Bromine partitioning in the tropical tropopause layer:
implications for stratospheric injection

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Abstract

Very short-lived (VSL) bromocarbons are produced at a prodigious rate by ocean biology and
these source compounds (SGVSL), together with their inorganic degradation products (PGVSL),
are lofted by vigorous convection to the tropical tropopause layer (TTL). Using a state-of-the-
art photochemical mechanism within a global model, we calculate annual average
stratospheric injection of total bromine due to VSL sources to be 5 pptv, with ~3 pptv entering
the stratosphere as PGVSL and ~2 pptv as SGVSL. The geographic distribution and partitioning
of VSL bromine within the TTL, and its consequent stratospheric injection, is highly
dependent on the oceanic flux, the strength of convection and the occurrence of
heterogeneous recycling reactions. Our calculations indicate atomic Br should be the
dominant inorganic species in large regions of the TTL during daytime, due to the low ozone
and cold conditions of this region. We propose the existence of a “tropical ring of atomic
bromine” located approximately between 15 and 19 km and 30ºN to 30ºS. Daytime Br/BrO
ratios of up to ~4 are predicted within this inhomogeneous ring in regions of highly
convective transport, such as the tropical Western Pacific. Then, we suggest experimental
programs designed to quantify the bromine budget of the TTL and the stratospheric injection
of VSL biogenic bromocarbons should include a strategy for the measurement of atomic Br during daytime and HOBr or BrCl during nighttime.

1 Introduction

Bromine compounds produced by biogenic processes in the oceans may influence the composition of the global atmosphere via their decomposition and the subsequent influence of the resulting bromine radicals on the tropospheric and stratospheric ozone budget, oxidation of elemental mercury and dimethyl sulphide (DMS) (Saiz-Lopez and von Glasow, 2012). Contributions of these biogenic sources, commonly known as very-short lived (VSL) bromocarbons, to inorganic bromine (Br$_r$) in the stratosphere (Ko et al., 1997; Pfeilsticker et al., 2000; Salawitch et al., 2005; Salawitch, 2006) would have important climatic effects on ozone photochemistry and trends (Dorf et al., 2006; Sinnhuber et al., 2009), and have been proposed as the missing contribution required to reconcile measurements and model simulations of stratospheric bromine chemistry (Montzka et al., 2011).

Considerable effort is presently being extended towards quantifying the oceanic production (Carpenter and Liss, 2000; Quack and Wallace, 2003; Pyle et al., 2011; Leedham et al., 2013; Liu et al., 2013) and convective transport (Ashfold et al., 2012; Fuhlbrügge et al., 2013) of biogenic VSL bromocarbons, as well as understanding the chemical transformations of these compounds to inorganic bromine species in the atmosphere (Yang et al., 2005; Salawitch, 2006; Hossaini et al., 2010; Parrella et al., 2012; Saiz-Lopez et al., 2012; Sommariva and von Glasow, 2012; Aschmann and Sinnhuber, 2013; Liang et al., 2014). Several global halocarbon emission inventories have been developed (Warwick et al., 2006; Liang et al., 2010; Ordóñez et al., 2012; Ziska et al., 2013) and compared (Hossaini et al., 2013) to address the contribution of VSL bromocarbons to the photochemistry of the troposphere and the stratospheric bromine injection. The short photochemical lifetime of many biogenic VSL bromocarbons (i.e. weeks to months, see Table 1) results in photochemical breakdown and release of reactive bromine atoms in the troposphere (Montzka et al., 2011), mainly in the tropical tropopause layer (TTL). The quantification of brominated organic source gas (SG) and inorganic product gas (PG) distributions in the TTL is of great importance for determining the impact of the VSL bromocarbons on the ozone budget of the free troposphere and lowermost stratosphere (Salawitch et al., 2005; Montzka et al., 2011; Aschmann and Sinnhuber, 2013; Liang et al., 2014).
While many global model simulations indicate that the SG injection or carbon-bonded portion (SG\text{VSL}) is the dominant contribution of VSL bromocarbons to stratospheric bromine loading (Aschmann et al., 2009; Hossaini et al., 2010, 2012; Aschmann and Sinnhuber, 2013), the combination of heterogeneous recycling processes and rapid gaseous cycling of the inorganic PG species (PG\text{VSL}, also referred hereafter as Br\text{y}) increases the lifetime of tropospheric bromine against washout by shifting the balance to the more hydrophilic inorganic portion (Parrella et al., 2012; Aschmann and Sinnhuber, 2013). Hence, the amount of inorganic bromine injected to the stratosphere is critically dependent on the balance between the efficiency of Br\text{y} removal by aerosol-, water- and ice- uptake (followed by particle sedimentation) and the heterogeneous recycling reactions that produce photolabile PG\text{VSL} species back to the gaseous phase (Sinnhuber and Folkins, 2006; Liang et al., 2010; Aschmann et al., 2011; Ordóñez et al., 2012; Aschmann and Sinnhuber, 2013). Additionally, the local convection would directly affect the timescale of the vertical transport (Aschmann et al., 2009; Hossaini et al., 2010; Ashfold et al., 2012; Liang et al., 2014), which also influences the SG\text{VSL} and PG\text{VSL} lifetimes.

Experimental campaigns are underway to measure the abundance and speciation of a host of organic bromine species and the dominant inorganic compounds within the TTL (Dorf et al., 2008; Brinckmann et al., 2012; Wisher et al., 2013; ATTREX, 2014; CAST, 2014; CONTRAST, 2014), because this region is critical for quantifying the fate of biogenic bromine decomposition products and their impact on atmospheric chemistry (Hossaini et al., 2012; Saiz-Lopez et al., 2012; Aschmann and Sinnhuber, 2013; Liang et al., 2014). Here we highlight some of the unique aspects of the TTL that drive the partitioning of inorganic bromine species by means of combined global- and box- modeling simulations. Based on our results, we discuss implications for the understanding of inorganic bromine chemistry and transport within the TTL, and assess its relevance on estimations of stratospheric bromine injection from VSL bromocarbon sources.

1.1 The tropical tropopause layer

The tropopause transition layer or tropical tropopause layer is the region between the lapse rate minimum at ~12 km (i.e., with a zero radiative heating potential temperature $\theta_{\text{ZRH}} \approx 350$K) and the cold point tropopause ($\theta_{\text{CPT}} \approx 380$ K, at ~17 km; see Gettelman and Forster, 2002). The TTL definition used here has been taken from the 2010 WMO/UNEP report (Montzka et al., 2011). Differences on the upper and lower TTL limits with respect to other
studies (Fueglistaler et al., 2009; Randel and Jensen, 2013) has no bearing on the scientific results of this study. Most gases that enter the stratosphere are first transported upwards by deep convection that detrains within the TTL (Sinnhuber and Folkins, 2006), followed by large-scale ascent.

Chemically, the TTL is characterized by low ozone and cold conditions. Very low ozone mixing ratios between 25 and 50 ppbv are routinely observed for vast regions of the TTL (Thompson et al., 2003), usually in the lower TTL. Folkins and Braun (2002) noted that air parcels in the TTL with ozone mixing ratios < 25 ppbv are “likely to be of recent boundary layer origin”, and they used low ozone as a proxy for recent, deep convection. Murphy et al. (1993) showed profiles of O$_3$ in the TTL with mixing ratios of ~26 ppbv, similar to surface values, consistent with a long photochemical lifetime for ozone and smoothing of the profile via vertical mixing. At the coldest regions of the TTL (i.e., overshooting convection), temperature can fall well below 200 K (Gettelman and Forster, 2002).

2 Model description

The 3-D chemistry climate model CAM-Chem (Community Atmospheric Model with Chemistry, version 4.0), included into the CESM framework (Community Earth System Model, version 1.1.1) had been used for this study (Lamarque et al., 2012). The development of the benchmark CAM-Chem chemical mechanism is based on MOZART-4 (Emmons et al., 2010). The current configuration includes an improved representation of stratospheric chemistry, considering heterogeneous processes for halogen species in polar stratospheric clouds from MOZART-3 (Kinnison et al., 2007; Wegner et al., 2013). CAM-Chem includes all of the physical parameterizations of CAM4 (Neale et al., 2013) and a finite volume dynamical core (Lin, 2004) for the tracer advection.

A model spin-up of 15 years considering constant boundary conditions representative of the 2000 decade have been performed to ensure stratospheric stabilization of halogen sources, and the last of a 3 years-long simulation was used to compute the bromine atmospheric burden for all sensitivity runs. A set of 5 global sensitivity simulations was performed with CAM-Chem to identify the contribution of each individual VSL species to the PG$_{VSL}$ and SG$_{VSL}$ stratospheric injection and the impact of transport within different tropical regions.
2.1 CAM-Chem setup

The setup used here considers a horizontal grid resolution of 1.9º (latitude) × 2.5º (longitude) and 26 hybrid vertical levels from the surface to approximately 40 km (Ordóñez et al., 2012; Saiz-Lopez et al., 2012). The model can be configured to perform either climatic simulations in “free running” mode (FR), where the physical state of the atmosphere is allowed to evolve independently following the hydrostatic equations, or in specified dynamics mode (SD), where offline meteorological fields are input into the model instead of calculated online. The SD procedure can allow for more precise comparisons between measurements of atmospheric composition and model output, and also allow to compare different sensitivity simulations where the chemical perturbations can be distinguished from the transport changes related to the physical state of the atmosphere (Lamarque et al., 2012).

To avoid dynamical perturbations, all sensitivities were performed in SD mode considering a high frequency meteorological input from a previous CAM-Chem FR climatic simulation. In this way, our CAM-Chem setup implies that we force the system to evolve as if it was a CTM (Chemical Transport Model), resolving the complexity of the novel tropospheric/stratospheric halogen-chemistry mechanism while using the meteorological fields from a standard chemistry-climate simulation. Then, the horizontal wind components, air temperature, surface temperature, surface pressure, sensible and latent heat flux and wind stress are read from the input meteorological dataset every 3-6 hs, while for time-steps between the reading times, all fields are linearly interpolated to avoid jumps. Even when meteorologically CAM-Chem is used as a CTM, the model is allowed to proceed with an independent inter-annual chemical evolution of all tropospheric and stratospheric constituents, and a direct comparison of the oxidative capacity of different types of atmospheres (e.g. with tropospheric bromine chemistry and without it) can be addressed (see Lamarque et al. (2012) for a complete description of this specific type of configuration). The model was configured with prescribed sea surface temperatures (SST) and ice-coverage for the 2000 decade (Rayner, 2003), so results are not representative of the meteorology of any specific year (i.e., the SD mode used here does not reproduce the specific characteristics of any of the El-Niño/La-Niña type events within the modeled period).
2.1.1 Bromine chemistry scheme

The model version used here includes a state-of-the-art organic and inorganic halogen (i.e. chlorine and bromine) photochemistry mechanism, considering both natural and anthropogenic sources, heterogeneous recycling, dry and wet deposition, both in the troposphere and stratosphere (Ordóñez et al., 2012). Rate constants have been updated to JPL-2010 (Sander et al., 2011) and IUPAC-2008 (Atkinson et al., 2007, 2008). The chemical solver for all simulations was initialized with identical chemical boundary conditions for all species, and all the atmospheric oxidants were computed online at all times (i.e., without considering prescribed monthly OH fields as done in previous studies). As major improvements with respect to previous studies are related to heterogeneous recycling over ice-particles and wet deposition processes, in Sects. 2.1.3 and 2.1.4 we present details of the implementation of these methods in CAM-Chem. For further details on bromine reactions with organic compounds, please see Table S1 in Ordóñez et al. (2012).

2.1.2 VSL bromocarbon emissions inventory

Geographically and seasonally distributed natural oceanic sources of VSL bromo- and iodo-carbons have been included and validated in the model (Ordóñez et al., 2012). The top-down emissions inventory includes seasonality and geographical emission distributions based on parameterizations of chlorophyll-α satellite maps within the tropical oceans (20ºN–20ºS) with a 2.5 coast-to-ocean emission ratio. In the extra tropics (20º–50º and 50º–90º) a seasonal latitudinal variation is also considered. The current setup includes bromocarbon sources of CHBr₃, CH₂Br₂, CH₂BrCl, CHBr₂Cl and CHBrCl₂, while iodocarbons species include CH₃I, CH₂I₂, CH₂ICl and CH₂IBr. Hereafter SG₉SL = CHBr₃, CH₂Br₂, CH₂BrCl, CHBr₂Cl, CHBrCl₂ and CH₂IBr. The anthropogenic halogen lower boundary conditions are based on prescribed surface volume mixing ratios (vmr) of long-lived species (CFCs, CH₃Cl, CH₃Br and Halons H1301, H1211, H1202, H2402) representative of year 2000. The long-lived inventory is based on Meinshausen et al. (2011), and has been recently used in IPCC AR5 hindcast and projection simulations (Marsh et al., 2013).

The global Br mass flux from all VSL bromocarbons is approximately 630 Gg Br yr⁻¹, comparable to our previous model estimates (Ordóñez et al., 2012). Table 1 presents the individual emission contribution from each VSL considered, as well as their respective photochemical lifetime and the annual tropical surface mixing ratios that the inclusion of the
emissions inventory generates. The geographical and seasonal representation of VSL sources constitutes a major improvement in CAM-Chem relative to studies performed with other models, where constant surface mixing ratios of VSL halocarbons are imposed at the lower level (Hossaini et al., 2012; Aschmann and Sinnhuber, 2013). Indeed, the current CAM-Chem setup includes constant surface mixing ratios only for long-lived species, because their long lifetimes ensure tropospheric homogenization.

2.1.3 Heterogeneous recycling

Mechanistic improvements with respect to previous works (Ordóñez et al., 2012; Saiz-Lopez et al., 2012) are mainly based on the implementation of heterogeneous reactions for HBr, HOBr and BrONO$_2$ on tropospheric ice-crystals and stratospheric aerosol surfaces. The inclusion of these reactions in CAM-Chem is based on the stratospheric implementation of heterogeneous recycling processes of Kinnison at al. (2007). Table S1 in the supplement presents the full set of chlorine and bromine heterogeneous reactions occurring on different types of surfaces [i.e., stratospheric sulfate aerosols (SULF), nitric acid tri-hydrate (NAT), ice crystals (ICE) and sea-salt (SSLT) aerosols] with their respective reactive uptake coefficients ($\gamma$). For the case of liquid sulfate aerosols in the stratosphere, a physically-based treatment of heterogeneous kinetics ($\gamma_{\text{SULF}}$) following the framework of Hanson et al. (1994) is applied. All heterogeneous reactions occurring over ice surfaces with the exception of the ones for HBr were considered in our previous model configurations, although a different latitudinal/vertical efficiency below the tropopause was applied. Reactive uptake coefficients from Crowley et al. (2010), Ammann et al. (2013) and JPL-2010 (Sander et al., 2011) were used, unless stated otherwise.

Heterogeneous recycling reactions of HOBr, BrNO$_2$ and BrONO$_2$ (and their equivalent chlorine compounds) are also considered to proceed on sea-salt aerosol surfaces ($SA_{\text{SSLT}}$), considering that the rate limiting step is the uptake of halogen species on the aerosol surface (McFiggans et al., 2000). Within this approach we assume that the initial chloride and bromide existent in the bulk of the aerosol is large enough for the heterogeneous reaction to occur until the sea-salt aerosol is removed by washout. The sea-salt aerosol scheme considers four size bins for SSLT based on Mahowald et al. (2006). The formation of sea-salt aerosol in each size bin is a function of wind speed and humidity. The surface area density ($SA_{\text{SSLT}}$) used in this work is derived from the NaCl mass in each size bin and the effective radius of that bin. Sea-salt is lost through dry deposition (including gravitational settling) and wet
removal. Overall the lifetime of sea-salt is less than 1 day (see Mahowald et al. (2006) for details).

Note that this process represents an additional source of inorganic bromine and chlorine in the troposphere, independent from the oceanic flux of VSL halocarbons described above (i.e. these are non-stoichiometric reactions releasing Cl and Br atoms from sea-salt aerosols to the gas phase). An average depletion efficiency of 0.65 and 0.35 for bromide and chloride, respectively, has been considered following Ordóñez et al. (2012) (see Table S1). The annual SSLT bromine source yields 2.9 Tg yr\(^{-1}\) (53% in the Southern Hemisphere) of which only 1.3 Tg yr\(^{-1}\) are released within the tropics. The contribution of SSLT heterogeneous reactions to the Br\(_y\) loading for different geographical and vertical regions is analyzed in Sect. 3.4.

### 2.1.4 Washout and ice-uptake

The removal of halogen species in CAM-Chem occurs via washout and scavenging in water and ice clouds (Lamarque et al., 2012; Ordóñez et al., 2012), treating each of the Br\(_y\) species independently. Both nucleation scavenging (rain-out) and impaction scavenging (below-cloud washout) are implemented in the wet-removal schemes (see Lamarque et al. (2012) for details). The Henry Law’s coefficients (k\(_H\)) for all bromine and chlorine species included in the model are shown in Table S2. Values are taken from the compilation of Henry’s Law constants of Sander (1999).

As nighttime reservoir species are more likely to be adsorbed by water/ice particles and removed from the gaseous phase (Crowley et al., 2010; Ammann et al., 2013), the modeled Br\(_y\) loading in the TTL depends on the efficiency of the wet deposition schemes (such as in-cloud washout or removal by ice-uptake) relative to the heterogeneous recycling (Aschmann et al., 2011; Aschmann and Sinnhuber, 2013). CAM-Chem has been updated to include ice-uptake removal of halogen species following an equivalent procedure to that used by Neu and Prather (2012) for HNO\(_3\). The distinguishing features of the Neu and Prather scheme are related to the partitioning between in-cloud and below-cloud scavenging, the treatment of soluble gas uptake by ice and the overlap of condensate and precipitation within a column (Lamarque et al., 2012). Several sensitivity studies showed that an extremely efficient washout of Br\(_y\) was artificially introduced in the case of considering ice-uptake for HBr, HOBr and other bromine acids. Then, the halogen removal scheme in CAM-Chem was implemented with ice-uptake turned on for all chlorine species and for BrONO\(_2\), BrNO\(_2\), BrCl
and Br$_2$, considering that the removal of this reservoir halogen species occurs at liquid, ice, and mixed-phase clouds in the troposphere (Neu and Prather, 2012). For the rest of Br$_y$ species (BrO, HOB, HBr) the removal scheme considered only washout due to liquid clouds. An effective Henry’s Law washout efficiency considering the acid dissociation constant has been used for HBr, the most abundant bromine reservoir in the lower troposphere ($K_{H}^{\text{eff}}$(HBr)=7.2×10$^{13}$ M atm$^{-1}$, see Table S2). Uptake of HBr on liquid surfaces regulates the bromine scavenging within the MBL and FT in agreement with previous studies (Yang et al., 2005; Parrella et al., 2012).

Several studies have determined that the efficiency of HBr recycling on ice particles greatly surpass the net uptake into the condensed phase, and ice-mediated HBr losses within the TTL were estimated to represent at most 2% of the actual loss rate when a full chemistry scheme is considered (Aschmann et al., 2011). Then, even when inorganic bromine removal in the lower troposphere is highly dependent on $k_{H}$(HBr), its impact decreases within the TTL. Our approach is consistent with the modelling framework of Aschmann and Sinnhuber (2013), who report that in their reference setup the organic VSL sources “contribute completely to stratospheric bromine thus ruling out dehydration (within the TTL) as an efficient loss process for bromine”. Indeed, due to the very fast photochemical time constants for the Br$_y$ system that rapidly equilibrate the steady-state abundance of all bromine species, bromine scavenging from the gaseous phase can proceed via the washout of other species besides HBr. For example, the ice-uptake for BrONO$_2$, one of the most abundant nighttime reservoirs in the absence of heterogeneous recycling, is considered to be infinitely efficient (i.e. it is instantaneously removed from the gaseous phase once it collides with an ice-particle, see Sander and Crutzen, 1996; Sander, 1999) and controls the bromine burden in the upper TTL.

### 2.2 Box-model configuration

We have used a photochemical box-model (Salawitch et al., 2005) to compute the concentration of inorganic bromine species throughout a daily cycle, subject to the constraint that production and loss of each species is balanced over a 24 hour period (diurnal photochemical steady state). The model includes a representation of 36 species constrained by specified values of p, T, O$_3$, Br$_y$, Cl$_y$, NO$_y$, H$_2$O, CH$_4$, CO, as well as particle surface area. Reaction rates and absorption cross-sections are from JPL-2010 (Sander et al., 2011). With the exception of the ozone sensitivity study (Sect. 3.3.1), all box-model simulations had been performed with an ambient O$_3$ concentration of 25 ppbv, which is the expected ozone value...
existent in the TTL within strong convective regions. The abundance of H$_2$O is set at 12.5 ppm, the saturation mixing ratio for T=200 K and p=130 hPa. Values of NO$_y$ are based on observations (Murphy et al., 1993). The abundance of CO is based on measurements in the subtropical upper troposphere (Marcy et al., 2004) and the value of CH$_4$ is set to contemporary surface levels. Surface Area for sulfate (SA$_\text{SULF}$) had been estimated from SAGE II measurements, obtained during July 2005. SA$_\text{ICE}$ was found assuming a condensation of 12.5 ppmv of H$_2$O onto 10 particles cm$^{-3}$, resulting in 7.5 µm radius particles. Clearly, SA$_\text{ICE}$ in the TTL is highly variable. Our calculations are designed to represent the fact that typically, SA$_\text{ICE} >>$ SA$_\text{SULF}$ in the TTL (see Fig. 10). Reactive uptake coefficients for the heterogeneous recycling reactions were taken from JPL-2010 (Sander et al., 2011), based on the laboratory measurements of Abbatt (1994). Unless stated otherwise, all simulations were performed for Lat=10ºN and Solar Decl=20º, and numerical values for all model inputs are as follow: p=130 hPa, T=200 K, H$_2$O=12.5 ppmv, CH$_4$=1.77 ppmv, CO=60 ppbv, Br$_y$=4 pptv, NO$_x$=400 pptv and O$_3$=25 ppbv. Model output for the box-model simulations is shown as [X]/[Br$_y$], where [X] represents the mixing ratio of specific inorganic bromine species. We have conducted simulations for Br$_y$=4 pptv, but all results shown here are extremely insensitive to the particular choice of Br$_y$, for values ranging from at least 1 to 10 pptv. Box model results are insensitive to assumptions regarding values of H$_2$O, SA$_\text{ICE}$, and CO in the TTL, provided some reasonable value is used.

3 Results and discussions

All the CAM-Chem results shown here include an ocean mask below 3 km altitude to avoid considering grids above land that could produce artificially reduced vertical profiles within the marine boundary layer (MBL). Besides the standard 24 hour averaged streaming, time dependent output for day and night has been generated considering the noon (11:30−12:30) and midnight (23:30−00:30) local time, respectively, for all latitudes and longitudes. Tropical averages have been computed between 20º N − 20º S, while the Western Pacific (WP) warm pool area is defined by the equator (0º) and the 20º N parallels, and the 120ºE and 165ºE meridians (see black rectangle in Fig. 5).

3.1 The partitioning of inorganic and organic bromine

The vertical distribution of annually averaged inorganic bromine species within the tropics (20ºN−20ºS) at noon and midnight is shown in Figs. 1A and 1B, respectively. Figure 1C...
illustrates the 24h-mean inorganic and organic contributions to the total bromine budget of the tropical atmosphere. Our model shows a total \( (\text{Br}_{\text{total}} = \text{Br}_{\text{org}} + \text{Br}_y) \) stratospheric bromine loading of \( \sim 21 \) pptv, within the range of values \( (19.5–24.5 \) pptv) inferred from measurements of stratospheric \( \text{BrO} \) reported in the last WMO ozone assessment report (Montzka et al., 2011). Below the tropopause, \( \text{Br}_{\text{org}} \) (defined as \( \text{Br}_{\text{Halons}} + \text{CH}_3\text{Br} + \text{SG}_{\text{VSL}} \)) is the dominant form of bromine because the abundant long-lived bromocarbons have not yet decomposed. Within this organic portion, \( \sim 4.3 \) pptv of \( \text{SG}_{\text{VSL}} \) reaches the lower TTL, where 2.3 pptv are transformed to active \( \text{Br}_y \) before stratospheric injection (see Fig. 2 and Table 1 for details on the individual \( \text{SG}_{\text{VSL}} \) partitioning). In strong convective regions the faster transport increase the total amount of \( \text{SG}_{\text{VSL}} \) reaching the lower TTL relative to the tropical annual average: our model results show an enhancement of 0.8 pptv or \( \sim 20\% \) for the WP region in agreement with recent measurements performed in the upper troposphere (Sala et al., 2014). In the middle and upper stratosphere (i.e., above \( \sim 25 \) km altitude), inorganic bromine dominates because of photodissociation of any organic compounds that cross the tropopause (Fig. 1C).

\( \text{HBr} \) is the dominant inorganic bromine species in the MBL, free troposphere (FT) and lower TTL both during daytime and at night (Fig. 1A, B). Then, the removal rate of \( \text{HBr} \) controls the total amount of \( \text{Br}_y \) that reaches the lower TTL and can subsequently be injected into the stratosphere (Yang et al., 2005; Sinnhuber and Folkins, 2006; Parrella et al., 2012; Aschmann and Sinnhuber, 2013). In the upper TTL, \( \text{HBr}/\text{Br}_y \sim 0.1 \) due to fast heterogeneous recycling over ice-crystals that releases reactive bromine back to the gas phase, so bromine washout is controlled by the scavenging of nighttime reservoirs. On an annual average, noontime \( \text{BrO} \) values in the FT are \( \sim 0.2 \) pptv within the tropics (Fig. 1A) and in the range \( 0.2–0.4 \) pptv for mid-latitudes (Fig. 6), with higher values of up to a few pptv calculated for certain coastal locations within the MBL, in agreement with previous studies (Yang et al., 2005; Parrella et al., 2012; Saiz-Lopez et al., 2012). The modeled tropical vertical profiles of \( \text{BrO} \) are in excellent agreement with balloon measurements performed in the tropical regions where \( \text{BrO} \) concentrations of \( (2.0\pm1.5) \) pptv for the local tropopause (17 km) and \( (3.2\pm1.6) \) pptv at 18 km were observed, with very low, or even negligible, \( \text{BrO} \) concentrations (<1ppt) for the lower, middle and upper troposphere (Dorf et al., 2008).

During nighttime, \( \text{BrCl}, \text{HOBr} \) and \( \text{BrONO}_2 \) dominate the inorganic bromine budget in the TTL and stratosphere, while \( \text{HBr} \) dominates in the FT and MBL (Fig 1B). Large volume mixing ratios of the diatomic \( \text{Br}_2 \) and \( \text{BrCl} \) species are maintained in the first few kilometres
above the oceans due to the occurrence of heterogeneous recycling reactions over sea-salt aerosols (see Table S1). As the average depletion for bromide is larger than for chloride, the calculated abundance of Br$_2$ surpasses that of BrCl in the MBL. Their abundance decrease rapidly with increasing altitude following the vertical profiles of SA$_{SSLT}$ (see Fig. 10a). The abundance of BrCl in the upper TTL increases due to the efficient heterogeneous recycling of inorganic chlorine reservoirs (i.e., HCl), which turns out to be more efficient than the Br$_2$ production via HOBr + HBr (see Sect. 3.3). The potential importance of BrCl is quite uncertain due to two factors: the contribution of VSL chlorocarbons to Cl$_y$ in the upper TTL is not well constrained by observations but likely is between 0 and 50 pptv (Montzka et al., 2011); the particle habitat is highly variable and the ClO/Cl$_y$ ratio is quite sensitive to temperature. A box-model sensitivity analysis of the recycling processes affecting the partitioning of nighttime bromine reservoirs is given in Sect. 3.3.

The tropical annual vertical profiles for all VSL bromocarbons and their overall SG$_{VSL}$ contribution are shown in Fig. 2A. The equivalent profiles computed for February within the WP region are presented in Fig. 2B. The Br$_y$ loading within the TTL depends primarily on the release of reactive bromine from CHBr$_3$ and CH$_2$Br$_2$, which are the most abundant species and represent more than 85% of SG$_{VSL}$ in the lower TTL (Montzka et al., 2011). CHBr$_3$ is removed mainly by photolysis (it has the smallest lifetime of VSL bromocarbons), releasing an annual average of 1.7 pptv Br atoms into the gaseous inorganic phase before being transported to the stratosphere (see Table 1). In comparison, only ~17% of the bromine contained in CH$_2$Br$_2$ that reaches the lower TTL is released below the cold point tropopause (~17 km). Within the WP region, where the strength of convective transport is increased, 5.1 pptv of SG$_{VSL}$ reach the lower TTL. Also, due to the faster vertical transport, the photochemical loss of CHBr$_3$ within the TTL is reduced by ~35%, releasing only ~1.1 pptv Br before stratospheric injection occurs. Comparatively, only 5% of CH$_2$Br$_2$ is decomposed before stratospheric injection.

3.2 The tropical ring of atomic bromine

Daytime levels of atomic Br increase significantly within the TTL due to the low abundance of ozone and cold conditions, which restrict formation of BrO by the Br $+$ O$_3$ reaction. As a result we simulate a daytime “tropical ring of atomic bromine” extending from 30ºN to 30ºS (Fig. 3), where Br atoms constitute a major portion of Br$_y$. Within this tropical ring, annual average Br maximizes at ~1.3 pptv near the cold point tropopause, representing up to 30% of
the total annually-averaged Br$_y$ within the tropics (see contour lines in Fig. 3).

The tropical ring of atomic Br is a photochemical phenomenon. It exists only on the illuminated portion of the earth, so it circles the tropics with the sun. The tropical ring extends approximately from 15 to 19 km, driven by vertical variations in ozone and temperature. The height thickness is delimited by temperatures colder than ~200 K, with O$_3$ mixing ratios ranging between ~25 ppbv and ~500 ppbv at the bottom and top boundaries, respectively. Below the lower limit of the ring, the ambient temperature is high enough (e.g. > 220 K) for the highly exothermic Br + O$_3$ reaction (Sander et al., 2011) to proceed efficiently. Above the ring, in the lower stratosphere, ozone abundances of a few ppmv and warmer conditions lead to the rapid formation of BrO through Br + O$_3$. It is worth recalling that due to its strong ozone and temperature dependence, the proposed tropical ring of atomic bromine should occur as an inhomogeneous ring of Br that follows the illuminated portion of the earth with variable mixing ratios at various longitudes, latitudes and altitudes.

The existence of the atomic Br ring is a direct consequence of the O$_3$/T dependences described above, and as such, presents a marked seasonality and geographical distribution (see Fig. 3C and 5A). The maximum abundances of atomic Br within the tropical ring are coincident with low ozone regions, i.e., in geographical areas of extremely strong convection such as the tropical WP warm pool (Ashfold et al., 2012; Brinckmann et al., 2012), where O$_3$-poor air masses originating from the MBL are rapidly transported to the lower TTL (Folkins and Braun, 2002). When O$_3$ falls below a critical level, loss of atomic bromine occurs primarily via reaction with the formaldehyde (HCHO). Our results indicate atomic Br levels up to 3 pptv may occur in the WP region during periods of vigorous convection (Fig. 4). Due to the fast vertical transport during convective events (hours to days), SG$_{VSL}$ have not completely decomposed, and an additional source of Br$_y$ in the lower TTL of the WP region arise from the detrainment of bromine-rich air masses entrained in the MBL and lower troposphere. Model sensitivities (see Table 4) indicate that during strong convection within the WP region the efficient sea-salt recycling occurring up to a height of ~3 km increase significantly the Br$_y$ loading in the lower troposphere (see Fig. 11), and approximately half of the Br$_y$ released in the MBL can reach the lower TTL (Br$_y^{MBL}$ ~ 7.9 pptv and Br$_y^{12\text{ km}}$ ~ 3.6 pptv). The increase in atomic bromine levels within the middle and upper troposphere directly influences the lifetime of elemental mercury against oxidation via Hg + Br (Holmes et al., 2006, 2010).
3.2.1 The Br/BrO ratio in the TTL

Even though BrO is the most abundant species throughout the tropics on an annual average (Fig. 1A), a ratio of Br/BrO > 1 will occur in key areas. The abundance of atomic Br surpasses BrO in regions of the TTL where temperatures and O$_3$ levels range from 190 K to 220 K and from 50 ppbv to 25 ppbv, respectively (see Figs. 4 and 5). As Br and BrO establish a photochemical steady-state, their ratio can be approximated by the BrO photolysis ($J_{\text{BrO}}$) and the Arrhenius type Br + O$_3$ reaction ($k_{\text{Br+O}_3}$), i.e., $[\text{Br}]/[\text{BrO}]=J_{\text{BrO}}/(k_{\text{Br+O}_3}[\text{O}_3])$. A box-modeling analysis of the sensitivity of the partitioning between Br and BrO to ozone is given in Sect. 3.3.1.

The Br/BrO ratio peaks in regions with O$_3$ < 40 ppbv and T < 190 K, while for regions with O$_3$ > 60 ppb and T > 200 K within the tropics a ratio of Br/BrO < 1 is found. The altitude where the Br/BrO ratio peaks is usually located just below the maximum level of atomic Br, as a consequence of a compromise between the decrease of temperature and the increase of O$_3$ towards the lower stratosphere.

The magnitude of the Br/BrO ratio peak strongly depends on season and geographical region, due to the changes in oceanic sources and vertical transport. Figures 5A-C show the modeled geographic distributions of Br, BrO and Br/BrO ratio at the top of the TTL (~17 km) for February. At this altitude, monthly averaged values of Br/BrO ~2.5 are found for the WP region (see Fig. 4). The modeled ozone vertical profiles in the WP present a very good agreement with ozonesondes measurements of Rex et al. (2014), who reported that ozone loss in the boundary layer combined with convectively driven vertical mixing in the WP region is the most likely explanation for the very low O$_3$ mixing ratios found in the upper stratosphere.

The WP is a critical region: during Northern Hemisphere winter, a majority of air parcels that are lofted to the $\theta_{\text{CPT}}=380$ K level by rapid convection (which marks likely future ascent to the stratosphere) detrain in the TTL (Bergman et al., 2012). For individual events of vigorous convection within the WP, atomic Br represents ~60% of daytime Br$_3$ and a maximum Br/BrO ratio of ~4 is found (see Fig. 4B).

Figure 6 shows the zonal average vertical distribution of BrO abundances between 60°N–60°S, similar to Fig. 3A for atomic Br. The characteristic latitudinal variation of the tropopause is clearly defined by the change in BrO abundances: within the tropics, the altitude of the tropopause is ~17 km or ~100 hPa while for the mid-latitudes remains between 10–12 km (~200 hPa). Within the FT, BrO abundances remain below 0.4 pptv on an annual average,
while in the stratosphere up to 70% of Br is due to BrO, with a clear stratification as we move up to the higher levels. At the top of the model (~35 km), daytime BrO reaches 16 pptv, and together with BrONO₂ and HOBr are the most abundant species in the sunlit stratosphere.

Even when the geographic distributions of Br and BrO differ (Figs. 5A and 5B), the Br/BrO ratio follows the spatial patterns of O₃ and temperature (Figs. 5D and 5E) and defines the inhomogeneous extension of the tropical ring. This confirms that the Br/BrO ratio is nearly independent of the total amount of Br (Fig. 5F), as expected due to the rapid photochemical time constants of the Br system (1, 23). Notwithstanding uncertainties in heterogeneous reactions and removal processes, all the global sensitivity simulations presented in Sect. 3.4 predict the existence of the tropical ring of atomic bromine and a pronounced Br/BrO peak within the TTL. Hence, measurements of the abundance of daytime Br, in addition to that of BrO, would be valuable for quantifying the bromine budget in the TTL and the injection of Br to the stratosphere.

3.3 Sensitivity to heterogeneous recycling: A box-modelling approach

The species HBr, HOBr and BrONO₂ are soluble and likely to be absorbed by aerosol or liquid cloud particles (Iraci et al., 2005; Abbatt et al., 2012; Saiz-Lopez and von Glasow, 2012), and the reactive uptake of HBr, HCl and HOBr has been observed to proceed over ice (Abbatt, 1994). Depending on the local microphysical environment, sedimentation of these particles and dehydrogenation could represent an efficient sink for Br, preventing the decomposition products of biogenic bromocarbons from reaching the lower stratosphere (Sinnhuber and Folkins, 2006) or global troposphere (Yang et al., 2005). However, the efficiency of aerosol/cloud washout of Br is likely altered by heterogeneous reactions that release bromine to the gas phase (Iraci et al., 2005; Salawitch, 2006; Sinnhuber and Folkins, 2006), and several recent global modeling studies have included HBr heterogeneous recycling with different efficiencies and highlighted its importance (Yang et al., 2005; Aschmann et al., 2011; Parrella et al., 2012; Aschmann and Sinnhuber, 2013). Here we go a step forward and, instead addressing the bromine removal efficiency, we focus on the important changes in the nighttime partitioning of bromine reservoirs due to the inclusion (or not) of heterogeneous recycling reactions. Table 2 presents a complete set of kinetic model sensitivities aimed at evaluating the uncertainties of such processes for atmospheric conditions representative of the cold point tropopause of the tropical WP. Results are shown for various model runs, with different assumptions for particle surface area (SA) and total inorganic chlorine (Cl₃)
affecting the heterogeneous reactions

\[
\text{HOBr} + \text{HBr} \rightarrow \text{Br}_2 + \text{H}_2\text{O}. 
\] (R1)

\[
\text{HOBr} + \text{HCl} \rightarrow \text{BrCl} + \text{H}_2\text{O}. 
\] (R2).

Once inorganic bromine is produced from the decomposition of biogenic bromine compounds, we expect the chemistry to equilibrate rapidly to a state similar to one of those shown in Fig. 7. In all cases, the nighttime reservoirs are converted almost entirely to Br and BrO during each sunlit cycle, with HBr being the only other species potentially present during daytime. The particular state will be determined by local ambient levels of O\(_3\), temperature, Cl\(_y\), as well as particle surface area.

Figure 7B shows the diurnal variation of Br\(_y\) species for Run\(_1a\), which assumes the presence of only sulfate aerosols (SA\(_\text{SULF}\)) and that Cl\(_y\)=0. The calculated abundance of BrONO\(_2\) displays a rapid rise during evening twilight, because the nighttime buildup of NO\(_2\) occurs coincident with the presence of BrO. However, BrONO\(_2\) is converted to HOBr by heterogeneous hydrolysis, with a lifetime of several hours (Lary et al., 1996). Prior to sunrise, HOBr is the dominant nighttime reservoir, while the Br\(_2\) produced by R1 constitutes about 20% of total Br\(_y\). If the recycling reaction is turned off (Run\(_0\), where \(\gamma_{\text{HOBr+HBr}}=0\)), significant levels of HBr are present at all times (Fig 7A) and the lowest values of atomic Br are found. On the other hand, in the presence of large areas of ice particles (SA\(_\text{ICE}\)), the heterogeneous reaction R1 acts as a rapid sink of HBr, resulting in nearly complete nighttime conversion of all Br\(_y\) to HOBr (Fig. 7C). R1 reaction has been observed to proceed on ice (Abbatt, 1994) and sulfate surfaces (Iraci et al., 2005) under laboratory settings, so it is likely that the conditions of Run\(_0\) are unrealistic. Hence, very low abundances of HBr are expected to be present in the TTL. Figure 7D shows calculated lifetimes for photochemical loss of HBr; the longest time constant is for Run\(_0\), with HBr removal becoming progressively faster for Run\(_1a\) and Run\(_1b\).

The nighttime partitioning of Br\(_y\) species is also sensitive to the possible presence of trace chlorine. Background chlorine levels in the FT and lower TTL are < 10 pptv due to the long photolytic lifetimes of CFCs and CH\(_3\)Cl, and the small contributions from VSL chlorocarbons (Montzka et al., 2011). Despite this, stratospheric intrusion events are frequent within the TTL, and observations of Cl\(_y\) as high as 50 pptv have been reported in the subtropical upper troposphere (Marcy et al., 2004). Additionally, Mébarki et al. (2010) have determined a
background HCl mixing ratio of ~20–30 pptv in the upper TTL not influenced by tropospheric nor stratospheric air. The presence of this much Cl\textsubscript{y} results in rapid conversion of HOBr to BrCl, due to R2. As shown in Figs. 7E and 7F, BrCl will likely be the dominant nighttime Br\textsubscript{y} reservoir whenever Cl\textsubscript{y} \gg Br\textsubscript{y} (for Run\textsubscript{2a}, Br\textsubscript{2} is the other nighttime reservoir). Abundances of Cl\textsubscript{y} large enough to drive nighttime Br\textsubscript{y} almost entirely into BrCl might be supplied either by decomposition of VSL chlorocarbons, or by irreversible mixing of a small portion of stratospheric air into the upper troposphere (Marcy et al., 2004).

3.3.1 Dependence of Br\textsubscript{y} partitioning on ozone and temperature

Figure 8 shows the box-model calculated noontime abundance of Br\textsubscript{y} species, as a function of O\textsubscript{3} for Run\textsubscript{1a}. The vertical dashed lines denote O\textsubscript{3} mixing ratios of 20 and 30 ppbv, values commonly present near convective outflow in the TTL of the WP region (Rex et al., 2014). For these levels of O\textsubscript{3}, atomic Br is the dominant inorganic species during daytime, representing at least 50% of the total inorganic Br\textsubscript{y}. For values of O\textsubscript{3}=25 ppbv, the Br/Br\textsubscript{y} and HBr/Br\textsubscript{y} ratios display a slight sensitivity to total NO\textsubscript{y} (400 pptv), but Br and BrO remain the dominant daytime species for all values of NO\textsubscript{y} reported by Murphy et al. (1993) for the TTL. In regions with higher ozone abundances (i.e. > 100 ppbv), as in the lower stratosphere, the diurnal Br-BrO steady-state is displaced to favor the BrO production.

The sensitivity of noontime Br/Br\textsubscript{y}, BrO/Br\textsubscript{y}, and HBr/Br\textsubscript{y} to temperature is shown in Fig. 9. For these calculations, O\textsubscript{3}=25 ppbv and H\textsubscript{2}O is set to its temperature dependent saturation mixing ratio, although nearly identical results are found if H\textsubscript{2}O is kept constant at 12.5 ppmv. The simulations show that for temperatures less than ~200 K, atomic Br is expected to constitute at least half of total Br\textsubscript{y}. Model results shown by the solid lines are for Run\textsubscript{1a} conditions. The temperature sensitivity is driven by the Br+O\textsubscript{3}, Br+H\textsubscript{2}CO, Br+HO\textsubscript{2}, and BrO+NO rate constants, each of which acts to increase the Br/Br\textsubscript{y} ratio and decrease the HBr/Br\textsubscript{y} ratio, with decreasing temperature. The uncertainty in the calculated abundances of Br, BrO, and HBr due to gas phase and heterogeneous processes is shown by the dashed and dotted lines in Figs. 9A-C. For gas phase processes, the uncertainty of individual kinetic parameters is based on JPL-2010 (Sander et al., 2011). The uncertainty due to heterogeneous processes considers cases Run\textsubscript{0} and Run\textsubscript{2b} (see supplementary material).
3.4 Global model sensitivities to processes and sources

Table 3 describes a set of 5 global sensitivity simulations that were performed with CAM-Chem to quantify the contribution of each source gas to the $P_{VSL}$ and $S_{VSL}$ bromine burden. Besides including the surface oceanic flux of only a unique VSL source gas or turning ON/OFF the heterogeneous reactions, the rest of the parameterizations considered in the chemical scheme are identical to those used for the baseline simulation ($cam_{Full\_Br}$). To avoid extraneous chemical perturbations related to differences in stratospheric halogen loading of bromine and chlorine, all simulations considered identical boundary conditions for long-lived halocarbon species (i.e., CFCs, CH$_3$Br and halons).

Table 4 presents values of $P_{VSL}$ and $S_{VSL}$ as well as the total bromine loading due to VSL sources ($\Sigma Br$) for the lower TTL (12 km) and upper TTL (17 km), for various regions and averaging periods. Note that for the $cam_{NoVSL}$ simulation, $S_{VSL}$ includes the contribution of all long-lived bromocarbons, while the $P_{VSL}$ portion represents the Br$_y$ decomposition products released by photolysis of CH$_3$Br below the tropopause. Table 4 also presents the R quotient for each CAM-Chem sensitivity simulation, computed as:

$$R = \frac{P_{VSL}}{S_{VSL}},$$

As CAM-Chem includes a complete treatment of halogen chemistry within the troposphere, the term $P_{VSL}$ in the numerator is the sum of: i) the decomposition products from the $S_{VSL}$ entering the lower TTL, ii) and the vertical transport of Br$_y$ reaching the lower TTL from the FT and MBL. For the $cam_{CHBr3}$, $cam_{CH2Br2}$ and $cam_{MinorVSL}$ runs, $P_{cam_{NoVSL}}$ is subtracted from the numerator to account for stratospheric injection of product gases resulting from tropospheric degradation of CH$_3$Br below the coldest point tropopause. Several global modeling studies oriented to determine the magnitude of the stratospheric bromine injection due to VSL do not consider a detailed treatment of the tropospheric halogen chemistry, and then approximate the entrainment of Br$_y$ from the lower troposphere.

CAM-Chem sensitivities indicate that on an annual average, the Br$_y$ abundance in the upper TTL can be up to one order of magnitude greater than $S_{VSL}$ if only CHBr$_3$ sources are considered ($R \sim 8$). In comparison, when only CH$_2$Br$_2$ is taken into account, $R \sim 0.2$ due to the greater photochemical lifetime of this source (see Table 4) and only 0.3 pptv of bromine is released from the $S_{VSL}$ to the inorganic portion below the tropopause. During convective events, the photo-degradation of bromocarbons within the TTL account only for a small
portion of the $Br_y$ loading: between 12 and 17 km, the $SG_{VSL}$ reduction is $\sim 0.06$, 0.1 and 1.0 pptv for $cam_{MinorVSL}$, $cam_{CH2Br2}$ and $cam_{CHBr3}$, respectively. This highlights the importance of considering the different dynamical timescales of large-scale ascent and deep convective events, as well as a detailed tropospheric chemical mechanism, when computing the partitioning between $SG_{VSL}$ and $PG_{VSL}$ of short-lived species.

The entrainment of $Br_y$ into the TTL from the lower troposphere constitutes an important portion ($\sim 1.0$ pptv when all VSL sources are considered for the tropical annual average) of the $PG_{VSL}$ component to stratospheric injection (see Table 4). Photodegradation of CHBr$_3$ represents $\sim 90\%$ of the annual $PG_{VSL}$ reaching the lower TTL even when heterogeneous recycling on sea-salt aerosol is not considered. Indeed, integrated in the tropics SSLT recycling can account, at most, for about 10\% of the $Br_y$ levels at 12 km, and only 3\% of the total $PG_{VSL}$ at 17 km. The situation changes considerably during events of vigorous convection within the WP region. There, the $Br_y$ loading at the lower TTL is significantly reduced from 3.6 pptv for $cam_{Full_{Br}}$ to 0.5 pptv for $cam_{NoSSLT}$. Then, in strong convective regions the $Br_y$ loading in the TTL is dominated by the entrainment of product gases from the FT and MBL instead of being controlled by VSL degradation (see below).

**3.4.1 Implications of heterogeneous recycling reactions on the composition of the TTL**

Figure 10 shows vertical profiles for the main variables affecting the heterogeneous rates for reactions R1 (HBr + HOBr) and R2 (HCl + HOBr) considering the baseline $cam_{Full_{Br}}$ simulation. The vertical variation of the surface area density of ice particles ($SA_{ICE}$), liquid droplets ($SA_{LIQ}$), stratospheric sulfate aerosols ($SA_{SULF}$) and sea-salt aerosols ($SA_{SSLT}$) between $20^\circ N$–$20^\circ S$ is presented in Fig. 10A. Within the TTL, $SA_{ICE}$ densities are at least one order of magnitude greater than those for $SA_{SULF}$, therefore ice-mediated heterogeneous reactions are expected to predominate within the TTL in agreement with other modeling studies (Aschmann and Sinnhuber, 2013). The $SA_{ICE}$ density for the tropical annual average (solid lines) can be up to a factor of 10 smaller than for the WP region during February (dashed lines). This indicates that in regions of strong convection the efficiency of heterogeneous reactions within the TTL can be considerably enhanced. Indeed, the rapid uplift of air-masses and the colder temperatures increase the total amount of ice-cloud droplets where heterogeneous reactions occur (Williams et al., 2009).
Our box-model sensitivities indicate that the other important factor controlling the efficiency of heterogeneous recycling of HBr and HOBr is the total amount of inorganic chlorine available as HCl. Figure 10B shows the vertical profile abundance of HCl and Cl\textsubscript{y} for the tropical annual average and the WP region during February. Chlorine levels up to 50 pptv on an annual average are modeled within the tropics in agreement with previous reports (Marcy et al., 2004; Mébarki et al., 2010), although WP abundances remain below 10 pptv within the TTL, due to the faster vertical transport of VSL-chlorocarbons that decreases the release of chlorine atoms.

In the absence of solar radiation, BrCl is the dominant bromine nocturnal reservoir at 17 km within the WP region (see Fig. 11), indicating that the current implementation of heterogeneous recycling in CAM-Chem is more sensitive to the increase in SA\textsubscript{ICE} than to the decrease in Cl\textsubscript{y} levels within the high convective regions. Note in Fig. 10C that the efficiency of R1 and R2 within the TTL is at least one order of magnitude greater for the reactions occurring over SA\textsubscript{ICE} than over SA\textsubscript{SULF}, being the latter the dominant substrate for recycling only above 20 km. As the reaction rate for HCl recycling surpass that for HBr recycling, and the HCl abundance rapidly increase towards the upper TTL and above (Mébarki et al., 2010), BrCl shows a pronounced night-time peak at ~17 km that rapidly decrease following the SA\textsubscript{ICE} profile. Note that there still are very large uncertainties on both the vertical and geographical representation of SA\textsubscript{ICE} fields in chemical climate models (Williams et al., 2009; Aschmann and Sinnhuber, 2013) and the reactive uptake efficiencies of heterogeneous processes for halogen species (Crowley et al., 2010; Ammann et al., 2013). Hence, field measurements of BrCl, HOBr and other bromine nighttime reservoirs, as well as laboratory measurements of SA\textsubscript{ICE} reactions would shed some light in the understanding of these fundamental processes.

### 3.4.2 Impact of convective transport on the bromine burden and partitioning

Figure 11 shows the distribution of the day-/night- time inorganic and organic vertical profile partitioning within the tropical WP during February, in a manner equivalent to Fig. 1, which showed equivalent model results for an annual average of the entire tropics. Several aspects of the sources and processes controlling the bromine partitioning in the TTL are evident comparing Figs. 1 and 11. First, there is a clear predominance of atomic Br as the most abundant Br\textsubscript{y} species during daytime, which corresponds to a localized enhancement of the
bromine atoms forming the inhomogeneous tropical ring, as well as a marked increase in the 
Br/BrO ratio in the WP (see Fig. 4). Second, BrCl becomes the most abundant bromine 
species during midnight, in agreement with the box-model sensitivity studies performed for 
the Run_2a and Run_2b cases (see also Fig. 10c). The role of heterogeneous recycling in the 
TTL highlights the importance of reducing the uncertainties related to the efficiency and 
temperature dependence of these reactions, and their competition relative to complete 
washout or ice-removal from the atmosphere (Sinnhuber and Folkins, 2006; Aschmann and 
Sinnhuber, 2013). Third, both the organic and inorganic bromine burden of the TTL is 
enhanced within this region. Then, within the WP region during strong convection periods, 
the ratio Br/BrO > 1 throughout the TTL, and atomic Br represents up to 60% of total Br. It 
is worth noting that for the cam_Full_Br scheme, the total bromine (i.e., SG_{VSL} + PG_{VSL}) 
loading in the upper TTL is slightly larger than in the lower TTL (see Table 4). This increase 
for the monthly average can be attributed to different local transport pathways within our 
definition of the WP region (0° < Lat < 20°N and 120°E < Lon < 165°E): i) transport from 
other latitudes and longitudes outside the defined WP; and ii) intrusion events or subsidence 
from the Br, rich stratosphere.

Our results show that in the MBL and lower troposphere of the WP region, the total Br, 
abundance is up to a factor of 3 larger relative to the abundance modeled for the entire tropics 
(Figs. 1c and 11c). This substantial change can be attributed to the larger occurrence of sea-
salt recycling reactions in coastal regions with strong convection, and their rapid vertical 
transport to the TTL: cam_NoSSLT sensitivity indicate that total PG_{VSL} in the MBL is reduced 
~7.5 pptv when sea-salt reactions are turned off, decreasing the inorganic bromine reaching 
the coldest point tropopause by ~ 2.2 pptv. This uncertainty to Br, loading in the TTL due to 
SSLT recycling is of similar magnitude to the uncertainties recently found by Liang et al. 
(2014) due to changes in convection strength (~2.6 pptv), although they modeled a reduction 
in stratospheric bromine injection for their maximum convection conditions. It is worth noting 
that Liang et al. (2014) did not considered heterogeneous reactions for HBr nor HOBr, and 
compared different sensitivities where not only the initiation and relaxation of convection was 
altered, but also the evaporation of rain and cloud formation was modified, i.e. the washout 
efficiency between sensitivities was changed. In contrast, all of the CAM-Chem sensitivities 
considered identical removal schemes and parameterizations, and we have focused on 
addressing the changes in Br, loading when we shift from the dominant large-scale ascent
(tropical annual average) to the periods and regions where strong convective events are frequent (tropical Western Pacific).

3.5 Implications for stratospheric injection and atmospheric ozone

Our simulations suggest that the proportion of inorganic Br input to the stratosphere is higher (PG_{VSL} \approx 3 \text{ pptv}) than the contribution from the carbon-bonded portion (SG_{VSL} \approx 2 \text{ pptv}), resulting in a total stratospheric bromine injection of \sim 5 \text{ pptv}. Tropospheric degradation of the long-lived bromine species CH_{3}Br makes an important contribution, 0.7 \text{ pptv}, to the total amount of inorganic bromine (PG_{VSL} \approx 3 \text{ pptv}) that is transported across the tropopause (see Table 4). CH_{3}Br is not commonly considered to be a VSL compound because its atmospheric removal lifetime of \sim 9.6 months is slightly longer than the 6 month cut off used to denote very short lived species (Montzka et al., 2011). However, we consider stratospheric injection of Br_{y} compounds produced following the tropospheric decomposition of CH_{3}Br to be part of PG_{VSL} because the ozone destruction potential of bromine in the lowermost stratosphere (LMS) depends critically on the amount of active bromine above the tropopause, regardless of origin (Salawitch et al., 2005). This highlights the need for climate models to incorporate a complete treatment of tropospheric bromine chemistry that includes both VSL and long-lived bromocarbons.

Our global simulations reveal distinctive features in the partitioning of SG_{VSL} and PG_{VSL} between air masses injected to the stratosphere within the WP region and the average tropics (see Figs. 1 and 11, and Table 4). First, VSL bromocarbons are produced at a remarkable rate by ocean biology within the WP, primarily in shallow coastal waters, and are subsequently transported by strong convection to the TTL. As expected, the faster the air masses are transported, the smaller the photolytic release of atomic bromine from VSL bromocarbons and the stronger the injection of the organic SG_{VSL} fraction (Figs. 2B and 11C). Second, the greater SA_{SSLT} existent in the WP region compared to the average Tropics (Fig. 10A) enhances the sea-salt aerosol recycling flux of Br_{2} and BrCl. The resulting Br_{y} abundances within the MBL of the western pacific show a \sim 3-fold increase for the cam_Full_Br scheme (Br_{y}^{Tropics} \approx 3.3 \text{ pptv} and Br_{y}^{WP} \approx 10.6 \text{ pptv}), mostly due to the recycling of bromine released from the shortest lived of all the VSL species considered (CH_{3}IBr). Third, the rapid transport of Br_{y}-rich air masses from the MBL to the lower TTL reduces the wet-deposition of PG_{VSL}
species, additionally increasing the Br\textsubscript{y} abundance in the WP TTL. This result in a local enhancement of the total bromine loading at $\theta_{\text{CPT}}$ of 7.7 pptv for the WP region during February (with both SG\textsubscript{VSL} and PG\textsubscript{VSL} coincidentally reaching 3.85 pptv) compared to the annual average injection of 5 pptv. Third, during the fast uplifting event that results in the Br/BrO peak shown in Fig. 4, the SG\textsubscript{VSL} (4.6 pptv) > PG\textsubscript{VSL} (3.0 pptv) suggests that the tropical ring of atomic bromine exists even when R < 1. Finally, note that in the case of the tropical annual average the Br\textsubscript{y} loading in the TTL is controlled by SG\textsubscript{VSL} decomposition (mostly CHBr\textsubscript{3}, Fig. 2A) and the sea-salt contribution represents < 3% of the PG\textsubscript{VSL} reaching the upper TTL. Then, the additional inorganic source released from sea-salt only impact on the stratospheric bromine injection within strong convective regions, while due to the longer timescales of the dominant large scale ascent, sea-salt recycling reactions do not affect the average tropical PG\textsubscript{VSL} injection on the global scale.

4 Summary and conclusions

Our estimate of the stratospheric injection of bromine compounds, naturally emitted from the oceans, suggests that the inorganic bromine injected to the stratosphere is larger (~3pptv) than that from very short-lived bromocarbons (~2pptv). Then, accounting for the contributions from all VSL bromocarbons, an overall quotient $R = \text{PG}_{\text{VSL}}/\text{SG}_{\text{VSL}} \sim 1.5$ (dominated by the CHBr\textsubscript{3} partitioning) is obtained. This result differs from previous model studies that reported that the SG\textsubscript{VSL} injection is the dominant contribution (Hossaini et al., 2012; Aschmann and Sinnhuber, 2013), or that SG\textsubscript{VSL} and Br\textsubscript{y} contributions are equivalent (Liang et al., 2010), although those reports considered either constant surface mixing ratios of VSL bromocarbons or a simplified representation of inorganic bromine chemistry and removal processes.

The current CAM-Chem scheme includes geographically and seasonally varying VSL emissions as well as a detailed representation of the multi-phase photochemistry from the ocean surface to the upper stratosphere. Then, our model captures regions where surface halocarbon fluxes are relatively large (i.e., coastal areas) and represents the rapid, deep convective updraft of these air masses to the lower TTL (i.e. WP region). The independent representation of removal processes for each species (HOBr, HBr, BrONO\textsubscript{2}, etc.) allows improved consideration of the various deposition processes (i.e., scavenging in-cloud precipitation, below-cloud washout, large-scale removal and re-evaporation, etc.) that affect the upper atmosphere inorganic bromine loading. In addition, the consideration of sea-salt
recycling reactions within the MBL and its convective transport, allow us to account for an additional geographically varying natural source of Br$_y$ to the lower TTL.

As shown above, a substantial portion of the bromine supplied by VSL bromocarbons is available in active form before reaching the LMS. Low ozone and temperature conditions make bromine atoms the dominant inorganic species during the sunlit tropical upper troposphere, generating a natural ring of atomic bromine that circles the tropics with the sun. If confirmed by observation, the tropical ring of atomic bromine would be indicative of direct supply of inorganic bromine to the LMS by biogenic VSL bromocarbons. The inorganic bromine injection to the LMS has the most important effect on stratospheric ozone during times of high sulfate aerosol loading after major volcanic eruptions (3), following the convection injection of water vapor (Anderson et al., 2012), or under proposed scenarios to artificially increase the stratospheric aerosol burden to mitigate climate change (Tilmes et al., 2012). Then, the supply of inorganic bromine from VSL would likely render the ozone layer more sensitive to future changes in ocean biogeochemistry than current predictions, since most chemistry-climate models presently neglect oceanic bromocarbons sources or detailed tropospheric bromine chemistry.

Heterogeneous reactions of HBr and HOBr are key uncertainties in quantifying the fate of inorganic products produced following the decomposition of biogenic bromocarbons. The fingerprint of heterogeneous reactions is revealed by the behavior of nighttime reservoirs and the abundance of daytime HBr and atomic Br. Therefore, speciated measurements of atomic bromine during the day and nighttime BrCl and/or HOBr in the TTL would provide insight into the nature of heterogeneous chemical processes, which are profoundly important because they potentially regulate the efficiency of Br$_y$ removal by particle sedimentation.

Our modeled contribution of VSL bromocarbons and their breakdown products to stratospheric Br$_y$ represents an increment of ~25% to the stratospheric bromine burden supplied by long-lived sources (CH$_3$Br and halons). This percentage will likely rise in the future as the atmospheric burden of long-lived sources declines due to a near cessation of anthropogenic emission. If the production and transport of VSL bromocarbons is found to be sensitive to climatically driven changes in the state of the world’s oceans (atmospheric winds, sea surface temperature, upwelling, and/or nutrient supply), then the tropical Br ring represents a potentially important new link between climate change and atmospheric ozone. All of this highlights the importance of including a detailed description of the interaction
between surface fluxes, convective transport and halogen photochemistry in chemistry-climate model simulations.

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Table 1. Global annual oceanic flux, SGVSL abundances and atmospheric lifetime of VSL bromocarbons

<table>
<thead>
<tr>
<th>Source Gas</th>
<th>Global Flux (Gg Br yr(^{-1}))</th>
<th>Lifetime(^{$}) (days)</th>
<th>Surface vmr(^{†}) (pptv of Br)</th>
<th>Lower TTL(^{†}) (CAM-Chem)</th>
<th>Lower TTL (WMO-2010)</th>
<th>Upper TTL(^{†}) (CAM-Chem)</th>
<th>Upper TTL (WMO-2010)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHBr(_3)</td>
<td>505.6</td>
<td>17</td>
<td>5.5</td>
<td>1.89</td>
<td>(0.9-3.3)</td>
<td>0.20</td>
<td>(0.03-0.87)</td>
</tr>
<tr>
<td>CH(_2)Br(_2)</td>
<td>61.9</td>
<td>130</td>
<td>2.2</td>
<td>1.82</td>
<td>(1.54-2.30)</td>
<td>1.5</td>
<td>(0.86-1.66)</td>
</tr>
<tr>
<td>CH(_2)BrCl</td>
<td>6.2</td>
<td>145</td>
<td>0.3</td>
<td>0.21</td>
<td>(0.13-0.16)</td>
<td>0.19</td>
<td>(0.1-0.12)</td>
</tr>
<tr>
<td>CHBr(_2)Cl</td>
<td>15.1</td>
<td>56</td>
<td>0.4</td>
<td>0.24</td>
<td>(0.12-0.30)</td>
<td>0.06</td>
<td>(0.02-0.22)</td>
</tr>
<tr>
<td>CHBrCl(_2)</td>
<td>11.0</td>
<td>46</td>
<td>0.2</td>
<td>0.14</td>
<td>(0.18-0.22)</td>
<td>0.05</td>
<td>(0.11-0.14)</td>
</tr>
<tr>
<td>CH(_2)IBr</td>
<td>31.6</td>
<td>0.1</td>
<td>0.01</td>
<td>~ 0</td>
<td>Not reported</td>
<td>~ 0</td>
<td>Not reported</td>
</tr>
<tr>
<td>Total</td>
<td>631.4</td>
<td>-</td>
<td>8.5</td>
<td>4.30</td>
<td>(2.86-6.5)</td>
<td>2.0</td>
<td>(1.1-3.2)</td>
</tr>
</tbody>
</table>

Adapted from Table 3 in Ordoñez et al. (2012) and Table 1-7 from WMO-2010 (Montzka et al., 2011). The vmr for each species have been multiplied by their respective atomicity in order to represent the contribution to Br\(_y\).

\(^{\$}\) Lifetimes for each species have been computed considering tropospheric loss by photolysis and reaction with OH.

\(^{†}\) Values are in pptv of Br, and have been averaged for the tropical latitudinal band (20° N–20° S)
<table>
<thead>
<tr>
<th>Simulation</th>
<th>Particle Type</th>
<th>Surface Area (SA) ($\text{cm}^2/\text{cm}^3$)</th>
<th>$\gamma_{\text{HOBr+HBr}}$ (pptv)</th>
<th>$\text{Cl}_y$ (pptv)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run_0</td>
<td>Sulfate</td>
<td>$2.9 \times 10^{-8}$</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Run_1a</td>
<td>Sulfate</td>
<td>$2.9 \times 10^{-8}$</td>
<td>0.25</td>
<td>0.0</td>
</tr>
<tr>
<td>Run_1b</td>
<td>Ice</td>
<td>$1.8 \times 10^{-5}$</td>
<td>0.1</td>
<td>0.0</td>
</tr>
<tr>
<td>Run_2a</td>
<td>Sulfate</td>
<td>$2.9 \times 10^{-8}$</td>
<td>0.25</td>
<td>50.0</td>
</tr>
<tr>
<td>Run_2b</td>
<td>Ice</td>
<td>$1.8 \times 10^{-5}$</td>
<td>0.1</td>
<td>50.0</td>
</tr>
</tbody>
</table>
Table 3. Description of the CAM-Chem setup used for different global sensitivity simulations.

<table>
<thead>
<tr>
<th>Simulation name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>cam_Full_Br</td>
<td>Considers all SG\textsubscript{VSL} and Long-Lived (LL) sources</td>
</tr>
<tr>
<td>cam_NoVSL</td>
<td>Considers only LL sources and no SG\textsubscript{VSL}</td>
</tr>
<tr>
<td>cam_CHBr3</td>
<td>Considers only CHBr\textsubscript{3} and LL sources</td>
</tr>
<tr>
<td>cam_CH2Br2</td>
<td>Considers only CH\textsubscript{2}Br\textsubscript{2} and LL sources</td>
</tr>
<tr>
<td>cam_MinorVSL</td>
<td>Considers CH\textsubscript{2}BrCl, CHBr\textsubscript{2}Cl, CHBrCl\textsubscript{2}, CH\textsubscript{2}IBr and LL sources</td>
</tr>
<tr>
<td>cam_NoSSLT</td>
<td>Same sources as cam_Full_Br. Recycling reactions over Sea-salt aerosols had been turned OFF\textsuperscript{§}</td>
</tr>
</tbody>
</table>

\textsuperscript{§} Heterogeneous reactions HSS0–HSS5 of Table S1 have not been considered.
Table 4. Partitioning of source gas (SG\textsubscript{VSL}) and product gas (PG\textsubscript{VSL}) species within the TTL for different CAM-Chem simulations.

<table>
<thead>
<tr>
<th>Region (Period)</th>
<th>Tropical Average (Annual) ( ^\dagger )</th>
<th>Western Pacific (February) ( ^\ddagger )</th>
<th>Vigorous Event (3days) ( ^\S )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity</td>
<td>Upper TTL (17 km)</td>
<td>Upper TTL (17 km)</td>
<td>Upper TTL (17 km)</td>
</tr>
<tr>
<td>cam_Full_Br</td>
<td>PG\textsubscript{VSL} 3.0, SG\textsubscript{VSL} 2.0, ( \Sigma\text{Br} ) 5.0, R 1.5</td>
<td>PG\textsubscript{VSL} 3.86, SG\textsubscript{VSL} 3.86, ( \Sigma\text{Br} ) 7.7, R 1.0</td>
<td>PG\textsubscript{VSL} 3.0, SG\textsubscript{VSL} 4.6, ( \Sigma\text{Br} ) 7.6, R 0.6</td>
</tr>
<tr>
<td>cam_NoVSL( ^\E )</td>
<td>PG\textsubscript{VSL} 0.8, SG\textsubscript{VSL} 15.5, ( \Sigma\text{Br} ) 16.4, R 0.05</td>
<td>PG\textsubscript{VSL} 0.4, SG\textsubscript{VSL} 16.3, ( \Sigma\text{Br} ) 16.8, R 0.03</td>
<td>PG\textsubscript{VSL} 0.3, SG\textsubscript{VSL} 16.3, ( \Sigma\text{Br} ) 16.6, R 0.02</td>
</tr>
<tr>
<td>cam_CHBr3</td>
<td>PG\textsubscript{VSL} 2.5, SG\textsubscript{VSL} 0.2, ( \Sigma\text{Br} ) 2.7, R 8.0</td>
<td>PG\textsubscript{VSL} 3.1, SG\textsubscript{VSL} 1.3, ( \Sigma\text{Br} ) 4.4, R 2.0</td>
<td>PG\textsubscript{VSL} 2.5, SG\textsubscript{VSL} 2.0, ( \Sigma\text{Br} ) 4.5, R 1.1</td>
</tr>
<tr>
<td>cam_CH2Br2</td>
<td>PG\textsubscript{VSL} 1.1, SG\textsubscript{VSL} 1.5, ( \Sigma\text{Br} ) 2.6, R 0.2</td>
<td>PG\textsubscript{VSL} 0.7, SG\textsubscript{VSL} 1.9, ( \Sigma\text{Br} ) 2.6, R 0.1</td>
<td>PG\textsubscript{VSL} 0.5, SG\textsubscript{VSL} 1.9, ( \Sigma\text{Br} ) 2.4, R 0.1</td>
</tr>
<tr>
<td>cam_MinorVSL</td>
<td>PG\textsubscript{VSL} 1.1, SG\textsubscript{VSL} 0.3, ( \Sigma\text{Br} ) 1.5, R 0.9</td>
<td>PG\textsubscript{VSL} 2.1, SG\textsubscript{VSL} 0.6, ( \Sigma\text{Br} ) 2.7, R 2.7</td>
<td>PG\textsubscript{VSL} 1.8, SG\textsubscript{VSL} 0.7, ( \Sigma\text{Br} ) 2.5, R 2.3</td>
</tr>
<tr>
<td>cam_NoSSLT</td>
<td>PG\textsubscript{VSL} 2.9, SG\textsubscript{VSL} 2.0, ( \Sigma\text{Br} ) 4.9, R 1.4</td>
<td>PG\textsubscript{VSL} 1.8, SG\textsubscript{VSL} 3.8, ( \Sigma\text{Br} ) 5.6, R 0.5</td>
<td>PG\textsubscript{VSL} 0.8, SG\textsubscript{VSL} 4.6, ( \Sigma\text{Br} ) 5.4, R 0.2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Region (Period)</th>
<th>Lower TTL (12 km)</th>
<th>Lower TTL (12 km)</th>
<th>Lower TTL (12 km)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity</td>
<td>Upper TTL (17 km)</td>
<td>Upper TTL (17 km)</td>
<td>Upper TTL (17 km)</td>
</tr>
<tr>
<td>cam_Full_Br</td>
<td>PG\textsubscript{VSL} 1.0, SG\textsubscript{VSL} 4.3, ( \Sigma\text{Br} ) 5.3, R 0.2</td>
<td>PG\textsubscript{VSL} 2.5, SG\textsubscript{VSL} 5.1, ( \Sigma\text{Br} ) 7.5, R 0.5</td>
<td>PG\textsubscript{VSL} 3.6, SG\textsubscript{VSL} 5.7, ( \Sigma\text{Br} ) 9.3, R 0.6</td>
</tr>
<tr>
<td>cam_NoVSL( ^\E )</td>
<td>PG\textsubscript{VSL} 0.1, SG\textsubscript{VSL} 16.3, ( \Sigma\text{Br} ) 16.4, R 0.003</td>
<td>PG\textsubscript{VSL} 0.4, SG\textsubscript{VSL} 16.4, ( \Sigma\text{Br} ) 16.8, R 0.02</td>
<td>PG\textsubscript{VSL} 0.3, SG\textsubscript{VSL} 16.5, ( \Sigma\text{Br} ) 16.8, R 0.02</td>
</tr>
<tr>
<td></td>
<td>0.9</td>
<td>1.9</td>
<td>2.8</td>
</tr>
<tr>
<td>----------------</td>
<td>-----</td>
<td>-----</td>
<td>-----</td>
</tr>
<tr>
<td>$cam_{CHBr3}$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$cam_{CH2Br2}$</td>
<td>0.1</td>
<td>1.8</td>
<td>1.9</td>
</tr>
<tr>
<td>$cam_{MinorVSL}$</td>
<td>0.2</td>
<td>0.6</td>
<td>0.8</td>
</tr>
<tr>
<td>$cam_{NoSSLT}$</td>
<td>0.9</td>
<td>4.3</td>
<td>5.1</td>
</tr>
</tbody>
</table>

All values with exception of the R quotient are reported in parts per trillion by volume (pptv).

$\sum Br = SG_{VSL} + PG_{VSL}$ is the total bromine loading from VSL sources.

$R = PG_{VSL}/SG_{VSL}$ for the $cam_{Full_{Br}}$, $cam_{NoVSL}$ and $cam_{NoSSLT}$ runs whereas $R = (PG_{VSL} - PG_{cam_{NoVSL}})/SG_{VSL}$ for the $cam_{CHBr3}$, $cam_{CH2Br2}$, and $cam_{MinorVSL}$ runs. $PG_{cam_{NoVSL}}$ is subtracted from $PG_{VSL}$ to account for stratospheric injection of product gases resulting from tropospheric degradation of CH$_3$Br, which was present in the $cam_{CHBr3}$, $cam_{CH2Br2}$, and $cam_{MinorVSL}$ simulations.

*For $cam_{NoVSL}$ sensitivity, $PG_{VSL}$ and $SG_{VSL}$ represent the inorganic and organic contributions from Long-Lived sources.

† 24-hs annual tropical average has been computed for the (20° N−20° S) latitudinal band.

‡ 24-hs monthly averages for February within the WP region outlined in black in Fig. 5.

§ Noontime 3 days average for a vigorous event within the WP region.
Figure 1. Annual vertical profile abundances of organic and inorganic bromine species within the tropics (20°N–20°S): (A) Main Br$_y$ species at noon; (B) Main Br$_y$ species at midnight; and (C) 24 hour average contribution of organic and inorganic species to total bromine. The horizontal dotted lines represent the approximate location of the TTL.
Figure 2. Vertical profile abundances of the five VSL bromocarbons considered in CAM-Chem for the tropical annual average (panel A) and the Western Pacific region (panel B). All halocarbon abundances are multiplied by their respective atomicity in order to represent their contribution to Br\textsubscript{y} after photolysis or reaction with OH. The total organic SG\textsubscript{VSL} contribution is also shown by the filled diamonds.
Figure 3. The “tropical ring of atomic bromine”: (A) annual zonal average; (B) annual meridional average abundance of atomic Br within the tropics (20°N–20°S); and (C) seasonal evolution of the zonally-averaged atomic Br ring within the tropics. The color scale represents noontime volume mixing ratios (pptv) while black contour lines show the percentage contribution of atomic Br to Br$_y$. 
Figure 4. Vertical noontime profiles of Br and BrO vmr (panel A), the Br/BrO ratio (panel B), and T and O₃ (panel C) in the tropics. Vertical profiles have been averaged over different regions and time periods: (solid) annual tropical averages (20ºN−20ºS); (dashed) WP region during February; (dotted) at the midpoint of a strong convective cell within the WP during a 3-day period in February.
Figure 5. Average noontime geographical distribution of atmospheric bromine, ozone and temperature at 17 km during February: (A) atomic Br; (B) BrO; (C) Br/BrO ratio; (D) ozone; (E) temperature; and (F) Br$_y$. The location of the WP region, also considered to compute the vertical profiles of Figs. 4 and 11 is outlined in black.
Figure 6. Annual zonal average BrO distributions between 60°N–60°S. The color scale represents absolute values (pptv), while black contour lines show the percentage contribution of BrO to Br$_2$. 
Figure 7. Calculated abundance of inorganic bromine species as a function of local solar time for different sensitivity simulations: (A) Run_0 (SA_SULF; $\gamma_{\text{HOBr+HBr}}=0$; Cl$_y=0$); (B) Run_1a (SA_SULF; $\gamma_{\text{HOBr+HBr}}=0.25$; Cl$_y=0$); (C) Run_1b (SA_ICE; $\gamma_{\text{HOBr+HBr}}=0.1$; Cl$_y=0$); (D) Photochemical lifetime of HBr for all 5 sensitivity simulations; (E) Run_2a (SA_SULF; $\gamma_{\text{HOBr+HBr}}=0.25$; Cl$_y=50$ pptv); (F) Run_2b (SA_ICE; $\gamma_{\text{HOBr+HBr}}=0.1$; Cl$_y=50$ pptv).
Figure 8. Box-model partitioning of Br$_y$ species as a function of ozone mixing ratio. The abundances (normalized to total Br$_y$) are from diurnal photochemical steady state conditions at local noon. Inputs for aerosol surface area, inorganic chlorine, and $\gamma_{\text{HOBr+HBr}}$ are for Run_1a.
Figure 9. Temperature dependence of the calculated abundance of atomic Br, BrO and HBr at local noon. The baseline simulation (solid line) assumes conditions for SA, Cl, and $\gamma_{\text{HOBr+HBr}}$ from Run_1a: (A) Atomic Br uncertainties considering only gas phase processes (dashed lines) and combined uncertainty considering both heterogeneous and gas phase processes (dotted lines). Uncertainties for BrO and HBr are shown in panels (B) and (C), respectively.
Figure 10. Vertical variation of CAM-Chem atmospheric constituents affecting the rate of heterogeneous reactions within the tropics (solid lines) and WP region (dotted lines): (A) Surface area for ice particles (SA_{ICE}), liquid droplets (SA_{LIQ}), stratospheric sulphate aerosols (SA_{SULF}) and sea-salt aerosols (SA_{SSLT}); (B) Average vertical profiles of HCl, HOCl, ClONO$_2$ and total inorganic chlorine (Cl$_y$); (C) heterogeneous reaction rates over different types of surfaces.
Figure 11. Vertical profile abundances of organic and inorganic bromine species within the WP region during February: (A) Main Br\textsubscript{y} species at noon; (B) Main Br\textsubscript{y} species at midnight; and (C) 24 hour average contribution of organic and inorganic species to total bromine.