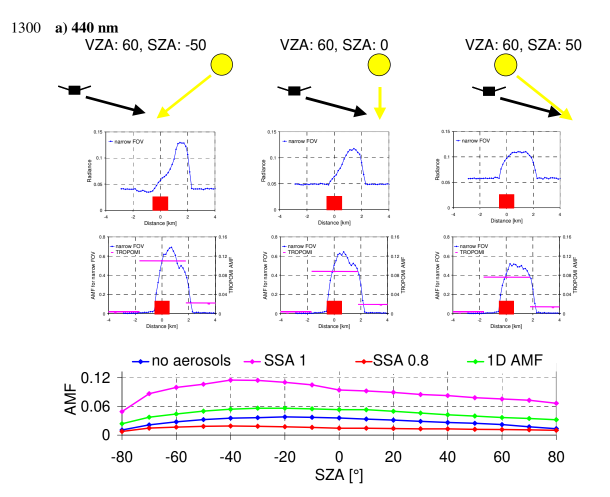


Fig. 16 AMFs for TROPOMI observations of elevated plumes for different combinations of SZA and VZA. The blue lines show the AMFs for observations with a narrow FOV (~ 0.014°) (right y axis); the bright blue symbols represent simulations with a TROPOMI FOV (left axis). All simulations are for IO at 440 nm and plume sizes of 1 x 1 x 1 km³. Results for 313 nm (SO₂) and 340 nm (BrO) are shown in Figs. A5.1 and A5.2 in appendix A5.

number of pixels above the detection limit for retrieved fraction of the plume content for the the weak SO₂ scenario (313 nm) weak SO₂ scenario (313 nm) VZA 0 content pixels above detection limit VZA 40° RAA 0° VZA 0 VZA 40° RAA 0° VZA 40° RAA 180° 3.0 VZA 40° RAA 180° VZA 60° RAA 0° VZA 60° RAA 0° VZA 60° RAA 180° VZA 60° RAA 180 number of Retrieved 0.2 SZA number of pixels above the detection limit for retrieved fraction of the plume content for BrO BrO (340 nm) (340 nm) VZA 0 VZA 0 content 8.0 number of pixels above detection limit VZA 40° RAA 0° VZA 40° RAA 0° VZA 40° RAA 180 VZA 40° BAA 180 VZA 60° RAA 0° the plume VZA 60° RAA 0° VZA 60° RAA 180 VZA 60° RAA 180 fraction of 60 SZA SZA number of pixels above the detection limit for retrieved fraction of the plume content for IO IO (440 nm) (440 nm) VZA 0 content 8.0 above detection limit VZA 40° RAA 0° VZA 60° BAA 0° /ZA 60° RAA 180° of the 0.6 fraction 0.4 of pixels VZA 0 VZA 40° RAA 0° number 0.2 VZA 40° RAA 180 VZA 60° RAA 0° VZA 60° RAA 180°

Fig. 17 Number of satellite pixels with SCD above the detection limit (left) and the retrieved fraction of the plume content (right) for TROPOMI observations of SO_2 (weak, 313 nm)), BrO (340 nm), and IO (440 nm) with different viewing and solar zenith angles. The plume has an extension of 1 x 1 x 1 km³ and is located at 5 – 6 km altitude. The red circles indicate scenarios with the sun and the satellite in the same direction for which the smearing effect is smallest. RAA indicates the relative azimuth angle between the viewing direction and the sun. Here a RAA of 180° represents cases where the sun shines in the same direction of the satellite view.



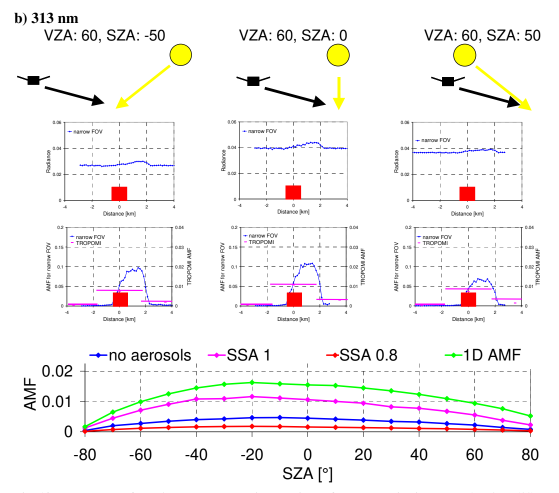


Fig. 18 Dependence of the AMF and normalised radiance for a slant viewing angle (VZA = 60°) on the illumination angle for IO at 440 nm (a) and SO₂ at 313 nm (b). The upper panels show sketches of the investigated viewing and illumination angles (SZA of -50°, 0°, 50°). The middle panels show the normalised radiances and AMFs for these scenarios (for plumes with scattering aerosols) as function of the relative distance from the plume (1 x 1 x 1 km³, red boxes). Note the different y-axes for the AMFs calculated for a narrow FOV (left) or TROPOMI FOV (right). The plume is located directly above the surface (0 – 1 km). The bottom panels show the AMFs for a TROPOMI FOV as function of the SZA (also for a VZA of 60°). The different lines represent 3D-AMFs for plumes without aerosols (blue), with scattering (magenta) and absorbing (red) aerosols as well as the 1D-AMF (without aerosols).

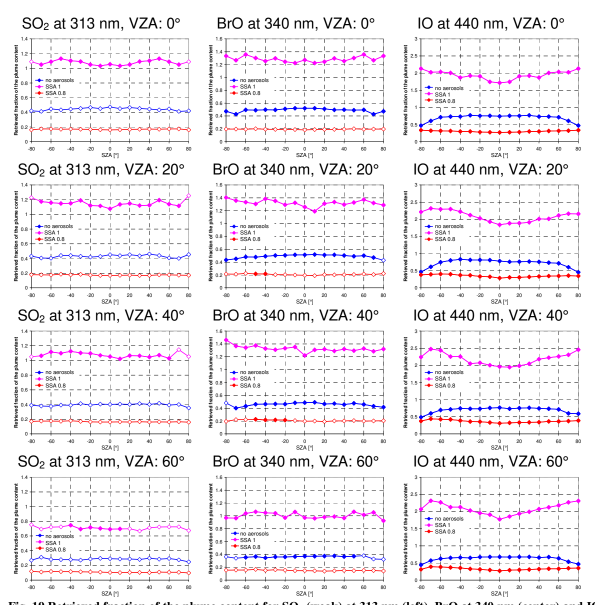


Fig. 19 Retrieved fraction of the plume content for SO₂ (weak) at 313 nm (left), BrO at 340 nm (center) and IO at 440 nm (right) as function of the SZA for different viewing angles (blue: plume without aerosols; magenta: plume with scattering aerosols; red: plume with absorbing aerosols). Symbols in full colour represent results above the detection limit.

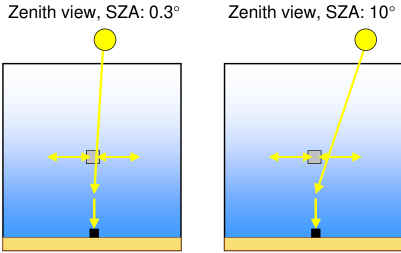
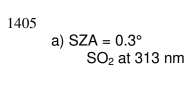
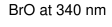
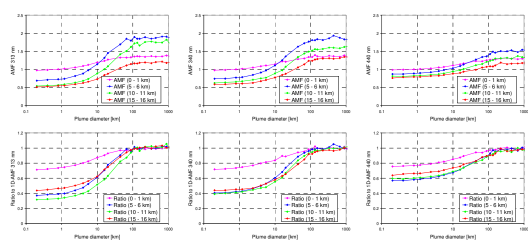


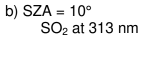
Fig. 20 Ground based observations of a narrow plume in zenith direction. For almost overhead sun (SZA of 0.3°, left), the direct sun light traverses the plume before it is scattered into the field of view of the instrument. For larger SZA (right), direct sun light is scattered into the field of view of the instrument without having crossed the plume. In both scenarios no direct sun light is observed by the instruments.

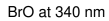




IO at 440 nm









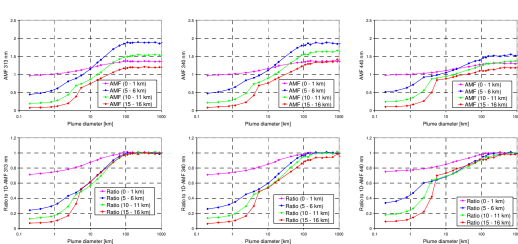


Fig. 21 Dependence of the AMF on the horizontal plume extension for ground based observations (FOV: 0.014° , VZA = 0° , SZA = 0.3° (a) and SZA = 10° (b). In the upper panels the AMFs for different plume altitudes and wavelengths are shown. In the bottom panels the AMFs normalised by the corresponding 1D AMFs are shown.

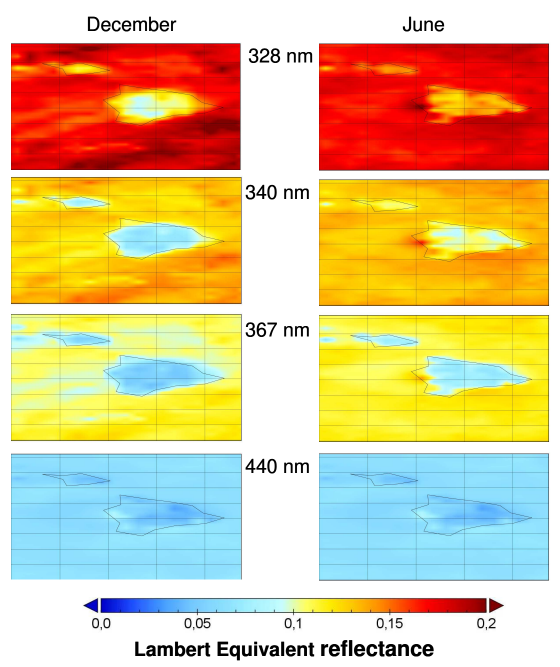
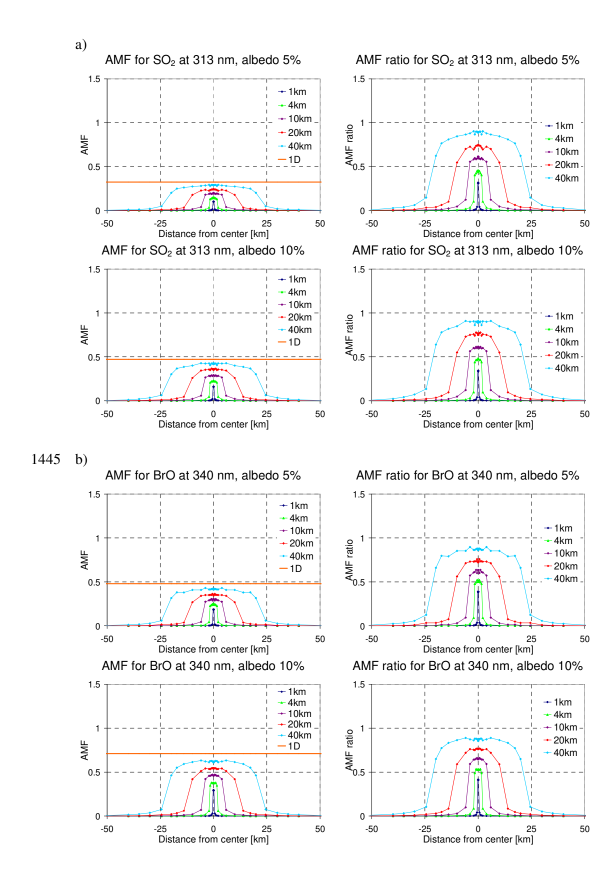


Fig. A1.1 Minimum Lambertian equivalent albedos over Hawaii derived from TROPOMI observations for December (left) and June (right). Data obtained from the TEMIS data base (https://www.temis.nl/), see also Tilstra et al. (2021).



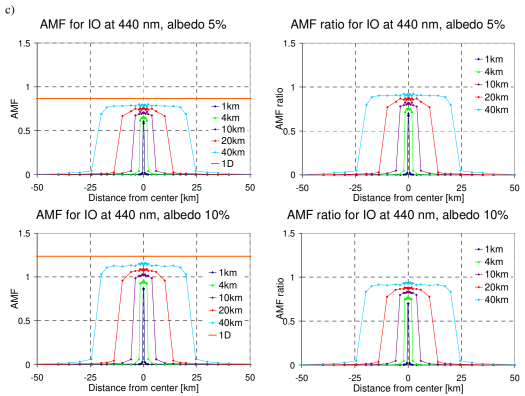


Fig. A1.2 AMFs (left) and corresponding ratios to the 1D-AMFs (right) for plume scans in near-nadir viewing geometry (SZA=0°, VZA=0°) for (a) SO_2 at 313 nm, (b) BrO at 340 nm, and (c) IO at 440 nm. It is assumed that the satellite scans the plume with a narrow FOV (~ 0.014°). The different colours represent AMFs for plumes at 5-6 km altitude and with different horizontal extensions (from 1 x 1 km² to 40 x 40 km²). Results are shown for a surface albedo of 5% (as used in general in this study) and 10%. The ratios of the AMFs for narrow plumes to the corresponding 1D-AMFs hardly depend on the surface albedo.

Appendix A2 Differential absorption cross sections for the selected trace gases and wavelengths

The following table shows a summary of the differential absorption cross sections used in this study to relate the trace gas SCDs to the corresponding optical depths.

The values were determined by convolution of the original cross section with a Gaussian slit function with FWHM of 0.5 nm (similar to TROPOMI). In addition to the IO absorption cross section also those of H₂O and NO₂ are shown, which could also be analysed in the same spectral range around 440 nm. Note that the absorption cross sections depend on temperature. Also the values from different studies differ, especially for IO and H₂O. Finally, the differential absorption cross sections will depend on the spectral resolution of the instrument. Thus the values in Table A.1.1 can serve as useful reference (e.g. in this study), but might differ for other applications.

Table A2.1 Differential absorption cross sections for the trace gases considered in this study.

Trace gas, wavelength	Differential absorption cross section after
	convolution with Gaussian instrument line
	shape with FWHM of 0.5 nm.
SO ₂ , 313 nm (Bogumil et al., 2000)	1 · 10 ⁻¹⁹ cm ²
SO ₂ , 324 nm (Bogumil et al., 2000)	4 · 10 ⁻²¹ cm ²
SO ₂ , 332 nm (Bogumil et al., 2000)	4 · 10 ⁻²² cm ²
SO ₂ , 370 nm (Bogumil et al., 2000)	5 · 10 ⁻²² cm ²
BrO, 340 nm (Wilmouth et al., 1999)	1.7 · 10 ⁻¹⁷ cm ²
IO, 440 nm (Hönninger, 1999)	2 · 10 ⁻¹⁷ cm ²
NO ₂ , 440 nm (Vandaele et al., 1997)	3 · 10 ⁻¹⁹ cm ²
H ₂ O, 440 nm (Rothman et al., 2005)	3.4 · 10 ²⁶ cm ²

1500 Appendix A3 Simulation results for different plume altitudes

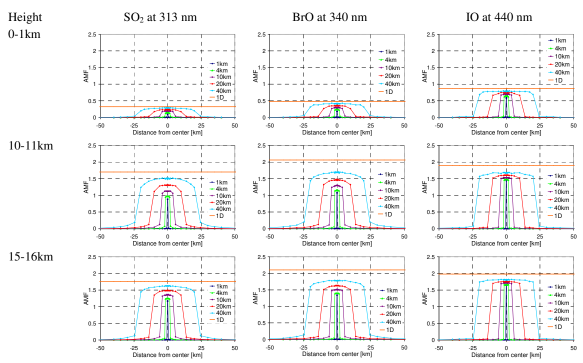
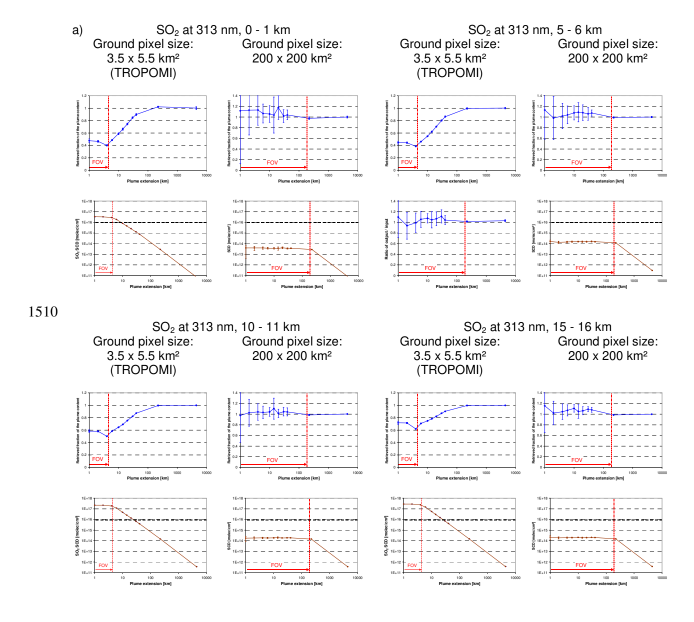
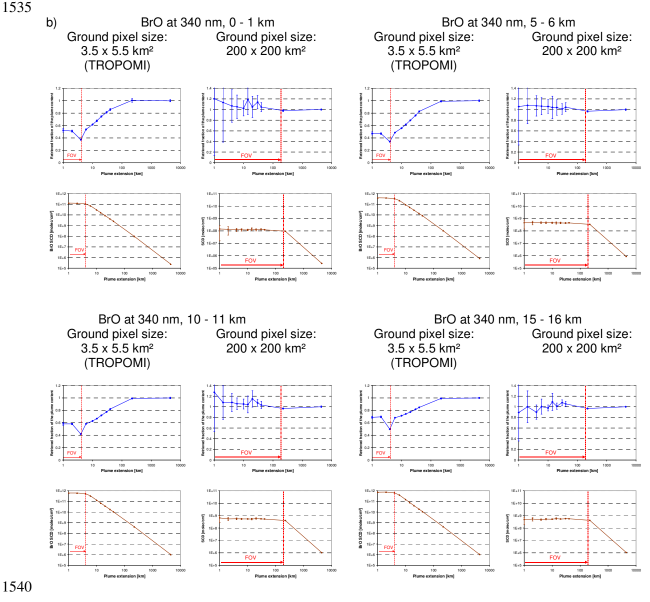


Fig. A3.1 Same as Fig. 5, but for additional plume heights: AMFs for plume scans in near-nadir viewing geometry (SZA=0°, VZA=0°) at different wavelengths. It is assumed that the satellite scans the plume with a narrow FOV ($\sim 0.014^{\circ}$). The different colours represent AMFs for plumes with different horizontal extensions (from 1 x 1 km² to 40 x 40 km²).







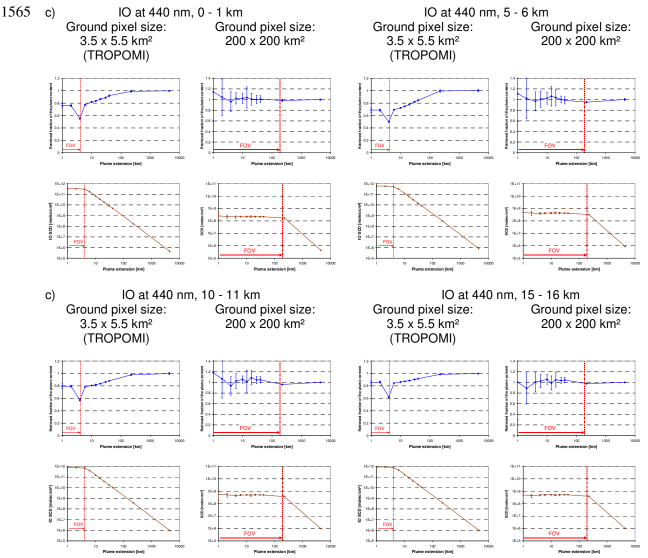
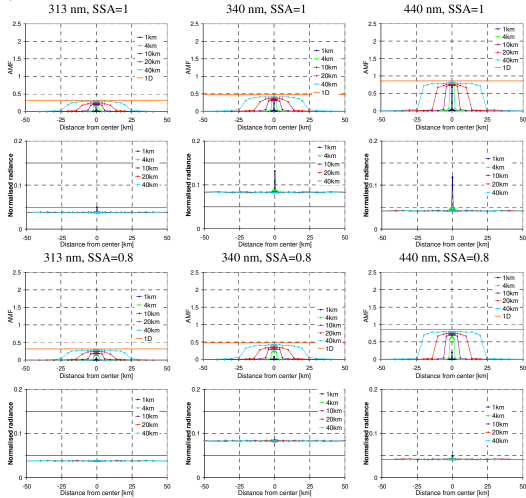
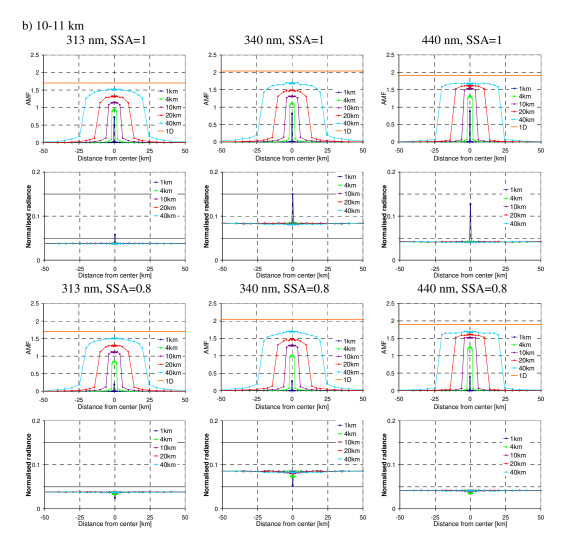


Fig. A3.2 Same as Fig. 8, but for additional plume heights and wavelengths: Top: Retrievd fraction of the plume cintent as a function of the horizontal plume extension (note the logarithmic scale). Bottom: measured trace gas SCDs (for a 1 x 1 km² pixel). The horizontal dashed line indicates the detection limits (see section 2.1). Left: results for a TROPOMI ground pixel; right: results for a large ground pixel of 200 x 200 km². Note that in the case of plume size > ground pixel size, only the fraction of the plume within the ground pixel size is considered. Simulations for VZA=0, and SZA=0°. The error bars represent the standard deviation calculated from 40 individual simulations.







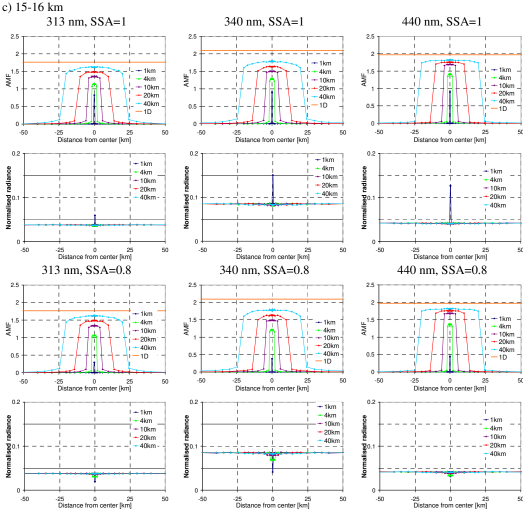
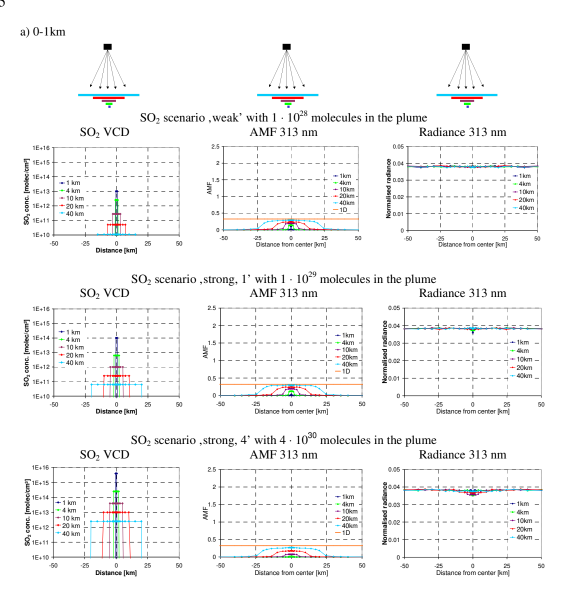


Fig. A3.3 Same as Fig. 11, but for additional plume heights: AMFs and normalised radiances for plume scans in near-nadir viewing geometry (SZA=0°) for different wavelengths and aerosol contents. The satellite scans the plume with a narrow FOV ($\sim 0.014^{\circ}$). The different colours represent AMFs for plumes with different horizontal extensions (from 1 x 1 km² to 40 x 40 km²). The rather low radiance at 313 nm (in spite of the high probability of Rayleigh scattering) is caused by the stratospheric ozone absorption.



1720

1725

1730

1735

b) 10-11 km SO_2 scenario ,weak' with $1 \cdot 10^{28}$ molecules in the plume AMF 313 nm Ra SO₂ VCD Radiance 313 nm 1E+16 SO² conc. [molec/cm³] 1E+14 1E+13 1E+12 1E+11 +10km +20km + 4 km + 10 km + 20 km + 40 km -40km SO_2 scenario, strong, 1' with $1 \cdot 10^{29}$ molecules in the plume AMF 313 nm SO₂ VCD Radiance 313 nm 1E+16 E 1E+15 1E+14 + 1km + 4km + 10km + 1km + 4km + 10km + 20km 1E+12 20km 40km g 1E+11 -50 0 Distance [km] -25 0 25 Distance from center [km] -25 0 25 Distance from center [km] SO_2 scenario, strong, 4' with $4 \cdot 10^{30}$ molecules in the plume SO₂ VCD AMF 313 nm Radiance 313 nm 1E+16 ፪ 1E+15 0/21E+14 1E+13 + 1 km + 4 km + 10 km + 20 km + 40 km + 1km - 4km - 10km **5** 1E+12 g 1E+11 0.5 -25 0 25 Distance from center [km] 0 Distance [km] -25 0 25 Distance from center [km]



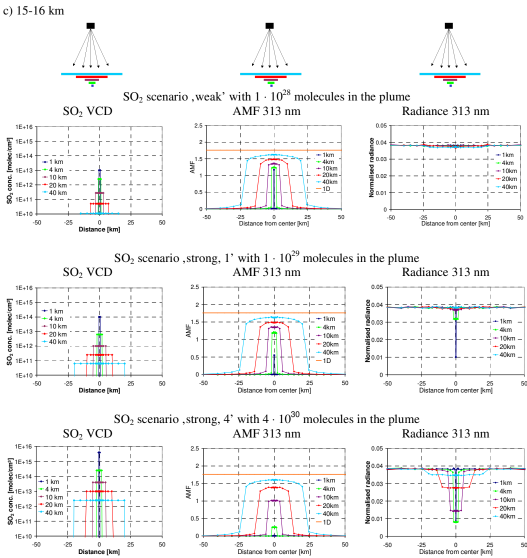
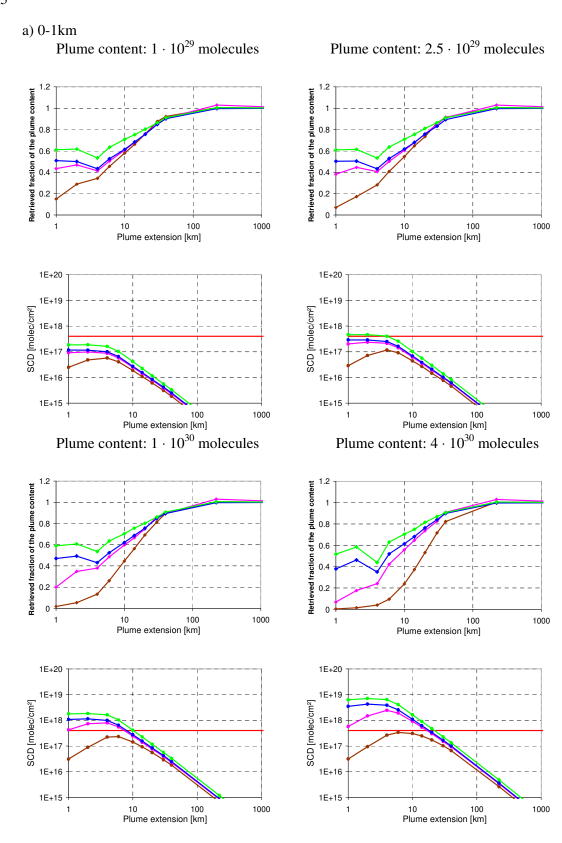


Fig. A3.4 Same as Fig. 13, but for additional plume heights: Plume scans for SO_2 plumes with different amounts of molecules and different horizontal extensions (for the different SO_2 scenarios, see table 3). Left: SO_2 VCDs of the plumes, middle: AMFs at 313 nm; right: normalised radiances at 313 nm.



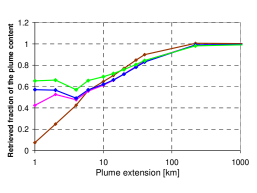
0.8 0.6

0.4

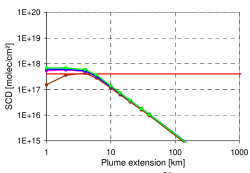
0.2

0

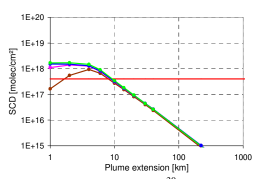
b) 10-11km Plume content: $1 \cdot 10^{29}$ molecules Retrieved fraction of the plume content



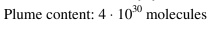
Plume content: $2.5 \cdot 10^{29}$ molecules

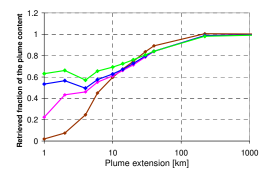


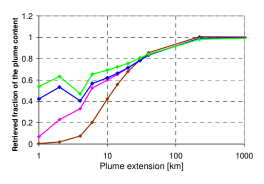
10 100 Plume extension [km]

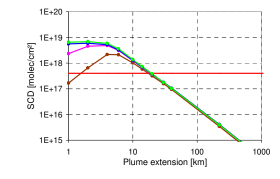


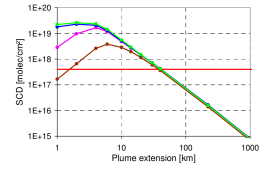
Plume content: $1 \cdot 10^{30}$ molecules











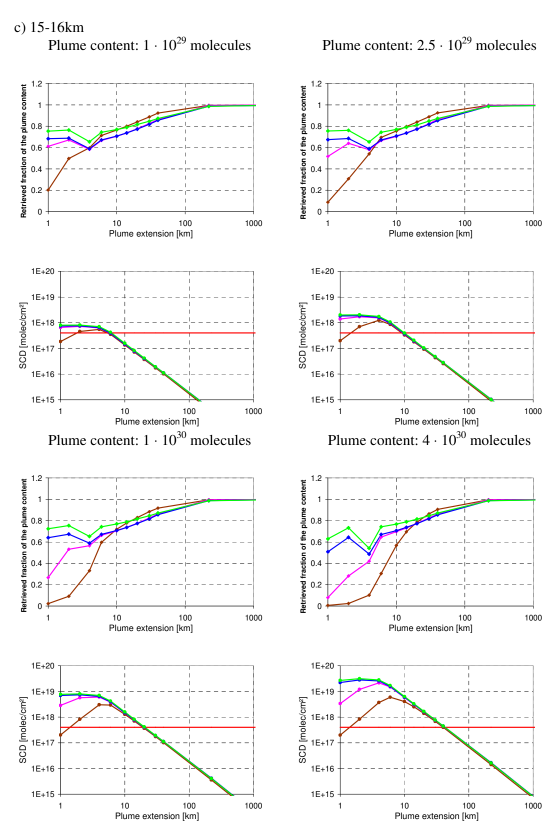
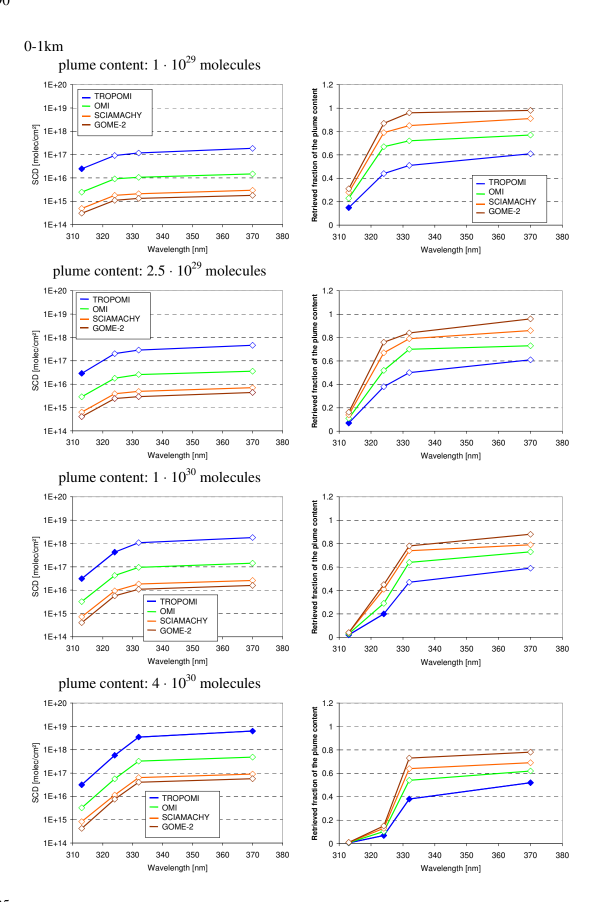
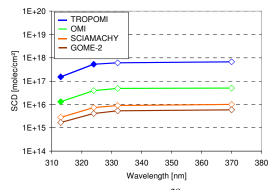


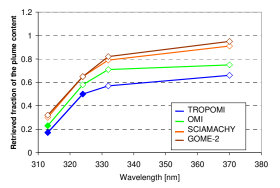
Fig. A3.5 Same as Fig. 14, but for additional plume heights: retrieved fraction of the plume content (top) and SO₂ SCDs (bottom) from TROPOMI measurements at different wavelengths as function of the plume size and for different amounts of molecules in the plume. The red horizontal line indicates the threshold, above which the operational SO₂ retrieval switches from the standard fit window to the first alternative fit window (at longer wavelengths).



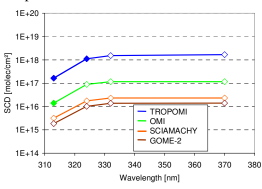
10-11km

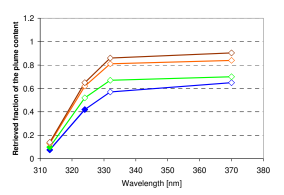




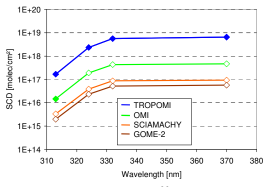


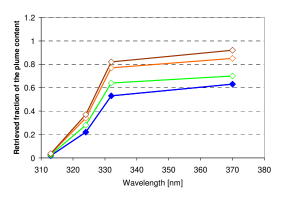
plume content: $2.5 \cdot 10^{29}$ molecules



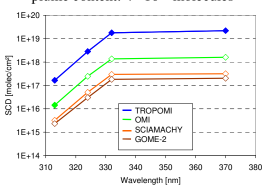


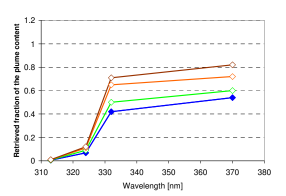
plume content: $1 \cdot 10^{30}$ molecules





plume content: $4 \cdot 10^{30}$ molecules





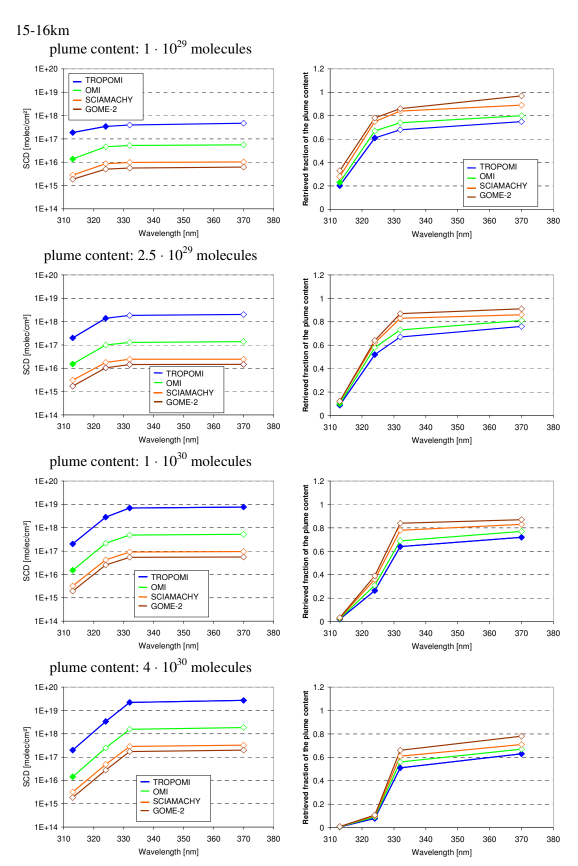


Fig. A3.6 Same as Fig. 15, but for additional plume heights: Retrievd fractions of the plume content and SO_2 SCDs for different satellite instruments and for plume size of 1 x 1 km² as function of the wavelength. The full symbols represent measurements with SCDs above the detection limit. Note that for the rather large ground pixel sizes of SCIAMACHY and GOME-2 the errors are between 5 and 10%.

Appendix A4 Effect of the wavelength dependence of the AMF

- 1815 In this section we quantify the effect of the wavelength dependence of the AMF for selected representative scenarios by the comparison of the results of two simulations:
 - a) monochromatic AMFs (only affected by the saturation effect)

SO₂ VCD (used as input for the respective scenario) to yield the ,true' AMF.

- b) ,true' AMFs (affected by the saturation effect and the wavelength dependence of the AMF)
- The ,true' AMFs are calculated in the following way (see also Marquard et al., 2000): first a measured spectrum for a selected SO₂ scenario is simulated at 'high' spectral resolution (~0.22 FWHM, the spectral resolution of the SO₂ cross section). In order to minimise the computational effort for Monte Carlo simulations (and also to minimise the noise), the spectra are calculated in the following way: first a wavelength close to the center of a SO₂ fit range is chosen. For this wavelength, the radiance is simulated as function of the absorption cross section. Note that in order to minimise the noise the same photon trajectories are used for the simulations with different absorption cross sections. Based on the derived relationship between the radiance and the absorption cross section, a high resolution spectrum for the selected SO₂ scenario is calculated and convolved by the instrument slit function (here we assume a FWHM of 0.5 nm). In the next step, the simulated spectra are analysed by DOAS similar to the measured spectra. Finally, the derived SO₂ SCD is divided by the
- The upper part of Figure A4.1 shows the ,true AMFs' derived for the standard fit window around 313 nm and the corresponding monochromatic AMFs for single wavelengths for 4 different plume scenarios. Additional results for 332 nm are shown in the bottom part of the figure. Besides the AMFs, also the ratios of the monochromatic AMFs and the true AMFs are shown. For scenarios with high AMFs (monochromatic AMFs > about 0.6), the difference between the monochromatic AMFs and the true AMFs is small (<10%), but for smaller AMFs, the difference increases (up to more than 50% for monochromatic AMF below about 0.4). In Fig. A4.2, the ratios of the true AMFs and monochromatic AMFs as function of the vertical optical depth of the SO₂ absorption are shown. For the same fit window, a similar dependence is found, almost independent from the SO₂ scenario. However, a different relationship is found for a different fit window indicating the specific spectral patterns of the SO₂ absorption in the different fit windows.

These findings indicate that for an accurate quantitative assessment, true AMFs for selected plume scenarios and wavelengths have to be individually calculated. Nevertheless, for the wavelengths considered here, the underestimation is < 5% if the vertical OD is below about 2.5%.

It should be also noted that for scenarios with high optical depth, the ,optical definition' of the AMF (equation 1) should be used for the interpretation of the measurements. For such cases, the AMFs calculated based on geometric photon trajectories (e.g. box-AMFs) will differ from those calculated by equation 1 in Pukite and Wagner (2016).

1845

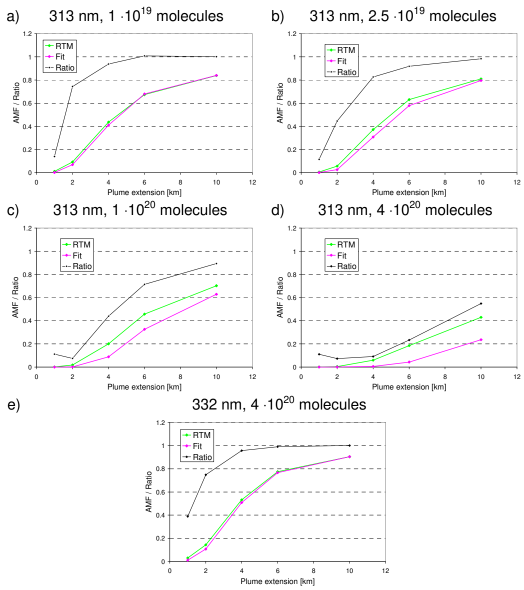


Fig. A4.1 True AMFs (magenta lines) and monochromatic AMFs (green lines) as function of the horizontal plume extension for different SO_2 scenarios. Also the ratios of both AMFs are shown (black lines). The AMFs and AMF ratios depend on the wavelength and the SO_2 amount.

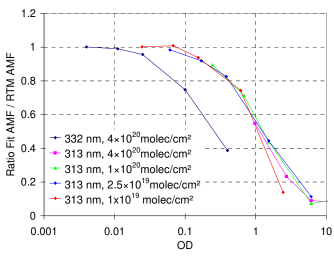


Fig. A4.2 Ratio between true AMFs and monochromatic AMFs as function of the vertical optical depth of SO_2 in the plume.



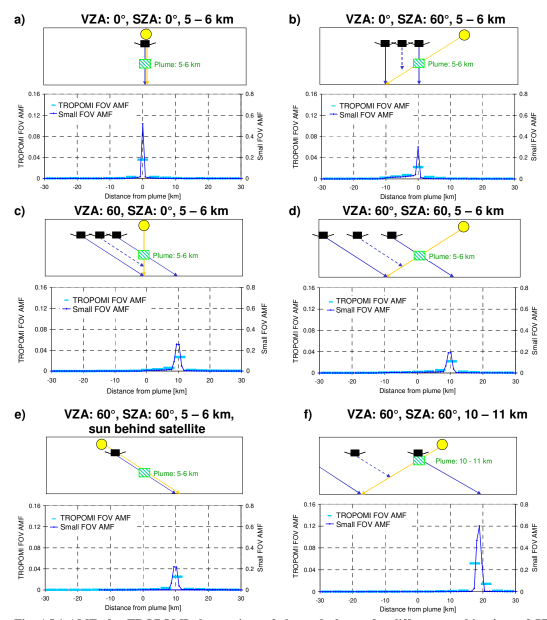


Fig. A5.1 AMFs for TROPOMI observations of elevated plumes for different combinations of SZA and VZA. The blue lines show the AMFs for observations with a narrow FOV ($\sim 0.014^{\circ}$) (right y axis); the bright blue symbols represent simulations with a TROPOMI FOV (left axis). All simulations are for SO₂ (weak) at 313 nm and plume sizes of 1 x 1 x 1 km³.

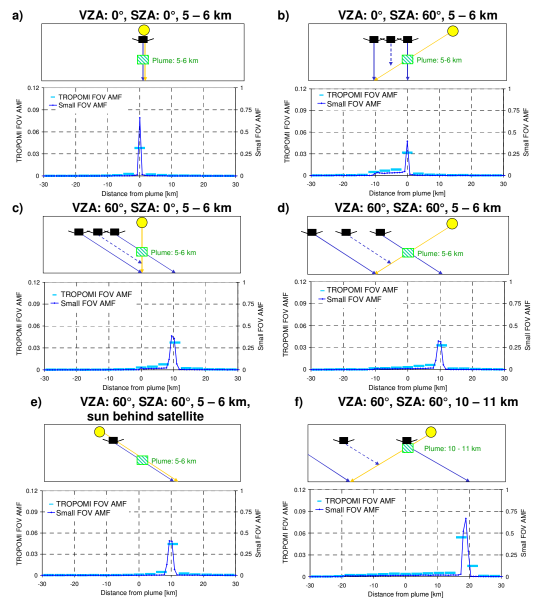


Fig. A5.2 AMFs for TROPOMI observations of elevated plumes for different combinations of SZA and VZA. The blue lines show the AMFs for observations with a narrow FOV ($\sim 0.014^{\circ}$) (right y axis); the bright blue symbols represent simulations with a TROPOMI FOV (left axis). All simulations are for BrO at 340 nm and plume sizes of 1 x 1 x 1 km³.