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Chlorinated very short-lived substances offset the long-term reduction of inorganic stratospheric chlorine



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Over the past few decades, a reduction in chlorinated long-lived ozone-depleting substance emissions due to the regulations imposed by the Montreal Protocol has led to a global decrease in stratospheric chlorine. At the same time, emissions of chlorinated Very Short-Lived Substances, which are unregulated, have increased. Here we show that observed changes of inorganic stratospheric chlorine are inconsistent with changes in the tropospheric abundances of long-lived ozone-depleting substances. Satellite observations of stratospheric chlorine species from the Atmospheric Chemistry Experiment—Fourier Transform Spectrometer during 2004–2020 reveal that the observed decrease in inorganic stratospheric chlorine is 25%–30% smaller than expected based on trends of long-lived ozone-depleting substances alone. At mid-latitudes in the lower stratosphere, this can be explained by the chlorinated Very Short-Lived Substances increase, which offsets the long-term reduction of stratospheric chlorine by up to 30%.

The stratospheric ozone layer protects life on Earth by absorbing harmful solar ultraviolet (UV) radiation. The discovery of ozone (O₃) destruction by halogen-containing Ozone-Depleting Substances (ODSs), such as chlorofluorocarbons (CFCs), in the 1980s was thus of major concern [e.g. refs. 1,2]. The 1987 Montreal Protocol and its subsequent amendments and adjustments successfully reduced anthropogenic emissions of long-lived ODSs³, leading to 21st century increases in stratospheric O₃ concentrations in the middle and upper stratosphere [e.g. refs. 4–6]. However, O₃ depletion can also be caused by Very Short-Lived Substances (VSLs), which are not regulated by the Montreal protocol [e.g. refs. 7,8]. VSLs are trace gases that have lifetimes of less than 6 months and less uniform tropospheric concentrations compared to longer-lived species such as CFCs³. Brominated and iodinated VSLs have mostly natural sources, such as algae, seaweed, and phytoplankton [e.g. refs. 9–11], but the most important chlorinated VSLs (Cl-VSLs, such as dichloromethane (CH₂Cl₂), chloroform (CHCl₃) and 1,2-dichloroethane (C₂H₄Cl₂)) are mainly emitted by anthropogenic activities [e.g. refs. 12,13]. The contribution of Cl-VSLs to the total inorganic stratospheric chlorine (Cl_y) is small, but growing, as emissions of chlorinated long-lived ODSs decrease and emissions of Cl-VSLs increase^{14,15}. The global mean total

tropospheric Cl-VSL abundance increased from 103 ppt in 2016 to about 113 ppt in 2020, a rate of 2.5 ppt/year³. This can impact the rate of O₃ recovery, potentially delaying it to a later date than projected by models that neglect VSLs^{16,17}.

Previous studies have calculated the Cl-VSL trend in the upper troposphere/lower stratosphere (UTLS) using a Chemical Transport Model (CTM)^{14,18} or a Chemistry-Climate Model (CCM)¹⁹ constrained by observed surface abundances of Cl-VSLs. Using a CTM resulted in a Cl-VSL trend at the tropical tropopause of 3.8 ± 0.3 ppt/year for 2004–2017¹⁴, and using a CCM resulted in similar Cl-VSL trend of 3.7 ± 0.5 for the same period¹⁹. Cl-VSL measurements in the UTLS are available from aircraft campaigns [e.g. refs. 20,21], and there is reasonable agreement between modelled and observed Cl-VSL concentrations near 17 km in the tropics between 2004 and 2016¹⁴. At higher altitudes in the stratosphere it is only possible to measure the chlorinated product gases, such as HCl (also formed from long-lived ODSs), and not the Cl-VSLs themselves, which are no longer present in detectable amounts at these altitudes. Modelled stratospheric HCl trends were shown to agree better with HCl trends in observations from the Atmospheric Chemistry Experiment—Fourier Transform Spectrometer (ACE-FTS)²² when Cl-VSLs were included in the

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Fig. 1 | Cl-EESC for 3-year-old air (red) and 5.5-year-old air (blue), calculated following the method of ref. 23. Trends are from a simple linear fit (dashed black lines) to the Cl-EESC for 2004 to 2020.

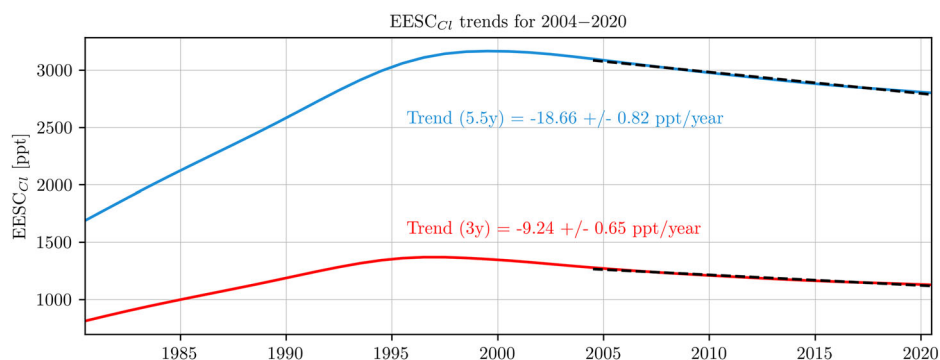
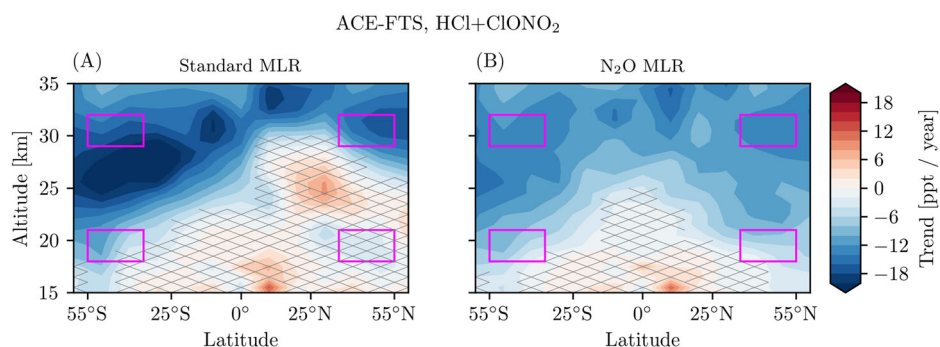


Fig. 2 | Trends in HCl+ClONO₂ from ACE-FTS as a function of latitude and altitude. Trends are for 2004–2 to 2020–12. The magenta boxes mark the approximate regions of 3- and 5.5-year-old air at mid-latitudes (35°–55° in each hemisphere; older air is at higher altitudes for a fixed latitude). The height of each box is 3 km, corresponding to the vertical resolution of the AoA observations. **A** Trends calculated with a standard MLR. **B** Trends calculated with the N₂O MLR, which uses N₂O as a proxy for dynamical variability. Hatching denotes statistically insignificant trends at the 2σ level.



simulations^{14,19}, but the specific impact of Cl-VSLs on stratospheric Cl_y trends was not quantified.

Here we use ACE-FTS observations of HCl and ClONO₂ to determine the stratospheric Cl_y trend at mid-latitudes for 2004–2020. The observed Cl_y trend is derived by scaling the ACE-FTS HCl+ClONO₂ trend by the modelled ratio of the Cl_y trend to the HCl+ClONO₂ trend. The resulting ACE-FTS Cl_y trends are shown to be inconsistent with the trend in the chlorine component of the Equivalent Effective Stratospheric Chlorine (Cl-EESC)²³, which is used as a proxy for the trend in chlorinated long-lived ODSs. The stratospheric Cl_y trends are 25% – 30% smaller than expected based only on decreases in long-lived ODSs. Cl-VSLs can explain most of this difference. In particular, for younger air, the combined Cl-VSLs+Cl-EESC trend is equivalent to the observed stratospheric Cl_y trend in both hemispheres.

Results

Trends in chlorinated long-lived ODSs

We use the trend in the chlorine component of EESC to represent the trend in long-lived chlorinated ODSs. EESC is a metric for the combined effects of long-lived chlorinated and brominated ODSs on the stratosphere, based on abundances of the tropospheric source gases^{23–25}. EESC is derived by propagating tropospheric ODS abundances into the stratosphere according to the age spectrum described by a Green's function for air with a given mean age²³. Typically 3 and 5.5 years are used for the mean ages, chosen to roughly correspond to the lower stratosphere at mid-latitudes and polar latitudes, respectively²⁵. The age spectra are further modified to account for chemical loss, in addition to transport. Details of the EESC calculation are provided in ref. 23. We use the same method here to calculate the chlorine contribution to the total EESC: the Cl-EESC. The tropospheric abundances of ODS used in the Cl-EESC calculation are from the WMO 2022 Ozone Assessment³, based primarily on measurements from the National Oceanic and Atmospheric Administration (NOAA) [e.g. ref. 26] and the Advanced Global Atmospheric Gases Experiment (AGAGE) [e.g. ref. 27].

Figure 1 shows the resulting Cl-EESC from 1980 to 2020 for 3- and 5.5-year-old air. Since ~2000, the Cl-EESC mixing ratio has decreased at a

steady rate due to the phase-out of long-lived ODSs, so the trend in the Cl-EESC can be approximated using a simple linear regression. The trend is calculated for 2004–2020: the start year is limited by the ACE-FTS record, and the end year is limited by the availability of ODS abundance measurements that are needed for the Cl-EESC calculation. The dashed black lines in Fig. 1 show the resulting fit for 3- and 5.5-year-old air. The corresponding trends for 2004–2020 are -9.24 ± 0.65 ppt/year for 3-year-old air and -18.66 ± 0.82 ppt/year for 5-year-old air.

Trends in inorganic stratospheric chlorine

For our initial assessment, we use the sum of ACE-FTS HCl observations and ClONO₂ observations to represent the stratospheric inorganic chlorine reservoir. Trends in stratospheric trace gas concentrations are typically calculated using a multiple linear regression (MLR) model that includes proxies for phenomena that are known to contribute to the observed variability, such as the solar cycle and the quasi-biennial oscillation (QBO). By applying this standard MLR to monthly zonal mean (MZM) ACE-FTS observations of HCl+ClONO₂ we find a distinct hemispheric asymmetry in the trends for 2004–2020: below 30 km the inorganic stratospheric chlorine decreased in the SH by up to 20 ppt/year and increased by an insignificant amount in the NH (Fig. 2A). This pattern was observed previously in HCl trends^{16,28–30} and it was attributed to a slowdown of the Brewer-Dobson circulation (BDC) in the NH relative to the SH²⁸. We would expect to see HCl decrease in both hemispheres if the trend was only due to surface emissions of chlorine-containing substances. The HCl+ClONO₂ trends are therefore caused by the combined effect of transport and emission changes. Using the sum of HCl and ClONO₂ also ensures that the trends are not significantly impacted by the 2020 Australian wildfires, which occurred near the end of the time period used for the trend calculation. The associated anomalies in both gases following the wildfires compensate for each other: HCl decreased and ClONO₂ increased³¹.

To determine the trend solely due to changes in chlorine-containing substance emissions, it is necessary to account for BDC changes in the regression. This can be done using N₂O observations as a proxy for dynamical variability³². N₂O is a long-lived trace gas with a tropospheric source

and a known surface trend, so it provides a good representation of transport anomalies throughout the stratosphere. Later work further demonstrated that using ACE-FTS N_2O observations as a proxy for dynamical variability could explain nearly all of the inter-hemispheric asymmetry in ACE-FTS HCl trends over 2004–2018³⁰. Fig. 2B shows the trend in ACE-FTS HCl+ClONO₂ from this version of the regression, which we call the N_2O MLR (see data and methods section for details). Both hemispheres have a significant negative HCl+ClONO₂ trend, which suggests that hemispherically asymmetric changes in the BDC were properly controlled for in this version of the trend calculation.

To compare the ACE-FTS HCl+ClONO₂ trends to the Cl-EESC trends, it is necessary to have the HCl+ClONO₂ trends for air of the same mean ages (3 and 5.5 years). The ACE-FTS MZM age of air profiles³³ are used to determine the mean altitude of each age in four 10° latitude bins centred at 50°S, 40°S, 40°N, and 50°N, which we refer to as mid-latitudes. The focus is on mid-latitudes as the fraction of the source gases that has been converted to Cl_y in the tropics is low, and the polar vortex regions are highly variable. Mid-latitude Cl_y trends are also a more representative measure of stratospheric chlorine trends than tropical Cl_y trends since many more transport pathways are integrated into mid-latitude air masses.

The ACE-FTS HCl and ClONO₂ are interpolated to the mean altitudes of 3- and 5.5-year-old air at mid-latitudes (19.0 km and 29.5 km, respectively), and the trends are then calculated with the N_2O MLR. The resulting HCl+ClONO₂ trends are nearly equivalent at all four latitudes for 5.5-year-

old air (Fig. 3). There are some small differences between the trends at each latitude for 3-year-old air, with the values varying from -7.63 ± 1.82 ppt/year to -5.34 ± 1.39 ppt/year. This suggests that while the N_2O MLR can significantly reduce inter-hemispheric differences in the HCl+ClONO₂ caused by BDC changes compared to the standard MLR, there are still some small differences that the method does not account for at lower altitudes.

The mean HCl+ClONO₂ trends in each hemisphere are used as estimates of the overall mid-latitude NH and SH chlorine trends for 3- and 5.5-year-old air. The resulting NH HCl+ClONO₂ trend is -6.97 ± 1.72 ppt/year for 3-year-old air and -13.04 ± 1.80 ppt/year for 5.5-year-old air. The SH HCl+ClONO₂ trend is -6.42 ± 1.59 ppt/year for 3-year-old air and -13.03 ± 1.88 ppt/year for 5.5-year-old air. The absolute HCl+ClONO₂ decrease is greater for older air because the total HCl+ClONO₂ abundance is larger: in older air, there has been more time for chlorinated source gases to be converted to stratospheric inorganic chlorine reservoirs. This can be seen in Supplementary Fig. 1: the mean HCl+ClONO₂ trend is -0.4% /year for both 3- and 5.5-year-old air.

To assess how well the HCl+ClONO₂ trend captures the total Cl_y trend, we use results from the Canadian Middle Atmosphere Model (CMAM)^{34,35}. Fig. 4A shows the trend in CMAM HCl+ClONO₂, and Fig. 4B shows the trend in CMAM Cl_y . The trends in the two cases are very similar below ~ 25 km, but the Cl_y decrease is greater than the HCl+ClONO₂ decrease at higher altitudes. The ratio of the Cl_y trend to the HCl+ClONO₂ trend is given in Fig. 4C: the ratio is close to 1 at lower altitudes, and reaches up to ~ 1.15 at 35 km. This shows that the HCl+ClONO₂ trend can fully explain the Cl_y trend at lower altitudes, but is missing part of the Cl_y trend at higher altitudes, in particular contributions from ClO. It should also be noted that the inconsistency between the HCl+ClONO₂ trend from CMAM in Fig. 4 and from ACE-FTS in Fig. 2 is not important, as we are only using the model results to get the relationship between HCl+ClONO₂ and Cl_y , not the actual trend values, and this relationship largely depends on the model chemistry.

The ACE-FTS HCl+ClONO₂ trends are converted to Cl_y trends using a scale factor from CMAM. At mid-latitudes, the Cl_y /HCl+ClONO₂ trend ratio is 1.02 for 3-year-old air and 1.05 for 5.5-year-old air. Multiplying the ACE-FTS HCl+ClONO₂ trends by this ratio gives an ACE-FTS Cl_y trend in the NH of -6.53 ± 1.59 ppt/year for 3-year-old air and -13.97 ± 1.93 ppt/year for 5.5-year-old air during 2004–2020. The corresponding Cl_y trends in the SH are -6.59 ± 1.63 ppt/year for 3-year-old air and -13.50 ± 1.95 ppt/year for 5.5-year-old air.

To test the ability of CMAM to accurately represent the ratio of Cl_y to HCl+ClONO₂ trends, we also considered scale factors from six other models using runs that follow the same REF1 scenario as the CMAM runs. Five of the models have realistic Cl_y amounts and partitioning and can be compared to CMAM. Scaling factors from all of these models are very similar, which suggests that CMAM provides an accurate representation of the chlorine partitioning in these regions. Further discussion and the scaling factors for each model are provided in the Supplementary Information document.

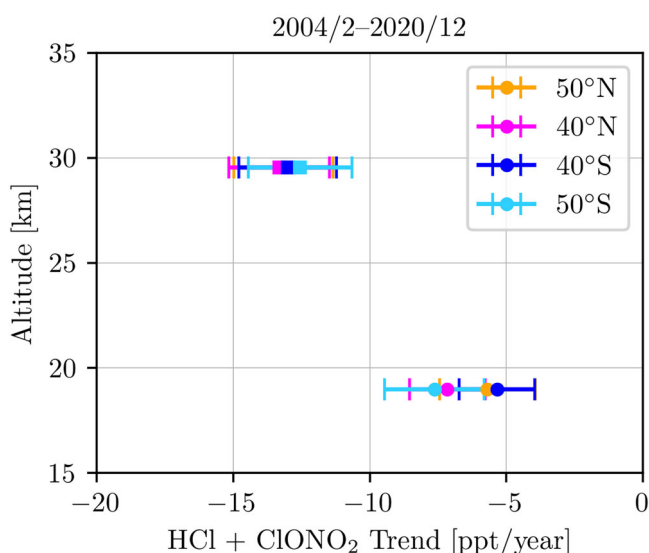
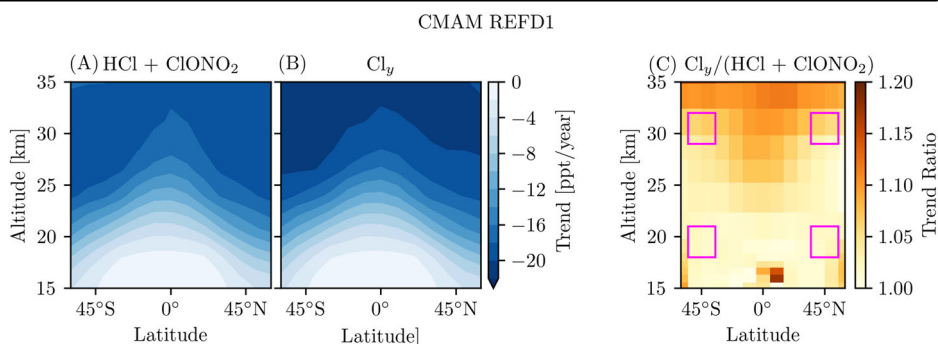


Fig. 3 | Trends in HCl+ClONO₂ from ACE-FTS at mid-latitudes and the mean altitudes of 3-year-old air (circles) and 5.5-year-old air (squares). Trends are for 2004–2 to 2020–12. Trends are calculated with the N_2O MLR, and error bars are the 2σ uncertainty in the calculated trends.

Fig. 4 | Trends for 2004–1 to 2018–12 in the mean of 5 ensemble members from the CMAM REF1 simulations. **A** Trends in HCl+ClONO₂. **B** Trends in Cl_y . **C** The ratio of the trends, Cl_y /(HCl+ClONO₂). The magenta boxes mark the approximate regions corresponding to 3- and 5.5-year-old air at mid-latitudes. The height of each box is 3 km, corresponding to the vertical resolution of the AoA observations. Trends are calculated with the N_2O MLR.



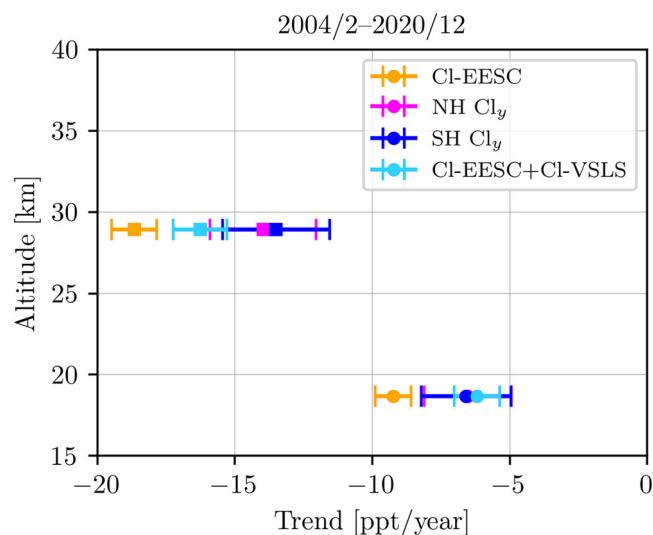


Fig. 5 | Trends in NH and SH Cl_y , CI-EESC, and CI-VSLs at mid-latitudes and the mean altitudes of 3-year-old air (circles) and 5.5-year-old air (squares). Trends are for 2004-2 to 2020-12. Error bars are the 2σ uncertainty in the calculated trends.

Discussion and conclusion

The recovery of the stratospheric ozone layer depends upon a reduction in anthropogenically emitted halocarbons. While atmospheric abundance of long-lived chlorinated ODSs have indeed decreased over the past two decades due to regulations imposed by the Montreal Protocol, stratospheric concentrations of Cl_y have not gone down at a similar rate: if long-lived chlorinated ODSs were the only source of stratospheric chlorine, we would expect the trend in the inorganic stratospheric chlorine reservoir to be the same. However, we find that the observed decrease in Cl_y is 25–30% smaller than the CI-EESC trend (Fig. 5). This difference cannot be explained by the uncertainties in the CI-EESC and Cl_y trend calculations.

To see if the discrepancy between long-lived ODS trends and total stratospheric chlorine trends can be explained by rising emissions in unregulated CI-VSLs, we also compare the Cl_y trends to the combined trend in CI-EESC+CI-VSLs. The CI-VSLs trends that we use are 3.0 ± 0.5 ppt/year for 3-year-old air and 2.4 ± 0.5 ppt/year for 5.5-year-old air. The CI-VSLs trends are from simulations done with the Met Office's Unified Model coupled to the United Kingdom Chemistry and Aerosol (UM-UKCA)³⁶, which are constrained to observed surface values of CI-VSLs¹⁹. The combined CI-EESC+CI-VSLs trend agrees with the stratospheric Cl_y trend for 3-year-old air in both hemispheres, showing the CI-VSLs have a significant impact on stratospheric chlorine in the lower stratosphere: CI-VSLs offset the decrease in stratospheric chlorine loading by ~30% in this region (Fig. 5). For 5.5-year-old air the agreement is less strong, but the combined CI-EESC+CI-VSLs trend is nonetheless more similar to the total stratospheric Cl_y trend than the CI-EESC trend is on its own, particularly in the NH.

The fact that our Cl_y trend estimates based on stratospheric satellite measurements are consistent with the CI-EESC + CI-VSLs trend estimates based on modelling combined with ground-based observations provides convincing evidence that stratospheric chlorine trends can only be explained if all sources of chlorine are taken into account. It should be noted that the presented uncertainties are based on the trend-calculation method and describe unexplained variations of the quantities over time. Other sources of uncertainty, such as the potential impact of satellite sampling patterns on zonal mean stratospheric chlorine estimates, are not taken into account, but could bias the trends^{37,38}. Furthermore, transport variations that would impact N_2O differently than HCl or ClONO_2 will not be removed from the chlorine trends by our N_2O proxy method. However, the very good agreement between the Cl_y trends in the NH and the SH in Fig. 3 suggests that our trend-calculation method successfully removes the dominant first-

order effects of the well-known hemispheric asymmetries of the circulation changes. Finally, CI-VSLs trend estimates can be impacted by uncertainties in the observed circulation and its trends, alongside its representation in the different reanalysis products and models¹⁹, and uncertainties in the spatial distribution of VSLs emissions^{13,39} and their interactions with the atmospheric transport pathways that may not be adequately represented by using latitudinally uniform lower boundary conditions.

Overall, our results show that rising CI-VSLs levels in the stratosphere have a significant impact on the total inorganic stratospheric chlorine concentrations. CI-VSLs increases are causing stratospheric Cl_y to decrease at a slower rate than anticipated based on reductions in long-lived ODS emissions. These findings are important to consider when evaluating the progress of the Montreal Protocol and predicting the ozone recovery date.

Data and methods

Observations and model results

Observations from ACE-FTS^{22,40} are used to determine HCl and ClONO_2 trends, as well as the altitude corresponding to a given mean age. We use HCl, ClONO_2 , and N_2O profiles from version 5.2 of the ACE-FTS retrieval⁴¹. All trace gas profiles are filtered using the data quality flags developed by ref. 42 before any further analysis is performed. Trends are calculated in the area-weighted monthly zonal mean ACE-FTS observations for 10° latitude bins. The MZM age of air derived from v3.6 of the ACE-FTS SF_6 retrieval³³ is used to determine the altitude level corresponding to a given mean age. SF_6 was used to derive the AoA because it has a long chemical lifetime in the stratosphere and a well-defined tropospheric trend.

The relationship between HCl+ ClONO_2 changes and Cl_y changes is determined using model simulations from CMAM^{34,35}. CMAM is a vertically extended version of the Canadian Centre for Climate Modelling and Analysis third-generation atmospheric general circulation model [AGCM3, ref. 43] that also includes interactive stratospheric chemistry. The CMAM results used here follow the REF1 scenario, which includes forcing from observed sea surface temperatures, greenhouse gases, ozone-depleting substances, and volcanic aerosol⁴⁴. Results from five ensemble members, each with slightly different initial conditions, are considered. The CMAM configuration does not include VSLs. Including VSLs is not expected to change the scale factors, as the impact of VSLs would affect the Cl_y and HCl + ClONO_2 trends in a consistent way.

The values for the CI-VSLs trends are based on simulations from the UM-UKCA³⁶ chemistry-climate model run in a specified-dynamics configuration¹⁹, with temperatures and zonal and meridional winds nudged to European Centre for Medium-Range Weather Forecasts (ECMWF) ERA5 reanalysis⁴⁵. The model configuration includes the explicit treatment of four of the most important CI-VSLs: CH_2Cl_2 , CHCl_3 , C_2Cl_4 , and $\text{C}_2\text{H}_4\text{Cl}_2$ ⁴⁶. The modelled CI-VSLs are constrained at the surface by annual mean observations in five latitude bands. The CH_2Cl_2 and C_2Cl_4 surface values are from NOAA, the CHCl_3 surface values are from AGAGE, and the $\text{C}_2\text{H}_4\text{Cl}_2$ surface values are estimated from the HIPPO aircraft campaign^{14,46,47}. Trends are calculated in the modelled stratospheric source gas injection (SGI) plus the product gas injection (PGI) of chlorine from CI-VSLs, which corresponds to the simulated concentrations at 17 km and from 20°S – 20°N . We use the trend from 1999–2015 for comparison to 5.5-year-old air at mid-latitudes and the trend from 2001 to 2017 for the comparison to 3-year-old air at mid-latitudes to account for the transit time from the tropical tropopause region.

Trend calculations

Trends are calculated with a generalised least squares method using the Long-term Ozone Trends and Uncertainties in the Stratosphere (LOTUS) regression code⁴⁸. Three different sets of proxies are considered, each of which is described in this section. In all cases, the uncertainties are defined as the 2σ error of the trend parameter estimate in the regression (95% confidence level). The uncertainties account for first-order autocorrelation in the residuals⁴⁹.

The uncertainty in the mean Cl_y trend presented in Fig. 5 is calculated as

$$\sigma = \frac{1}{N} \sqrt{\sum_{i=1}^N \sum_{j=1}^N \sigma_i \sigma_j} \quad (1)$$

where $N = 4$, corresponding to the four latitude bins that are considered.

The Cl-EESC and Cl-VSLs trends are calculated with a simple linear regression, defined by the equation

$$y(t) = \beta + \beta_{\text{trend}} \times \text{linear}(t) + R(t), \quad (2)$$

where $y(t)$ is the mixing ratio of either Cl-EESC or Cl-VSLs as a function of time, β is the constant term, β_{trend} is the trend, and $R(t)$ is the residual. The $\text{linear}(t)$ variable represents a linear function of time that is fit to the observations (the trend is the slope of this line).

Trends in $\text{HCl} + \text{ClONO}_2$ are calculated in two ways: with the standard MLR and with the N_2O MLR. The standard MLR is defined as

$$y(t) = \beta^{(2)} + \beta_{\text{trend}} \times \text{linear}(t) + \beta_{\text{qboa}}^{(2)} \times \text{QBO}_a(t) + \beta_{\text{qbob}}^{(2)} \times \text{QBO}_b(t) + \beta_{\text{solar}} \times F_{10.7}(t) + \beta_{\text{enso}} \times \text{ENSO}(t) + R(t). \quad (3)$$

In Eq. (3) each β_i defines a regression coefficient $\text{QBO}_a(t)$ and $\text{QBO}_b(t)$ are the first two principal components of the Singapore zonal winds, $F_{10.7}(t)$ is the solar flux at 10.7 cm, and $\text{ENSO}(t)$ is the multivariate El-Niño Southern Oscillation (ENSO) index. The superscripts specify the number of seasonal harmonics included for a given term. Therefore, the coefficient for the constant term expands to

$$\beta^{(2)} = \beta^0 + \sum_{k=1}^2 \left(\beta^{2k-1} \sin \frac{2\pi}{365.25} kt + \beta^{2k} \cos \frac{2\pi}{365.25} kt \right). \quad (4)$$

$\beta_{\text{qboa}}^{(2)}$ and $\beta_{\text{qbob}}^{(2)}$ are expanded similarly. There are 18 regression coefficients in total: one corresponding to each of the trend, $\text{ENSO}(t)$, and $F_{10.7}(t)$ terms and five coefficients from each of the constant, $\text{QBO}_a(t)$, and $\text{QBO}_b(t)$ terms. Further details on this version of the regression model, the data sources for the proxies, and examples showing how the regression model can be used to calculate stratospheric ozone trends are provided in refs. 48,49.

The N_2O MLR equation is

$$y(t) = \beta^{(2)} + \beta_{\text{trend}} \times \text{linear}(t) + \beta_{\text{solar}} \times F_{10.7}(t) + \beta_{\text{N}_2\text{O}} \times \text{N}_2\text{O}(t) + R(t). \quad (5)$$

All terms are the same as in Eq. (3), except in this case the $\text{N}_2\text{O}(t)$ term represents all sources of dynamical variability, including ENSO, the QBO, and the BDC. The N_2O proxy is the anomaly in the MZM ACE-FTS N_2O mixing ratio, with the trend due to surface emissions of N_2O removed. The proxy is therefore different in each latitude and altitude bin. Further details on the N_2O proxy and MLR are provided in ref. 30.

Altitudes for a specified mean age

The ACE-FTS MZM age of air profiles are used to determine the altitude corresponding to each mean age. First, a cubic spline is fit to each MZM age of air profile, and then that curve is interpolated to the altitude corresponding to the specified mean age. The result is an MZM time series of the altitude corresponding to each mean age. This time series is then filtered by removing monthly values that are more than 5 median absolute deviations from the median. The mean altitude of the remaining values is taken as the altitude of 3- and 5.5-year-old air at each latitude.

For 3-year-old air, the altitudes are 18.5 km, 19.6 km, 20.1 km, 19.1 km for the latitudes 50°S, 40°S, 40°N, 50°N. For a 5.5-year-old air, the altitudes are 29.3 km, 30.2 km, 30.4 km, 29.6 km for the latitudes 50°S, 40°S, 40°N, 50°N. The ACE-FTS AoA has a vertical resolution of 3 km, so the difference in the altitude for each mean age across the four latitudes is smaller than the

resolution of the data. We therefore present our results in terms of the average altitude across all four latitudes for each mean age: 19.0 km for 3-year-old air and 29.5 km for 5.5-year-old air.

Conversion of $\text{HCl} + \text{ClONO}_2$ trends to Cl_y trends

We convert the ACE-FTS $\text{HCl} + \text{ClONO}_2$ trend to a Cl_y trend using a scale factor from CMAM:

$$\text{Cl}_y \text{ trend}[\text{ACE}] = \text{HCl} + \text{ClONO}_2 \text{ trend}[\text{ACE}] \times \frac{\text{Cl}_y \text{ trend}[\text{CMAM}]}{\text{HCl} + \text{ClONO}_2 \text{ trend}[\text{CMAM}]} \quad (6)$$

The CMAM $\text{HCl} + \text{ClONO}_2$ and Cl_y trends are calculated with the N_2O MLR to be consistent with the ACE-FTS trends. The ACE-FTS pressure profiles that are retrieved with each scan are used to determine the average pressure corresponding to the altitude of 3- and 5.5-year-old air, and then the scale factors are calculated from CMAM trends interpolated to this pressure level.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

ACE-FTS ClONO_2 , HCl , and N_2O observations are available by registration at <https://database.scisat.ca/level2/>⁵⁰. ACE-FTS data quality flags are available from <https://doi.org/10.5683/SP3/NAYNFE>⁵¹. ACE-FTS age of air is available from <https://doi.org/10.5683/SP3/5AC1F0>⁵². CMAM REFD1 simulations are available by registration from the CEDA archive, <https://catalogue.ceda.ac.uk/uuid/3f06d743521f44a5ba8e666e11f66d6b/>⁵³.

Code availability

The LOTUS regression code and documentation are available at <https://github.com/usask-arg/lotus-regression>⁴⁸.

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Author contributions

K.D. performed the analysis and wrote the first draft of the manuscript. S.T. conceptualised and supervised the project. A.E. provided the code for the CI-EESC calculation. J.L. provided the ODS mole fractions needed to calculate the EESC. L.S. provided the ACE-FTS age of air dataset. R.H. and E.B. provided the CI-VLS trends. All authors provided significant feedback on the method and the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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