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Partitioning and budget of inorganic and organic chlorine species observed by MIPAS-B and TELIS in the Arctic in March 2011

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Abstract. The Arctic winter 2010/2011 was characterized by a persistent vortex with extremely low temperatures in the lower stratosphere above northern Scandinavia leading to a strong activation of chlorine compounds (ClO_x) like Cl, Cl₂, ClO, ClOOCl, OClO, and HOCl, which rapidly destroyed ozone when sunlight returned after winter solstice. The MIPAS-B (Michelson Interferometer for Passive Atmospheric Sounding) and TELIS (TErahertz and submillimeter LImb Sounder) balloon measurements obtained in northern Sweden on 31 March 2011 inside the polar vortex have provided vertical profiles of inorganic and organic chlorine species as well as diurnal variations of ClO around sunrise over the whole altitude range in which chlorine has been undergoing activation and deactivation. This flight was performed at the end of the winter during the last phase of ClO_x deactivation. The complete inorganic and organic chlorine partitioning and budget for 31 March 2011, assumed to be representative for the late-winter Arctic stratosphere, has been derived by combining MIPAS-B and TELIS simultaneously observed molecules. A total chlorine amount of 3.41 ± 0.30 parts per billion by volume is inferred from the measurements (above 24 km). This value is in line with previous stratospheric observations carried out outside the tropics confirming the slightly decreasing chlorine amount in the stratosphere. Observations are compared and discussed with the output of a multi-year simulation performed with the chemistry climate model EMAC (ECHAM5/MESSy Atmospheric Chemistry). The simulated stratospheric total chlorine amount is in accordance with the MIPAS-B/TELIS observations, taking into account the fact that some chlorine source gases and very short-lived species are not included in the model.

1 Introduction

The discovery of the Antarctic stratospheric "ozone hole" in the 1970s (Farman et al., 1985) strongly intensified research to unravel the reason for this ozone depletion. Chemically active chlorine (ClO_x) species like Cl, Cl₂, ClO, ClOOCl, OCIO, and HOCI are part of total inorganic chlorine Cl_{v} $(ClO_x + HCl + ClONO_2)$. They play a dominant role in the catalytic destruction of stratospheric ozone during polar winter when low temperatures and heterogeneous chemical reactions on polar stratospheric cloud (PSC) particles have previously enabled active chlorine compounds (mainly Cl₂) to be produced from its reservoir species ClONO₂, HCl, and HOCl (e.g. Molina and Rowland, 1974; Solomon et al., 1986; Molina et al., 1987; Solomon, 1999; Crutzen and Oppenheimer, 2008). Due to the Montreal Protocol and successive agreements, emissions of dominant halocarbons were reduced such that total tropospheric (organic) chlorine has been decreasing since 1994 after reaching a peak value of nearly 3.7 parts per billion by volume (ppbv) (O'Doherty et al., 2004; WMO, 2011). The stratospheric total chlorine peak occurred several years later, because it took this time for emitted air masses to propagate into the stratosphere (Engel et al., 2002; WMO, 2011; Kohlhepp et al., 2012). The amount of equivalent effective stratospheric chlorine (chlorine and bromine halogens) is predicted to return to 1980 values around 2050 at mid-latitudes (Stolarski et al., 2010; WMO, 2011).

To assess and monitor the partitioning and budget of chlorine, a number of measurements of its individual compounds have been carried out to calculate the amount of inorganic (Cl_{v}) , organic (CCl_{v}) , and finally total chlorine (Cl_{total}) . An early observation based on data from the Atmospheric Trace Molecule Spectroscopy (ATMOS) instrument was published by Zander et al. (1992). A mean stratospheric total chlorine volume mixing ratio (VMR) of 2.58 ± 0.10 ppbv was observed at 30° N in 1985. Significantly enhanced values between 3.4 and 3.5 (± 0.4) ppbv in the 1992 Arctic lower stratosphere were estimated using retrieved data from the balloon-borne Michelson Interferometer for Passive Atmospheric Sounding (MIPAS-B) in combination with in situ measurements (von Clarmann et al., 1995). A further slightly enhanced value of 3.53 ± 0.10 ppbv was detected during the ATMOS/ATLAS-3 November 1994 mission at northern midlatitudes, also demonstrating the strong increase of stratospheric total chlorine before regulating measures could alter this linear trend of 0.10 ppbv per year in the time period from northern spring 1985 to autumn 1994 (Zander et al., 1996). The same trend has been deduced between 1991 and 1995 by estimating total chlorine with the help of HCl observations from the HALogen Occultation Experiment (HALOE; Russell III et al., 1996). A further increased Cl_{total} value of 3.7 ± 0.2 ppbv was derived from MkIV balloon measurements carried out in the 1997 Arctic summer (Sen et al., 1999). This measurement took place close to the turnover of the total stratospheric chlorine amount. Chlorine data obtained by the Atmospheric Chemistry Experiment Fourier Transform Spectrometer (ACE-FTS) in combination with in situ measurements from the Stratospheric Aerosol and Gas Experiment (SAGE) III Ozone Loss and Validation Experiment (SOLVE) campaign (Schauffler et al., 2003; Nassar et al., 2006) were used to estimate Cl_{total} between February 2004 and January 2005 in five latitude zones. A mean stratospheric Cl_{total} value of 3.65 ± 0.13 ppbv was determined for both the northern and southern midlatitudes. This start of a temporal decrease of stratospheric chlorine was confirmed by observations from the Microwave Limb Sounder (MLS) from August 2004 until January 2006 (Froidevaux et al., 2006). A Cl_{total} value of 3.60 ± 0.13 ppbv at the end of this time period was inferred from HCl measurements and a decrease of about 43 pptv in the stratospheric chlorine loading within this 18-month period was detected.

The long term trend of stratospheric inorganic chlorine was investigated by using data of multiple space-borne sensors like ACE-FTS, ATMOS, MLS, CLAES (Cryogenic Limb Array Etalon Spectrometer), CRISTA (CRyogenic Infrared Spectrometer and Telescope for the Atmosphere), and HALOE (Lary et al., 2007). This time series confirms that stratospheric Cl_y peaked in the late 1990s and started to decrease as expected due to the changing concentrations of tropospheric source gases and related transport times from the troposphere to the stratosphere. More recently published observations of Cl_{total} were performed by ACE-FTS covering the years 2004 until 2009. Nine chlorine containing species have been directly measured by the satellite instrument (Brown et al., 2011, 2013). These data were supplemented by a number of further trace gases calculated using the SLIMCAT 3-dimensional chemical transport model (Chipperfield, 2006). Global mean stratospheric chlorine was found to decrease by 0.46 % per year in the time period under investigation.

The purpose of this paper is to assess the partitioning and budget of inorganic and organic stratospheric chlorine inside the late-winter Arctic vortex. Winter 2010/2011 was characterized by a cold vortex defining a strong transport barrier until approximately mid-April (Manney et al., 2011; Sinnhuber et al., 2011). Temperatures were below the threshold associated for chlorine activation ($\sim 196 \,\mathrm{K}$) for more than 100 days between about 15 and 23 km. Consequently, an unprecedented Arctic ozone loss was observed which could be described for the first time as an Arctic ozone hole since ozone profiles in late March resembled typical Antarctic latewinter profiles (Manney et al., 2011; Sinnhuber et al., 2011). Trace gas profiles of individual chlorine compounds were retrieved from limb-emission spectra recorded during a balloon flight of MIPAS-B and the TErahertz and submillimeter LImb Sounder (TELIS) on 31 March 2011 inside the polar vortex. A description of the instruments, data analysis, and chemical modelling is given in Sect. 2. A discussion of the observed chlorine partitioning and budget follows in Sect. 3 and in the conclusions, together with a comparison of the combined measured data to simulations of the chemistry climate model EMAC (ECHAM5/MESSy Atmospheric Chemistry model).

2 Instruments, data analysis, and modelling

The MIPAS-B/TELIS flight took place on 31 March 2011 over northern Scandinavia inside the Arctic vortex at the end of the chlorine deactivation period that started slowly in early March and accelerated towards the end of this month (Manney et al., 2011; Sinnhuber et al., 2011). The balloon gondola was launched from Esrange near Kiruna (Sweden, 67.9° N, 21.1° E) and reached its float level at about 35 km. Recorded limb sequences of MIPAS-B and TELIS are depicted in Fig. 1.

2.1 MIPAS-B instrument and data analysis

The balloon-borne limb-emission sounder MIPAS-B is a cryogenic Fourier transform spectrometer which operates in

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Figure 1. Potential vorticity (PV) field (in 10^{-6} K m² s⁻¹ kg⁻¹) from European Centre for Medium-Range Weather Forecasts (ECMWF) analysis on 31 March 2011, 00:00 UTC. MIPAS-B tangent points are plotted as black solid circles and TELIS tangent points as yellow solid circles (altitude range: 9.1 to 35.4 km). Both instruments look in the same direction. The vortex boundary which represents the strongest PV gradient (Nash et al., 1996) is shown as black dashed line. The insert (top left) shows the approximate measurement region (yellow marker) in relation to the position of the whole polar vortex.

the mid-infrared spectral range between about 4 and 14 µm. The maximum optical path difference of 14.5 cm of the beam in the interferometer allows a high unapodized spectral resolution of 0.0345 cm^{-1} (about 0.07 cm^{-1} after apodization with the Norton and Beer (1976) "strong" function), which allows the separation of individual spectral lines from continuum-like emissions in combination with a high radiometric accuracy of typically 1 %. Values of the noise equivalent spectral radiance (NESR) are typically within 1×10^{-9} and 7×10^{-9} W(cm² sr cm⁻¹)⁻¹ for a single calibrated spectrum. Averaging over *n* spectra ($n \le 5$) per single elevation scan reduces the spectral noise by a factor of $1/\sqrt{n}$. The instrument is characterized by a high performance and flexibility of the pointing system with a knowledge of the tangent altitude of better than 50 m at the 1σ confidence limit. A comprehensive overview and description of the instrument together with processing of recorded interferograms to calibrated spectra including phase correction, Fourier Transformation to the spectral domain, and two-point calibration of the spectra from arbitrary to radiance units is given by Friedl-Vallon et al. (2004) and references therein. This includes instrument characterization in terms of the instrumental line shape, field of view, NESR, line of sight of the instrument, detector non-linearity (Kleinert, 2006), and the error budget of the calibrated spectra.

Forward radiance calculations were performed with the Karlsruhe Optimized and Precise Radiative transfer Algo-



Figure 2. Temporal evolution of chlorine monoxide (ClO) mixing ratios (ppbv) as observed by MIPAS-B (top) and TELIS (bottom) on 31 March 2011 above northern Scandinavia between 02:00 and 04:38 UTC inside the Arctic vortex within the latitude/longitude sector shown in Fig. 1. The black solid line marks the sunrise terminator. A residual activation of chlorine is visible between 16 and 22 km with slightly enhanced ClO values up to 0.4 ppbv.

rithm (KOPRA; Stiller et al., 2002), which is a line-by-line and layer-by-layer model to simulate the infrared radiative transfer through the atmosphere. Molecular spectroscopic parameters for the calculation of limb-emission spectra were taken from the high-resolution transmission molecular absorption database (HITRAN; Rothman et al., 2009) and a MIPAS dedicated spectroscopic database (Raspollini et al., 2013). KOPRA also calculates derivatives of the radiance spectrum with respect to atmospheric state and instrument parameters and thus provides the Jacobians for the retrieval procedure KOPRAFIT (Höpfner et al., 2002). Since the vertical scan distance of adjacent tangent altitudes ranges between 1 and 1.5 km, the retrieval grid was set to 1 km up to the balloon float altitude. Above this level, the vertical spacing increases gradually to 10 km at the top altitude at 100 km. Considering the smoothing of the vertical part of the instrumental field of view, the retrieval grid is finer than the achievable vertical resolution of the measurement for a large part of the altitude region covered (especially above the observer altitude). To avoid retrieval instabilities due to this oversampling of the vertical retrieval grid, a Tikhonov–Phillips regularization approach (Phillips, 1962; Tikhonov, 1963) was applied with a constraint with respect to a first derivative of the a priori profile x_a of the target species:

$$\boldsymbol{x}_{i+1} = \boldsymbol{x}_i + [\mathbf{K}_i^T \mathbf{S}_y^{-1} \mathbf{K}_i + \mathbf{R}]^{-1}$$
$$[\mathbf{K}_i^T \mathbf{S}_y^{-1} (\boldsymbol{y}_{\text{meas}} - \boldsymbol{y}(\boldsymbol{x}_i)) - \mathbf{R}(\boldsymbol{x}_i - \boldsymbol{x}_a)], \qquad (1)$$

where \mathbf{x}_{i+1} is the vector of the desired state parameters for iteration i + 1; \mathbf{y}_{meas} is the measured radiance vector and $\mathbf{y}(\mathbf{x}_i)$ the calculation of the radiative transfer model using state parameters of iteration number i; **K** is the Jacobian matrix containing partial derivatives $\partial \mathbf{y}(\mathbf{x}_i)/\partial \mathbf{x}_i$, while \mathbf{S}_y^{-1} is the inverse noise measurement covariance matrix and **R** a regularization matrix with the first derivative operator and a regularization strength parameter.

In a first step, a temperature retrieval was performed using appropriate CO₂ lines of two separate bands around 810 and $950 \,\mathrm{cm}^{-1}$ and a priori pressure-temperature information from European Centre for Medium-Range Weather Forecasts (ECMWF) analyses together with a CO₂ VMR profile updated with data from NOAA ESRL GMD (National Oceanic and Atmospheric Administration, Earth System Research Laboratory, Global Monitoring Division; Montzka et al., 1999). The temperature retrieval 1σ accuracy is estimated to be within about 0.7 K. Then, VMR profiles of the target species are individually retrieved in selected spectral regions (see Table 1). Profiles of species interfering with the target molecule were adjusted simultaneously during the retrieval procedure. An overview of the analysis of spectra with regard to chlorine- and nitrogen-containing molecules is given in von Clarmann et al. (1995) and Wetzel et al. (2002, 2010). The error estimation of the target parameter consists of random and systematic errors that were added in quadrature to yield the total error, which refers to the 1σ confidence limit. Random errors include spectral noise as well as covariance effects of the simultaneously fitted parameters. Systematic errors mainly comprise spectroscopic data inaccuracies (band intensities), uncertainties in the line of sight, and gain calibration errors. The altitude resolution is calculated from the number of degrees of freedom of the retrieval, which corresponds to the trace of the averaging kernel matrix. Typical values for the retrieved parameters are given in Table 1.

Many trace gases measured by the MIPAS-B instrument have been involved in a large number of validation activities and cross-comparisons of satellite sensors like MIPAS, ILAS/ILAS-II (Improved Limb Atmospheric Spectrometer), and SMILES (Superconducting Submillimeter-Wave Limb-Emission Sounder). For species used in this work we explicitly mention for evaluation the following studies: CIONO₂ (Höpfner et al., 2007; Wetzel et al., 2008, 2013), CFC-11 and CFC-12 (Wetzel et al., 2008, 2013), ClO (Sagawa et al., 2013), and N_2O (Wetzel et al., 2008; Payan et al., 2009).

2.2 TELIS instrument and data analysis

The cryogenic heterodyne balloon sounder TELIS was developed in a collaboration of three partners: the German Aerospace Centre (DLR), Rutherford Appleton Laboratory (RAL), United Kingdom, and the Netherlands Institute for Space Research (SRON). Each institute generated one channel: 1.8 THz (DLR), 500 GHz (RAL), and 480–650 GHz (SRON). A comprehensive description of the instrument is given by Birk et al. (2010) and de Lange et al. (2012). The HCl and ClO results presented here were derived from spectra in the 480-650 GHz channel with a tunable superconducting integrated receiver developed and characterized by de Lange et al. (2010, 2012). A local oscillator (LO) reference signal is mixed with the atmospheric signal in a nonlinear mixer. The measured spectrum is the superposition of two spectra covering the frequency ranges $v_{LO} + v_{IF}$ and $v_{\rm LO} - v_{\rm IF}$, where $v_{\rm IF}$ is the measured difference (intermediate) frequency (IF).

The analysis of the TELIS spectra is carried out in a similar way as for the MIPAS-B retrieval procedure. A forward line-by-line model is used to model the radiative transfer along the line of sight of the instrument. Spectroscopic parameters are also taken from the HITRAN database (Rothman et al., 2009). An instrument model to account for the specifics of the TELIS instrument is included in the forward algorithm. Further details on the forward model are described by de Lange et al. (2009) and references therein. The forward model is inverted with a Gauss–Newton iteration scheme in combination with a Tikhonov–Phillips regularization approach (Phillips, 1962; Tikhonov, 1963) as described in the previous section.

The HCl retrievals are performed for both chlorine isotopes H³⁵Cl and H³⁷Cl. The total amount of HCl can be determined by taking into account the isotope abundance of $H^{35}Cl$ (75.76%) and $H^{37}Cl$ (24.23%). While the random error of HCl is very small (~ 0.01 ppbv), the systematic error estimate yields between 0.05 and 0.4 ppbv, resulting in a total error of about 10 to 15 % in the region of the VMR maximum. Systematic error sources are instrumental uncertainties such as instrumental line shape and side band ratio inaccuracies, detector non-linearity, and calibration and pointing errors. Furthermore, errors in the atmospheric pressuretemperature profile as well as spectroscopic data errors are taken into account. The largest uncertainty stems from the non-linear behaviour of the detector. This holds also for the ClO retrievals. The overall accuracy of the ClO measurement is almost entirely determined by systematic error sources. Similar to HCl, the total error for the species ClO typically remains within 10 and 15 % in the altitude region of its VMR maximum. An overview of the characteristics of the retrieved species is given in Table 2.

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Table 1. Set-up for MIPAS-B trace species retrievals and typical (1σ) errors. Results are given for different state parameters in corresponding
spectral windows together with the retrieval altitude resolution (Alt. reso.).

Species	Spectral range (cm $^{-1}$)	Noise error ^a (%)	Total error ^a (%)	Alt. reso. (km)
ClONO ₂	779.7–780.7	2–3	5–6	4–5
ClO	821.0-841.5	10–25 ^b	20–30 ^b	5-8
CFC-11	840.0-860.0	2–3	5–6	3–4
CFC-12	918.0-924.0	2–3	5–6	3–4
HCFC-22	828.0-830.0	3–10	8-15	3–6
CFC-113	813.0-830.0	3–10	20-25	3–6
CCl ₄	792.0-806.0	1–5	10-20	4–6
CH ₃ Cl	742.5-755.0	1–5	8-15	9–13
HOCl	1215.0-1265.0	10–15	35-50	6–8
N ₂ O	1227.8-1303.1	2–3	5–6	2–4

 $^{\mathrm{a}}$ In the altitude region around the VMR maximum; $^{\mathrm{b}}$ daytime errors.

Table 2. Set-up for TELIS HCl and ClO retrievals with typical (1σ) errors and retrieval altitude resolution (Alt. reso.).

Species	Spectral line (GHz)	Noise error* (%)	Total error* (%)	Alt. reso. (km)
H ³⁵ Cl	625.9	< 1	10–15	2–5
H ³⁷ Cl	624.8	< 1	10-15	2–5
ClO	501.3	< 1	10–15	2–4

* In the altitude region around the VMR maximum.

TELIS HCl and ClO observations have been evaluated using MLS measurements (de Lange et al., 2012). ClO was additionally compared to SMILES observations (Sagawa et al., 2013). The Sagawa study also includes a cross-comparison with MIPAS-B ClO observations.

2.3 Model calculations

Measured data are compared to simulations performed with the chemistry climate model EMAC, which is a numerical chemistry and climate simulation system that includes submodels describing tropospheric and middle atmosphere processes (Jöckel et al., 2010). It uses the second version of the Modular Earth Submodel System (MESSy2) to link multiinstitutional computer codes. The core atmospheric model is the fifth-generation European Centre Hamburg general circulation model (ECHAM5, Roeckner et al., 2006). For the present study we applied EMAC (ECHAM5 version 5.3.02, MESSy version 2.50) in the T42L39MA-resolution, i.e. with a spherical truncation of T42 (corresponding to a quadratic Gaussian grid of approximately 2.8° by 2.8° in latitude and longitude) with 39 vertical hybrid pressure levels from the ground up to 0.01 hPa. The applied model set-up comprised, among others, the submodels MECCA (Sander et al., 2005) for the calculation of gas-phase chemistry and the submodel MSBM (Kirner et al., 2011) for the simulation of polar stratospheric clouds and the calculation of heterogeneous reaction rates. The PSC scheme was validated with the help of HNO₃, ClO, and O_3 data from the MLS instrument (Kirner et al., 2015).

A Newtonian relaxation technique of the prognostic variables temperature, vorticity, divergence, and surface pressure above the boundary layer and below 1 hPa towards the ECMWF reanalysis ERA-Interim (Dee et al., 2011) has been applied to the model to simulate realistic synoptic conditions (van Aalst, 2005). Boundary conditions for greenhouse gases, chlorofluorocarbons (CFCs), and halons are adapted from observations (WMO, 2011; Meinshausen et al., 2011). Halogenated hydrocarbons are included according to the WMO-A1 scenario (WMO, 2011). Chlorine-containing tropospheric source gases considered in EMAC are CFC-11, CFC-12, HCFC-22, CFC-113, CCl₄, CH₃Cl, and CH₃CCl₃. Photolysis rates of HCFC-22 and CFC-113 are the same as for CFC-12. The simulation includes a comprehensive chemistry set-up from the troposphere to the lower mesosphere with 104 gas-phase species, 234 gas-phase reactions, 67 photolysis reactions, and 11 heterogeneous reactions on liquid aerosols, nitric acid trihydrate, and ice particles. Rate constants of gas-phase reactions are taken from Atkinson et al. (2007) and Sander et al. (2011). The model output data were saved every 10 min. The temporally closest model output to the MIPAS-B measurements has been interpolated in space to the observed geolocations.

3 Chlorine partitioning and budget

The combination of two different sensors, MIPAS-B and TELIS, working in different spectral regions (mid-infrared and microwave), enables the simultaneous measurement of virtually all relevant inorganic and organic chlorine molecules. The amount of inorganic chlorine $[Cl_y]$ is defined as

$$[Cly] = [ClOx] + [HCl] + [ClONO2],$$
(2)

where active chlorine $[ClO_x]$ is calculated via

$$[ClO_x] = [ClO] + [HOCl] + 2[ClOOCl].$$
(3)

The amount of organic chlorine $[CCl_y]$ is composed of

$$[CCl_y] = 2[CFC-12] + 3[CFC-11] + [HCFC-22] + 3[CFC-113] + 4[CCl_4] + [CH_3Cl].$$
(4)

Total chlorine [Cl_{total}] is given as the sum of both budgets:

$$[Cl_{total}] = [Cl_y] + [CCl_y].$$
(5)

Constituents, which are of minor importance for the Arctic stratospheric chlorine budget (like Cl₂, Cl, OClO, CH₃CCl₃, CFC-114, CFC-115, HCFC-141b, HCFC-142b, Halon-1211; see e.g. Prinn et al., 2000) are neglected here. All the quantities defined in Eqs. (2) to (5) can be deduced from TELIS (measuring ClO and HCl) and MIPAS-B (measuring all gases except HCl) observations. However, the chlorine monoxide dimer ClOOCl is only measurable by MIPAS-B under activated chlorine conditions ([ClOOCl] > 0.5 ppbv) without any PSC emissions in the recorded spectra (Wetzel et al., 2010). On 31 March 2011, no PSC signatures are visible in the MIPAS-B spectra but ClOOCl concentrations are below the detection limit. However, [ClOOCl] can be estimated from [ClO] with the following relation (Wetzel et al., 2012):

$$[\text{ClOOCl}] = ([\text{ClO}_{\text{noon}}] + 2[\text{ClOOCl}_{\text{noon}}] - [\text{ClO}])/2, \quad (6)$$

while the amounts of $[ClO_{noon}]$ and $[ClOOCl_{noon}]$ which correspond to noon maximum and minimum values respectively can be both taken from EMAC simulations if the modelled ClO is constrained to the measured one.

MIPAS-B spectra have been recorded from night until day. The sunrise took place between 02:38 UTC at 36 km and 03:10 UTC at 9 km altitude. Figure 2 shows the measured ClO cross section from 02:00 to 04:38 UTC, corresponding to 64.0° N, 30.1° E and 63.5° N, 28.9° E. A temporal variation of ClO is visible. The concentration of this species is a measure of whether the air masses sounded are still chlorineactivated or not. After sunrise the mixing ratio of ClO increases in a layer between 16 and 22 km from nighttime values below 0.05 ppbv to daytime mixing ratios up to 0.4 ppbv. During periods of strong chlorine activation, significantly higher values around 2 ppbv are observed (see e.g. Santee



Figure 3. $CIONO_2$ mixing ratios (ppbv) as seen by MIPAS-B above northern Scandinavia on 31 March 2011 inside the late-winter Arctic vortex. The black solid line marks the sunrise terminator.

et al., 2003; Wetzel et al., 2012). The CIO increase is shown by both instruments, MIPAS-B and TELIS. The latter instrument measured not only with higher vertical resolution but also with higher temporal resolution compared to MIPAS-B; hence the TELIS data were transferred to the coarser temporal grid of MIPAS-B for better comparability. At altitudes above 26 km, MIPAS-B CIO temporal retrieval fluctuations are visible due to the large spectral noise error in this altitude region. As a consequence, the TELIS CIO data were used for calculating the chlorine partitioning and budget in the whole altitude range.

The decreasing ClO_x at the end of the Arctic winter in the lower stratosphere due to rising temperatures followed by shrinking ClO_x production from heterogeneous chemical reactions is in line with high amounts of ClONO2 in this altitude region. The reaction of ClO with NO₂ produces the reservoir species ClONO₂. The measured time evolution of this molecule is displayed in Fig. 3. Measured ClONO₂ data exhibit high values that are typical for observations in the late Arctic winter (see e.g. Oelhaf et al., 1994; von Clarmann et al., 1997, 2009; Wetzel et al., 2002). Only in an atmospheric layer around 19 km is the vertical mixing ratio gradient small, since ClONO₂ values (around 19 km) are slightly lower than they would be if chlorine was completely deactivated. This observed signature is in line with the enhanced ClO amounts around 19 km as seen in Fig. 2. A significant diurnal temporal variation is not visible in the ClONO₂ data.

The mean measured chlorine partitioning and budget for early morning is displayed in Fig. 4. A spectral noise error weighted averaging was applied to calculate the mean profiles, although statistical errors of the individual species profiles are similar. The molecules CIO and CIOOCI exhibit a temporal variation over the measured time period. However, since their mixing ratios are very low at this time in the year, vertical profiles of these species have also been avG. Wetzel et al.: Partitioning and budget of inorganic and organic chlorine species



Figure 4. Partitioning and budget of inorganic, organic, and total chlorine as measured by TELIS (HCl and ClO) and MIPAS-B (all other species) in the Arctic stratosphere on 31 March 2011 (see legend for line style and note non-linear abscissa). The reservoir species HCl and ClONO₂ dominate the stratospheric inorganic chlorine budget. Cl_y^* deduced from observed N₂O data with the help of a N₂O–Cl_y correlation (see Eq. 7) is shown for comparison. Note that for the calculation of the chlorine budgets the atomic content for each species has to be considered (some error bars have been omitted for clarity).

eraged over the observed time period, with almost no consequence. To obtain a proxy of total inorganic chlorine, a N₂O-Cl_v correlation was derived from air samples collected with the balloon-borne cryogenic whole air sampler BONBON in the Arctic between 2009 and 2011 according to the method described in Engel et al. (2002) and Wetzel et al. (2010). Cl_{ν} from the cryosampler measurements is calculated as the difference between total chlorine and observed organic chlorine from the source gases CFC-11, CFC-12, CFC-113, CH₃CCl₃, CCl₄, HCFC-22, HCFC-141b, and HCFC-142b. In addition, an input of 50 pptv of chlorine from short-lived source gases is taken into account, which is assumed to be transformed immediately to inorganic chlorine. Total chlorine from the gases is propagated into the stratosphere in the same way as an inert tracer, as described in Engel et al. (2002), using global mean observation data from NOAA ESRL. The proxy inorganic chlorine $[Cl_{y}^{*}]$ is calculated with the following dependence on the amount of $[N_2O]$, both given in ppbv:

$$[Cl_{y}^{*}] = 3.2008346 + 8.7786479 \times 10^{-6} [N_{2}O] - 2.9132361 \times 10^{-5} [N_{2}O]^{2}.$$
(7)

This correlation has been applied to MIPAS-B measured N_2O and yields up to 3.20 ppbv Cl_v^* in the stratosphere. The amount of inorganic chlorine is dominated by the chlorine reservoir species ClONO₂ and HCl, the latter one especially above 24 km. Above this altitude, where the Cl_v VMR is (vertically) approximately constant, the mean observed Cl_{ν} amounts to 3.25 ± 0.30 ppbv, which is in agreement with the deduced Cl_{ν}^{*} within the error bars although there is a tendency towards a small positive deviation in the observations compared to the Cl_{ν}^{*} reference according to Eq. (7). The deviation between Cl_{y} and Cl_{y}^{*} below 21 km is caused by different degrees of subsidence of the air masses in the case of the discussed balloon flight and the Cl^{*}_v reference, which results in different N₂O mixing ratios in a specific altitude. Cl_{y} species play by far the largest part in the total chlorine budget from the lower to the upper Arctic winter stratosphere. From about 17 km downwards, the amount of organic chlorine gets increasingly dominant in the total chlorine budget. Source gases that contribute to CCl_{y} are visible in Fig. 4: CFC-12 (CCl₂F₂), CFC-11 (CCl₃F), HCFC-22 (CHClF₂), CFC-113 $(C_2Cl_3F_3)$, CCl_4 , and CH_3Cl . The mean amount of Cl_{total} is calculated as 3.41 ± 0.30 ppbv above 24 km. From the ratio Cly to Cltotal it follows that about 95% of total chlorine is inorganic in this altitude region.

The mean chlorine partitioning and budget as simulated by EMAC is shown in Fig. 5. The principal vertical profile shape of the measured chlorine species is well reproduced by the model. However, some differences in detail between simulated and observed data are visible. The modelled HCl VMR maximum appears slightly broader than the measured one. Below about 20 km, the simulation shows significantly lower values compared to the observation by TELIS. A difference is visible in the case of ClONO₂. The model underestimates this reservoir species and deviates by 0.8 ppbv (42%) from the MIPAS-B data in the region of the VMR maximum at 22 km, although simulated and measured NO_{ν} and NO₂ (a necessary reactant in the production of ClONO₂ via NO2 plus ClO) agree in this altitude region. Since simulated HCl and ClO_x (near 22 km) are in agreement with the observed data, the simulated Cl_{y} deviation from the measurement can be attributed to the ClONO₂ deficit in EMAC. Around 19 km, the difference in simulated and measured Cl_{y} is largest due to very low HCl values in EMAC compared to the HCl seen by TELIS. The amount of available Cl_v below about 24 km is dependent on the degree of downwelling of the air masses inside the polar vortex. In EMAC, the subsidence of the air masses in the course of the winter was underestimated such that we find higher values of tracers like N₂O and CFCs at a given altitude of the lower stratosphere compared to the measurements. These higher N₂O values are connected with lower Cl_{y} values according to the compact N_2O-Cl_v relationship, resulting in an underestimation of the chlorine reservoir species (especially ClONO₂). So, at least part of the ClONO₂ deficit in EMAC can be explained by the underestimation of the subsidence in the model.



Figure 5. Partitioning and budget of inorganic, organic, and total chlorine as simulated with the chemistry climate model EMAC on 31 March 2011 (see legend for line style). Cl_y^* has been calculated from the simulated N₂O data according to Eq. (7). The budgets ClO_x , Cl_y , CCl_y , and Cl_{total} are calculated as listed in Eqs. (2) to (5) respectively. The shaded region of the budgets takes into account all minor chlorine species contained in EMAC (Cl_2 , Cl_2 , Cl_3) that were not measured by MIPAS-B and TELIS.

The simulated Cl_y reaches its maximum VMR in the quasi-altitude-constant region above 24 km with a mean value of 3.16 ppbv, which is slightly lower than the measured one and close to the simulated value of Cl_y^* (deduced from EMAC), which gives 3.19 ppbv. Below this altitude region, a similar bias between Cl_y and Cl_y^* , as in the case of the observations, is visible.

The mean amount of Cl_{total} in the model run is calculated as 3.21 ppbv above 24 km, which is 0.20 ppbv lower than the observed one. About half of this simulated chlorine deficit can be explained by the fact that some minor CFCs (e.g. CFC-114 and CFC-115) and HCFCs (e.g. HCFC-141b and HCFC-142b) as well as halons are not included in the EMAC model. Their contribution to Cl_{total} is not more than 1% above 24 km (Brown et al., 2013). The remaining deficit can be explained by very short-lived chlorine species which altogether amount to about 0.1 ppbv (Mébarki et al., 2010; WMO, 2011) and which are also not contained in the model simulation. However, the chlorine amount of these missing species is implicitly contained in the HCl measurement (since the short-lived chlorine species are converted to HCl after being photolysed) and hence included in the observed chlorine budget. In the altitude region above 24 km, about 98% of total chlorine in EMAC is inorganic. The shaded region of the budget profiles of ClO_x , Cl_y , CCl_y , and Cl_{total} shown in Fig. 5 takes into account all available chlorine species in EMAC that were not measured by MIPAS-B and TELIS. These molecules comprise Cl, Cl₂, OCIO (belonging to ClO_x and Cl_y), and CH_3CCl_3 (belonging to CCl_y) and add up to 0.1 ppbv at 16 km to the total chlorine budget (mainly due to Cl₂ and OCIO). However, at altitudes between 22 and 36 km, contributions of these gases to the chlorine budget are insignificant.

4 Conclusions

Observations from MIPAS-B/TELIS were performed at the end of the cold 2010/2011 stratospheric winter that was characterized by a persistent polar vortex enabling strong chlorine activation and ozone loss. The chlorine partitioning measured on 31 March 2011 reveals that in the outer part of the polar vortex (above Finland) the recovery of active chlorine (ClO_x) into the reservoir species (mainly $CIONO_2$) is almost completed by the end of March, only a few days before the cold period had finished (Manney et al., 2011). This is verified by low amounts of daytime ClO of up to 0.4 ppbv around 19 km. The observed total stratospheric chlorine amounts to 3.41 ± 0.30 ppbv above 24 km (see Table 3). This is in accordance with the EMAC simulation (3.21 ppbv), taking into account the fact that some chlorine source gases and very short-lived species are not included in the model. The horizontal Cl_{total} distribution in EMAC (above 24 km) exhibits virtually no variation inside the polar vortex. The variation inside/outside vortex is no larger than 0.1 ppbv. That is smaller than the estimated Cl_{total} measurement accuracy of 0.3 ppbv so that the observations can be treated as representative at least for the geographical region of the Arctic vortex. Mean Cl_{total} values deduced from spectra recorded by the ACE-FTS instrument (Brown et al., 2013) give 3.44 ± 0.18 ppbv (morning occultations) and 3.50 ± 0.13 ppbv (evening occultations) for northern mid-latitudes and the Arctic in 2009. Extrapolating these data to 2011 with the chlorine trend (between 2004 and 2009) obtained from these ACE-FTS observations (about -0.4 % per year) yields Cl_{total} values of 3.41 ppbv (morning occultations) and 3.47 ppbv (evening occultations) comparable to the MIPAS-B/TELIS data. The accumulated amount of minor species (not measured by MIPAS-B/TELIS) like CFC-114, CFC-115, HCFC-141b, HCFC-142b, and Halon-1211 was estimated to be about 0.7 % (\sim 0.02 ppbv) of total chlorine at 30 km (Brown et al., 2013). Hence, the MIPAS-B/TELIS Cl_{total} value is in line with the data obtained from ACE-FTS solar occultations and is consistent with the decreasing amount of stratospheric chlorine. Considering the 2005 mean global tropospheric Cl_{total} from in situ data of AGAGE (Advanced Global Atmospheric Gases Experiment) and NOAA ESRL databases, as compiled in WMO (2011), and transferring this value to 30 km, taking into account a typical time lag of 6 years of stratospheric mean age of air (Engel et al., 2002, 2009; Stiller et al., 2008; WMO, 2011),

Table 3. Mean stratospheric chlorine budgets (ppbv, including 1σ total errors) as measured by MIPAS-B/TELIS and simulated by EMAC in comparison to ACE-FTS observations (Brown et al., 2013) and in situ data from AGAGE and NOAA ESRL databases (WMO, 2011).

Budget	MIPAS-B/TELIS	EMAC	ACE-FTS	In situ
Cl _{total}	3.41 ± 0.30^a	3.21 ^a	3.41/3.47 ^b	3.40 ^c
Cl_y	3.25 ± 0.30^a	3.16 ^a	-	-
Cl_y^*	3.19 ± 0.002^a	3.19 ^a	-	_

^a Mean value between 25 and 36 km; ^b mean value (for morning/evening occultations) between 30 and 70° N for 2011, extrapolated from 2009 with trend between 2004 and 2009; ^c mean global tropospheric value from 2005 corresponding to a stratospheric value of 2011 assuming a stratospheric mean age of air of 6 years.

we get an estimated Cl_{total} value of 3.40 ppbv for the year 2011, which is very close to the MIPAS-B/TELIS result (3.41 ppbv).

We finally conclude that the stratospheric total chlorine as deduced from Arctic MIPAS-B/TELIS observations on 31 March 2011 confirms previously published total chlorine assessments and their related trends. A recently published study by Mahieu et al. (2014) shows a HCl concentration increase between 2005/2006 and 2010/2011 in large parts of the northern hemispheric lower stratosphere in combination with an increase in the mean age of stratospheric air of up to 0.4 years. However, in the Arctic above 24 km, ascertained changes of mean age of stratospheric air are small (Mahieu et al., 2014) and, therefore, do not alter the findings above.

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