The Michelson Interferometer for Passive Atmospheric Sounding global climatology of BrONO$_2$ 2002–2012: a test for stratospheric bromine chemistry

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Abstract. We present the first observational dataset of vertically resolved global stratospheric BrONO$_2$ distributions from July 2002 until April 2012 and compare them to results of the atmospheric chemical climate model ECHAM/MESSy Atmospheric Chemistry (EMAC). The retrieved distributions are based on space-borne measurements of infrared limb-emission spectra recorded by the Michelson Interferometer for Passive Atmospheric Sounding (MIPAS) on Envisat. The derived vertical profiles of BrONO$_2$ volume mixing ratios represent 10$^\circ$ latitude bins and 3 d means, separated into sunlit observations and observations in the dark. The estimated uncertainties are around 1–4 pptv, caused by spectral noise for single profiles as well as for further parameter and systematic errors which may not improve by averaging. Vertical resolutions range from 3 to 8 km between 15 and 35 km altitude.

All leading modes of spatial and temporal variability of stratospheric BrONO$_2$ in the observations are well replicated by the model simulations: the large diurnal variability, the low values during polar winter as well as the maximum values at mid and high latitudes during summer. Three major differences between observations and model results are observed: (1) a model underestimation of enhanced BrONO$_2$ in the polar winter stratosphere above about 30 km of up to 15 pptv, (2) up to 8 pptv higher modelled values than observed globally in the lower stratosphere up to 25 km, most obvious during night, and (3) up to 5 pptv lower modelled concentrations at tropical latitudes between 27 and 32 km during sunlit conditions. (1) is explained by the model missing enhanced NO$_x$ produced in the mesosphere and lower thermosphere subsiding at high latitudes in winter. This is the first time that observational evidence for enhancement of BrONO$_2$ caused by mesospheric NO$_x$ production is reported. The other major inconsistencies (2, 3) between EMAC model results and observations are studied by sensitivity runs with a 1D model. These tentatively hint at a model underestimation of heterogeneous loss of BrONO$_2$ in the lower stratosphere, a simulated production of BrONO$_2$ that is too low during the day as well as strongly underestimated BrONO$_2$ volume mixing ratios when loss via reaction with O($^3$P) is considered in addition to photolysis. However, considering the uncertainty ranges of model parameters and of measurements, an unambiguous identification of the causes of the differences remains difficult.

The observations have also been used to derive the total stratospheric bromine content relative to years of stratospheric entry between 1997 and 2007. With an average value of 21.2 ± 1.4 pptv of Br$_y$ at mid latitudes where the modelled adjustment from BrONO$_2$ to Br$_y$ is smallest, the MIPAS data agree with estimates of Br$_y$ derived from observations of BrO as well as from MIPAS-Balloon measurements of BrONO$_2$. 

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Introduction

Besides chlorine, bromine is the major halogen constituent, with anthropogenic and natural sources affecting stratospheric ozone (e.g. Sinnhuber et al., 2009; Engel et al., 2018). After Wofsy et al. (1975) had described the possible relevance of bromine for ozone, the important role of bromine nitrate (BrONO$_2$) within stratospheric bromine chemistry was proposed by Spencer and Rowland (1978). They noticed the much faster photolysis of BrONO$_2$ compared to ClONO$_2$, which is an important prerequisite for the effectiveness of bromine ozone destruction cycles compared to those of chlorine (Lary, 1997; Klobas et al., 2020).

BrONO$_2$ is produced via the termolecular reaction (Burkholder et al., 2019, and references therein)

\[
\text{BrO + NO}_2 \xrightarrow{\text{M}} \text{BrONO}_2. \tag{R1}
\]

Due to its relatively short lifetime, the BrONO$_2$ concentration is strongly coupled to changes in NO$_2$ (Lary, 1996). The 1σ uncertainty factor of the reaction rate (Reaction R1) as provided by Burkholder et al. (2015, 2019) is 1.2 (i.e. 20% uncertainty) at 298 K, increasing to ~1.9 at a stratospheric temperature of 220 K.

The main loss process of BrONO$_2$ during the day is photolysis (Burkholder et al., 2019, and references therein):

\[
\text{BrONO}_2 + h\nu \rightarrow \text{Products}, \tag{R2}
\]

in which the products are Br + NO$_3$ and BrO + NO$_2$. The recommended quantum yields at wavelengths above 300 nm, being most important in the lower stratosphere, are 0.85 and 0.15, respectively. While in Sander et al. (2011) a combined uncertainty in cross sections and quantum yields of 1.4 is provided, the most recent evaluations (Burkholder et al., 2015, 2019) assign one wavelength-independent uncertainty factor of 1.2 (2σ) to the cross sections.

Further loss of BrONO$_2$ is due to atomic oxygen (Soller et al., 2001):

\[
\text{BrONO}_2 + \text{O}^3\text{P} \rightarrow \text{BrO} + \text{NO}_3, \tag{R3}
\]

which occurs, like Reaction (R2), only during sunlit conditions due to the necessary presence of O$^3\text{P}$. The 1σ uncertainty factor for the reaction coefficient varies between 1.25 at room temperature and 1.3 at 220 K (Burkholder et al., 2019). However, independent confirmation of the reaction parameters of Reaction (R3) is pending (Burkholder et al., 2019).

Finally, heterogeneous reactions can affect BrONO$_2$ concentrations, like hydrolysis in sulfuric acid aerosols (Burkholder et al., 2019, and references therein).

\[
\text{BrONO}_2 + \text{H}_2\text{O(s, l, H}_2\text{SO}_4 \cdot \text{nH}_2\text{O}) \rightarrow \text{HOBr} + \text{HNO}_3, \tag{R4}
\]

or in combination with halogens at surfaces, like

\[
\text{BrONO}_2 + \text{HCl(H}_2\text{O(s, H}_2\text{SO}_4 \cdot \text{nH}_2\text{O})} \rightarrow \text{BrCl} + \text{HNO}_3,
\]

where typical uncertainty factors of the gas–surface reaction probabilities are in the range of 2–4 (Burkholder et al., 2019).

Given the relatively large uncertainties in most of these leading reactions involving BrONO$_2$, comparison of observations to model calculations can be helpful for verification or even for suggesting improvements. For example, Kreyéc et al. (2013) analysed stratospheric balloon observations and concluded that the ratio $J_{\text{BrONO}_2}/k_{\text{BrO}+\text{NO}_2}$ should be increased to fit their data. Such investigations can be useful, first, to improve model simulations of stratospheric ozone loss and, second, to aid the analysis of the total stratospheric bromine (Br$_r$) content from observations of one species, such as BrO (e.g. Harder et al., 2000; Dorf et al., 2006a, 2008; Millán et al., 2012; Stachnik et al., 2013; Kreyéc et al., 2013; Werner et al., 2017; Engel et al., 2018).

Anthropogenic and natural emissions both contribute roughly equally to the present-day stratospheric bromine loading: Engel et al. (2018) give a best estimate of the total stratospheric bromine loading for 2016 of 19.6 pptv, of which natural sources contribute slightly more than 10 pptv. Brominated very short-lived substances (VSLs), such as bromofrom (CHBr$_3$) and dibromomethane (CH$_2$Br$_2$), contribute about 5 pptv to the stratospheric bromine loading, but their precise current contribution, any possible long-term changes, and the additional influx of inorganic product gases (product gas injection, PGI) are still uncertain (e.g. Sinnhuber et al., 2009; Aschmann and Sinnhuber, 2013; Falk et al., 2017). In this context, the observation of BrONO$_2$ provides an additional independent approach to determine total Br$_r$ (Wetzel et al., 2017) and, in consequence, to estimate the relative contribution of brominated VSLs.

Due to its spectral lines in the microwave and UV-vis, remote-sensing observations of BrO, the major inorganic bromine species in the lower stratosphere during sunlit hours, are common from the ground (e.g. Solomon et al., 1989; Carroll et al., 1989; Fish et al., 1995; Theys et al., 2007; Hendrick et al., 2008), from aircraft (e.g. Koenig et al., 2017; Werner et al., 2017; Rotermund et al., 2021), from balloons (e.g. Harder et al., 2000; Pundt, 2002; Dorf et al., 2006a, 2008; Stachnik et al., 2013; Kreyéc et al., 2013), and from satellites (e.g. Sinnhuber et al., 2005; Livesey et al., 2006; Kovalenko et al., 2007; McLinden et al., 2010; Rozanov et al., 2011; Millán et al., 2012; Parrella et al., 2013). In contrast, BrONO$_2$, the most important night-time reservoir of bromine, was detected in infrared limb-emission observations by the Michelson Interferometer for Passive Atmospheric Sounding (MIPAS) instrument on board the Envisat satellite only a decade ago (Höpfner et al., 2009). At that time the retrieval of altitude profiles was complicated by uncertainties in the infrared absorption cross sections of BrONO$_2$. In the meantime, Wagner and Birk (2016) provided an improved infrared spectroscopic database covering stratospheric conditions. On the basis of these new data, Wetzel...
et al. (2017) analysed the diurnal variation of BrONO$_2$ during three flights of the MIPAS-Balloon instrument.

In this paper we introduce the first day- and night-time climatology of stratospheric BrONO$_2$ as derived for the ~10-year lifetime of MIPAS/Envisat. We compare the results to global model simulations and discuss major differences by use of 1D photochemical modelling. Finally, the total stratospheric Br$_y$ content is estimated.

2 Methods

2.1 MIPAS instrument and data analysis

Flying on the polar-orbiting satellite Envisat, the limb sounder MIPAS recorded infrared spectra of the atmospheric thermal emission from 2002 until 2012 (Fischer et al., 2008). MIPAS was operated in two major modes: during period 1 (P1), between July 2002 and March 2004, the spectral resolution, defined here as $0.5 \times$ (maximum optical path difference)$^{-1}$, was 0.025 cm$^{-1}$, and during period 2 (P2) between January 2005 and April 2012, the resolution was set to 0.0625 cm$^{-1}$. During P1 the spectra of the “nominal” viewing modes as used in this work were taken at 17 tangent points between 7 and 72 km with 3 km steps up to 42 km and somewhat larger steps above. During P2, 27 spectra were recorded per limb scan with latitude-dependent tangent altitudes ranging from 5–70 km at the poles to 12–77 km over the Equator, with steps increasing with height from 1.5 to 4.5 km. The along-track sampling distance between each limb scan was ~550 km during P1 and ~420 km during P2. The local solar Equator-crossing time at the position of the tangent points is around 10:10 for the descending node and 22:20 for the ascending node of the Sun-synchronous orbit.

Since the first stratospheric detection of BrONO$_2$ (Höpfner et al., 2009), retrievals from averaged MIPAS spectra have been established for species with very weak signatures, such as SO$_2$ and NH$_3$ (Höpfner et al., 2013, 2016). The retrieval of vertical profiles of BrONO$_2$ volume mixing ratios as applied for the current dataset follows closely the procedure described in Höpfner et al. (2009). Here we briefly describe the retrieval scheme as well as the applied improvements with respect to Höpfner et al. (2009).

For the selection of spectra to be averaged zonally as well as temporally, the cloud filter method by Spang et al. (2004) has been applied to sort out any measurements affected by tropospheric as well as polar stratospheric clouds. Further, only spectra above about 15 km tangent altitude have been used for averaging so as to concentrate mainly on the stratosphere. We have applied a constrained nonlinear multiparameter least-squares fitting procedure to each limb sequence of averaged spectra to derive profiles of trace gas volume mixing ratios at 1 km-spaced vertical levels. Here we have used the same spectral interval (801–820 cm$^{-1}$) and atmospheric parameters simultaneously fitted with BrONO$_2$ (O$_3$, ClONO$_2$, NO$_2$, CFC$-$22, HNO$_4$, COF$_2$, HNO$_3$, ClO, CCl$_4$, CFC$-$113, PAN, T) as in Höpfner et al. (2009). We have applied a first-order smoothing constraint (Steck, 2002; Tikhonov, 1963) to dampen oscillations in the retrieved profiles. The regularization strength for each of the simultaneously derived species has been adjusted separately, and the related a priori profile for the target species BrONO$_2$ was set to 0.1 pptv, while for the other species climatological profiles have been used.

Major improvements and updates compared to Höpfner et al. (2009) are the following.

1. The most recent version (V8.03) of level-1B calibrated limb radiances by the European Space Agency (ESA) has been used (https://earth.esa.int/web/sppa-mission-performance/esa-missions/envisat/mipas/products-availability/level-1/level1-8.03, last access: 14 December 2021).

2. To simulate the spectral feature of BrONO$_2$, the new pressure- and temperature-dependent infrared spectroscopic database by Wagner and Birk (2016) has been used.

3. The spectroscopy of the interfering gases has been taken from the high-resolution transmission molecular absorption database (HITRAN) 2016 (Gordon et al., 2017) with the exception of HO$_2$NO$_2$. For this gas, the infrared cross sections in HITRAN for 220 K by May and Friedl (1993) have been extended by the ones of Friedl et al. (1994), which were measured at 298 K, to account for different atmospheric temperatures by two-point interpolation (Wetzel et al., 2017).

4. While in Höpfner et al. (2009) retrievals have been performed based on monthly mean spectra of September 2002 and 2003 within few coarse latitude bands, here we have subdivided the MIPAS measurements into 18 latitude bands of 10$^\circ$ spacing with a temporal binning of 3 d over the whole observational period 2002–2012.

We have estimated altitude-dependent errors of the BrONO$_2$ retrieval by applying assumptions about single error sources to two randomly selected periods in March and June for the years 2003 and 2009, i.e. during P1 and P2, respectively. The results are shown in Fig. 1 together with the total error profile calculated by quadratic combination of single error components. Instrumental uncertainties are estimated at 3% for the instrument line shape expressed as linear loss of modulation efficiency toward the maximum optical path difference of the interferometer (ILS), 1% for radiometric gain calibration (RadGain), and 300 m for tangent height knowledge (Htang). The uncertainty of European Centre for Medium-Range Weather Forecasts (ECMWF) temperatures (Temp) has been set to values of 2 K below and 5 K above 35 km altitude. The uncertainty of the BrONO$_2$ spectroscopy has been assumed to be 5%, which is on the conservative
side considering the 2 % (1σ) error estimation given in Wagner and Birk (2016). Further errors refer to the spectroscopic parameters of interfering gases. For those, we have assumed uncertainties of 5 % for species described by cross sections (SpecXitf), 5 % for intensities (SpecINTitf) and 10 % for the half-widths of the line parameters (SpecHHWt). These assumptions are within the typical errors provided in the database (Gordon et al., 2017). A further error term describing the retrieval from averaged spectra (NonLin) has been estimated on the basis of dedicated retrieval simulations as detailed in Höpfner et al. (2009). For this estimate the values used for the tangent altitude scatter of single observations were set to 400 and 300 m (1σ) during P1 and P2, respectively.

The total error estimate as calculated by quadratic combination of the single components is given by the blue lines in Fig. 1. Around this total error estimate, the blue shading indicates the variability of the estimated errors for all latitude bands. In general, the estimated total errors vary between about 1 and 4 pptv, independent of day- or night-time observations. They appear to be slightly smaller during P1 compared to P2, which is probably due to the better spectral resolution during P1. Further, it is evident from Fig. 1 that, below 20–23 km, total parameter errors and spectral noise are the dominant contribution to the total error, while at larger altitudes it is mostly dominated by spectral noise.

As a further diagnostic measure of the retrieval, Fig. 2 shows the vertical resolution as a function of altitude. It has been calculated by dividing the retrieval grid width of 1 km by the diagonal elements of the averaging kernel matrices (Rodgers, 2004). The vertical resolution is about 3 km at 15 km altitude and becomes coarser with altitude, reaching 8 km at 35 km altitude. The vertical resolution is generally finer at the tangent altitudes and coarser at retrieval levels between the tangent altitudes. Conversely, the retrieval noise is larger at the tangent altitudes and smaller at altitude levels in between. These effects are only visible when a retrieval set-up is chosen where the retrieval grid is finer than the tangent altitude spacing. Since in period P1 the tangent altitude grid is fixed, this effect survives the averaging, leading to a zigzag profile of the vertical resolution. In contrast, in P2 the tangent altitude grid varies with latitude, and the zigzag features of vertical resolution average out.

2.2 Atmospheric modelling

We have compared the MIPAS BrONO₂ dataset with a multi-annual simulation from the chemical climate model ECHAM/MESSy Atmospheric Chemistry (EMAC) (Jöckel et al., 2010). Within EMAC, the interface Modular Earth Submodel System (MESSy) links the sub-models describing tropospheric and middle atmospheric processes to the dynamical core, the fifth-generation European Centre Hamburg general circulation model ECHAM5 (Roelckner et al., 2006). We have used EMAC (ECHAM5 version 5.3.02, MESSy version 2.52) at the T42L90MA resolution with 90 vertical hybrid pressure levels from the ground up to 0.01 hPa (~80 km) and a horizontal resolution of ∼2.8° × 2.8° latitude × longitude. The sub-models MECCA (Sander et al., 2005) and MSBM (Kirner et al., 2011) simulate gas-phase chemistry and polar stratospheric clouds including heterogeneous reaction rates, respectively. To reproduce realistic conditions for comparison with the observations, the model run was nudged towards the ECMWF reanalysis ERA-Interim (Dee et al., 2011) by a Newtonian relaxation technique of surface pressure, temperature, vorticity, and divergence above the boundary layer and below 1 hPa (van Aalst, 2005). We have applied a comprehensive chemistry set-up from the troposphere to the lower mesosphere with more than 100 species involved in gas-phase, photolysis, and heterogeneous reactions on liquid sulfate aerosols, nitric acid trihydrate (NAT), and ice particles. Rate constants for gas-phase reactions have been taken mainly from Atkinson et al. (2007) and the Jet Propulsion Laboratory (JPL) compilation (Sander et al., 2011). Photochemical reactions of short-lived bromine-containing organic compounds CHBr₃, CH₂Br₂, CH₂ClBr, CHClBr₂, and CHCl₂Br are included in the model set-up (Jöckel et al., 2016). Boundary conditions for CH₃Br and the bromine-containing halons were taken from Meinshausen et al. (2011) and extended with the RCP6.0 scenario as suggested by Eyring et al. (2013). We have used scenario 5 of Warwick et al. (2006) to describe the surface emissions of these organic bromine species. During the MIPAS measurement periods, from the model output first all data within 1 h around 10:00 and 22:00 LT were selected. Depending on their latitude, longitude and altitude, they were then assigned to sunlit and dark conditions and averaged over the observational bins of 10° latitude and 3 d periods. In the extreme cases of twilight conditions at high latitudes, this might induce differences between modelling and observations. However, in the discussion below we only refer to situations which are not affected by twilight conditions.

For specific sensitivity investigations at low latitudes, we have applied a 1D photochemical stacked box model. The chemical mechanism of the 1D model is based on the SLIMCAT model (Sinnhuber et al., 2005, and references therein). The 1D model runs have been initialized with equatorial mean profiles of the EMAC simulation of all inorganic Br species and NO₂. For the other species as well as pressure and temperature, equatorial profiles from the MIPAS reference database have been used (Remedios et al., 2007). A comparison of the parameters of several bromine reactions between EMAC, the 1D baseline model run and the JPL2019 compilation (Burkholder et al., 2019) is provided in Table D1.
Figure 1. Retrieval error estimates from four 3 d periods, two during P1 (a, b) and two during P2 (c, d), for both dark (a, c) and sunlit (b, d) conditions. Considered error sources are the uncertainties of the instrumental line shape and radiometric gain calibration (ILS, RadGain), the pointing knowledge (Htang), assumed temperature profiles (Temp), spectroscopic errors of BrONO$_2$ absorption cross sections (SpecBrONO$_2$) and errors in cross sections (SpecXitf), line half-widths (SpecHWitf) and line intensities (SpecINTitf) of interfering species, as well as the error due to the applied technique of retrievals from averaged spectra (NonLin) and the spectral noise of the instrument (Noise). The total error (Total err) has been determined by quadratic combination of all single error components, while the combined parameter and systematic error (Tot paraerr) considers all uncertainties except the spectral noise. The blue-filled space around the total error curves indicates the areas into which 90% of the total error profile estimates fall.

3 The MIPAS dataset in comparison to EMAC model results

3.1 Overview of the measurements

We provide overviews of the MIPAS BrONO$_2$ volume mixing ratio datasets at full temporal resolution in the left-hand-side panels of Figs. 3 and 4 for observations during dark (night) and sunlit (day) conditions, respectively. White spaces indicate regions where no measurements are available, such as observations in the dark during high-latitude summer as well as sunlit measurements during winter. A measurement gap due to instrumental issues of MIPAS happened between April 2004 and January 2005, and in the subsequent years, observations were ramped up through to
about 2007. From then on, quasi continuous coverage exists until April 2012. The coverage at lower altitudes is determined by the lower limit of 15 km, chosen to confine the retrievals primarily to the stratosphere, and by the presence of high-altitude clouds and the scan pattern of MIPAS (which is mainly a factor in the tropics). Some additional data gaps exist at high southern latitudes during winter, when thick polar stratospheric clouds (PSCs) obscured the observations.

From Figs. 3 and 4, the major features of the stratospheric BrONO$_2$ variability can be discerned in our measurements.

1. The diurnal variability (Fig. 3 versus Fig. 4) as a manifestation of the fast photolysis during the day (Reaction R2) versus the production (Reaction R1).

2. The annual recurrence of low values during night at high latitudes (Fig. 3) due to the lack of NO$_x$ as a supply for the production (Reaction R1) in combination with heterogeneous loss due to the presence of PSC particles (Reactions R4 and R5).

3. The annual maxima of BrONO$_2$ volume mixing ratios at high and mid latitudes during day- and night-time observations in summer caused by the annual variability of NO$_2$.

4. The lack of a similarly clear seasonal variability at tropical latitudes.

We have reduced the dataset to annual views by averaging over the whole MIPAS observational period in order to provide a clearer picture of intra-annual variabilities. The related horizontal and vertical cross sections are presented in Figs. 5–8. During dark conditions, maximum mean mixing ratios of around 22 pptv are reached mainly at mid latitudes at altitudes of 25–30 km during the summer months, while in winter only about 18 pptv is observed. During sunlit conditions, the largest BrONO$_2$ mixing ratios of up to about 11 pptv appear at high latitudes at 20–25 km altitude during summer. Negative retrieved mean mixing ratios in the subtropics and tropics below about 18–22 km altitude are discussed more in detail below.

3.2 Measurement–model comparisons

The results of the EMAC model run are presented in the second column in each of Figs. 3–8. To take into account the limited vertical resolution of the measurements for these comparisons, we have applied the averaging kernel matrix of each retrieved profile of BrONO$_2$ (see Sect. 2.1) to the related modelled profile. Differences between model results with and without averaging kernel application are generally below 1 pptv, with only sporadic exceptions at the highest altitudes near polar latitudes, where differences reach up to 3–4 pptv (Figs. A1 and A2).

From comparing measured and modelled distributions of BrONO$_2$ volume mixing ratios in Figs. 3–8, it is evident that the model reproduces all major modes of variability which are present in the observations as described in the previous section. Despite this agreement, there are a few areas where systematic deviations are prominent. The most obvious differences appear (1) at higher altitudes during polar winter, (2) in the lower stratosphere mainly at mid and low latitudes during the entire year as well as (3) at altitudes around 30 km in the tropics during sunlit conditions. We discuss these differences one by one below.

3.2.1 Low modelled polar winter BrONO$_2$

One disagreement between modelled BrONO$_2$ and MIPAS observations can best be observed in Figs. 5 and 7: at altitudes of around 30 km and above for latitudes south of 70$^\circ$S, the model predicts values smaller than 5 pptv from May until September. The corresponding measurements, however, reach values of 10 to 15 pptv. While not as pronounced, this feature is also visible in the Arctic wintertime stratosphere, with model estimates of 5–10 pptv and measurements of 10–15 pptv. We explain these low model values of BrONO$_2$ by an underestimation of NO$_2$ as visible in Fig. B1. In this figure, the MIPAS NO$_2$ distributions are compared to the EMAC model results of NO$_2$. The missing NO$_2$ in the simulations is due to an insufficient supply of NO$_x$ through downward propagation from the upper mesosphere and lower thermosphere. This stratospheric enhancement of
Figure 3. Horizontal cross sections (latitude versus time) of measured (left) and modelled (right) BrONO₂ volume mixing ratios at selected altitudes over the whole time period of MIPAS observations during dark conditions. Retrieval averaging kernels have been applied to model data. White areas indicate the absence of measurements.
Figure 4. Same as in Fig. 3 but for sunlit measurements.
Figure 5. Horizontal cross sections (latitude versus time) of the annual development of measured (left) and modelled (right, with averaging kernels applied) BrONO$_2$ volume mixing ratios during dark conditions at selected altitudes calculated as average values over the whole MIPAS dataset in weekly bins (see Fig. 3).
Figure 6. Same as Fig. 5 but for sunlit measurements (see Fig. 4).
Figure 7. Bimonthly averaged cross sections (altitude versus latitude) of BrONO₂ volume mixing ratios for dark conditions. Left: measurements, middle: model with retrieval averaging kernels applied, right: model minus MIPAS.
Figure 8. Same as Fig. 7 but for sunlit conditions.
NO\(_x\) through production by energetic particle precipitation and downwelling during polar winter has been investigated e.g. on the basis of MIPAS observations by Funke et al. (2005, 2014).

3.2.2 High modelled night-time BrONO\(_2\) at lower altitudes

As visible e.g. in Figs. 5 and 7, the model overestimates BrONO\(_2\) volume mixing ratios with respect to the MIPAS results by up to 8 pptv in the lower stratosphere at altitudes up to about 25 km during the night. Such differences are also present during the day, albeit to a smaller absolute extent (up to about 4 pptv; see Fig. 8). Relative differences between measurements and model results during the day and night are more similar, reaching e.g. 50 % at around 20 km at mid latitudes and 22–25 km in the tropics. Below, the day-time relative differences become much larger due to the very small mixing ratios of BrONO\(_2\) and are difficult to compare.

The black line in panel “night” of the top row in Fig. 9 represents the measured mean night-time profile over the whole period at 5° S. The related EMAC model result is shown in red. Maximum differences between both are about 8 pptv at around 20–22 km altitude. As possible explanations for these differences we will discuss below (1) measurement errors, (2) wrong modelling of the release of Br\(_x\) from its source gases, (3) wrong partitioning of Br\(_x\) between its main constituents in the model simulations, and (4) wrong modelling of NO\(_2\) mixing ratios.

1. The degree of discrepancy between measurement and model of up to 8 pptv over an altitude range of around 5 km cannot reasonably be explained given the errors estimated for the MIPAS BrONO\(_2\) retrieval (Fig. 1). This would allow for discrepancies of around 2–3 pptv, especially considering that the “noise” error term is strongly reduced by the temporal averaging over all equatorial measurements. Still, one cannot rule out unequivocally any unidentified additional systematic error source in the measurements contributing to these differences.

2. Dorf et al. (2008) report on balloon observations of BrO in the framework of an ENVISAT validation campaign in Teresina, Brazil (5.1° S, 42.9° W), on 17 June 2005. In the altitude region of 20–22 km they observed volume mixing ratios of BrO of around 6–10 pptv, which agrees with our EMAC model results during day time (see Fig. 9, second row, red line in panel “day”) that indicate 8–11 pptv of BrO in the same altitude range. Similar mixing ratios of BrO at these altitudes in the tropics have also been reproduced by other model simulations (e.g. Theys et al., 2009) as well as by satellite observations (Sinnhuber et al., 2005; Sioris et al., 2006; Rozanov et al., 2011; Parrella et al., 2013). A possible contribution to a model overestimation of BrONO\(_2\) might be the emission scenario used for organic bromine species, taken from Warwick et al. (2006). As has e.g. been shown by Keber et al. (2020), this scenario probably leads to an overestimation of brominated VSLS by up to 2 pptv, which is, however, not enough to explain the observed differences. Thus, it is highly unlikely that the inorganic bromine content at 20–25 km is strongly overestimated in the EMAC model calculations.

3. The modelled partitioning of Br\(_x\) at those altitudes during night is essentially determined by the heterogeneous conversion of BrONO\(_2\) into HOBr through sulfate aerosols (Reaction R4). For a more detailed investigation, we have performed sensitivity simulation with the 1D model, the results of which are shown in Fig. 9. The rate coefficient for heterogeneous reactions is proportional to the aerosol surface area density (SAD) and the reaction probability \( \gamma \). For the baseline 1D calculation, we have applied a vertical profile of aerosol surface area densities which was derived from the mean tropical aerosol volume densities as available from MIPAS (Güntner et al., 2018) assuming a lognormal size distribution with a number density of 10 cm\(^{-3}\) and a width of 1.8. Figure 10 provides a comparison of the resulting “MIPAS mean” surface area densities with SAGE II profiles in the tropics between July 2002 and August 2005 (Damadeo et al., 2013) as well as in situ measurements at mid latitudes over Laramie (41° S, 105° W) between 2002 and 2012 (Deshler et al., 2019). The reaction probabilities for the 1D baseline run (dashed pink curve in Fig. 10b) were determined according to Hanson (2003) as reported by Burkholder et al. (2019).

A test without consideration of heterogeneous conversion of BrONO\(_2\) into HOBr (dotted orange line in Fig. 9) results in a night-time increase in BrONO\(_2\) by up to 5 pptv as well as the corresponding decrease in HOBr. To replicate the observations of BrONO\(_2\), we performed two tests by adjusting the profile of aerosol surface area density: one (a) where we kept the altitude-dependent reaction probabilities (dashed pink curves in Figs. 9 and 10) and one (b) where the reaction probabilities were set to 1 over the entire altitude range (dashed-dotted purple curves in Figs. 9 and 10). As can be seen from Fig. 10, in test (a) one would require surface area densities enhanced by factors of more than 3 to deplete BrONO\(_2\) to such an extent that the observations in the altitude range 20–26 km are met. For the idealized case of \( \gamma = 1 \), the aerosol surface area densities had to be adjusted up to 24 km only, however, still by a factor of more than 3 at 20 km altitude.

4. As the formation of BrONO\(_2\) is strongly linked to NO\(_2\), we have inspected its modelled mixing ratios in comparison to the observed ones. As can be seen in Figs. B1 and B3, the model calculations agree well with the mea-
measurements, especially at the lower altitudes where the observed discrepancies in BrONO$_2$ occur. Thus, we exclude a wrong modelling of NO$_2$ as an explanation for these differences.

In summary, we have to conclude that there is no compelling evidence for any of the explanations above causing the observed differences between measurements and model results. While (3) would imply an increase in aerosol surface area density contradicting related observations, (2) would oppose observations of BrO, (4) provides no hint of a discrep-
Finally, under (3), we will briefly address differences between not considered in the present implementation of EMAC. Further, we will address the effect of Reaction (R3), which is uncertainties in the parameters of Reactions (R1) and (R2).

3.2.3 Low modelled day-time BrONO$_2$ at low latitudes

The EMAC model calculations fit the observed night-time BrONO$_2$ maximum at 29–30 km altitude in the tropics quite well (see Fig. 7). However, during day time (Fig. 8), BrONO$_2$ is almost entirely depleted in the EMAC simulations at these altitudes, while MIPAS still detects maximum values of up to 5 pptv. Possible reasons for this discrepancy discussed below might be (1) measurement errors or (2) the impact of uncertainties in the parameters of Reactions (R1) and (R2). Further, we will address the effect of Reaction (R3), which is not considered in the present implementation of EMAC. Finally, under (3), we will briefly address differences between measured and modelled NO$_2$ mixing ratios.

1. In the top row of Fig. 11, the mean profile of BrONO$_2$ measured at 5° S is again shown together with the EMAC model results in panel “day”. As in the case of the night-time discrepancy, the values of BrONO$_2$ observed during sunlit conditions cannot be explained by the estimated retrieval errors, which are about 1–2 pptv at the 30 km region (Fig. 1). However, as can be seen in Fig. 11, the mean measured profile values below about 24 km are negative, indicating an unresolved issue in the retrieval at these altitudes. Still, the negative values are in the range of the estimated uncertainties. To investigate whether the positive values above where the discrepancy with the model becomes apparent might be due to an oscillatory feature caused by the negative values below, we have tested different retrieval options (increasing the regularization strength, performing retrievals of log(vmr) instead of vmr, skipping tangent altitudes below). In all the tests, the maximum values at around 30 km still appeared in the retrieved profiles, indicating them as a robust feature.

2. Sensitivity calculations using the 1D model are also shown in Fig. 11. As a baseline for these simulations, we have applied the setting “SAD adapted” so that the calculations during night-time also fit to the observations at altitudes below 27 km. The resulting profile from the 1D model run also shows values which are smaller than the observed BrONO$_2$ mixing ratios by about 3 pptv at around 30 km altitude. Though about 1 pptv larger than the EMAC simulations at those altitudes, the 1D results are also not compatible with the measurements.

To test the sensitivity with modelled BrONO$_2$ to its production via the three-body Reaction (R1), we have used the JPL2019 formulation (Burkholder et al., 2019) instead of that by Atkinson et al. (2007) that was applied in the EMAC and 1D baseline runs (Table D1). This led to an increase in BrONO$_2$ vmr values by about 1pptv; see the dash-dotted green curve in Fig. 11. Further increasing these rate coefficients by a factor of 2, which is well covered by the 2σ uncertainty factor of 3.7 for that reaction at 220 K (Burkholder et al., 2019), led to an additional increase by about 3pptv (dash-dotted olive curves in Fig. 11). While these results coincide now with the observed day-time abundances of BrONO$_2$ at around 30 km, the increase at lower altitudes does not correspond well to the observations there. Moreover, around 30 km during night, the calculations overestimate the measurements by up to 2pptv.

To test the sensitivity with respect to the photolysis of BrONO$_2$ (Reaction R2), we have divided the photolysis rate by 1.2, the 2σ uncertainty as provided by Burkholder et al. (2019). The result is illustrated by the blue dashed curves in Fig. 11. The resulting increase of only around 0.5 pptv is too small compared to the observations. Moreover, the increase appears over a larger altitude range (from 22 up to 34 km) compared to the more confined region between about 27 and 33 km where the increased day-time values are observed.

The grey dotted curve in Fig. 11 illustrates the effect on the simulated mixing ratios when the loss Reaction (R3) is included in the 1D model by using the reaction coefficients from the JPL2019 compilation; see Table D1 (Burkholder et al., 2019). With a reduction of up to

![Figure 10](https://example.com/figure10.png)

**Figure 10.** (a) Vertical profiles of surface area densities (SADs). Blue: mean MIPAS tropical profile (Günther et al., 2018) (see text for details), orange: mean profile from in situ observations over Laramie (41° S, 105° W) between 2002 and 2012, green: mean SAGE II profiles in the tropics between 2002 and 2005 (Damadeo et al., 2013). The dashed pink and dash-dotted purple lines show the adapted SAD profiles corresponding to the 1D model results indicated by the same line styles in Fig. 9. (b) Reaction probability profiles for hydrolysis of BrONO$_2$ (Reaction R4). The dashed pink profile refers to the standard one determined in the 1D model (Hanson, 2003; Burkholder et al., 2019), and the dash-dotted purple curve has been used in the sensitivity analysis to adjust the SAD (a and Fig. 9).
1.5 pptv, this obviously drives the concentrations further away from the observations.

3. In the tropics at around 30 km, the mixing ratios of NO\textsubscript{2} from the measurements are around 10 \%-20 \% smaller than modelled by EMAC, with the results of the 1D model lying in between (see Figs. B2 and B4). These deviations cannot account for the model underestimation of BrONO\textsubscript{2} since they would even imply stronger modelled production of BrONO\textsubscript{2}.

In conclusion, we have found no unequivocal explanation for the high measured day-time mixing ratios of BrONO\textsubscript{2} at around 30 km over the tropics/subtropics: (a) the differences between models and observations are outside the estimated measurement errors, (b) the uncertainties of the cross sections for photolysis (Reaction R2) of BrONO\textsubscript{2} are by far too small, (c) the error estimates for Reaction (R1) would allow a sufficient increase in BrONO\textsubscript{2} mixing ratios but over an overly large vertical extent, and (d) any inclusion of Reaction (R3) opens the gap between simulated and observed BrONO\textsubscript{2} concentrations even more.

4 Estimation of total stratospheric Br\textsubscript{y}

The MIPAS dataset of BrONO\textsubscript{2} allows us to determine the total stratospheric content of bromine (Br\textsubscript{y}) by using the so-called inorganic method. It is a linear correction of the observed bromine species by multiplication by the ratio between total modelled bromine (Br\textsubscript{y}\textsubscript{mod}) and the modelled bromine species observed. This procedure has often been applied in case of day-time observations of BrO (e.g. Dorf et al., 2006a, b). Wetzel et al. (2017) were the first to apply this method to night-time observation of BrONO\textsubscript{2}:

\[
Br_y = BrONO_2^{\text{meas}} \times \frac{Br_y^{\text{mod}}}{BrONO_2^{\text{mod}}}. \tag{1}
\]

Using night-time observations of BrONO\textsubscript{2} instead of BrO measured during the day to derive Br\textsubscript{y} should have the advantage that BrONO\textsubscript{2} during the night makes up a larger fraction of Br\textsubscript{y} than BrO does during the day, due to the continuous production of BrONO\textsubscript{2} (Reaction R1). The modelled night-time ratio between BrONO\textsubscript{2}\textsubscript{mod} and Br\textsubscript{y}\textsubscript{mod} has been examined to decide which region (in terms of altitude and latitude) to use for determination of Br\textsubscript{y} from the MIPAS dataset (cf. Fig. C1). Values clearly exceeding 90 \% are simulated at mid latitudes, mainly during spring/summer/autumn, and centred at altitudes around 26 km. Thus, for the analysis we have chosen data at 25–26 km altitude and 40–60° latitude from October to March in the Southern Hemisphere and from April to September in the Northern Hemisphere. Additionally, due to the low seasonal variability of BrONO\textsubscript{2} and to capture relatively young air masses, we also considered tropical data at 29–30 km in the analysis. While not exceeding 90 \%, the ratio BrONO\textsubscript{2}\textsubscript{mod}−Br\textsubscript{y} is still larger than 85 \% (see Fig. C1).

For stratospheric measurements, the values of derived total inorganic bromine are generally assigned to their year of entry into the stratosphere (e.g. Engel et al., 2018, Fig. 1-16). Here we have used an update of the age-of-air dataset determined from MIPAS retrievals of SF\textsubscript{6} (Haenel et al., 2015; Stiller et al., 4–8 May 2020). Typical values of age of air in the case of the mid-latitude observations range between 5 and 6.5 years and in the tropics between 3.5 and 5 years, with errors of about 1 year.

Depending on the date of stratospheric entry, total Br\textsubscript{y} (red) estimated from the MIPAS observations through Eq. (1) in comparison to the original MIPAS observation (BrONO\textsubscript{2}\textsubscript{meas}, blue) is shown in Fig. 12. The solid lines represent weighted means of all data points, where for the weighting the a posteriori estimated random error (dotted black curves in Fig. 1) was applied. The resulting random error as indicated by the thickness of the line is very small (0.04–0.07 pptv) due to the large number of data points. The major part of the uncertainty in our estimation of total Br\textsubscript{y}, however, is due to the combined parameter and systematic error (the green line in Fig. 1), which is shown by the red shading in Fig. 12. It should be noted that this error term is a combination of a variety of estimated uncertainties, each of which might also partly be random in nature with different temporal correlation lengths (e.g. correlated over each of the two measurement phases). Thus, we have made a conservative assumption in considering all those to be systematic – but possibly also underestimate them by applying quadratic combination of the single error terms.

Our estimates of total Br\textsubscript{y} vary from 21.0 ± 1.4 and 21.4 ± 1.4 pptv for the northern and southern mid-latitude regions, respectively (years of stratospheric entry: 1997–2006), to a maximum of 22.4 ± 1.7 pptv in the tropical stratosphere (years of stratospheric entry: 1998–2007) (Table 1). The values of Br\textsubscript{y} from the mid latitudes of both hemispheres coincide clearly within their uncertainty ranges. Since it is unlikely that the real values of total Br\textsubscript{y} vary strongly in the stratosphere, the differences of 1–1.4 pptv between tropical and mid-latitude estimates are more probably caused by uncertainties. These may either be caused by errors in the BrONO\textsubscript{2} concentrations derived from MIPAS or may be due to the calculation of total Br\textsubscript{y} from BrONO\textsubscript{2} through Eq. (1) (or a combination of both). The first explanation would require a retrieval error component varying with latitude, e.g. due to some temperature dependence, while the second one implies model uncertainty. Since the model adjustment of Br\textsubscript{y} from BrONO\textsubscript{2} is much larger in the tropical stratosphere (about 2.5 pptv) than at mid latitudes (about 0.5 pptv), the second explanation would affect more strongly our estimation of Br\textsubscript{y} in the tropics.

Br\textsubscript{y} obtained from MIPAS can be compared to data of Br\textsubscript{y} derived from observations of BrO, as summarized for exam-
Figure 11. Sensitivity 1D model simulations to test the day-time EMAC model underestimation of BrONO$_2$ around 29 km altitude for BrONO$_2$ (top row), BrO (middle row), and HOB (bottom row). Left: diurnal evolution of the “1D base” run for a period of 2 d. The white dashed lines indicate the local solar times of the MIPAS observations. The columns “night” and “day” contain the averaged night-time and day-time profiles of MIPAS at 5° S as black solid lines along with error bars indicating the 2σ estimated measurement uncertainty. The mean EMAC results are provided in red. The MIPAS and EMAC curves are the same as in Fig. 9. The other curves illustrate the results of the 1D model simulations and the columns “diff night” and “diff day” show the differences between the 1D sensitivity runs and the “1D base” run.

In Fig. 1-16 of Engel et al. (2018). To provide an easy way of comparison, we have replicated the single values of those datasets in each panel of our Fig. 12 and have collected their respective mean values in Table 1, limited to the period of stratospheric entry from the MIPAS dataset. Notably, all of these observations are compatible with the values derived from MIPAS and lie clearly within the uncertainty estimates of MIPAS data. The balloon-borne observations ranging from about 20.4 to 21.3 pptv are more in line with the mid-latitude values of MIPAS, as are the ground-based observations from Harestua with 21.0 pptv. The Br$^y$ value of 22.4 pptv from the ground-based observations in Lauder fits,
Figure 12. Series of averaged MIPAS BrONO$_2$ measurements (dark blue dots) and derived total stratospheric Br$_2$ (red dots) for different altitude and latitude bands over the time of stratospheric entry. Dark blue and red lines indicate the related time-averaged mean values over the whole period and the red shading indicates the estimated 1σ uncertainty. The other data points and error bars are estimates of Br$_2$ from observations of BrO taken from Fig. 1-16 in Engel et al. (2018) as well as updates of the MIPAS-Balloon observations of BrONO$_2$ (Wetzel et al., 2017). Coarse locations (south of 20° S: top; 20° S–20° N: middle; north of 20° N: bottom) of the non-MIPAS/Envisat observations are indicated by the larger symbol size. See also Table 1.

however, more to the higher tropical MIPAS estimates. Further, the Br$_2$ estimates from BrONO$_2$ measurements during two balloon flights of the MIPAS-B instrument (21.6 and 22.7 pptv, respectively) agree with both mid-latitude and tropical MIPAS/Envisat values.

5 Conclusions

We have presented the first global dataset of BrONO$_2$ volume mixing ratio profiles for day and night derived from 10°-zonally and 3-daily averaged MIPAS spectra covering the whole period of observations from 2002 to 2012. A comparison with EMAC model simulations confirms overall our current understanding of the chemical processes influencing the global zonal mean distribution as well as the diurnal and seasonal variations of BrONO$_2$ in the stratosphere. Still, remaining differences indicate uncertainties in modelled processes as well as in boundary conditions.

One deviation, the underestimation of BrONO$_2$ concentrations by the model at high latitudes during winter, could be explained. It is caused by the missing additional NO$_x$ source in the model located in the mesosphere and lower thermosphere. The energetic-particle-produced NO$_x$ is transported downwards by polar winter subsidence, thereby contributing to the production of BrONO$_2$ — a process which we could observe here for the first time. In future, modelling efforts are envisaged to study this effect on the high-latitude bromine budget as well as its impact on stratospheric ozone.

Two further inconsistencies between model and measurement are more difficult to unravel, and final explanations remain open. First, a globally present disparity is the higher simulated values in the lower stratosphere, especially at night. Sensitivity calculations with our 1D model indicate as the only possible means to decrease BrONO$_2$ concentrations a more efficient heterogeneous loss of BrONO$_2$, e.g. via Reaction (R4). However, to reach values compatible with
Table 1. Mean values, standard deviation, the standard error of the mean, and the estimated accuracy of total stratospheric Br$_y$ as derived from MIPAS, from Fig. 1-16 of Engel et al. (2018) as well as single observations by the MIPAS-Balloon experiment (Wetzel et al., 2017) during the stratospheric entry years 1997–2007; see also Fig. 12.

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Year of stratospheric entry</th>
<th>Br$_y$ derived from</th>
<th>Br$_y$ mean (pptv)</th>
<th>Br$_y$ Std (pptv)</th>
<th>Br$_y$ Std (mean) (pptv)</th>
<th>Est. accuracy (pptv)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIPAS$^a$ (40°–60° S)</td>
<td>1997–2006</td>
<td>BrONO$_2$</td>
<td>21.37</td>
<td>3.0</td>
<td>0.07</td>
<td>1.4</td>
</tr>
<tr>
<td>MIPAS$^a$ (20° S–20° N)</td>
<td>1998–2007</td>
<td>BrONO$_2$</td>
<td>22.37</td>
<td>3.62</td>
<td>0.04</td>
<td>1.7</td>
</tr>
<tr>
<td>MIPAS$^a$ (40–60° N)</td>
<td>1997–2006</td>
<td>BrONO$_2$</td>
<td>20.98</td>
<td>3.69</td>
<td>0.07</td>
<td>1.4</td>
</tr>
<tr>
<td>Balloon$^{b,c}$ (occultation)</td>
<td>1998–2000</td>
<td>BrO</td>
<td>21.25</td>
<td>0.00</td>
<td>0.00</td>
<td>2.5</td>
</tr>
<tr>
<td>Balloon$^{b,c}$ (occultation, diff. instr.)</td>
<td>Nov 2006</td>
<td>BrO</td>
<td>20.5</td>
<td></td>
<td></td>
<td>3.5</td>
</tr>
<tr>
<td>Balloon$^b$ (Langley)</td>
<td>1998–2004</td>
<td>BrO</td>
<td>20.36</td>
<td>0.65</td>
<td>0.13</td>
<td>2.5</td>
</tr>
<tr>
<td>Ground-based$^{b,d}$ (Harestua)</td>
<td>1997–2007</td>
<td>BrO</td>
<td>20.97</td>
<td>0.48</td>
<td>0.04</td>
<td>3.8</td>
</tr>
<tr>
<td>Ground-based$^{b,d}$ (Lauder)</td>
<td>1997–2001</td>
<td>BrO</td>
<td>22.35</td>
<td>0.36</td>
<td>0.07</td>
<td>4.0</td>
</tr>
<tr>
<td>MIPAS-Balloon$^e$ (Kiruna)</td>
<td>Apr 2005</td>
<td>BrONO$_2$</td>
<td>21.6</td>
<td></td>
<td></td>
<td>2.2</td>
</tr>
<tr>
<td>MIPAS-Balloon$^e$ (Timmins)</td>
<td>Sep 2009</td>
<td>BrONO$_2$</td>
<td>22.7</td>
<td></td>
<td></td>
<td>1.9</td>
</tr>
</tbody>
</table>

$^a$ This work, $^b$ WMO (2018), $^c$ Dorf et al. (2006b), $^d$ Hendrick et al. (2008), $^e$ update of Wetzel et al. (2017).
dataset of BrONO$_2$ from MIPAS can be combined e.g. with the simultaneous day-time BrO observations from the SCIAMACHY instrument on Envisat to investigate the revealed issues about possible deficiencies in our understanding of stratospheric bromine chemistry as well as to gain more insight into possible uncertainties in the observations.
Appendix A: Application of averaging kernels to EMAC model results

The EMAC model data without and with application of the respective averaging kernels from the BrONO\(_2\) retrieval are shown in Figs. A1 and A2.

**Figure A1.** Same as in Fig. 3 but showing the pure model results of BrONO\(_2\) volume mixing ratios in dark conditions on the left-hand side in comparison to the model results with the retrieval averaging kernels applied on the right-hand side.
Figure A2. Same as in Fig. A1 but for sunlit conditions.
Appendix B: NO₂

Figures B1 and B2 show the results of NO₂ volume mixing ratio profiles simultaneously retrieved with BrONO₂ in comparison with EMAC model data.

**Figure B1.** Bimonthly averaged cross sections (altitude versus latitude) of NO₂ volume mixing ratios for dark conditions. Left: measurements, middle: model with retrieval averaging kernels applied, right: model minus MIPAS.

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Figure B2. Same as Fig. B1 but for sunlit conditions.
**Figure B3.** Same as Fig. 9 but for NO$_2$.

**Figure B4.** Same as Fig. 11 but for NO$_2$.
Appendix C: BrONO₂–Brᵢ ratio

In Fig. C1 the ratio between BrONO₂ and total inorganic Brᵢ from the EMAC model data is shown.

Figure C1. Modelled ratio of BrONO₂ to Brᵢ volume mixing ratios averaged over the whole measurement period per month.
## Appendix D: Reaction parameters

**Table D1.** Major bromine reaction parameters as used in the EMAC and the 1D baseline model runs compared to the JPL2019 compilation.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>EMAC</th>
<th>JPL2019</th>
<th>1D base</th>
</tr>
</thead>
<tbody>
<tr>
<td>BrO + NO₂ → BrONO₂</td>
<td>k₃rd_iupac(k₀, n, kₘ, m, f₉)ᵇ,c</td>
<td>k₃rd(k₀, n, kₘ, m, f₉)ᵈ</td>
<td>EMAC</td>
</tr>
<tr>
<td>BrONO₂ + O(³P) → BrO + NO₃</td>
<td>–</td>
<td>1.9 × 10⁻¹¹, −21⁵ᵉ</td>
<td>–</td>
</tr>
<tr>
<td>BrONO₂ + H₂O(s, l, H₂SO₄ · nH₂O)</td>
<td>Hanson et al. (1996)</td>
<td>Hanson (2003)</td>
<td>JPL2019</td>
</tr>
<tr>
<td>BrO + NO → Br + NO₂</td>
<td>1.7 × 10⁻¹¹, 80⁰ᵇ</td>
<td>1.6 × 10⁻¹¹, 78⁰</td>
<td>EMAC</td>
</tr>
<tr>
<td>BrO + NO₂ → Br + NO₂</td>
<td>8.7 × 10⁻¹², −26⁰ᵇ</td>
<td>8.8 × 10⁻¹², −26⁰</td>
<td>EMAC</td>
</tr>
<tr>
<td>BrO + HO₂ → HOBr + O₂</td>
<td>4.5 × 10⁻¹², −50⁰ᵇ</td>
<td>4.5 × 10⁻¹², −46⁰</td>
<td>EMAC</td>
</tr>
<tr>
<td>HOBr + hv → Products</td>
<td>Ingham et al. (1998)</td>
<td>Ingham et al. (1998)</td>
<td>EMAC</td>
</tr>
</tbody>
</table>

ᵃ Burkholder et al. (2019),ᵇ Atkinson et al. (2007),ᶜ k₀(T) = k₀(300K/T)ⁿ, kₘ(T) = kₘ(300K/T)ⁿ, kₘ = kₘ(T)/kₘ(T), N = 0.75 − 1.27log₁₀(f₉).

k₃rd_iupac = \left(\frac{[M]k₀(T)}{k₃rd} \right)^{\left(1+\frac{100\log(\text{ration})}{N}\right)^{2}} + 1, ² k₃rd = \left(\frac{[M]k₀(T)}{k₃rd} \right)^{\left(1+\frac{100\log(\text{ration})}{N}\right)^{2}} + 1, ³ A, E/R, k(T) = A \exp(-E/RT).
Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/acp-21-18433-2021-supplement.

Author contributions. MH performed the MIPAS data retrieval with input from GW, GS and TVC. JO advised on spectroscopy, OK, RR, BMS, GW, SJ and MH performed and supported simulations with EMAC and the 1D model. FH and GS provided age-of-air data from MIPAS. All the authors contributed to the scientific discussion. MH prepared the manuscript with support from all the co-authors.

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

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Data availability. MIPAS level-1b data are provided by ESA (https://earth.esa.int/web/sppa/mission-performance/esa-missions/ENVISAT/mipas/products-availability/level-1/level1-8.03, last access: 14 December 2021; European Space Agency, 2021). SAGE II data were obtained from the NASA Langley Research Center Atmospheric Science Data Center https://asd.larc.nasa.gov/project/SAGE%20II/SAGE2_AEROSOL_03_NO2_H2O_BINARY_V7.0 (last access: 14 December 2021; NASA Langley Research Center, 2021). The peer-review process was guided by an independent editor, and the authors also have no other competing interests to declare. The article processing charges for this open-access publication were covered by the Karlsruhe Institute of Technology (KIT).

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