Analysis of HCl and ClO time series in the upper stratosphere using satellite data sets


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Abstract. Previous analyses of satellite and ground-based measurements of hydrogen chloride (HCl) and chlorine monoxide (ClO) have suggested that total inorganic chlorine in the upper stratosphere is on the decline. We create HCl and ClO time series using satellite data sets extended to November 2008, so that an update can be made on the long term evolution of these two species. We use the HALogen Occultation Experiment (HALOE) and the Atmospheric Chemistry Experiment Fourier Transform Spectrometer (ACE-FTS) data for the HCl analysis, and the Odin Sub-Millimetre Radiometer (SMR) and the Aura Microwave Limb Sounder (Aura-MLS) measurements for the study of ClO. Altitudes between 35 and 45 km and two mid-latitude bands: 30°S–50°S and 30°N–50°N, for HCl, and 20°S–20°N for ClO and HCl are studied. ACE-FTS and HALOE HCl anomaly time series (with QBO and seasonal contributions removed) are combined to produce all instrument average time series, which show HCl to be reducing from peak 1997 values at a linear estimated rate of −5.1% decade⁻¹ in the Northern Hemisphere and −5.2% decade⁻¹ in the Southern Hemisphere, while the tropics show a linear trend of −5.8% per decade (although we do not remove the QBO contribution there due to sparse data). Trend values are significantly different from a zero trend at the 2 sigma level. ClO is decreasing in the tropics by −7.1% ± 7.8% decade⁻¹ based on measurements made from December 2001 to November 2008. The statistically significant downward trend found in HCl after 1997 and the apparent downward ClO trend since 2001 (although not statistically significant) confirm how effective the 1987 Montreal protocol objectives and its amendments have been in reducing the total amount of inorganic chlorine.

1 Introduction

Stratospheric ozone depletion is the result of anthropogenic release of chlorofluorocarbons (CFCs) in the troposphere. The lifetimes of the CFCs in the troposphere are very long allowing for a build up of concentrations of these gases, which are slowly transported to the stratosphere. CFCs are eventually photolysed in the stratosphere due to intense ultra violet radiation, where active chlorine (Cl) is released and is available for the destruction of ozone (Molina and Rowland, 1974). However, due to the 1987 Montreal protocol agreement (and the subsequent adjustments), ozone depleting substance (ODS) emissions have slowly been phased out. It has been estimated that the total inorganic loading of tropospheric Cl peaked in 1993 (Rinsland et al., 2003) and has decreased by ∼5% through early 2002 (O’Doherty et al., 2004). As it is estimated that it takes 3–6 years for air to be transported into the middle and upper stratosphere (Stiller et al., 2008), then peak stratospheric values are suggested to be somewhere between 1996 and 1999 (WMO, 2006). Based on assumptions that the international agreements are adhered to, it is proposed using the equivalent effective stratospheric chlorine (EESC) parameter that total CI values will reach pre 1980 levels around 2040 (Newman et al., 2007).

Two gases that are important contributors to the Cly family (Cly = Cl + ClO + HCl + ClONO₂ + 2Cl₂O₂ + HOCl + OClO + BrCl + 2Cl₂) are hydrogen chloride (HCl) and chlorine monoxide (ClO). HCl is important to the gas phase chemistry of ozone depletion because of its role as a reservoir species for Cl. ClO on the other hand, carries most of the reactive chlorine in the stratosphere during day time. Past studies of these gases have suggested that there has been a slow down and even a decrease in the accumulation of ClO in the stratosphere using ground based measurements and satellite data sets (Considine et al., 1997, 1999; Anderson et al., 2000; Newchurch et al., 2003; Froidevaux et al., 2006; Solomon et al., 2006).
This study intends to examine the long term evolution of HCl and ClO using satellite data sets of the Halogen Occultation Experiment (HALOE), the Atmospheric Chemistry Experiment Fourier Transform Spectrometer (ACE-FTS), the Aura Microwave Limb Sounder (Aura-MLS), and the Odin Sub-Millimetre Radiometer (SMR). We examine altitudes from 35 to 45 km and mid-latitudes from $30^\circ$S–$50^\circ$S and $30^\circ$N–$50^\circ$N for HCl and $20^\circ$N–$20^\circ$S for ClO and HCl. The choice for investigating this altitude region is because ClO and HCl together make up more than 95% of the chlorine budget here (WMO, 2006). Additionally, it is in this altitude range where ozone is expected to show signs of a recovery first (Jucks et al., 1996). The analysis will make trend estimates based on monthly residual averaged data. By showing the current state of total inorganic chlorine in the stratosphere, we can ascertain if the Montreal protocol objectives are still working.

2 HCl and ClO data sets

2.1 HALOE

The HALOE instrument aboard the Upper Atmosphere Research Satellite (UARS) was operational from September 1991 to November 2005. HALOE employed a solar occultation instrument measuring many trace gases including HCl. Observations were made in the infrared part of the electromagnetic spectrum (between 2.45 and 10 µm) (Russell et al., 1993). The HALOE occultation instrument was highly sensitive and obtained 15 occultation measurements during each sunrise/sunset by comparing the solar spectra to the spectra obtained whilst scanning through the atmosphere. This produced in essence a self calibrating instrument with long term stability. A vertical profile resolution of approximately 2–4 km was typically obtained for HCl, between 10 and 65 km. Global coverage from about 80$^\circ$S and 80$^\circ$N was achieved in approximately six weeks. As solar occultation relies on the sun, it means there will be no measurements made during polar night in the winter hemisphere. Moreover, it should be noted that the coverage of HALOE degraded over time and there were fewer measurements during the later years of the mission. We suggest the reader to consult (http://haloe.gats-inc.com/home/index.php) for more details. For the trend analysis, we ignore data for 1991 and 1992 to stay consistent with the work done by Newchurch et al. (2003). The reason for their analysis not to include these two years was due to the fact that the slope of a linear fit will be greatly affected by the beginning and ending data values. Because the HALOE data start during the Pinatubo period, they were very cautious to obtain a robust regression. Hence, they omitted any suspicious data. Data used in this analysis are from the HALOE v19 and can be obtained from http://haloe.gats-inc.com/home/index.php. The combined systematic and random uncertainties in the upper stratosphere are typically between 12 and 15%, while correlative measurements with other instrument data show agreement between 8 and 19% (Russell et al., 1996).

2.2 ACE-FTS

Another infrared solar occultation instrument yielding vertical profiles of HCl concentrations is the ACE Fourier Transform Spectrometer (ACE-FTS) (Bernath et al., 2005). The SCISAT platform maintains a low Earth repeating orbit with a geographical coverage between approximately 85$^\circ$S–85$^\circ$N, providing the opportunity for ACE-FTS to make typically 15 sunrise and 15 sunset solar occultations per day. Unlike HALOE, the ACE-FTS orbit does not change on a year to year basis, meaning that typically the same latitudes are sampled for a given time period.

Vertical profiles are retrieved from observed solar spectra using a non-linear least-squares global fit technique (Boone et al., 2005). Currently, HCl v2.2 VMRs are from $\sim$10–50 km with a vertical resolution of approximately 3–4 km. Comparisons of ACE-FTS HCl to various other satellite and ground based instrumentation show that there is a good agreement of generally better than 5–10% above 20 km (Mahieu et al., 2008). At present, no error analysis for ACE-FTS atmospheric products has been made. For this study, ACE-FTS v2.2 data are filtered according to various criteria: that the associated measurement uncertainty is not greater than 100% of the corresponding VMR value. Data used in this analysis are the ACE-FTS measurements from February 2004–November 2008.

2.3 MLS

Stratospheric ClO is studied using measurements of the Microwave Limb Sounder (Aura-MLS), which was launched in July 2004 on the NASA Aura satellite (from here on, we refer to Aura-MLS as MLS). This is the second MLS instrument, whereas the previous instrument, on board the UARS platform (hence, UARS MLS), operated for $\sim$8 years between 1991 and 1999, but is not included in this analysis. MLS is a limb scanning instrument, observing thermal emission at millimetre and sub-millimetre wavelengths (Waters et al., 2006). MLS has the ability to measure at night and is not affected by stratospheric clouds. The Aura satellite maintains a suborbital track, covering 82$^\circ$S to 82$^\circ$N with an ascending node of 13:45 h local solar time (LST) and makes approximately 3500 scans per day.

MLS v2.2 CIO measurements are provided on pressure surfaces where the recommended levels to use are between 100 and 1 hPa. The vertical resolution and profile precision of the ClO measurements in this range are typically 3–4.5 km and $\sim$0.1 ppbv respectively. Comparisons to correlative measurements show the largest systematic uncertainties are found below 22 hPa for both night time and day time measurements when compared directly to co-located ground and
satellite measurements (Santee et al., 2008). However, this is of little relevance here as we examine MLS ClO measurements between 6.5 and 1.6 hPa (∼35–45 km) where systematic uncertainties are small.

MLS ClO data are screened using only profiles that have a zero status flag so that profiles with possible ambiguities are removed. Data are also only used if the quality flag is greater than 0.8. Moreover, we reject values that have a poor convergence criterion such that it is larger than 1.5. Finally, values that possess a negatively flagged uncertainty (precision) are also rejected. Please see Santee et al. (2008) for more details.

2.4 Odin/SMR

The Odin satellite was launched in the beginning of 2001 and is a joint initiative between Sweden, Canada, Finland and France. This small satellite comprises two instruments, the Sub-Millimetre Radiometer (SMR) and the Optical Spectrograph InfraRed Imager System (OSIRIS). The Odin satellite is polar orbiting (82.5° S to 82.5° N) and is sun synchronous with a descending node at 06:00 h and ascending node at 18:00 h LST. The SMR instrument makes measurements of various species including stratospheric ClO by observing thermal emission in the microwave region during day and night (Murtagh et al., 2002; Frisk et al., 2003). SMR consists of five receivers, where we use ClO from the 501.8 GHz band. We remove profiles if the quality flag is not equal to zero or for any observation with a measurement response less than 0.75.

The newest version of SMR, data version 2.1 (produced at the Chalmers University of Technology, Sweden), is used for the analysis of stratospheric ClO. The study by Santee et al. (2008) compares coincident MLS and SMR measurements, and it is found that there is an excellent agreement for altitudes between 35 and 45 km (within 0.1 ppbv). An overview of the SMR ClO measurements can be found in Urban et al. (2005, 2006). Note that SMR v2.1 and v2.0 ClO products are virtually identical. The vertical resolution for the ClO product at 501.8 GHz is 2–4 km in the stratosphere.

3 Methodology

Here, we describe the methods used in order to derive long term trends of stratospheric HCl and ClO. We use separate methodologies for each species, although a linear regression analysis is used in each case to derive the resulting trends.

3.1 HCl

As the main focus of the HCl analysis is on the upper stratosphere, we firstly filter both HALOE and ACE-FTS data between 35 and 45 km into three latitude bands: 30° S–50° S, 20° S–20° N, and 30° N–50° N. Monthly averages are calculated for the filtered data. Each data set comprises a large number of measurements, implying that the stochastic error should be minimised when measurements are averaged. We have chosen these altitude ranges firstly because the first signs of ozone recovery are expected to be seen in the upper stratosphere (Jucks et al., 1996) and since this upper altitude range is also studied by other analyses and hence the results found here can easily be compared (for example, Newchurch et al. (2003)). Figure 1 illustrates the monthly averaged time series for each data set for the three latitude bands mentioned above. Due to the SCISAT orbital parameters and due to the nature of the viewing geometry, ACE-FTS measurements are sparse in the tropics, and eventual trend estimates in the tropics have to be treated with more care compared to the mid-latitude cases. In all three cases there is an apparent ∼0.2–0.4 ppb positive ACE-FTS bias compared to the HALOE observations during overlapping measurement periods. This technique does not follow the conventional comparison method of profile to profile comparisons, but rather compares time series of average measurements over a long period of time as they are used in this analysis. Mismatches are thus possible in terms of time and space. This result agrees with similar analyses made by Mahieu et al. (2008), who show that there is ∼10–20 % ACE-FTS bias compared to HALOE for altitudes between 20 and 55 km. Work by Froidevaux et al. (2008) also show that the biases between MLS and HALOE are of a similar magnitude in the same altitude region. Moreover, MLS and ACE-FTS HCl data agree better than either versus HALOE. Additionally, it has also been shown that HALOE is biased low versus other correlative data, based on the Russell et al. (1996) HALOE HCl validation paper. Interesting features are the peak in HALOE values until around 1997 after which values start to slowly decrease until instrument termination at the end of 2005. However, there is a large degree of variability in the values between 1997 and 2002, which is also seen in previous findings for HALOE HCl observations at 55 km (Waugh et al., 2001; WMO, 2006), although these features are not fully understood. The ACE-FTS time series exhibits strong seasonal variations, especially in the Southern Hemisphere (lower panel) and tropics (middle panel). The tropics also exhibit smaller interannual variations due to the semi-annual oscillation (SAO), typically present at these altitudes, although data here are limited as occultation geometry does not continuously cover tropical latitudes. The seasonal and inter seasonal variations are less apparent in the Northern Hemisphere (top panel) compared to the Southern Hemisphere. This difference could be a consequence of stronger wave activity in the Northern Hemisphere in comparison to the Southern Hemisphere. If this is the case, then the end result is that the seasonal variations in the Northern Hemisphere will be masked due to the increased turbulent mixing in the surf zone of the northern mid-latitudes. HALOE values exhibit more noise in the measurements as they have a lower sampling rate compared to ACE-FTS (HALOE typically 50–350 profiles globally a month, ACE-FTS typically...
>300–500 profiles a month). A typical example of spatial coverage for all four instruments studied in this analysis is illustrated in Fig. 2. Here, the HALOE and ACE-FTS orbital paths represent the typical spatial coverage completed during one month, while the SMR and MLS orbital paths are the typical spatial coverage for observations made during one day.

### 3.1.1 Removing known sources of variation

The variations of each individual time series are cyclic in nature and are associated mainly to seasonal cycles (including SAO) and the quasi-biennial oscillation (QBO). HCl’s response to variations in solar intensity are small between 35 and 45 km (Egorova et al., 2005) and is not accounted for in this analysis. By being able to separate the relative contributions of known variation (by the process of linear regression), we will be left with the unexplained variability of the monthly mean signal to which a linear trend can be applied. We follow similar approaches to that of Newchurch et al. (2003), and Steinbrecht et al. (2004, 2006), where we firstly remove the seasonal cycle from the HCl time series. This is simply done for each instrument by finding the difference between each monthly mean value from their corresponding average (climatological) annual cycle. For example, the HALOE mean January value calculated for all Januarys during 1993–2005 is subtracted from each individual HALOE January VMR value. Hence, the result HALOE and ACE-FTS time series will be a monthly mean HCl time series with the seasonal components removed, but still containing contributions from the QBO and other sources of variation.

In order to model the HCl anomalies from HALOE and ACE-FTS time series (with the seasonal component already removed), we apply a linear regression model to each set of anomalies independently, accounting for fluctuations related to the QBO.

\[
\Delta \text{HCl}(t) = b + at + \sum_{i=1}^{QBO}(c_i \cos \left( \frac{2\pi t}{P_{\text{QBO}}(i)} \right) + d_i \sin \left( \frac{2\pi t}{P_{\text{QBO}}(i)} \right)) + N_i, \tag{1}
\]

where \(\Delta \text{HCl}(t)\) are the HCl anomalies calculated for each month, \(t\). The first and second components in the model represent a linear trend, where \(b\) is a constant when time \(t = 0\) and \(a\) is a linear fit coefficient. The third set of terms represents the QBO cycle, which is a combination of cosine and sine wave functions, which use harmonics, \(P_{\text{QBO}}(i)\), that best represent the behaviour of the QBO cycle. The last term in the model, \(N_i\), is the residual or autocorrelated error term, which is an autoregressive process such that, \(N_i = \theta N_{i-1} + e_i\), where \(\theta\) is the autocorrelation (lag 1 month) and \(e_i\) are the white noise residuals with zero mean and common variance. As we have already removed the seasonal cycle prior to this step, there is no need to include a seasonal cycle component. We have chosen to do this as we treat the tropical bin slightly different since we do not fit for the QBO parameter. The reason for this is due to the sparse contribution of ACE-FTS HCl data between 20° S–20° N, which causes difficulties in estimating reliable harmonics representing the cyclic behaviour of the QBO.

The linear fit, \(a\), calculated for each independent set of anomalies is the estimated fit over each entire time series. However, \(a\) is solely a product of our estimation of the cyclic behaviour of each full time series and is only used for calculating the QBO contribution. This makes sense since the HALOE time series, for example, does not maintain a linear trend for its entirety, as can be seen from Fig. 1.

It should also be noted that subtraction of the monthly mean values (for deseasonalisation) will be reflected in the axis intercept \(b\) (compared to a (possible) fit to the original HCl time-series).
The terms \(a, b, c\) and \(d\) need to be calculated so that the total summed contribution of the QBO can be determined. Hence, Eq. (1) can be solved for pre-determined QBO cycles \(P_{QBO}(i)\) by least squares analysis such that the sum of the squares of the residuals is minimised. Thus, the remaining HCl monthly anomalies are from sources of variability that cannot be explained by seasonal or QBO behaviour. The fundamental period of the QBO in the tropics is around 30 months, but there are also smaller periods modulating around 12 months, especially relevant for the mid-latitudes (Baldwin et al., 2001). Furthermore, Steinbrecht et al. (2004) showed statistically significant (at 90\% ) periods found in ground-based stratospheric ozone data, to be between 7.5 and 32.2 months. Here, we estimate the QBO signals by examining the HCl deseasonalised time series directly, when passed through a fast Fourier transform (FFT). Large peaks observed between 7–9, 16–22, and 26–32 months in the resulting FFT power spectrum give an estimate to those cycles that are due to the QBO and are used as \(P_{QBO}(i)\) in the regression model. We then compare the summed QBO component to zonal wind data provided at 10 hPa over Singapore to ensure that the modeled QBO component has the correct degree of phasing, such that there will be a phase shift in the modeled QBO output due to the delay in the transport of air from the tropics (typically 6–18 months Stiller et al., 2008).

### 3.1.2 Applying a trend to anomaly time series after seasonal and QBO components are removed

Examination of the HALOE anomalies (with the season and QBO contributions removed) for the southern mid-latitude bin in Fig. 3, shows little difference in structure for the same bin to that exhibited in Fig. 1. The Northern Hemisphere mid-latitude and tropics bin show similar structures, (not shown here), hence we assume that the turnaround time for HCl to be 1997 for both bins, which also agrees with EESC peak times in this altitude range (Newman et al., 2007; WMO, 2006; Schoeberl et al., 2005). In order to quantify the HALOE anomalies, we would like to try and estimate a linear trend before and after 1997. This can be done by applying the change in linear trend method suggested by Reinsel et al. (2002). Here, a piece-wise linear trend is defined over the total time series period, \(T\), such that there is a change in trend magnitude at a given time, \(t = T_0\) and is described here as

\[
[\Delta_{HALOE\text{HCl}}] = b + a_1 X_{1t} + a_2 X_{2t} + N_t, \quad (2)
\]

where \(\Delta_{HALOE\text{HCl}}(t)\) are the HALOE HCl anomalies to be modeled with seasonal and QBO parameters removed. \(X_t\) is a linear trend function, \(b\) is the intercept at \(t = 0\), while \(a_1\) is the trend at time \(t = 0\), and \(a_2\) signifies the magnitude of a possible change in trend at time \(t = T_0\). Thus

\[
X_{1t} = \begin{cases} \frac{0}{1000000} & 0 < t < T_0 \\ \frac{(t - T_0)}{1000000} & T_0 < t < T \end{cases}
\]

\[
X_{2t} = \begin{cases} \frac{0}{1000000} & 0 < t < T_0 \\ \frac{1}{1000000} & T_0 < t < T \end{cases}
\]

Hence, Eq. (2) can be simplified to

\[
[\Delta_{HALOE\text{HCl}}] = b - a_2 T_0 + t (a_1 + a_2) \quad (3)
\]

\(a_2 = 0\) where \(t < T_0\). From this linear regression model two different trends will be calculated: a trend \((a_1)\) before a turning point and one that implies a change in trend \((a_2)\) after the turning point, such that the new trend after the turning point, \(a = a_1 + a_2\). In order for this method to work, one assumes that there is a change in trend and that the transition is continuous. Hence, for the case in Fig. 3, it is possible to model linear fits to the HALOE anomalies such that there is a change in trend in January 1997 and this methodology is applied to each HALOE anomaly time series from each bin.

The change in trend method is not needed for the ACE-FTS anomalies as the data set started in 2004 and the decline in HCl is approximately linear. However, we can use the ACE-FTS anomalies to extend the HALOE time series, which finishes in 2005. As the HALOE and ACE-FTS anomalies both fluctuate around zero, we need to apply an offset to the ACE-FTS anomalies so that they align with the HALOE anomalies. By using the post 1997 linear fit line calculated from the HALOE anomaly time series, we can apply such an offset to the ACE-FTS anomalies so that they fit relative to the HALOE anomalies. As HALOE terminated in November 2005, it means that there is only 15 months worth of data overlap between ACE-FTS and HALOE. We make an important assumption that the post 1997 HALOE fit line can be extrapolated to cover the whole period covered by ACE-FTS (as shown in Fig. 3). As the ACE-FTS anomalies show no large changes in trend after HALOE’s termination, there is good reason to believe that this assumption is valid. The offset is simply determined by calculating the mean value of the HALOE fit line during the overlapping time of the ACE-FTS anomalies. For the case shown in Fig. 3 the ACE-FTS...
anomalies are offset by $-1.15\%$ (as indicated by the green star in Fig. 3). It should be noted that due to the measurement noise and the natural variability of the atmosphere there will be a residual offset uncertainty, which contributes to the overall correlated error term in the trend uncertainty.

When the ACE-FTS anomaly time series is aligned correctly with the HALOE anomalies, we can then create an all instrument average where both instrument anomaly time series are combined and averaged. Hence, where there is data overlap between ACE-FTS and HALOE an average of the two is calculated. Due to the influence of the ACE-FTS anomalies in the all instrument average, it means that the original estimated trends calculated only for the HALOE time series using Eq. (2) will now need updating. Hence, we make new trend calculations based on the all instrument average produced in each bin, $\Delta_{\text{All Inst}}^{\text{HCl}}(t)$, before and after 1997 using Eq. (2).

3.2 ClO

Similarly to the HCl analysis, we have decided to investigate ClO in the upper stratosphere, from 35 to 45 km, but only for the tropics between $20^\circ$ S to $20^\circ$ N. As MLS makes retrievals on pressure surfaces, we filter data by using approximate pressure surfaces that closely match the geometric altitude zones used here: 6.4–1.6 hPa $\sim 35$–$45$ km. As ClO has a strong diurnal variation, it is important to consider the local solar time (LST) of the measurements of both SMR and MLS. As SMR has an ascending node around 06:00 h and descending node around 18:00 h, we expect the satellite to pass between $20^\circ$ S to $20^\circ$ N somewhere between 05:00–07:00 h and 17:00-19:00 h. A similar principle applies to MLS, which observes ClO in the ascending node from 12:45–14:45 h and descending node from 00:45-02:45 h for this latitude range. Hence, data must be also filtered according to time of day.

On examination of the SMR measurements, it was found that the ascending and descending node LSTs have been slowly drifting forwards since Odin’s launch. The reason for this is because the Odin satellite has slowly descended from its initial position during its mission. Figures 4 and 5 illustrate the effect that this has on the monthly averaged ClO measurements for both dawn and dusk, illustrating the ascending and descending nodes respectively. As a result of increasing LST, it also means that the solar zenith angle (SZA) has slowly decreased over time during morning scans and increased during evening scans. The overall concentrations observed by SMR have therefore slowly increased (morning) or decreased (evening) over time. This causes problems if a trend analysis is to be made using the SMR data set as the induced trend in the measurements from the increased LST will be the dominating feature. For the record, we do not see such problems using MLS for either day or night time measurements since the Aura satellite has orbit stabilization.
the modelling of the HALOE gases are reduced. Moreover, by modelling the same day consequentially 15 times or longer ensures that all the chemical abundances in the model converge, such that subsequent model runs produce insignificant changes for all modeled species.

In this study, we use the box model to simulate diurnal ClO values for 1 km altitude steps from 35–45 km at the equator and during the day of equinox (spring, autumn would produce the same results). Essentially, each ClO diurnal cycle can be expressed as a function of SZA, which in itself is a function of time. The purpose of doing this is to calculate scaling factors, allowing for the binned SMR profiles to be scaled to a specific SZA, \( \theta \). This means for the equivalent satellite SZA, \( \theta_{obs} \), the model values will yield a scale factor, \( sf \), which is then applied to the satellite measurement,

\[
Y_{scl}(\theta_{mod},z) = \frac{Y_{mod}(\theta_{mod},z)}{Y_{mod}(\theta_{obs},z)} Y_{obs}(\theta_{obs},z)
\]

where \( Y_{obs} \) is the observed concentration profile, \( Y_{mod} \) represents the model values, \( Y_{scl} \) is the scaled concentration profile, and \( z \) is the vertical coordinate. For this study we chose \( \theta_{mod} = 90^\circ \) am (i.e. morning). In order to make a fair comparison, we also treat the MLS data in the same manner, such that both data sets contain profiles corrected for diurnal variation (according to Eq. 4) from 35 to 45 km that can be used to produce monthly means based solely for when \( \theta_{mod} = 90^\circ \) am. More importantly, it removes the LST problem concerning the SMR measurements and time of day no longer is an issue. A graphical illustration of the scaling process is given in Fig. 6.

When making a trend analysis the absolute values of the scaled monthly means are irrelevant as we only consider relative units by firstly removing the seasonal cycle (similar to that of the HCl method). A linear regression analysis is made (according to Eq. 1) on each set of scaled deseasonalised anomalies. We do not fit for the solar cycle since the time series length of each data set is too short. Mean ClO changes due to 11-year solar forcing are typically 0–2 % between 35 and 45 km (Egorova et al., 2005). Thus the remaining scaled anomalies will have the QBO and seasonal cycles removed, but will still include the solar cycle contribution. We apply an offset to the MLS anomalies so that they fit relative to the SMR anomalies. An all instrument average is created by combining the overlapping MLS (after applied offset) and SMR anomalies. Finally, a linear trend line is fitted to the all instrument average.

For reference, we have also analysed measurement scale factors based on other times of the year, including the winter and summer solstices. It means that the observations are scaled differently owing to different conditions compared to those at equinox. Such conditions are related to parameters like solar intensity (as a result of the alternating distance from the Earth to the Sun during an annual cycle) and variations in the maximum SZA during an annual cycle. However, we do not find any significant differences in the overall trend magnitudes using simulated diurnal cycles from the winter and summer solstice cases, therefore the choice of using a ClO diurnal cycle for equinox is justified.

4 Results

4.1 HCl

Figure 7 shows the all instrument average for combined HALOE and ACE-FTS anomalies in the three bins analysed. Each bin shows an increase in HCl values monotonically until around 1997. Values slowly start to decrease after 1997 and by mid 2008 they are typically 5–6 % lower than the 1997 turn around anomaly estimates.

Concerning the mid-latitudes, we find that in each bin HCl values increase by 27 % decade\(^{-1}\) in the Northern Hemisphere and 32.4 % decade\(^{-1}\) in the Southern Hemisphere until 1997. In each case the trend magnitudes are statistically significant from a zero trend at 2\(\sigma\) (or at 95 % confidence level). Error estimates are calculated using an algorithm suggested by Weatherhead et al. (1998). The trends calculated here are larger than those found by Newchurch et al. (2003), who examined HCl for the same altitude region using HALOE data from 1993–2003. This could be a
result of the “change in linear trend method”. Newchurch et al. (2003) only apply a linear trend from 1993 until 1997, thus they only consider the anomalies during this time, while the change in linear trend method looks for a specific temporal path through the whole data set. The difference being in this case that the method used here produces a less steep fit to the data in comparison to the Newchurch et al. (2003) method. Trend values after 1997 all show consistent declines in HCl: −5.2 % ± 2.2 % decade\(^{-1}\) in the southern mid-latitudes and −5.1 % ± 2.1 % decade\(^{-1}\) in the northern hemispheric mid-latitudes. These values are all statistically significant from a zero trend at 2\(\sigma\). It is also seen that the sudden increase in values after 2001 (most apparent in 30° S–50° S) reported in Fig. 1 is still present, implying that according to this analysis it is neither a related feature of seasonal or QBO behaviour. This feature may be related to possible anomalous atmospheric dynamics in the Southern Hemisphere, such as the Antarctic ozone hole split in 2002 (Manney et al., 2005; Charlton et al., 2005) and/or changes in tropical upwelling due to enhanced wave driving in 2000–2001 (Dhomse et al., 2008). The tropics bin from 20° S–20° N shows a more negative trend of 5.8 % decade\(^{-1}\) since 1997 compared to the mid-latitudes. However, we have not removed the QBO contribution from HALOE or ACE time series, suggesting the calculated trend could be different if the QBO contribution is removed.

4.2 ClO

A summary of the scaled SMR and MLS measurements is presented in Fig. 8. The upper panel illustrates AM and PM scaled monthly means for both SMR and MLS. It can be seen that there is an excellent agreement between the average SMR AM and PM scaled values. There is however about a 0.1–0.2 ppbv difference between MLS AM and PM scaled values. The MLS AM measurements are made typically when the LST is between 01:00 h and 03:00 h, a time when ClO concentrations are close to zero ppbv below 40 km. As a result, small systematic errors of the measurements may be amplified by the diurnal correction factors, yielding in this case unrealistically high scaled measurement values. The end result is a larger layer average. Due to this problem, we have decided to ignore the MLS night time measurements, thus we only consider MLS PM observations for the construction of the all instrument average (note, that the offset is applied to the MLS PM anomalies so that they fit relative to the SMR PM anomalies). The bottom panel of Fig. 8 shows the SMR scaled monthly means (combination of AM and PM) as well as the MLS PM scaled monthly means. There is a good agreement between the two scaled data sets, where each individual time series shows evidence for structural components due to natural variability, including the seasonal and QBO cycles. Differences between MLS and SMR are typically no larger than ~0.03 ppbv and there is also clear evidence that the LST trend in the SMR measurements is removed.

Examination of the ClO all instrument average based on the anomalies, with the seasonal and QBO cycles removed, is shown in Fig. 9. The top panel shows the SMR contribution to the all instrument average, while the bottom panel considers MLS. The trend calculated for the 2001–2008 period from both the combined data sets is −7.1 % ± 7.8 % decade\(^{-1}\), implying that it is not statistically significant from a zero trend at 2\(\sigma\).

5 Discussion of HCl and ClO results

The ClO trend value obtained here is considerably less negative than the analysis performed by Solomon et al. (2006), who used ground based millimetre wave measurements at Mauna Kea in Hawaii, USA, to observe ClO concentrations. They found a significant linear decline of ClO of −15 % ± 2 %/decade from 1995–2005 for altitudes between 35 and 39 km, although they estimate that about 30 % of this decline can be attributed to methane (CH\(_4\)) increments (a sink of Cl) in this altitude range, which have been measured by HALOE. A recent update of this analysis, where the time series was extended to 2007, shows a ClO trend of −9 % ± 2 % decade\(^{-1}\) (WMO, 2011). Our values are more consistent with the analysis by O’Doherty et al. (2004), who estimate that tropospheric Cl\(_2\) is decreasing between 6 and
Newchurch et al. (2003) examined HALOE observations from 30° S–30° N, but as HALOE had a sparse coverage due to the temporal progression of the scans as the satellite orbited from pole to pole, it means the total global coverage varied strongly from year-to-year. This implies that a bin covering a 60° span over the tropical zone is perhaps too large and there will be strong influences from different chemical and dynamical regimes. In order to test how the HALOE HCl behaves in the tropics, we have examined HALOE residuals in 5° latitude bands spanning 30° S–30° N. The HALOE residuals examined have only the seasonal component removed. As there is a sparse amount of data in each five degree bin, estimating a QBO component could be challenging, hence the QBO component is retained in the residuals. Using the analogy of Eq. (2), we apply the Reinsel et al. (2002) fitting technique to the HALOE residuals from January 1993 to November 2005 with a turnaround of January 1997. Table 1 shows the estimated trends before and after the turnaround date for the five degree latitude bins and four extra larger bins for a fuller comparison: 10° S–10° N, 15° S–15° N, 20° S–20° N, and 30° S–30° N.

From Table 1, it can be seen that the range of HCl trend values from each five degree latitude bin after 1997 are in the range of −4 % to −6 % decade$^{-1}$. It is also apparent that HCl increases at a higher rate prior to 1997 as one moves away from the pole, resulting in the QBO component being more clearly defined. The QBO component is retained in the residuals. Using the analogy of Eq. (2), we apply the Reinsel et al. (2002) fitting technique to the HALOE residuals from January 1993 to November 2005 with a turnaround of January 1997. Table 1 shows the estimated trends before and after the turnaround date for the five degree latitude bins and four extra larger bins for a fuller comparison: 10° S–10° N, 15° S–15° N, 20° S–20° N, and 30° S–30° N.

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From Table 1, it can be seen that the range of HCl trend values from each five degree latitude bin after 1997 are in the range of −4 % to −6 % decade$^{-1}$. It is also apparent that HCl increases at a higher rate prior to 1997 as one moves away from the equator. This is possibly due to a combination of stronger wave dynamics, which rapidly transport the tropical air towards the mid-latitudes. Moreover, the production of HCl is greater outside the tropics as CFCs are more rapidly broken down here due to longer residence times at the higher stratospheric altitudes. Consequently, HCl is destroyed faster as one moves away from the pole, resulting in the QBO component being more clearly defined. The QBO component is retained in the residuals. Using the analogy of Eq. (2), we apply the Reinsel et al. (2002) fitting technique to the HALOE residuals from January 1993 to November 2005 with a turnaround of January 1997. Table 1 shows the estimated trends before and after the turnaround date for the five degree latitude bins and four extra larger bins for a fuller comparison: 10° S–10° N, 15° S–15° N, 20° S–20° N, and 30° S–30° N.
from a peak and decline in CFC concentrations after 1997. Examination of the trends for the four larger bins (10° S–10° N, 15° S–15° N, 20° S–20° N, and 30° S–30° N) shows that the differences in trend magnitude between each bin is relatively small, indicating that the use of larger bins is acceptable, although caution is needed. We chose to use 20° S–20° N for representing the tropics in Fig. 7 so as to stay consistent with the ClO analysis. It should be noted that the trends for 20° S–20° N in Fig. 7 are slightly different to those in Table 1. This is simply because the ACE-FTS data has now been added to the all instrument average, which alters the time series properties when the Reinsel fit is made. The all instrument average for the tropics (and the tropical trend estimates from Table 1) are not dissimilar from the mid-latitude all instrument average trends shown in Fig. 7, although the QBO contributions are removed from the mid-latitude calculations and not for the tropics. It suggests that since 2005 until present, HCl has continued to decrease at a similar rate. Solomon et al. (2006) also investigated HALOE HCl between 35 and 39 km for 10° N–30° N, where they applied a fit to the HALOE data from 1997–2005, producing a HCl linear trend of $-5 \pm 1.2 \text{ % decade}^{-1}$, which is similar to the all instrument average values found in this study as well as the HALOE values from Table 1. Simulated CMAM monthly averaged HCl values show that from 1997 to 2008 HCl decreases by $-3.1 \text{ % decade}^{-1}$, for 35–45 km at northern mid-latitudes and $-4.1 \text{ % decade}^{-1}$ at Southern Hemisphere mid-latitudes, values which are slightly less than our estimated trend values. The difference in trend values can be attributed to differences in the choice of peak loading times of HCl assumed here and to that found in the CMAM model, where CMAM HCl peaks typically around 1999 and not 1997. Recent trend analyses concerning the upper stratosphere (~55 km) have been performed, using various satellite and ground-based data. Aura-MLS and ACE-FTS trend values show a decline in HCl to be $-6 \pm 1 \text{ % decade}^{-1}$ and $-9 \pm 2 \text{ % decade}^{-1}$ respectively, since 2004. Moreover, ground-based FTIR total column measurements of HCl at Jungfraujoch and Kiruna show similar trends from 1996–2009: $-9 \pm 1 \text{ % decade}^{-1}$ and $-8 \pm 2 \text{ % decade}^{-1}$ respectively (WMO, 2011). These reported trends are larger in magnitude compared to here. However these differences can be possibly explained due to contrasts in HCl abundances with respect to altitude. HCl is estimated to account for >95% of the total chlorine above 50 km compared to ~80–90% between 35 and 45 km (WMO, 2006). Moreover, there are different chemical and dynamical processes controlling the evolution of HCl throughout the stratosphere, hence we do not necessarily expect to see similar trend magnitudes.

Linear trend model uncertainties are mainly determined by three factors: the time series length, the autocorrelation of the residuals, and variance of the residuals factors (Weatherhead et al., 1998; Reinsel et al., 2002). In this analysis, we find the ClO all instrument average anomalies to have an autocorrelation of 0.51, which is moderate (the degree of correlation increases towards 1 or −1), while the standard deviation is 4.4%. The ClO anomalies of course still have variations due to the solar cycle included, which will account for some of the variance and autocorrelation in the modeled values. The variation of tropical ClO due to changes in solar intensity is estimated to be as much as 2% between 35 and 40 km, but it is almost negligible between 40 and 45 km according to model simulations (Egorova et al., 2005). The total length of the time series is also quite short, which means it is hard to differentiate between long and short term variations. Inevitably, the regression parameter errors will be reduced when the time series is extended (assuming that the other two influencing parameters remain constant). A longer time series will also mean that a reasonable solar cycle estimate can be made to help further reduce some of the uncertainty.

Ultimately, the apparent decline in chlorine compounds should mean an anti-correlation with ozone. Recent ozone trends calculated for 1997 onwards in this altitude region have shown signs of a small recovery (but not significantly different from a zero trend), although the magnitudes found are typically smaller than those of the chlorine species reported here. Jones et al. (2009) reported that ozone is increasing at $\sim 1.7 \% \left(\pm 2.0 \text{ % decade}^{-1}\right)$ at mid-latitudes based on linear trends from 1997–2008, while Steinbrecht et al. (2006) suggest a 1997–2005 trend for Hawaii to be increasing by $1.9 \pm 1.9 \text{ % decade}^{-1}$. Both of these examples show trend magnitudes to be less than the trend values estimated here using HCl and ClO. This implies that ozone’s recovery is also dependent on other atmospheric parameters and not just entirely on chlorine chemistry. For example, ozone is influenced by cycles involving, ClO, BrO, NOx, and HOx and thus depends on changes of the source gases of these families (for example, H2O and CH4 for the HOx family) (WMO, 2006).

6 Summary

We have analysed the long-term evolution of stratospheric HCl, using combined HALOE and ACE-FTS HCl measurements from 1993–2008. Furthermore, we have estimated a trend in stratospheric ClO, using measurements from the Aura-MLS and Odin/SMR over a shorter time period, 2001–2008. The analyses shown here utilise monthly averaged data in the 35–45 km altitude region for two mid-latitude bands (30° S–50° S and 30° N–50° N) for HCl, while ClO and HCl measurements are examined for the tropics (20° S–20° N).

We find that the SMR ascending and descending node LSTs are monotonically increasing in time since its launch. This has resulted in ClO measurements being significantly influenced by lower SZAs during morning scans and higher SZAs during evening scans. To remove the LST trend in the data, a 1-D atmospheric box model is used to determine scale factors, which are applied to the measurements so that they will represent a value when the SZA equals 90° during
sunrise. The same method is applied to the MLS data so that a comparison can be made. However, the computed scale factors for MLS night profiles are overestimated due to the very low night time ClO abundances below 40 km. As a result, only the MLS PM scaled values are considered in the trend analysis. SMR scaled values show consistency between AM and PM profiles, thus both sets of scaled values are used.

Anomalies of the monthly averages are calculated by removing the seasonal and QBO cycles. However, due to the sparse ACE-FTS HCl data around the tropics, we only remove the seasonal cycle in the tropics bin. The anomalies are then concatenated in each respective bin to produce all instrument averages, which are used for the trend analysis. HCl trend estimates after 1997 are \(-5.1 \pm 2.1\) % decade\(^{-1}\) (northern mid-latitudes), whilst \(-5.2 \pm 2.2\) % decade\(^{-1}\) (southern mid-latitudes) in the upper stratosphere, and \(-5.8 \pm 2.2\) % decade\(^{-1}\) in the tropics. The modeling uncertainties indicate that the estimated trends are significantly different from a zero trend at 2\(\sigma\). ClO trend estimates show slightly larger trend magnitudes since 2001, although not statistically significant at 2\(\sigma\) (7.1 \pm 7.8) % decade\(^{-1}\). It is expected that in the future uncertainties of the ClO analysis can be reduced by employing a longer time series. It will also allow for the possibility to estimate the contribution of the solar cycle. This shows the importance of time series length. The fact that SMR has only lived its 2 year expected life time is a bonus as it has provided one of the longest ClO data sets so far, hence it is important that the observations continue to be made until the satellite finally ceases to naturally operate. This is also certainly applicable for ACE-FTS, MLS, and other recent satellite instruments.

The results found here are in agreement with and extend on previous findings (for example, Considine et al., 1997, 1999; Anderson et al., 2000; Newchurch et al., 2003; Froidevaux et al., 2006; Solomon et al., 2006; WMO, 2006). Although the ClO trend is not statistically significant, it does imply a reduction over the past 7 years, while the HCl trend shows a clear statistically significant downward trend. These results illustrate that the Montreal protocol and the subsequent amendments are having an impact on the total chlorine concentration in the stratosphere and ultimately the recovery of ozone.

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