Impact of deep convection and dehydration on bromine loading in the upper troposphere and lower stratosphere

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Abstract

Stratospheric bromine loading due to very short-lived substances is investigated with a three-dimensional chemical transport model over a period of 21 years using meteorological input data from the European Centre for Medium-Range Weather Forecasts ERA-Interim reanalysis from 1989 to the end of 2009. Within this framework we analyze the impact of dehydration and deep convection on the amount of stratospheric bromine using an idealized and a detailed full chemistry approach. We model the two most important brominated short-lived substances, bromoform (CHBr$_3$) and dibromomethane (CH$_2$Br$_2$), assuming a uniform detrainment mixing ratio of 1 part per trillion by volume (pptv) for both species. The contribution of very short-lived substances to stratospheric bromine varies drastically with the applied dehydration mechanism and the associated scavenging of soluble species ranging from 3.4 pptv in the idealized setup up to 5 pptv using the full chemistry scheme. In the latter case virtually the entire amount of bromine originating from very short-lived source gases is able to reach the stratosphere thus rendering the impact of dehydration and scavenging on inorganic bromine in the tropopause insignificant. Furthermore, our long-term calculations show that the mixing ratios of very short-lived substances are strongly correlated to convective activity, i.e. intensified convection leads to higher amounts of very short-lived substances in the upper troposphere/lower stratosphere especially under extreme conditions like El Niño seasons. However, this does not apply to the inorganic brominated product gases whose concentrations are anti-correlated to convective activity mainly due to convective dilution and possible scavenging, depending on the applied approach.
1 Introduction

It is generally accepted that very short-lived substances (VSLS) contribute significantly to stratospheric bromine loading and ozone depletion in addition to the long-lived halons and methyl bromide (e.g., World Meteorological Organization, 2007, in the following WMO07). However, there are still significant uncertainties how and to what extent VSLS are transported into the upper troposphere/lower stratosphere (UTLS) and whether this will respond to possible changes under future atmospheric conditions.

Recent studies deduced the contribution of VSLS to stratospheric inorganic bromine ($Br_y$) via observations of BrO ranging from $\sim 3$ pptv (Sinnhuber et al., 2005) over $\sim 5$ pptv (McLinden et al., 2010) up to $\sim 8.4$ pptv (Sioris et al., 2006) for satellite measurements and $\sim 5.2$ pptv using a balloon-borne instrument (Dorf et al., 2008). Modeling studies to evaluate the contribution of brominated VSLS also give a wide range of estimates reaching from 0.8–3 pptv (Dvortsov et al., 1999; Nielsen and Douglass, 2001; Sinnhuber and Folkins, 2006; Kerkweg et al., 2008; Gettelman et al., 2009; Aschmann et al., 2009; Hossaini et al., 2010) to $\sim 5$ pptv (Liang et al., 2010) and 6–7 pptv (Warwick et al., 2006), although the differences in the underlying approaches and assumptions between each study make a direct comparison difficult. Commonly included VSLS are bromoform (CHBr$_3$) and dibromomethane (CH$_2$Br$_2$) which are according to the WMO07 report the most abundant very short-lived bromocarbons with an average mixing ratio in the marine boundary layer of 1.6 and 1.1 pptv, respectively. Being mainly of natural origin from maritime lifeforms, local mixing ratios can be much higher in coastal or upwelling areas (e.g., Carpenter and Liss, 2000; Quack and Wallace, 2003; Yokouchi et al., 2005; Butler et al., 2007) thus introducing a significant uncertainty in the actual source strength.

Furthermore, the transport and dehydration mechanisms in the tropical tropopause are not fully understood. Deep convection is thought to be the most important pathway into the stratosphere by enabling air parcels to overcome the level of zero clear sky radiative heating (e.g., Corti et al., 2005; Sinnhuber and Folkins, 2006; Fueglistaler et al.,
During this ascent air gets effectively dehydrated and thus soluble species are likely to be scavenged by falling water/ice particles. However, important details of the dehydration process remain unclear and are still subject to current discussions (e.g., Sherwood and Dessler, 2001; Holton and Gettelman, 2001; Fueglistaler et al., 2005, 2009). Moreover, dissolved or adsorbed halogenated species may undergo heterogeneous reactions which are able to release active radicals again into the gas phase (e.g., WMO07). Since a significant part of Br$_y$ at the base of the tropical tropopause layer (TTL) is actually HBr and therefore highly soluble (e.g., Lary, 1996; Sinnhuber et al., 2002; Yang et al., 2005, see also Fig. 4) the estimation of VSLS contribution to stratospheric bromine loading is highly sensitive to the implementation of these processes in a model. Note that due to the general absence of liquid water in the UTLS the terms “washout” and “scavenging” will be used in the following to refer to the uptake and loss of trace gases on falling ice particles rather on droplets, unless stated otherwise.

For this study we use a three-dimensional (3-D) chemistry transport model (CTM) to investigate the impact of VSLS on stratospheric bromine loading over a 21-year period. The isentropic model contains an explicit treatment of convection separated from the large-scale vertical transport which allows a realistic distribution of VSLS in the UTLS region (Aschmann et al., 2009). For the chemistry and dehydration implementation we used a twofold approach: On the one hand we have a simplified setup based on idealized tracers and processes acting as a sensitivity study. On the other hand we use a complex chemistry scheme with detailed treatment of gas phase and heterogeneous reactions, photochemistry and wet and dry deposition to assure a more realistic representation of the involved processes.
2 Model description

2.1 Three-dimensional chemical transport model

The model used in this study is an isentropic 3-D CTM with 29 levels between 330 and 2700 K (about 10 to 55 km) and a spatial resolution of 2.5° lat. × 3.75° lon. which was already utilized in our previous work (for details see Aschmann et al., 2009). It is driven by meteorological input data from ERA-Interim (EI) reanalysis of the European Centre for Medium-Range Weather Forecasts (ECMWF). A distinctive feature of our model is the separation of vertical transport into a slow large-scale heating-rate-driven component on the one hand and fast localized convective updrafts on the other hand which allows a more realistic representation of troposphere to stratosphere exchange. The convective transport is based on archived detrainment rates taken from the EI reanalysis.

For the tracer and chemistry configuration we use two approaches, an idealized setup based on simple assumptions and a complex setup including a more complete chemistry scheme which are described in detail in the following sections.

All runs were executed from 1989 to 2009 spanning almost the entire available EI dataset with a model timestep of 15 min. However, we found that in order to use as much as possible of the 21-year time period for our purpose we needed a realistic initial configuration. A recent similar study conducted by Liang et al. (2010) shows that their model (GEOS CCM Version 2) needs a spin-up time of 13 years to reach steady state for the zero-initialized brominated product gases. Our model needs significantly longer to reach equilibrium (about 20 years) in a comparable setup. The mean age of air as indicator for the circulation speed in our model agrees generally well with results from SF$_6$ observations (Stiller et al., 2008) but tends to be indeed up to two years older in higher altitudes/latitudes which may lead to the longer spin-up time. To reduce the impact of initialization artifacts on our study we therefore used the output of previous long-term runs as initial configuration.
2.2 Idealized setup

The simplified approach presented here is an expanded version of the setup described in Aschmann et al. (2009): two idealized tracers with a lifetime of 20 and 120 days (TT20, TT120) resemble the most important short-lived brominated source gases bromoform (CHBr$_3$) and dibromomethane (CH$_2$Br$_2$), respectively (WMO07). Convective upwelling is the only source for TT20/TT120 which detrain with a uniform detrainment mixing ratio ([TT20]$_c$, [TT120]$_c$) of 1 pptv each. These values are lower than the recommended mixing ratios in the marine boundary layer (1.6 and 1.1 pptv for CHBr$_3$ and CH$_2$Br$_2$, respectively, WMO07), however, as stated above the local emission strength of VSLS is highly uncertain and entrainment of tropospheric air into the convective updrafts will also decrease the detrainment mixing ratios of these species.

The brominated source gases eventually decay into inorganic bromine (Br$_y$), yielding three and two units of Br$_y$ for one unit of TT20 and TT120, respectively. Water vapor content is controlled by convection (detraining air is assumed to be fully saturated following Dessler and Sherwood, 2004), fallout of ice (if the relative humidity in a model grid box raises above 100% all surplus water is removed instantaneously) and a simple methane oxidation parametrization (details in Aschmann et al., 2009).

Inorganic bromine is represented in two variants, soluble and insoluble, both having a constant detrainment mixing ratio of 0 pptv. Whereas the insoluble Br$_y$ is inert with no additional sinks in our model the soluble version is subject to a simple washout mechanism: whenever ice forms in a model grid box (i.e. relative humidity exceeds 100%) all soluble Br$_y$ is removed completely. These two extreme cases allow the estimation of an upper and lower bound of bromine loading avoiding the details of cloud and aerosol microphysics.
2.3 Full chemistry setup

The complex chemistry scheme used here is based originally on the chemistry part of the SLIMCAT by Chipperfield (1999) and was updated and modified by Sinnhuber et al. (2003) and Winkler et al. (2008). It contains about 180 gas phase, heterogeneous and photochemical reactions for 59 tracers which are evaluated in every model timestep. Reaction rates and absorption cross sections utilized in the chemistry scheme correspond to the Jet Propulsion Laboratory (JPL) recommendations (Sander et al., 2006). Especially relevant for this study are the brominated species which are listed in Table 1. The loss of the included brominated VSLS (CHBr$_3$ and CH$_2$Br$_2$) by OH-reaction and photolysis is modeled explicitly. However, we ignore the detailed chemistry of the degradation products and assume that the VSLS directly decay into atomic bromine, which is according to the recent study of Hossaini et al. (2010) a valid approximation.

The SLIMCAT chemistry scheme incorporates an explicit treatment of uptake of halogenated species on aerosol/particle surfaces such as liquid sulfuric acid aerosols and solid nitric acid trihydrate. However, more relevant for the tropical UTLS is the uptake of halogenated species on ice particles (e.g., Lary et al., 1996) which may lead either to eventual scavenging by sedimentation or activation by heterogeneous reactions. Ice is formed in a model grid box whenever the actual water mixing ratio exceeds the saturation mixing ratio of water vapor over ice, i.e. if the relative humidity is higher than 100% all excess water is in solid phase. Then we model the uptake of gaseous HBr on ice using the partition coefficient $K_{\text{linC}}$ for HBr (4.14×10$^5$ cm) from the recent recommendation of the IUPAC Subcommittee on Gas Kinetic Data Evaluation for Atmospheric Chemistry (Crowley et al., 2010):

$$[\text{HBr}]_s = [\text{HBr}]_g^{0.88} \cdot K_{\text{linC}} \cdot A$$

(1)

Here $A$ denotes the surface area density which is calculated from the effective particle radius $r_e$ and an assumed number density $n$ of 10 particles per cubic centimeter (Chipperfield, 1999):
\[ A = 4\pi \cdot r_e^2 \cdot n \]  

The effective particle radius \( r_e \) is approximated from the total ice volume and the total number of ice particles in a particular model box. The relation of \( A \) to the fraction of adsorbed HBr is illustrated in Fig. 7. Sedimentation is calculated as well but is treated simply assuming a fixed mean particle radius of 10\( \mu \)m for ice to determine the fall velocity (about 1.5 km/day, Böhm, 1989). After one model timestep all dissolved and adsorbed species are released back into gas phase instantaneously.

Finally heterogeneous reactions on aerosol/particle surfaces are important to consider since they may release active halogen atoms back into the gaseous phase. The relevant implemented reactions for bromine are listed in Table 2.

The initial tracer configuration and detrainment mixing ratios are derived from previous 2-D model runs with a similar chemistry scheme (Sinnhuber et al., 2003). The detrainment mixing ratios of the brominated VSLS are set uniformly to 1 pptv analogous to the idealized setup. To capture the observed trend of long-lived halocarbons in our model we use time-dependent detrainment mixing ratios for these species according to the estimations in WMO07 (scenario A1). Further details of the chemistry scheme, the involved reactions and processes are given in Chipperfield (1999).

3 Results

3.1 Water vapor

A realistic distribution of water vapor content in the UTLS region is crucial for our modeling approach because uptake of soluble species and scavenging relies directly on an adequate representation of dehydration. Figure 1 shows the tape recorder signature (e.g., Mote et al., 1995) of tropical water vapor mixing ratio for the idealized and full chemistry setup and for satellite observations from HALOE (Groß and Russell, 2005).

Dry phase and speed of the tape recorder signal is generally well reproduced but both
modeling approaches show too much water (on average about 1 to 1.5 ppmv) in the UTLS region in boreal summer which is not supported by the HALOE observations. It is possible that our calculations tend to overestimate the water content of the tropical tropopause and therefore the influence of dehydration on stratospheric bromine loading. However, other measurements show indeed higher mixing ratios of water vapor in this region during boreal summer (e.g., Dessler and Sherwood, 2004; Hanisco et al., 2007; Jiang et al., 2010).

In our model, dehydration occurs mainly in the upper troposphere/tropopause region, predominantly in the tropics (Fig. 2) which is in agreement with previous studies (e.g., Gettelman et al., 2002a; Fueglistaler et al., 2005; Read et al., 2008). At high latitudes the ambient meteorological conditions allow the formation of ice even up to 25 km altitude, i.e. Type II polar stratospheric clouds (e.g., Solomon, 1999; Lowe and MacKenzie, 2008). Generally, the mixing ratios for ice are larger in the full chemistry setup since the idealized approach lacks a sedimentation mechanism, thus all ice is instantaneously removed from the corresponding grid box.

### 3.2 Stratospheric bromine loading due to VSLS

In general two processes control stratospheric bromine loading due to VSLS in both modeling approaches. The first one, convection, represented by the convective detrainment rate $d_c$, injects VSLS into the UTLS region and thus enables these species to contribute to stratospheric bromine abundance eventually. Therefore, the mixing ratio of Br$_y$ (or the corresponding inorganic bromine substances in the full chemistry setup, Table 1) is reduced by dilution since the detraining air is assumed to be free of Br$_y$, i.e. its detrainment mixing ratio is zero. The other important mechanism is the dehydration of air at the tropopause which may cause a significant removal of soluble Br$_y$ by falling ice depending on the implementation in the model.

As first step, we illustrate the impact of these processes on the stratospheric bromine loading by comparing both modeling approaches for a single year, 2006. We picked this particular year because it is in the last quarter of our covered time period minimizing the
effect of initialization artifacts. More important, 2006 is a year relatively unaffected by the El Niño Southern Oscillation (ENSO) that is typically accompanied by strong perturbations in sea surface temperature and convective activity (e.g., Ramanathan and Collins, 1991; Barsugli and Sardeshmukh, 2002; Chiang and Sobel, 2002; Guilyardi et al., 2009) that would otherwise influence our results (see Sect. 3.3).

3.2.1 Idealized setup

Figure 3 shows averaged tropical (20° N to 20° S) profiles for 2006 of brominated source and product gases illustrating typical distributions. The dashed profiles represent our earlier modeling study where we only incorporated TT20 (CHBr₃) as source gas (Aschmann et al., 2009). The suggested stratospheric bromine loading is in the range of 1.6 to 3 pptv assuming the detrainment mixing ratio of the only source, TT20, is 1 pptv. Taking into account that one unit of TT20 decays into three units of Brᵧ this means that at least 50% of all bromine originating from TT20 was unaffected by washout and is able to enter the stratosphere. Introducing TT120 (CH₂Br₂) as additional source gas with [TT120]c=1 pptv changes the previous estimation to 3.1 to 5 pptv (solid profiles). Consequently about 1.5 to 2 pptv is originating from TT120, i.e. at least 75% of the total available TT120 is not affected by washout assuming a production rate of 1:2 for TT120/Brᵧ.

Another aspect that influences our results of the idealized setup is the assumption regarding the partitioning of soluble and insoluble Brᵧ. Figure 4 shows the percentage of HBr to the total amount of Brᵧ for the full chemistry run. The fraction of soluble inorganic bromine is high only at the base of the tropical tropopause and diminishes rapidly with increasing altitude and latitude which is supported by earlier studies (e.g., Lary, 1996; Sinnhuber et al., 2002). Since we constructed the soluble Brᵧ tracer to be affected uniformly by washout this significantly overestimates the impact of washout especially near the south pole which is shown in Fig. 5: the top panel illustrates the drastic decrease of Brᵧ near the southern polar region in the standard idealized setup. The lower panel shows the result of a sensitivity calculation where the scavenging
of \( \text{Br}_y \) is restricted to within 40° N to 40° S. Switching off the scavenging process at higher latitudes increases our lower limit of the contribution of VSLS from 3.1 to 3.4 pptv (dotted profile in Fig. 3).

### 3.2.2 Full chemistry setup

The full chemistry run incorporates long-lived brominated source gases that define a bromine background in the stratosphere. Moreover, emissions of the long-lived source gases are changing over time which makes the assessment of the impact of VSLS on stratospheric bromine loading more complicated. To identify this background we conducted a reference run which does not include brominated VSLS but is identical to the full chemistry setup otherwise.

Figure 6 shows the averaged tropical profiles of brominated source and product gases in 2006 for the full chemistry runs. The mixing ratio of the long-lived source gases (\( \text{CH}_3\text{Br}, \text{CBrClF}_2, \text{CBrF}_3 \)) at the tropopause (\( \sim 14.7 \text{ pptv} \)) defines the abundance of inorganic bromine in the stratosphere (\( \sim 14.1 \text{ pptv} \) for the reference run) with a small discrepancy of 0.6 pptv. Significant loss of \( \text{Br}_y \) due to scavenging can be ruled out since the long-lived species are primarily photolyzed above altitudes where dehydration is important (e.g., Sander et al., 2006; Hossaini et al., 2010). What causes the difference is rather the variable “emission” (to be precise: the changing detrainment mixing ratio) of the long-lived brominated species. The relatively slow transport in the stratosphere compared to the troposphere introduces a time lag in the mixing ratio of \( \text{Br}_y \), i.e. some parts of the stratosphere still contain older air parcels with less \( \text{Br}_y \).

The full chemistry run with very short-lived source gases included shows a stratospheric abundance of \( \text{Br}_y \) of \( \sim 19 \text{ pptv} \), i.e. \( \sim 4.9 \text{ pptv} \) more than the reference run (Fig. 6). Considering that we added 5 pptv of additional bromine in the VSLS setup originally (detrainment mixing ratio of \( \text{CHBr}_3 \) and \( \text{CH}_2\text{Br}_2 \) is 1 pptv each) this means that almost the entire amount of bromine from short-lived substances is able to reach the stratosphere. Apparently the effect of dehydration and scavenging on stratospheric bromine loading is negligible in the full chemistry setup.
This has several reasons. First, the mixing ratio of HBr in the transient altitude range between the level of zero radiative heating and the cold point is relatively small (about 0.15 pptv, Fig. 4), the average fraction of HBr to Br$_y$ below 5%. Consequently, the majority of inorganic bromine at this altitude is actually insoluble with respect to ice in our model. Note that we have assumed here that only HBr is adsorbed on ice. We included heterogeneous reactions for HOBr and BrONO$_2$ on ice surfaces as well (Table 2), however, as it is currently not clear how and to what extent HOBr and BrONO$_2$ are adsorbed on ice (Crowley et al., 2010), we do not explicitly model the uptake as we do with HBr. The partitioning between HBr and Br$_y$ in the chemistry scheme is controlled mainly by the gas phase reactions in Table 3, the heterogeneous reactions in Table 2 and the uptake and sedimentation on ice.

Furthermore, the fraction of adsorbed HBr on ice is relatively small as well. Figure 7 shows the relation of available ice particle surface area density and the resulting fraction of adsorbed HBr to total HBr according to Eq. (1) and the relative frequency of occurrence of surface area density values in the tropical tropopause region for 2006: in 95% of all model grid boxes where ice forms the percentage of HBr on ice is significantly lower than 46% thus limiting the possible impact of scavenging directly.

In addition a major part of HBr on ice undergoes heterogeneous activation that releases reactive bromine back into gas phase. To investigate this effect we conducted another sensitivity run identical to the full chemistry setup with VSLS but without the heterogeneous reactions for HBr as listed in Table 2. Without the activation reactions the mixing ratio of HBr in the UTLS is up to four times higher than in the standard setup (Fig. 8). The resulting profile for Br$_y$ (Fig. 6) is about 0.3 pptv lower than the standard run. Consequently 4.6 out of 5 pptv of bromine originating from VSLS is able to enter the stratosphere when heterogeneous activation is ignored, 4.9 pptv otherwise.

Finally adsorbed HBr is not washed out instantaneously but sediments slowly downwards, i.e. it is possible that a significant part evaporates again and eventually reaches the stratosphere. In fact we likely overestimate the impact of sedimentation in assuming a fixed fall velocity based on an average particle radius of 10 µm.
According to Fig. 7 most ice particles are actually smaller (about 0.5 to 2 µm).

### 3.3 Development of stratospheric bromine loading over time

The balance between the processes controlling bromine loading is not static but rather varies significantly with ambient meteorological conditions. To illustrate the connection between the most important variables Figs. 9 and 11 show timeseries of tropical sea surface temperature (SST), detrainment rate $d_c$, brominated short-lived source gases, inorganic brominated product gases and total bromine abundance (i.e. the sum of source and product gases) and the corresponding monthly anomalies from 1990 to 2009 (1989 was discarded to reduce spin-up effects). The values for $d_c$ are averages over the altitude range being most relevant for bromine in the UTLS: from the level of zero clear sky radiative heating at approximately 350 K (15.5 km), which marks the transition from large-scale subsidence to large-scale upwelling (e.g., Corti et al., 2005; Sinnhuber and Folkins, 2006), to the cold point at about 380 K (17 km) where major parts of tropopause air get effectively dehydrated (e.g., Gettelman et al., 2002a; Fueglistaler et al., 2005). The cold point is also commonly regarded as a likely upper altitude limit for deep convection (e.g., Dessler, 2002). For the tracer timeseries we picked 380 K for the same reasons: almost all detrainment and also dehydration (in the tropics, Fig. 2) occurs below this level so tracers at this altitude are likely to enter the stratosphere with their current mixing ratio at 380 K.

#### 3.3.1 Idealized setup

Firstly, Fig. 9 illustrates the relation between sea surface temperature and convective activity in the ECMWF ERA-Interim reanalysis: as expected, higher SST lead in general to intensified convection denoted by higher values of $d_c$ (correlation coefficient between the two timeseries is $r=0.53$) which is in line with real world observations (e.g., Ramanathan and Collins, 1991). Secondly, our calculations show clearly a tight coupling of brominated source gas abundance in the UTLS and detrainment rate ($r=0.69$ for $d_c$/TT20 and $d_c$/TT120). Quantitatively this affects mostly the short-lived species
TT20 (CHBr$_3$), for example its mixing ratio at 380 K increases about 20% during the exceptional strong El Niño event 1997/98. For TT120 (CH$_2$Br$_2$) the magnitude of this effect is smaller (e.g., ∼6% during 1997/98) due to its relatively long lifetime.

However, for the product gas Br$_y$ the situation is more complex. The insoluble variant (not shown here) has no sink in our model but is diluted by detraining air ([Br$_y$]$_c$=0 pptv) leading to a strict anti-correlation to the detrainment rate ($r$=−0.70). Since the insoluble Br$_y$ tracer is inert and all brominated VSLS eventually end up as Br$_y$, the mixing ratio of total bromine converges to a fixed value, in the case of our setup 5 pptv. That means variations in convective activity cannot alter the amount of total bromine once it reaches its equilibrium value of 5 pptv. In contrast, the soluble Br$_y$ tracer is subject to washout as described in Sect. 2.2 which introduces an additional process that disrupts the strict anti-correlation to $d_c$ ($r$=−0.40 for soluble Br$_y$/$d_c$). Consequently total bromine abundance cannot reach equilibrium but varies with the amount of ice or scavenging efficiency. This process is due to our model setup mainly controlled by relative humidity in the UTLS region, i.e. water vapor content and ambient temperature. Actually there is weak correlation of total bromine with $d_c$ ($r$=0.44 for total Br/$d_c$) which may lead to the conclusion that the additional injection of brominated source gases during strong convection outweighs the decrease due to scavenging, however, there are also short periods where total bromine is anti-correlated to convective activity.

To illustrate the impact of an intense perturbation in convective transport Fig. 10 shows the spatial anomaly distribution of the aforementioned variables for the particular strong El Niño/La Niña seasons 1997/98 and 1999, respectively. For the El Niño season 1997/98 we observe an increase of SST in the equatorial East Pacific and to a lesser extent in the western part of the Indian Ocean. Convective activity generally raises in these areas although there is also a decrease of $d_c$ over the maritime continent. This dipole-type structure is also discernable in the distribution of the brominated source gases, TT20 and TT120, but the increase outweighs the loss considering the whole tropics. Interestingly the mixing ratios of both source gases are actually dropping over the Western Indian Ocean although $d_c$ shows a positive anomaly there.
The relative humidity as indicator for scavenging efficiency shows two zonal bands of strong positive anomalies at 20° N/20° S disrupted by a extensive negative anomaly over the Western Pacific stretching as equatorial band eastward. Mixing ratios of soluble Br\text{y} are significantly reduced in areas with positive anomalies of relative humidity exceeding the effect of Br\text{y} reduction due to dilution by detraining air. The patterns of relative humidity also appear distinctively in the distribution of total bromine: more bromine is available in the equatorial Pacific and Maritime Continent and less over the Indian Ocean. The La Niña season 1999 shows a similar picture with opposing sign. Lower SSTs lead to lower equatorial detrainment especially in the western Pacific with corresponding lows in source gas abundance. Relative humidity distribution lead to a split-up of total amount of bromine: a small increase in the Northern Hemisphere and a more distinctive loss in the south.

Both examples show the generally tight coupling between convection and the abundance of brominated source gases in the UTLS. However, the amount of bromine that actually advances further into the stratosphere is mainly controlled by the scavenging efficiency which is in turn determined by ambient temperature and water vapor content (relative humidity) in our idealized framework. Of course in reality convection and relative humidity are not independent but form a complex interaction (e.g., Sherwood and Dessler, 2001; Gettelman et al., 2002b; Jain et al., 2006; Fueglistaler et al., 2009; Tost et al., 2010) which cannot be captured in detail by our idealized approach. Our results suggest that in general the scavenging efficiency tends to be low in areas which show strong convective activity (Figs. 9 and 10) but the limitations of the utilized convective parametrization and water vapor treatment prohibit definite conclusions.

### 3.3.2 Full chemistry setup

The corresponding timeseries for the full chemistry setup are shown in Fig. 11. The picture for the brominated VSLS, bromoform and dibromomethane, is essentially the same as in the idealized setup. Both are correlated to $d_c$ ($r=0.59, 0.61$ for $d_c/\text{CHBr}_3$, $d_c/\text{CH}_2\text{Br}_2$) and bromoform is also more affected quantitatively than dibromomethane.
The pronounced difference to the idealized setup manifests in the mixing ratio of Br\textsubscript{y}/total bromine at 380 K. To remove the influence of the variable long-lived source gases we use for the timeseries of Br\textsubscript{y}/total bromine in Fig. 11 the difference between the reference run without VSLS and the standard full chemistry setup as already described in Sect. 3.2.2. As stated earlier our model suggests that virtually the entire amount of bromine originating from VSLS is able to reach the UTLS region at 380 K. Temporal variations in convective activity and washout efficiency do not have a significant effect; after about two years of spin-up time the mixing ratio of total bromine stays at 5 pptv. In return, Br\textsubscript{y} acts as the insoluble product gas tracer discussed in Sect. 3.3.1: since total bromine has reached the equilibrium value of 5 pptv Br\textsubscript{y} is strictly anti-correlated to convection (\textit{r} = \text{-0.62 for } \frac{d_c}{Br_y}) thus balancing out the changing amount of available source gases. Apparently the loss of soluble inorganic bromine species in the tropopause by uptake on ice is negligible in our full chemistry approach as stated in Sect. 3.2.2.

Results from the sensitivity calculation without heterogeneous activation presented in Fig. 12 show that uptake of soluble Br\textsubscript{y} on ice causes an average loss of total bromine of about 0.3 pptv that increases up to 0.5 pptv, for example, during the El Niño season 1997/98 which is roughly 25% to 30% of the Br\textsubscript{y} originating from VSLS at 380 K. Interesting to note is the time lag of the total bromine loss during the aforementioned El Niño season: in the idealized setup with instantaneous washout the major loss of total bromine occurs at December 1997 (Fig. 9) in contrast to the full chemistry (sensitivity) run where the minimum is reached around May 1998. Another notable difference to the idealized approach is revealed in the La Niña season 2000. The idealized run shows a small minimum in total bromine of about 0.1 pptv (Fig. 9) whereas the loss in the full chemistry run is doubled. Apart from these two major events the effect of dehydration on total bromine is generally smaller in the full chemistry sensitivity run than in the idealized setup.
The spatial anomaly patterns of the full chemistry run for the prominent El Niño/La Niña seasons 1997/98 and 1999 are presented in Fig. 13. For Br$_y$ and total bromine we show the sensitivity calculation without heterogeneous activation to emphasize the effect of ice uptake and sedimentation. For the standard run with heterogeneous activation the effect is much smaller. In addition, to remove the influence of changing long-lived source gases, for Br$_y$ and total bromine again the difference between the sensitivity run with VSLs and the reference run without VSLs is shown.

For the two VSLs and the relative humidity the spatial patterns and also the magnitude are very similar to the idealized setup (Fig. 10) during both seasons. For inorganic and total bromine there is also a certain resemblance in the patterns but the magnitude differs significantly due to the different treatment of dehydration compared to the idealized setup. In the El Niño season 1997/98 the loss of total bromine is not very distinctive during November 1997 to February 1998 as in the idealized run but occurs rather in May 1998 as shown in Fig. 11. On the other hand the amount of total bromine uniformly decreases during La Niña 1999 in contrast to the partial increase in the idealized framework.

4 Discussion and conclusions

In this study we explore the impact of deep convection and dehydration on stratospheric bromine loading due to VSLs over a 21-year period. The isentropic CTM framework we use was already evaluated with observations in a previous study (Aschmann et al., 2009). Utilizing this framework we apply two modeling approaches: a simplified setup with idealized tracers and a basic dehydration mechanism and a state of the art full chemistry scheme based on the original SLIMCAT model. In both modeling approaches we include the two most abundant brominated VSLs, bromoform and dibromomethane, with a uniform detrainment mixing ratio of 1 pptv each.

Our estimated contribution of these short-lived species to the stratospheric bromine loading is 3.4 to 5 pptv for the idealized setup and 4.9 to 5 pptv for the full chemistry
setup. These values are in agreement with observations ranging from 3 pptv (Sinnhuber et al., 2005) to 5 and 5.2 pptv (McLinden et al., 2010; Dorf et al., 2008) to 8.4 pptv (Sioris et al., 2006). Compared to other modeling studies our estimation is rather at the higher end. Considering only bromoform the estimated contributions to stratospheric bromine range from 0.8 to 1 pptv (Dvortsov et al., 1999; Nielsen and Douglass, 2001), 0.5 to 3 pptv (Sinnhuber and Folkins, 2006) to 1.6 to 3 pptv (Aschmann et al., 2009). When taking both bromoform and dibromomethane into account recent studies suggest a contribution of 2.4 to 2.8 pptv (Hossaini et al., 2010; Gettelman et al., 2009) up to 5 pptv (Liang et al., 2010). Warwick et al. (2006) suggest an even higher value of 6 to 7 pptv but considers five short-lived bromocarbons.

Aside from the uncertainties in the emission strength of short-lived bromocarbons which is not discussed here another key question is the impact of dehydration and scavenging on soluble inorganic bromine species. Depending on the representation of these processes in the model the estimated contribution of VSLS to bromine loading differ significantly. Hossaini et al. (2010) estimates a contribution of 2.4 pptv assuming a boundary layer mixing ratio of bromine atoms originally residing in CHBr₃ and CH₂Br₂ of 6 pptv and a constant washout lifetime for Brᵧ of 10 days up to 380 K (~17 km). This means about 60% of Brᵧ produced by VSLS was lost to scavenging. In contrast, Liang et al. (2010) give an estimation of 5 pptv assuming a boundary layer mixing ratio for bromine bound in VSLS of about 7 pptv utilizing a detailed wet and dry deposition scheme to calculate the loss of Brᵧ. They state that 30% of Brᵧ produced by VSLS is removed by wet scavenging and 85% of this wet removal occurs below 500 hPa (~5–6 km) in the troposphere; consequently about 5% of Brᵧ produced by VSLS is scavenged above 500 hPa. In our idealized approach the assumption that the soluble Brᵧ tracer is subject to total and instantaneous washout whenever ice is present in a model grid box leads to an estimation of 3.1 to 3.4 pptv of stratospheric bromine originating from VSLS (loss ratio 32–38%) considering a detrainment mixing ratio of bromine in VSLS of 5 pptv (Sect. 3.2.1). If we apply the detailed chemistry scheme but ignore the effect of heterogeneous activation the predicted contribution of VSLS
The modeled fraction of HBr to Br$_y$ is small in the tropical UTLS (∼5%).

2. The relatively small amount of adsorbed HBr on ice (6–8% of Br$_y$ originating from VSLS) is almost entirely released back into gas phase by heterogeneous activation.
3. Convective transport shows large inter-annual variation in relation to SST variations. In turn, the abundance of VSLS reaching the stratosphere is highly correlated to convective activity. However, the impact on total stratospheric bromine depends on the implementation of the dehydration process. Total bromine in the full chemistry scheme is nearly insensitive to dehydration whereas the idealized setup generally predicts an increased amount of total bromine in the UTLS during events of strong convective activity.

4. The fact that the detailed chemistry scheme indicates very little removal of Br$_y$ by falling ice means that almost all of the bromine from VSLS in areas of deep convection ultimately enters the stratosphere, largely independent of the details of the transport pathways. This important result should be confirmed by further refined studies of ice particle formation, sedimentation and uptake of brominated species (like, e.g., Grosvenor et al., 2007; Chaboureau et al., 2007; Read et al., 2008).

5. If removal of Br$_y$ by uptake and sedimentation is indeed not very efficient then the flux of VSLS bromine into the stratosphere will be largely controlled by the distribution of VSLS in the troposphere and the pathways by which VSLS enters deep convective uplift.

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References


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### Table 1. Brominated species in full chemistry setup.

<table>
<thead>
<tr>
<th>Category</th>
<th>Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>Source gases</td>
<td>CH$_3$Br, CBrClF$_2$, CBrF$_3$, CHBr$_3$, CH$_2$Br$_2$</td>
</tr>
<tr>
<td>Product gases (Br$_y$)</td>
<td>BrONO$_2$, BrCl, HOBr, HBr, Br, BrO</td>
</tr>
</tbody>
</table>

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Table 2. Heterogeneous activation reactions for bromine in full chemistry setup and the corresponding uptake coefficient $\gamma$ obtained from the JPL recommendations (Sander et al., 2006).

<table>
<thead>
<tr>
<th>Reactants</th>
<th>Products</th>
<th>$\gamma$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOBr+HBr</td>
<td>2Br+H$_2$O</td>
<td>0.12</td>
</tr>
<tr>
<td>HOBr+HCl</td>
<td>BrCl+H$_2$O</td>
<td>0.3</td>
</tr>
<tr>
<td>HOCl+HBr</td>
<td>BrCl+H$_2$O</td>
<td>0.06</td>
</tr>
<tr>
<td>BrONO$_2$+HBr</td>
<td>2Br+HNO$_3$</td>
<td>0.3</td>
</tr>
<tr>
<td>BrONO$_2$+H$_2$O</td>
<td>HOBr+HNO$_3$</td>
<td>0.3</td>
</tr>
<tr>
<td>ClONO$_2$+HBr</td>
<td>BrCl+HNO$_3$</td>
<td>0.3</td>
</tr>
</tbody>
</table>
**Table 3.** Gas phase reactions in the full chemistry setup that lead to the formation/destruction of HBr and the corresponding rate constant $k(T)$ obtained from the JPL recommendations (Sander et al., 2006). The reactions denoted by * are not included in the standard setup. Sensitivity calculations have shown that these reactions do not significantly affect the amount of HBr in the UTLS in our model.

<table>
<thead>
<tr>
<th>Reactants</th>
<th>Products</th>
<th>$k(T)$ [cm$^3$ molec$^{-1}$ s$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Br+CH$_2$O $\rightarrow$ HBr+HCO</td>
<td>$1.7 \times 10^{-11} \cdot \exp(-800/T)$</td>
<td></td>
</tr>
<tr>
<td>Br+HO$_2$ $\rightarrow$ HBr+O$_2$</td>
<td>$4.8 \times 10^{-12} \cdot \exp(-310/T)$</td>
<td></td>
</tr>
<tr>
<td>*Br+H$_2$O$_2$ $\rightarrow$ HBr+HO$_2$</td>
<td>$1.0 \times 10^{-11} \cdot \exp(-3000/T)$</td>
<td></td>
</tr>
<tr>
<td>*BrO+OH $\rightarrow$ HBr+O$_2$</td>
<td>$5.1 \times 10^{-13} \cdot \exp(+250/T)$</td>
<td></td>
</tr>
<tr>
<td>HBr+OH $\rightarrow$ Br+H$_2$O</td>
<td>$5.5 \times 10^{-12} \cdot \exp(+200/T)$</td>
<td></td>
</tr>
<tr>
<td>HBr+O $\rightarrow$ Br+OH</td>
<td>$5.8 \times 10^{-12} \cdot \exp(-1500/T)$</td>
<td></td>
</tr>
<tr>
<td>HBr+O$^1$D $\rightarrow$ Br+OH</td>
<td>$1.5 \times 10^{-10}$</td>
<td></td>
</tr>
</tbody>
</table>
**Fig. 1.** Comparison of tropical (20° N to 20° S) water vapor mixing ratios ("tape recorder") of idealized setup (top), full chemistry setup (middle) and HALOE observations (bottom, Grooß and Russell, 2005).
Fig. 2. Averaged equivalent gas phase volume mixing ratio of ice in idealized setup (top) and full chemistry setup (bottom) for 2006.
Fig. 3. Averaged tropical profiles (20° N to 20° S) of TT20 (CHBr₃), TT120 (CH₂Br₂) and soluble and insoluble Brₚ for 2006. The dashed profiles of soluble/insoluble Brₚ resulting from a setup where TT20 is the only brominated source gas (as in Aschmann et al., 2009) whereas the solid profiles denote the model run with both TT20 and TT120. The assumed detrainment mixing ratio for both VSLS is 1 pptv. The dotted blue profile marks soluble Brₚ from a sensitivity run where washout of Brₚ is restricted to within 40° N to 40° S.
**Fig. 4.** Average Br$_y$ partitioning for 2006 derived from the full chemistry run. Top panel: Ratio of the soluble species HBr to Br$_y$. Bottom panel: Profiles of Br$_y$ constituents in the tropical UTLS.
Fig. 5. Annual mean distribution of soluble Br\textsubscript{y} in 2006 for standard idealized run (top) and sensitivity calculation where washout of Br\textsubscript{y} is restricted to within 40° N to 40° S (bottom).
Fig. 6. Averaged tropical profiles (20° N to 20° S) of long-lived brominated source gases (CH$_3$Br, CBrClF$_2$ and CBrF$_3$), bromoform (CHBr$_3$), dibromomethane (CH$_2$Br$_2$) and inorganic brominated product gas (Br$_y$, see Table 1) for 2006. Both VSLS have an assumed uniform detrainment mixing ratio of 1 pptv. The Br$_y$ profile with filled and empty markers denote a model run with and without VSLS incorporated, respectively. The dotted Br$_y$ profile is derived from a sensitivity calculation where VSLS are included but the heterogeneous activation reactions (see Table 2) are switched off.
Fig. 7. Relation between available ice particle surface area density in a model box and the resulting fraction of adsorbed HBr on ice to total HBr derived from Eq. (1) (blue line, left ordinate). The red bars indicate the relative frequency of occurrence of surface area density values in the tropical tropopause (20° N to 20° S, 330 K to 380 K) for 2006 of the full chemistry run (right ordinate). The fraction of adsorbed HBr is lower than 46% in 95% of all calculated surface area densities.
Fig. 8. Tropical profiles of HBr from the standard full chemistry run (blue) and the sensitivity calculation without heterogeneous activation (red) for 2006.
**Fig. 9.** Timeseries of monthly mean sea surface temperature SST, detrainment rate $d_c$, idealized bromoform tracer TT20, idealized dibromomethane tracer TT120, soluble Br$_y$ and total bromine (left panels) and the corresponding anomalies (right panels) from 1990 to 2009 averaged over the tropics ($20^\circ$ N to $20^\circ$ S) at an altitude of approximately 17 km (380 K) for the tracers. The values for $d_c$ are an average of the range between the level of zero radiative heating and the cold point (350–380 K). The red lines denote the corresponding yearly average.
Fig. 10. Averaged anomalies of sea surface temperature SST, detrainment rate $d_c$, idealized bromoform tracer TT20, idealized dibromomethane tracer TT120, relative humidity RH, soluble Br$_y$ and total bromine. The small boxes denote the corresponding potential temperature level. Left: Averages over El Niño season November 1997 to February 1998. Right: Averages over La Niña season September 1999 to December 1999.
Fig. 11. Timeseries of monthly mean sea surface temperature SST, detrainment rate $d_c$, bromoform (CHBr$_3$), dibromomethane (CH$_2$Br$_2$), inorganic brominated product gases (Br$_y$, see Table 1) and total bromine (left panels) and the corresponding anomalies (right panels) from 1990 to 2009 averaged over the tropics (20° N to 20° S) at an altitude of approximately 17 km (380 K) for the tracers. The values for $d_c$ are an average of the range between the level of zero radiative heating and the cold point (350–380 K). The red lines denote the corresponding yearly average. Note: the values for Br$_y$ and total bromine are actually the difference of the standard full chemistry run and the reference run without VSLS to show the impact of the short-lived substances.
**Fig. 12.** Timeseries of Br$_y$, total bromine and the corresponding anomalies as in Fig. 11 but from the sensitivity run without heterogeneous activation.
Fig. 13. Averaged anomalies for sea surface temperature, detrainment rate, bromoform, dibromomethane, relative humidity and inorganic and total bromine for the El Niño season 1997/98 (left) and La Niña season 1999 (right) as in Fig. 10 but for the full chemistry setup. Note: the values for Br$_y$ and total bromine are actually the difference of the full chemistry run without heterogeneous activation and the reference run without VSLS to show the impact of dehydration and short-lived substances (see also Sect. 3.3.2).