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3	Satellite Observations of Stratospheric Hydrogen Fluoride and		
4	Comparisons with SLIMCAT Calculations		
5	by		
6	Jeremy J. Harrison ^{1,2} , Martyn P. Chipperfield ^{3,4} , Christopher D. Boone ⁵ , Sandip S.		
7	Dhomse ^{3,4} , Peter F. Bernath ⁶ , Lucien Froidevaux ⁷ , John Anderson ⁸ , and James Russell III ⁸		
8			
9	¹ Department of Physics and Astronomy, University of Leicester, Leicester LE1 7RH, United		
10	Kingdom.		
11	² National Centre for Earth Observation, University of Leicester, Leicester LE1 7RH, United		
12	Kingdom.		
13	³ Institute for Climate and Atmospheric Science, School of Earth and Environment, University		
14	of Leeds, Leeds LS2 9JT, United Kingdom.		
15	⁴ National Centre for Earth Observation, School of Earth and Environment, University of		
16	Leeds, Leeds LS2 9JT, United Kingdom.		
17	⁵ Department of Chemistry, University of Waterloo, Waterloo, Ontario N2L 3G1, Canada.		
18	⁶ Department of Chemistry and Biochemistry, Old Dominion University, Norfolk, Virginia		
19	23529, United States of America.		
20	⁷ Jet Propulsion Laboratory, Pasadena, California 91109, United States of America.		
21	⁸ Department of Atmospheric and Planetary Sciences, Hampton University, Hampton,		
22	Virginia 23668, United States of America.		
23 24 25 26 27	Number of pages = 30 Number of tables = 5 Number of figures = 11		
28 29 30 31 32 33 34 35 36	Address for correspondence:Dr. Jeremy J. HarrisonNational Centre for Earth ObservationDepartment of Physics and AstronomyUniversity of LeicesterUniversity RoadLeicester LE1 7RHUnited KingdomTel:(44)-116-2231943		
37 38	<i>e-mail</i> : jh592@leicester.ac.uk		

39 Abstract

The vast majority of emissions of fluorine-containing molecules are anthropogenic in nature, e.g. chlorofluorocarbons (CFCs), hydrochlorofluorocarbons (HCFCs), and hydrofluorocarbons (HFCs). Many of these fluorine-containing species deplete stratospheric ozone, and are regulated by the Montreal Protocol. Once in the atmosphere they slowly degrade, ultimately leading to the formation of HF, the dominant reservoir of stratospheric fluorine due to its extreme stability. Monitoring the growth of stratospheric HF is therefore an important marker for the success of the Montreal Protocol.

47 We report the comparison of global distributions and trends of HF measured in the Earth's atmosphere by the satellite remote-sensing instruments ACE-FTS (Atmospheric 48 49 Chemistry Experiment Fourier Transform Spectrometer), which has been recording 50 atmospheric spectra since 2004, and HALOE (HALogen Occultation Experiment), which 51 recorded atmospheric spectra between 1991 and 2005, with the output of SLIMCAT, a state-52 of-the-art three-dimensional chemical transport model. In general the agreement between 53 observation and model is good, although the ACE-FTS measurements are biased high by 54 ~10% relative to HALOE. The observed global HF trends reveal a substantial slowing down 55 in the rate of increase of HF since the 1990s: 4.97 ± 0.12 %/year (1991-1997; HALOE), 1.12 56 \pm 0.08 %/year (1998-2005; HALOE), and 0.52 \pm 0.03 %/year (2004-2012; ACE-FTS). In 57 comparison, SLIMCAT calculates trends of 4.01 %/year, 1.10 %/year, and 0.48 %/year, respectively, for the same periods; the agreement is very good for all but the earlier of the 58 59 two HALOE periods. Furthermore, the observations reveal variations in the HF trends with 60 latitude and altitude, for example between 2004 and 2012 HF actually decreased in the southern hemisphere below ~35 km. An additional SLIMCAT simulation with repeating 61 62 meteorology for the year 2000 produces much cleaner trends in HF with minimal variations 63 with latitude and altitude. Therefore, the variations with latitude and altitude in the observed 64 HF trends are due to variability in stratospheric dynamics on the timescale of a few years. 65 Overall, the agreement between observation and model points towards the ongoing success of 66 the Montreal Protocol and the usefulness of HF as a metric for stratospheric fluorine.

68

1. Introduction

69 The accumulation of fluorine in the Earth's atmosphere has resulted from 70 anthropogenic emissions of organic molecules such as chlorofluorocarbons (CFCs), 71 hydrochlorofluorocarbons (HCFCs), and hydrofluorocarbons (HFCs). The long atmospheric 72 lifetimes of such molecules allow them to reach the stratosphere, where they break down and 73 liberate fluorine in various inorganic forms. The most abundant of the emitted organic source 74 molecules are CFC-12 (CCl₂F₂), CFC-11 (CCl₃F), CFC-113 (CCl₂FCClF₂), which are all 75 now banned under the Montreal Protocol because they deplete stratospheric ozone, and 76 HCFC-22 (CHClF₂), the most abundant HCFC and a transitional substitute under the Protocol. Although long-lived, these molecules do degrade in the atmosphere at high 77 78 altitudes, ultimately to the long-lived stratospheric reservoir molecule hydrogen fluoride, HF; 79 the chemistry schemes are presented below. Monitoring HF as part of the atmospheric 80 fluorine family is important in closing the fluorine budget, particularly as anthropogenic emissions of fluorine species, many of which are ozone-depleting and all of which are 81 82 greenhouse gases, have varied substantially over time. Certainly, monitoring the growth of 83 stratospheric HF, which has slowed in recent years, is an important marker for the success of 84 the Montreal Protocol (in addition to monitoring stratospheric HCl).

For the three most abundant fluorine source gases, CFC-12, CFC-11, and HCFC-22,
atmospheric degradation proceeds with the breaking of a C-Cl (CFC-12 and CFC-11) or C-H
(HCFC-22) bond (Ricaud and Lefevre, 2006),

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89	$CCl_2F_2 + hv \rightarrow CClF_2 + Cl$	
90	$CCl_3F + hv \rightarrow CCl_2F + Cl$	
91	$CHClF_2 + OH \rightarrow CClF_2 + H_2O.$	(R1)

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93 Depending on their structure, the intermediates produced in R1 react further,

 $CClF_2 + O_2 + M \rightarrow CClF_2O_2 + M$ $CClF_2O_2 + NO \rightarrow CClF_2O + NO_2$ $CClF_2O + O_2 \rightarrow COF_2 + ClO_2$, (R2) 98 $CCl_2F + O_2 + M \rightarrow CCl_2FO_2 + M$ $CCl_2FO_2 + NO \rightarrow CCl_2FO + NO_2$ $CCl_2FO + O_2 \rightarrow COClF + ClO_2$. (R3) 102

For CFC-113, and minor sources such as HFCs (e.g. HFC-134a, HFC-152a), the reactionscheme is similar.

In Equations R2 and R3, carbonyl chloride fluoride (COCIF) and carbonyl fluoride 105 106 (COF₂), are important 'inorganic' reservoirs (the common terminology in atmospheric 107 science differs from that in chemistry) of fluorine in the stratosphere, with lifetimes of 1.6 108 (Fu et al., 2009) and 3.8 (Harrison et al., 2014) years, respectively; COF₂ is more abundant 109 than COCIF. The trends in these inorganic reservoirs over time are directly related to the 110 trends of the individual source gases. The decrease in the amounts of atmospheric CFC-11 111 and CFC-113, the principal sources of carbonyl chloride fluoride, has led to a decreasing 112 trend in this reservoir (Brown et al., 2011), whereas carbonyl fluoride is still slowly increasing over time due to the increase in HCFC-22, which more than compensates for the 113 114 decrease in the CFC-12 and CFC-113 source gases (Brown et al., 2011; Harrison et al., 115 2014).

116 COClF and COF₂ volume mixing ratios (VMRs) slowly increase with altitude 117 through the lower stratosphere until they reach their respective maxima, at $\sim 25 - 30$ km for 118 COClF (Fu et al., 2009) and $\sim 30 - 40$ km for COF₂ (Harrison et al., 2014). Above these 119 altitudes photolysis becomes more efficient, leading to the formation of fluorine atoms,

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121	$\text{COF}_2 + \text{hv} \rightarrow \text{FCO} + \text{F}$	
122	$\text{COClF} + \text{hv} \rightarrow \text{FCO} + \text{Cl}$	
123	$FCO + O_2 + M \rightarrow FC(O)O_2 + M$	
124	$FC(O)O_2 + NO \rightarrow FCO_2 + NO_2$	
125	$FCO_2 + hv \rightarrow F + CO_2.$	(R4)

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The liberated fluorine atoms then react with methane, water or molecular hydrogen, to form 127 the inorganic product hydrogen fluoride, HF. At the top of the stratosphere most of the 128 129 fluorine is present as HF (Brown et al., 2014), the dominant reservoir of stratospheric 130 fluorine due to its extreme stability. Note that due to this stability, F is not important for 131 catalytic stratospheric ozone loss. HF is removed from the stratosphere by slow transport to, 132 and rainout in, the troposphere, or by upward transport to the mesosphere, where it is 133 destroyed by photolysis (Duchatelet et al., 2010). Overall the amount of HF in the 134 atmosphere is increasing (e.g. Brown et al., 2014).

135 The first detection of HF in the Earth's stratosphere, based on solar spectra recorded from balloon and on the ground at Jungfraujoch, was made by Zander et al. (1977). 136 137 Measurements continue to be made at Jungfraujoch using a ground-based Fourier transform 138 spectrometer (FTS), (e.g. Duchatelet et al., 2010). There have also been measurements of HF 139 taken, for example, by the Atmospheric Trace MOlecule Spectrometry Experiment 140 (ATMOS) instrument which flew four times on NASA Space Shuttles between 1985 and 141 1994 (Irion et al., 2002) and the MkIV interferometer, a balloon-borne solar occultation FTS 142 (Velazco et al., 2011). Measurements taken by satellite-borne instruments, however, allow 143 HF to be observed with global coverage, and seasonal and latitudinal variability to be 144 investigated fully. The first global atmospheric distributions of HF were provided by the 145 HALogen Occultation Experiment (HALOE) instrument, onboard the Upper Atmosphere Research Satellite (UARS), which recorded atmospheric spectra between 1991 and 2005. 146 147 More recently, the Atmospheric Chemistry Experiment (ACE)-FTS, onboard the SCISAT 148 satellite, has been recording atmospheric spectra since 2004, carrying the mantle of HF 149 measurements into the second decade of the twenty-first century. In fact, the ACE-FTS is the 150 only satellite instrument currently taking measurements of HF.

151 This paper follows on from our recent work on the global distributions and trends of 152 COF₂, the most important 'temporary' stratospheric fluorine reservoir that directly leads to 153 the formation of HF. The aim of the present work is to understand the HF global distribution 154 and trends derived from satellite observations taken by the HALOE and ACE-FTS 155 instruments. To do this, we use the SLIMCAT model, a state-of-the-art three-dimensional 156 (3D) chemical transport model (CTM), one of the few to include stratospheric fluorine 157 chemistry. Additionally, we compare tracer-tracer correlations between some of the major 158 HF 'sources' for SLIMCAT and satellite observations as a further test of the model 159 chemistry.

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161 **2. Hydrogen fluoride datasets**

162 **2.1. ACE-FTS**

163 The ACE-FTS instrument, which covers the spectral region 750 to 4400 cm⁻¹ with a 164 maximum optical path difference (MOPD) of 25 cm and a resolution of 0.02 cm⁻¹ (using the 165 definition of 0.5/MOPD throughout), uses the sun as a source of infrared radiation to record 166 limb transmission through the Earth's atmosphere during sunrise and sunset ('solar 167 occultation'). Transmittance spectra are obtained by ratioing against exo-atmospheric 'high 168 sun' spectra measured each orbit. These spectra, with high signal-to-noise ratios, are 169 recorded through long atmospheric limb paths (~300 km effective length), thus providing a 170 low detection threshold for trace species. ACE has an excellent vertical resolution of about 171 3-4 km and can measure up to 30 occultations per day, with each occultation sampling the 172 atmosphere from 150 km down to the cloud tops (or 5 km in the absence of clouds). The 173 locations of ACE occultations are dictated by the low Earth circular orbit of the SCISAT 174 satellite and the relative position of the sun. Over the course of a year, the ACE-FTS records 175 atmospheric spectra over a large portion of the globe (Bernath et al., 2005), from which it is 176 possible to extract profiles of many fluorine-containing species, including CCl₃F (CFC-11), CCl₂F₂ (CFC-12), CHClF₂ (HCFC-22), CCl₂FCClF₂ (CFC-113), CH₃CCl₂F (HCFC-141b), 177 178 CH₃CClF₂ (HCFC-142b), CH₂FCF₃ (HFC-134a), CHF₃ (HFC-23), CF₄, COF₂, COClF, HF 179 and SF₆.

180 The atmospheric pressure and temperature profiles, the tangent heights of the 181 measurements, and the hydrogen fluoride VMRs were taken from version 3.0 (January 2004 182 until September 2010) and v3.5 (from October 2010) processing of the ACE-FTS data. Note 183 that the retrieval scheme is identical for both the v3.0 and v3.5 datasets, the difference being 184 in the meteorological data used as input for the pressure and temperature retrievals (the 185 lowest ACE-FTS levels use these data directly). Due to an error in these inputs, v3.0 data should only be used for measurements taken until the end of September 2010, while v3.5 is 186 valid for all ACE-FTS measurements. Details of the retrieval scheme for versions 3.0/3.5 187 processing have been explained elsewhere (e.g., Boone et al., 2013; Harrison et al., 2014). 188 189 Briefly, vertical profiles of trace gases (along with temperature and pressure) are derived 190 from the recorded transmittance spectra via an iterative Levenberg-Marquardt nonlinear 191 least-squares global fit to the selected spectral region(s) for all measurements within the 192 altitude range of interest. The microwindow set and associated altitude ranges for the HF 193 retrieval are listed in Table 1. The VMRs for molecules with absorption features in the 194 microwindow set (see Table 2) were adjusted simultaneously with the HF amount. All 195 spectroscopic line parameters were taken from the HITRAN 2004 database (Rothman et al., 196 2005), with HF parameters apparently unchanged since HITRAN 1992. The microwindow 197 set covers eight spectroscopic lines (P₁, P₂, P₃, P₄, R₀, R₁, R₂, R₃) from the fundamental (1-0) 198 band of HF. The HF retrieval extends from a lower altitude of 12 km up to altitudes corresponding to an atmospheric density of 9.0×10^{15} molecules cm⁻³, in practice ~50-55 km, 199 200 thus providing a variation in upper altitude with both latitude and season (see Table 1). The 201 HF spectral signal in ACE-FTS spectra recorded above the upper altitude retrieval limit is 202 generally below the noise level, so it is not possible to retrieve VMRs directly at these altitudes. Instead, the VMR profile above the highest analysed ACE measurement is
calculated by scaling the 'initial' VMR profile, taken from ATMOS measurements (Irion et
al., 2002), over these altitudes; the scaling factor is determined during the least-squares
fitting.

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208 **2.2. HALOE**

209 Like the ACE-FTS, the HALOE instrument (Russell et al., 1993) used the principle 210 of solar occultation to sound the middle atmosphere at sunset and sunrise (relative to the instrument). HALOE used broadband and gas-filter radiometry, with channels covering 211 212 selected portions of the spectrum between 2.45 and 10.04 µm, to determine the mixing ratios 213 of molecules related to the chemistry of stratospheric ozone and its destruction by CFCs. In particular, HALOE provided measurements of O₃, HCl, HF, CH₄, H₂O, NO, NO₂, aerosol 214 215 extinction, and temperature versus pressure, over an altitude range of ~ 15 to 60 - 130 km 216 depending on channel (HF, HCl, CH₄ and NO were measured using gas filter radiometry). 217 As with the ACE-FTS, the locations of HALOE occultations and hence the extent of its 218 global coverage were dictated by its orbit and the relative position of the sun. HALOE, with 219 an orbit inclination of 57° compared with 74° for the ACE-FTS, had a more even latitudinal 220 coverage and provided more data over tropical regions, for example, than the ACE-FTS 221 which takes most of its measurements at high latitude.

The atmospheric pressure, temperature, tangent heights, and hydrogen fluoride 222 223 VMRs were taken from version 19 processing of the HALOE data, which are available from 224 October 1991 to November 2005. The retrieval scheme incorporates a simple 'onion 225 peeling' approach stabilised at the top and bottom of the profile with a scalar optimal estimation formulation developed by Connor and Rodgers (1989). For the HF channel, the 226 spectral bandpass 5% relative response points are 4025 cm⁻¹ and 4135 cm⁻¹. The HF spectral 227 line parameters were taken from the HITRAN 1992 database (Rothman et al., 1992). The 228 229 instantaneous vertical field-of-view in the limb is ~1.6 km. Detailed validation studies for 230 HALOE HF measurements were published by Russell et al. (1996). Note that for internal consistency with previous work on the fluorine budget (Brown et al., 2014) and COF₂ 231 232 (Harrison et al., 2014), the vertical pressure grid has been interpolated onto the standard 1-km 233 grid used by the ACE-FTS.

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235 **2.3. GOZCARDS**

236 The ACE-FTS, HALOE and SLIMCAT model time series are also compared with 237 those of the GOZCARDS (Global OZone Chemistry And Related Datasets for the Stratosphere) HF data product in Section 5. GOZCARDS provides a global long-term 238 239 stratospheric Earth System Data Record (ESDR) for stratospheric ozone and related chemical 240 species, including HF. The HF data record was not ready in time to be included in the 241 original dataset provided for public usage (temperature, O₃, H₂O, HCl, N₂O, and HNO₃); we 242 are presenting the HF data for the first time here. Froidevaux et al. (2015) have described the 243 GOZCARDS data creation methodology and some stratospheric characteristics concerning 244 the latter five species. The constituent datasets are time series of monthly zonal means 245 versus latitude (in 10° latitude bins) taken from existing satellite datasets. In particular, the 246 GOZCARDS HF data product is derived by merging the v19 HALOE (1991 – 2005) and v2.2 ACE-FTS (2004 - 2010) datasets, with the relative bias between source datasets 247 248 removed by averaging them over the overlap period 2004 - 2005 and adjusting the series 249 accordingly; note that such a process does not account for any systematic biases in the 250 original datasets. All GOZCARDS datasets are provided on a vertical pressure grid. Again, 251 to be consistent with previous ACE work, this vertical pressure grid has been interpolated 252 onto the standard 1-km grid used by the ACE-FTS. Note that as this GOZCARDS dataset 253 uses v2.2 ACE-FTS data, the time series terminates in September 2010.

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3. Retrieval errors

256 **3.1. Infrared spectroscopy of hydrogen fluoride**

257 One of the major sources of systematic error for any retrieved atmospheric profile 258 arises from uncertainties in the laboratory spectroscopic data. A discussion of spectroscopic 259 errors is therefore appropriate. The ACE-FTS retrieval makes use of HF line parameters first 260 made available as part of the HITRAN 1992 database (and remaining unchanged until the 261 HITRAN 2012 release), with partition data taken from the Total Internal Partition Sums (TIPS) subroutine included in the HITRAN compilation. HITRAN simply provides error 262 263 codes for line parameters in the form of uncertainty ranges, but with no information as to how the parameters are correlated. For the HF line parameters used in this work, the errors 264 correspond to 0.0001 - 0.001 cm⁻¹ for the line wavenumber, v, 2 - 5 % for the line intensity, 265 S, and 1 - 2 % for the air-broadened half-width, γ_{air} . Errors are unreported for γ_{self} (self-266 267 broadened half-width), n_{air} (temperature-dependence exponent for γ_{air}), and δ_{air} (air pressure-268 induced line shift). Recently, and after v3.0 processing of the ACE data was complete, 269 HITRAN 2012 became available; it includes a complete re-evaluation of HF spectroscopy. The associated publication (Rothman et al., 2013) also explains that all the air-broadening parameters, γ_{air} , for the fundamental band of HF were fitted with the Galatry profile, not the Voigt profile, which is the lineshape of choice for the HITRAN database. Additionally, the Dicke narrowing parameters in the original analysis were simply neglected. For the purposes of this work, we assume a retrieval error of at most ~4 % arising from uncertainties in HF line parameters.

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277 **3.2. ACE-FTS**

The ACE v2.2 HF data product, which uses a slightly different microwindow set 278 279 from v3.0/v3.5 as well as an earlier version of the PT retrieval, has previously been validated, 280 for example, against measurements taken by HALOE and the MkIV interferometer (Mahieu 281 et al., 2008). It was found that ACE-FTS v2.2 HF measurements are biased high compared 282 to HALOE, with mean differences around 5-20% between 15 and 49 km. Comparison of ACE-FTS v2.2 HF with MkIV data is generally good, with relative differences above 19 km 283 284 within $\pm 10\%$. There have been no detailed comparisons in the literature between ACE-FTS 285 v2.2 and v3.0 HF datasets, although Duchatelet et al. (2010) suggest that ACE-FTS v3.0 HF VMRs have decreased by close to 5% relative to v2.2. However, a re-evaluation of filtered 286 v2.2 and v3.0/v3.5 HF data carried out as part of this work indicates good agreement within \pm 287 288 5 %, with no significant overall bias between the two datasets.

289 The 1 σ statistical fitting errors for a single ACE profile are typically ~5 % over most of the altitude range. These errors are random in nature and largely determined by the 290 measurement noise of the ACE-FTS. Averaged profiles tend to be dominated by systematic 291 errors, with random errors reduced by a factor of $1/\sqrt{N}$, where N is the number of profiles 292 Spectroscopic sources of systematic error predominantly arise from the HF 293 averaged. 294 HITRAN linelist (~4 %; see Section 3.1), with minor contributions from interfering species 295 that absorb in the microwindow regions; we assume that these contributions are small, at 296 most 1 %, due to the lack of systematic features in the spectral residuals (Harrison et al., 297 2014). Additional systematic errors arise from uncertainties in temperature, pressure, tangent 298 altitude (i.e. pointing) and instrumental line shape (ILS); these were estimated by running the 299 ACE-FTS retrieval for a subset of occultations, with each quantity (b_i) perturbed in turn by its assumed 1 σ uncertainty (Δb_i). The fractional retrieval error, μ_i , is defined as 300 301

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$$\mu_{j} = \left| \frac{\text{VMR}(b_{j} + \Delta b_{j}) - \text{VMR}(b_{j})}{\text{VMR}(b_{j})} \right|.$$
(3)

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304 Note that pressure, temperature and tangent height are not strictly independent quantities for 305 ACE-FTS retrievals; tangent heights are determined from hydrostatic equilibrium, and so 306 these quantities are strongly correlated. Therefore, only two of these three quantities are 307 altered: temperature is adjusted by 2 K and tangent height by 150 m (Harrison et al., 2014). 308 ILS uncertainty is determined by adjusting the field of view by 5 % (Harrison et al., 2014). A subset of 81 occultations measured between 65° and 70°N in July 2010 was selected for 309 310 The fractional value estimates of the systematic uncertainties, and their this analysis. 311 symbols, are given in Table 3. Assuming uncorrelated quantities, the overall systematic error 312 in the HF retrieval can be calculated as

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- 314
- 315

 $\mu_{systematic}^{2} = \mu_{spec}^{2} + \mu_{int}^{2} + \mu_{T}^{2} + \mu_{z}^{2} + \mu_{ILS}^{2}.$ (4)

The total systematic error contribution to the ACE-FTS HF retrieval is estimated to be $\sim 10 \%$ over the altitude range of the retrieval.

318 As discussed in Section 3.1, HF VMRs are not directly retrieved for ACE 319 measurements taken at tangent heights above the upper altitude limits listed in Table 1. In 320 the ACE-FTS HF retrieval, the calculated spectrum is generated from the sum of 321 contributions from the tangent layer up to 150 km. For the highest analysed measurement, 322 the retrieved VMR in the tangent layer is generated from the piecewise quadratic 323 interpolation scheme (Boone et al., 2013), while the VMR in every layer above that is 324 determined from scaling the 'initial' VMR profile, with the scaling factor determined during 325 the retrieval by forcing the calculated spectrum to match as best as possible the measured 326 spectrum for the highest analysed measurement. Since the 'initial' profile is fixed to a 327 constant VMR between 50 and 100 km altitude, and since this portion of the profile is scaled 328 based on the VMR of the highest analysed ACE measurement, this will likely introduce 329 systematic errors into the highest altitudes of the retrieved profile. However, since the 330 scaling factor errors are dominated by the 1σ statistical fitting errors of the highest analysed 331 ACE measurement, it is anticipated that the systematic errors at the top of the profiles are 332 reduced when they are averaged to create zonal means.

334 **3.3. HALOE**

335 As discussed in Section 3.2, HALOE v19 HF has been validated against ACE-FTS 336 v2.2, with the ACE measurements biased high by around 5-20% between 15 and 49 km 337 (Mahieu et al., 2008). Furthermore, HF data from the MkIV interferometer for three flights 338 (2003 - 2005) agree well with ACE-FTS, with relative differences above 19 km within 339 $\pm 10\%$, suggesting that there is a low bias in HALOE. Detailed HALOE HF error estimates 340 and validation studies have previously been conducted by Russell et al. (1996). The 341 estimated errors range from ~27% at 100 hPa to 15% at 1 hPa. Actual mean differences 342 between HALOE and balloon data from a series of nine FTS under-flights, five operating in 343 the far-IR and four MkIV comparisons in the near-IR, collectively ranged from more than 344 17% below 70 hPa where the mixing ratio is very low to <7% above that level, with no positive or negative bias implied. These HALOE data were produced using an early 345 346 algorithm version, but results have proven to be very stable for later versions.

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348 **4. TOMCAT/SLIMCAT 3D Chemical Transport Model**

349 SLIMCAT, an off-line 3D CTM, calculates the abundances of a number of 350 stratospheric trace gases from prescribed source-gas surface boundary conditions and a 351 detailed treatment of stratospheric chemistry, including the major species in the O_x, NO_y, 352 HO_x, F_y, Cl_y and Br_y chemical families (e.g. Chipperfield, 1999; Feng et al., 2007). The 353 model uses winds from meteorological analyses to specify winds and temperatures. This 354 approach gives a realistic stratospheric circulation (Chipperfield, 2006; Monge-Sanz et al., 355 2007). In the version used here the troposphere is assumed to be well-mixed.

356 For this study SLIMCAT was integrated from 1977 to 2013 at a horizontal 357 resolution of 5.6° \times 5.6° and 32 levels from the surface to 60 km. The model uses a σ -p 358 vertical coordinate (Chipperfield, 2006) and was forced by European Centre for Medium-359 Range Weather Forecasts (ECMWF) reanalyses (ERA-Interim from 1979 onwards). The 360 VMRs of source gases at the surface level were specified using datasets prepared for the 361 WMO/UNEP (2011) ozone assessment. These global mean surface values define the long-362 term tropospheric source gas trends in the model. Similarly, specification of the surface 363 VMRs of degradation products acts as a sink for these species. The model initialisation used 364 the estimated halocarbon loading for 1977, taken from the WMO/UNEP scenarios.

The SLIMCAT run makes use of the same chemistry scheme that was previously used for our work on COF₂ (Harrison et al., 2014), however in the present version the photolysis scheme has been updated to use latitudinally and monthly varying ozone profile

368 shapes in the photolysis look-up table. All source and degradation products related to 369 fluorine chemistry are listed in Table 4. COF₂ contributions arise from the degradation of 370 CFC-12, CFC-113, CFC-114, CFC-115, HCFC-22, HCFC-142b, HFC-23, HFC-134a, HFC-371 152a, Halon 1211, and Halon 1301, with COCIF production arising from the degradation of 372 CFC-11, CFC-113, and HCFC-141b (CH₃CCl₂F). Some HF is assumed to form directly 373 from the source gases (see Table 4), however this is almost negligible in practical terms (\sim 3 374 % in 2010, mainly arising from HFC-134a). The relative amounts of HF formed (in 2010) 375 via COClF and COF₂ are 30 and 67 %, respectively.

376 The SLIMCAT calculations reveal that at altitudes above the maximum COCIF and 377 COF₂ VMRs, there is net loss of these at all latitudes. The primary loss of COF₂ and COCIF 378 in the stratosphere occurs via photolysis, with an additional secondary loss mechanism through reaction with $O(^{1}D)$; SLIMCAT calculates relative contributions of 90 % and 10 %, 379 380 respectively, for COF₂ (Harrison et al., 2014) and 98 % and 2 % for COCIF. The SLIMCAT 381 outputs enable an estimation of the atmospheric lifetimes of COF₂ and COCIF by dividing 382 the total modelled atmospheric burden of each species by the total calculated atmospheric 383 loss rate. The total calculated mean atmospheric lifetimes are 3.9 years for COF₂, revised 384 upwards from the calculated value of 3.8 years presented by Harrison et al. (2014), and 1.7 years for COCIF. 385

386

387 **5.** Comparison between ACE-FTS/HALOE/SLIMCAT datasets

388 The ACE-FTS HF data were binned into five latitude bands by month; VMRs 389 outside six median absolute deviations (MAD) of the median VMR for each bin and altitude 390 were removed from the analysis. Once filtered to remove significant outliers, the data were 391 used to create monthly zonal means at each altitude within 5° latitude bins. In Figure 1 these 392 have been plotted next to SLIMCAT zonal means for the months September 2009 to August 393 2010, thereby revealing the seasonal variation in the HF distribution over this period. Note 394 that these dates have been chosen to match those used in the previous work on carbonyl 395 fluoride (see Figure 11 of Harrison et al., 2014). The HF profiles generally show an increase 396 in VMR with altitude, with the rate of this increase varying with latitude and time of year. 397 Note that ACE-FTS observations do not cover all latitude bins over a single month (see 398 Section 2.1), and that latitude bins containing fewer occultations are noisier in appearance. 399 Despite these caveats, Figure 1 reveals a good agreement between the ACE-FTS observations 400 and the model which reproduces very well the significant seasonal variation. For example, 401 note in particular the agreement for regions of low HF VMR (< 1000 ppt) at \sim 30 – 40 km 402 over the southern tropics in February 2010 and the northern tropics in August 2010; at 403 southern mid- to high-latitudes in December 2009, and March and August 2010; and at 404 northern mid- to high-latitudes in February and March 2010.

405 Plots of ACE-FTS and HALOE HF observations side-by-side with SLIMCAT HF 406 calculations for September 2004 to August 2005 are shown in Figure 2. As for Figure 1, the 407 agreement between observations and model is generally good and the significant seasonal 408 variation is well reproduced. Note that, as for the ACE-FTS, HALOE data do not cover all 409 latitude bins in a given month, although HALOE does take more measurements at lower 410 latitudes. One noticeable difference revealed in Figure 2 is the relative low bias of HALOE 411 measurements compared with ACE-FTS and SLIMCAT; this is most notable at the top of the 412 altitude range. Using the period of overlap between ACE and HALOE, we can estimate that 413 ACE v3.0 is biased high by about 10% relative to HALOE. Biases between observations and 414 SLIMCAT will be more fully addressed in Section 6.

415 The overall atmospheric distribution of HF is determined by a complicated 416 combination of its production and transport, which depends on the production and lifetimes of its 'sources' COF₂ and COCIF. Figure 3 shows the observed and modelled COF₂ and 417 418 COCIF zonal means for October 2009, and February and August 2010. Due to the upwelling of relatively organic-fluorine-rich air in the tropical regions, the largest VMRs of COF₂ and 419 COCIF are found over the tropics (Harrison et al., 2014; Fu et al., 2009), where solar 420 421 insolation is highest due to the small solar zenith angle, at altitudes of $\sim 30 - 40$ and $\sim 25 - 30$ 422 km. For COF₂ the model agrees well with the ACE observations in terms of magnitude and 423 spatial distribution. For COCIF the modelled distribution agrees with ACE but the peak 424 VMR is overestimated. Analysis of the SLIMCAT simulation shows that there is net loss of 425 COF₂ and COCIF at altitudes above those of the maximum VMRs, at all locations. There is 426 therefore a correlation between the stratospheric regions of low HF VMR (< 1000 ppt) above 427 \sim 20 km at the poles and \sim 25 km at the equator and those of peak COF₂ and COCIF VMRs.

428 Figure 1 reveals an asymmetry in the seasonal HF distribution between the two 429 hemispheres. This is attributable to asymmetries in the distributions of the 'sources' COF₂ / 430 COCIF and their precursors, due to differences in the meridional Brewer-Dobson circulation, 431 and to the stronger descent of air associated with the winter polar vortex in the southern 432 hemisphere; for example, compare the enhanced ACE-FTS HF VMRs near the southern pole 433 in August 2010 with those near the northern pole in February 2010 at ~25 km. An additional 434 source of asymmetry in the COF₂ distribution, which directly influences the HF distribution, 435 arises from the temperature-dependent loss reaction of the COF₂-precursor CHClF₂ (HCFC-

436 22) with OH, leading to a secondary COF_2 maximum at southern hemisphere high-latitudes 437 in the summer mid-stratosphere (~10 K warmer than the corresponding location in the 438 northern hemisphere) (Harrison et al., 2014); for example, compare the ACE-FTS HF 439 southern hemisphere VMRs at ~30-35 km in January 2010 with those in the northern 440 hemisphere in July 2010.

441 In addition to side-by-side comparisons of model and observation, the chemistry 442 scheme in SLIMCAT can be tested by comparing (chemically related) tracer-tracer 443 correlations for model and observation; only ACE measurements of fluorine-containing HF 444 'precursors' are available for this purpose. It is widely known that all long-lived species in 445 the stratosphere have compact correlations, even if there is no chemical link between them. 446 As explained by Plumb and Ko (1992), two tracers with lifetimes longer than quasi-447 horizontal mixing time scales should be in 'slope equilibrium' and produce a compact 448 correlation. Species with lifetimes longer than vertical transport time scales will also be in 'gradient equilibrium' and the compact correlation will be linear. Furthermore, relative 449 450 lower-stratospheric lifetimes of long-lived species (with stratospheric sinks via photolysis or 451 reaction with $O(^{1}D)$) under gradient equilibrium can be derived from the linear slope of the 452 tracer-tracer correlation (Chipperfield et al., 2014).

453 In the lower stratosphere COF₂ and COClF can be regarded as long-lived tracers 454 (local lifetimes of many years). Therefore, their tracer isopleths should follow the typical 455 tropopause-following contours of any long-lived tracer. In this sense, COF₂ and COCIF are 456 analogous to NO_y which is produced from N₂O. Figure 4 contains correlation plots between 457 COF₂ and its major source, CFC-12, over the two latitude bands 65-70°S and 65-70°N for 458 two months each over the period September 2009 – August 2010. Comparisons are made at 459 high latitudes, where ACE-FTS observations are more plentiful, and for individual months to 460 ensure that time trends in the source gas VMRs are minimised. The figure reveals that COF₂ is indeed long-lived enough to show a good anti-correlation with CFC-12 in the lower 461 462 stratosphere. Furthermore, agreement between the model and observations is good although 463 there are a few discrepancies around the region of maximum COF₂ VMR; these are due to 464 issues surrounding the scaled a priori used in the retrieval for this altitude region of the 465 profile where the spectral signal has dropped to within the noise level (refer to Harrison et al. 466 (2014) for more details).

Figure 5 contains correlation plots between COClF and its major source, CFC-11, for the same conditions as in Figure 4. Unlike for $COF_2 / CFC-12$, the agreement between model and ACE-FTS is particularly poor and the model overestimates the peak observed 470 values of COCIF; this can also be observed in Figure 3. There are several possible reasons 471 for this. Firstly, as the modelled VMRs are ~50 % higher than the ACE-FTS VMRs, the 472 modelled COCIF lifetime might be too long, i.e. the model underestimates the COCIF loss 473 processes. This would result in the calculated HF VMRs being slightly lower than they 474 should be, probably by less than a percent, but certainly by less than the uncertainty of the 475 ACE-FTS measurements. An additional SLIMCAT calculation with the COCIF lifetime 476 lowered by a third does improve the agreement with observations. Secondly, the COCIF 477 sources might be overestimated, however SLIMCAT calculations for CFC-11 reveal good 478 agreement with ACE-FTS observations, generally within 10% (Brown et al., 2011). 479 Additionally, the chemistry could be more complicated with additional destruction routes 480 missing from the model. Lastly, there could be a problem with the ACE-FTS retrieval itself. 481 The COCIF linelist used in the ACE-FTS retrieval was taken from the ATMOS database and 482 is described in the literature as 'very crude' (Perrin et al., 2011). At the time v3.0 data were 483 first released, this was the best linelist available, however a new and improved linelist has 484 subsequently been generated (Perrin et al., 2011), in which the band intensities are taken 485 from quantum-mechanical calculations. ACE-FTS COCIF retrievals for a handful of 486 occultations have been carried out using the new linelist, however there is no improvement in 487 the disagreement with SLIMCAT.

488 It is expected that the sum of all fluorine source gas VMRs (not including those 489 which have very long lifetimes compared with the period of observations, e.g. CF₄ and SF₆) 490 is anti-correlated with total F_y VMR (HF + 2COF₂ + COClF) in a conservative way (i.e. the 491 total adds up to a constant). As the ACE-FTS does not measure every source gas, and some 492 minor species have known biases (Brown et al., 2011), we only compare total F_y against the 493 sum of the major source gases, taken as CFC-11, CFC-12, and HCFC-22. The good 494 agreement between plots for SLIMCAT and ACE-FTS (see Figure 6) confirms that the 495 discrepancy in modelled and retrieved COCIF VMRs has a minimal effect on the overall 496 agreement between model and observation for HF.

497

498 **6. Trends**

Since HF has no chemical sink, with only minor losses arising from rainout in the troposphere and photolysis in the mesosphere, and since the atmosphere contains many longlived fluorine source gases, the overall HF atmospheric abundance has been increasing for many years and is expected to increase in the foreseeable future. In this section trends in ACE-FTS, HALOE, GOZCARDS and SLIMCAT time series are quantified as a function of altitude and latitude. A number of previous studies have quantified trends; for example, a linear trend of $8.5 \pm 1.0\%$ /year (1977 to 1986) (Zander et al., 1987) and $0.48 \pm 0.25\%$ /year (2000 to 2009) (Kohlhepp et al., 2012) in total columns measured at Jungfraujoch (46.5°N latitude, 8.0°E longitude), and 0.74 ± 0.2 %/year (between 30°S and 30°N) derived from ACE-FTS data for 2004 to 2010 (Brown et al., 2011).

509 Prior to the calculation of ACE-FTS, HALOE, and SLIMCAT trends, we derived 510 time series as a function of altitude (on the ACE-FTS grid) and latitude (in 10° bins). Figure 511 7 illustrates the ACE-FTS, HALOE and SLIMCAT time series for HF between 1991 and 512 2013 at selected altitudes for six of the latitude bins; for ease of viewing, error bars are not 513 shown. The annual cycle is clearly visible in each time series, a result of the seasonality of 514 the main 'source', COF₂ (Harrison et al., 2014), and careful inspection of Figure 7 reveals 515 that as expected the phase of this cycle is opposite in each hemisphere. The amplitude is 516 largest at high southern latitudes (note the maxima at 29.5 km for the 60° - 70°S plot), due to the descent of HF-rich air in southern winter polar vortices. Note also evidence of the quasi-517 518 biennial oscillation (QBO) signal in the tropical plots.

519 Overall the agreement between SLIMCAT and observations presented in Figure 7 is 520 good, however obvious biases are present. In Section 5 it was discussed that ACE v3.0 is biased high relative to HALOE by ~10%. HALOE VMRs are biased low relative to 521 SLIMCAT, generally by between ~5 and 15 %, although SLIMCAT is biased low relative to 522 523 HALOE by up to ~20 % between ~20 and 30 km in the 0-30°N region. Additionally, there is 524 a discrepancy in the observed and calculated annual cycle structure over the tropics, e.g. 10° -525 20°N at 34.5 km. In terms of bias, ACE-FTS v3.0/3.5 data generally agree with SLIMCAT 526 to within \pm 5 %, except over much of the lower stratosphere (below 30 km) where SLIMCAT 527 is biased low by at least ~5-15 %, peaking at ~20 % in the 0-30°N region and ~25-35 % at 528 the southern and northern high latitudes (poleward of 50°).

529 Figure 8 illustrates the GOZCARDS and SLIMCAT time series for HF plotted in the 530 same manner as Figure 7. Recall that the GOZCARDS HF data product is a merging of the 531 HALOE v19 and ACE-FTS v2.2 HF datasets, with the relative bias between the datasets 532 removed. The agreement between SLIMCAT and the HALOE component of GOZCARDS 533 above ~30 km is reasonably good, however at lower altitudes there are several regions in 534 which the low bias of SLIMCAT is significantly larger than presented in Figure 7, in 535 particular below 20 km near the south pole, and between ~ 20 and 30 km in the northern 536 hemisphere, where the bias peaks at ~35 % in the 0-30°N region. The ACE-FTS component 537 of GOZCARDS generally agrees with SLIMCAT to within \pm 5 % above 30 km at the tropics and above 25 km in the polar regions. At altitudes lower than these SLIMCAT is biased low, for example by $\sim 5 - 10$ % at latitudes above 50 °N and up to 25 % lower between 20 and 30 km in the 0-30°N region.

The GOZCARDS merging process for HF relied only on the relative bias between the HALOE v19 HF and ACE-FTS v2.2 HF datasets. In this study, it is not possible to comment on systematic or absolute biases. However, regardless of the absolute biases of the various datasets, it is clear that SLIMCAT tends to consistently underestimate HF VMRs at low altitudes (below 30 km at the tropics and 25 km at the poles) relative to those at higher altitudes.

547 HF trends for the ACE-FTS, HALOE, and SLIMCAT time series (trends were not 548 considered for GOZCARDS as this is a merged dataset directly related in a multiplicative 549 fashion to the original datasets) at each altitude within each latitude bin have been calculated 550 for three time periods from monthly percentage anomalies in HF zonal means, $C^{z,\theta}(n)$, 551 defined as

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$$C^{z,\theta}(n) = 100 \frac{VMR^{z,\theta}(n) - \sum_{m=1}^{12} \delta_{nm} \overline{VMR}^{z,\theta}(m)}{\sum_{m=1}^{12} \delta_{nm} \overline{VMR}^{z,\theta}(m)},$$
(7)

where *n* is a running index from month zero to month n-1, $VMR^{z,\theta}(n)$ is the corresponding 553 554 mixing ratio at altitude z and latitude θ , $\overline{VMR}^{z,\theta}(m)$ is the average of all zonal means for each 555 of the twelve months, m, and δ_{nm} , although not used in its strict mathematical sense, is one 556 when index n corresponds to one of the months m and is zero otherwise (Harrison et al., 557 2014). Such an approach essentially removes the annual cycle and the effect of biases in 558 VMRs; the trend, in units of %/year, is simply equated to the 'slope' of the linear regression 559 between $C^{z,\theta}(n)$ and the dependent variable n/12. The inclusion of additional terms such as 560 the annual cycle and its harmonics resulted in no additional improvement in the regression. 561 The three time periods considered are January 2004 to December 2012 (ACE-FTS, 562 SLIMCAT), October 1991 to December 1997 (HALOE, SLIMCAT), and January 1998 to November 2005 (HALOE, SLIMCAT). The HALOE time series was split into two periods 563 564 for which HF growth could be modelled linearly. Errors have been explicitly treated in the 565 linear regression of the ACE and HALOE data, but not the SLIMCAT outputs.

Figure 9 presents the trends in the growth of HF (percent per year) (January 2004 to December 2012) for ACE and SLIMCAT as a function of latitude and altitude, up to the top of the ACE-FTS retrieval range. The ACE-FTS plot in Figure 9 indicates that between 2004

569 and 2012, HF has increased most rapidly (2 - 3 %/year) at altitudes below ~25 km in the 570 northern hemisphere and at ~35 km near the equator. Similarly, HF has decreased most 571 rapidly in the southern hemisphere below \sim 35 km and in the northern hemisphere between \sim 572 30 and 35 km. The SLIMCAT plot in the second panel contains a number of features which 573 agree well with those in the ACE plot. In particular, note the region of negative trends in the southern hemisphere below $\sim 30 - 35$ km, peaking at -3.5 - -4.0 %/year, the region of high 574 positive trends in the northern hemisphere below ~ 30 km, peaking at 4.5 - 5.0 %/year, the 575 small region of positive trend at \sim 35 km near the equator, peaking at 2.0 – 2.5 %/vear, and 576 577 the slightly larger region of negative trend at $\sim 30 - 40$ km at 0° to 30°N, peaking at -1.0 - -578 1.5 %/year. However, the magnitudes of the SLIMCAT trends in the lower stratosphere are 579 biased high compared with the ACE measurements.

580 An additional SLIMCAT run has been performed with dynamics arbitrarily annually 581 repeating those for the year 2000; results from this run give a 'clean' HF signal without the complication of changes in stratospheric dynamics. Results from this run are included in the 582 583 times series plots in Figure 7; the annual repetition in dynamical structure reveals a clearer 584 signal of the long-term chemical changes. 'Clean' HF trends for 2004 – 2012 calculated in 585 the same manner as before are plotted in the lowest panel of Figure 9, revealing trends distributed relatively uniformly throughout the stratosphere with values between 0 and 1 586 %/year. This indicates that the variations in trends observed for the full SLIMCAT run result 587 from changes in stratospheric dynamics over the observation period. The information on 588 589 stratospheric circulation is provided solely by the analyses used to force the SLIMCAT 590 calculations. Similar changes due to stratospheric dynamics were observed for COF₂ 591 (Harrison et al., 2014). Moreover, Ploeger et al. (2015) used a Lagrangian chemical transport 592 model, also forced by ECMWF ERA-Interim reanalyses, to look at variations in stratospheric 593 age-of-air (AoA) over the period 1988-2013. They compared their model results with 594 estimates derived from MIPAS satellite observations for 2002-2012. During the period of 595 MIPAS observations they found that stratospheric AoA decreased in the lower stratosphere 596 but showed interhemispheric differences in the trend above about 20 km. Also, despite the 597 ongoing monotonic decrease of near-surface chlorine source gases, recent ground-based and 598 satellite remote-sensing measurements have shown a significant increase in hydrogen 599 chloride (HCl), the main stratospheric chlorine reservoir, in the lower stratosphere of the 600 Northern Hemisphere between 2007 and 2011 (Mahieu et al., 2014). By comparison to 601 similar SLIMCAT simulations as used here, this trend 'anomaly' was attributed to multiyear 602 variability in the stratospheric circulation and dynamics.

603 Together, the studies discussed above paint a consistent picture whereby variability 604 in stratospheric transport, which varies with altitude and hemisphere, significantly modifies 605 the observed trends in long-lived tracers. This variability seems to be well captured by 606 reanalysis products such as ERA-Interim. Even if these tracers have monotonic VMR trends 607 in the troposphere, this dynamical variability can lead to complicated behaviour in the 608 stratosphere and must therefore be accounted for when using observations to determine 609 underlying chemical trends. A detailed analysis of the changing stratospheric dynamics that 610 are responsible for the observed trends in HF and other species is beyond the scope of this 611 work and would require a coupled chemistry-dynamical model.

Trends have similarly been derived for the two HALOE periods. HALOE plots 612 613 corresponding to the ACE-FTS plots in Figure 9 can be found in Figures 10 (1991 – 1997) 614 and 11 (1998 – 2005). The HALOE trends in Figure 10 peaking in the northern hemisphere 615 between 0° and 40°N broadly agree with those calculated by SLIMCAT in the same region, however SLIMCAT calculates smaller trends at the lowest altitudes and generally 616 617 underestimates the trends in the southern hemisphere. The differences between the full and 618 fixed-dynamics SLIMCAT runs show the impact of dynamical variability; the fixed-619 dynamics run provides a clean chemical signal. Of the three periods considered, the 620 comparison between 1991 – 1997 HALOE trends and those calculated from SLIMCAT is the poorest. The North-South asymmetry in trends for the full SLIMCAT calculation, which 621 622 does not agree with observations, must be due to dynamical variability in the model, with the 623 dynamics imposed solely by the ECMWF analyses. Over the measurement period, the 624 quality of these analyses may vary depending on the available datasets used for the 625 assimilation, but it is very difficult to test how realistic the stratospheric transport is. There 626 are only a handful of other height-resolved datasets that test this aspect of the stratospheric 627 circulation, e.g. ozone (Dhomse et al., 2015). As evidenced by Figure 11, however, the HALOE trends for 1998 – 2005 agree better with SLIMCAT than for the 1991 – 1997 period, 628 629 with 'background' trends generally between 0.5 and 1.5 %/year. In fact, there is very little 630 variability over the majority of the plotted range.

Overall global trends in HF, weighted at each altitude and latitude by $\cos^2(\text{latitude}^\circ)$ and the average VMR, have been calculated from the three time series using errors determined from the linear regression; these trends are listed in Table 5. The observed HF trends reveal a substantial slowing down in the rate of increase of HF by ~90 % from the mid-1990s over the next 15 years, namely from 4.97 ± 0.12 %/year (1991-1997) to 1.12 ± 0.08 %/year (1998-2005) to 0.52 ± 0.03 %/year (2004-2012). In addition to direct

637 stratospheric ozone recovery (e.g. Chipperfield et al., 2015), this marked decline in the 638 growth rate of HF is a particularly important marker for the success of the Montreal Protocol, 639 and should drop even further once HCFC-22 is phased out in developing countries over the 640 coming years. Global trends calculated by SLIMCAT for the HALOE (1998 - 2005) and 641 ACE-FTS (2004 – 2012) time series, 1.10 %/year and 0.48 %/year, respectively, agree very well with observations, however for the 1991 – 1997 HALOE period the model produces a 642 643 value ~20 % lower (4.01 %/year). Again, the reason for this is not completely clear, but is 644 likely related to the ECMWF analysis used to drive the dynamics in the SLIMCAT 645 calculation.

646

647 **7. Conclusions**

648 Hydrogen fluoride (HF) is the most abundant fluorine reservoir in the stratosphere 649 with main sources arising from the atmospheric degradation of CFC-12 (CCl₂F₂), CFC-11 650 (CCl₃F), HCFC-22 (CHClF₂), and CFC-113 (CCl₂FCClF₂), ozone-depleting species whose 651 emissions are anthropogenic. Monitoring the growth of stratospheric HF is therefore an 652 important marker for the success of the Montreal Protocol.

Global distributions and trends of stratospheric HF have been determined from ACE-FTS (2004 –) and HALOE (1991 – 2005) data. Based on the overlap period between datasets, ACE-FTS HF measurements are biased high by ~10% relative to HALOE. The observations have been compared with the output of SLIMCAT, a three-dimensional CTM, and the agreement is generally good, although SLIMCAT tends to underestimate HF VMRs at low altitudes (below 30 km at the tropics and 25 km at the poles) relative to those at higher altitudes.

The observed global HF trends reveal a substantial slowing down in the rate of 660 increase of HF since the 1990s: 4.97 ± 0.12 %/year (1991-1997; HALOE), 1.12 ± 0.08 661 %/year (1998-2005; HALOE), and 0.52 ± 0.03 %/year (2004-2012; ACE-FTS), indicating 662 663 the effectiveness of the Montreal Protocol in phasing out the principal precursor species. For the same periods, SLIMCAT calculates trends of 4.01 %/year, 1.10 %/year, and 0.48 %/year, 664 respectively. The observations also reveal variations in the HF trends with latitude and 665 altitude, for example between 2004 and 2012 HF actually decreased in the southern 666 667 hemisphere below ~35 km. SLIMCAT calculations broadly agree with these observations, 668 most notably between 2004 and 2012. Such variations are attributed to variability in 669 stratospheric dynamics over the observation period.

The ACE-FTS is the only satellite instrument currently making measurements of HF, and continues to operate with only minor loss in performance since its launch. It will therefore be possible to extend the HF time series to the present day and beyond, and subsequently extend the comparison with SLIMCAT.

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675 Author contribution

J. J. Harrison devised the study and performed the data analysis. C. D. Boone and P.
F. Bernath provided the ACE-FTS data. J. Russell III provided the HALOE HF data. L.
Froidevaux and J. Anderson provided the HF GOZCARDS data. M. P. Chipperfield and S.
S. Dhomse ran the SLIMCAT model and provided additional explanation of the outputs. J. J.
Harrison prepared the manuscript with contributions from M. P. Chipperfield and the other
co-authors.

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693

694 **Figure Captions**

695

Figure 1. A comparison between ACE-FTS and SLIMCAT HF zonal means (September
2009 to August 2010). A full discussion of the seasonal variation in the HF distribution is
provided in the text.

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Figure 2. A comparison between ACE-FTS, HALOE and SLIMCAT HF zonal means
(September 2004 to August 2005). The ACE-FTS and HALOE time series of measurements
overlap during the period January 2004 to November 2005.

- Figure 3. A comparison of COF₂ and COClF zonal means from ACE-FTS and SLIMCAT
 for October 2009, and February and August 2010.
- 706

Figure 4. Correlation plots between coincident CFC-12 and COF₂ ACE-FTS observations
 and SLIMCAT calculations for November 2009 / July 2010 65-70°S and January / May 2010

- 709 65-70°N. The error bars represent the standard deviations in the ACE-FTS VMRs.
- 710
- Figure 5. Correlation plots between coincident CFC-11 and COCIF ACE-FTS observations
 and SLIMCAT calculations for November 2009 / July 2010 65-70°S and January / May 2010
- 713 65-70°N. The error bars represent the standard deviations in the ACE-FTS VMRs.
- 714
- **Figure 6**. Correlation plots between coincident ACE-FTS observations and SLIMCAT calculations of total 'major' organic fluorine, based on CFC-11, CFC-12, and HCFC-22, and total inorganic fluorine, F_y, for November 2009 / July 2010 65-70°S and January / May 2010
- 718 65-70°N. The error bars represent the standard deviations in the ACE-FTS VMRs.
- 719

Figure 7. The HALOE, ACE-FTS and SLIMCAT HF time series for selected altitude – latitude bin combinations. Observations are plotted between October 1991 and December 2012. Overlaid are the time series from a SLIMCAT run with dynamics arbitrarily annually repeating those for the year 2000; this provides a clearer signal of the long-term chemical changes without the complication of variations in stratospheric dynamics.

- 725
- Figure 8. The GOZCARDS and SLIMCAT HF time series for selected altitude latitude bin
 combinations. Observations are plotted between October 1991 and September 2010.
- 728
- Figure 9. Trends in the growth of HF (% yr⁻¹; January 2004 to December 2012) for ACEFTS and SLIMCAT as a function of latitude and altitude. A full discussion of these trends is
 provided in the text.
- 732

Figure 10. Trends in the growth of HF (% yr⁻¹; October 1991 to December 1997) for
HALOE and SLIMCAT as a function of latitude and altitude. A full discussion of these
trends is provided in the text.

Figure 11. Trends in the growth of HF (% yr⁻¹; January 1998 to November 2005) for
HALOE and SLIMCAT as a function of latitude and altitude. A full discussion of these
trends is provided in the text.

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918

920 Tables

Table 1: Microwindows for the v3.0/v3.5 ACE-FTS hydrogen fluoride retrieval.

Centre Frequency	Microwindow width	Lower altitude	Upper altitude
(cm^{-1})	(cm^{-1})	(km)	(km)
3787.60	1.60	40	2.00E+16 ^b
3788.28	0.60	12	40
3792.65 ^a	0.40	20	40
3833.70	0.80	16	40
3834.30	1.60	40	$9.00E+15^{b}$
3877.60	0.80	12	$9.00E+15^{b}$
3920.15	0.70	25	9.00E+15 ^b
4000.87	0.65	12	9.00E+15 ^b
4038.82	1.00	12	$9.00E+15^{b}$
4075.35	0.80	25	$9.00E+15^{b}$
4109.75	0.80	25	$2.00E+16^{b}$

923 ^{*a*} Included to improve results for interferer HDO.

 b Upper altitude given in atmospheric density units, molecules cm⁻³.

927 Table 2: Interferers in the v3.0/v3.5 ACE-FTS hydrogen fluoride retrieval.

Malagula	Lower altitude	Upper altitude limit
Wiolecule	limit (km)	(km)
H ₂ O	12	65
$H^{18}OH$	12	50
$H^{17}OH$	12	40
HDO	12	25
CO ₂	12	40
O3	12	38
CH4	12	30
$OC^{18}O$	12	20
N ₂ O	12	30

Table 3: Sources of systematic uncertainty in the ACE-FTS v3.0/v3.5 hydrogen fluorideretrieval.

Source	Symbol	Fractional value
HF spectroscopy	$\mu_{\scriptscriptstyle spec}$	0.04
Spectral interferers	$\mu_{ m int}$	0.01
Temperature	$\mu_{\scriptscriptstyle T}$	0.01
Altitude	μ_z	0.03
ILS	$\mu_{{\scriptscriptstyle I\!L\!S}}$	0.09

932

933 Table 4: Fluorine source gases in the SLIMCAT chemical scheme and their atmospheric

934 degradation products.

Source g	Product gases	
Commercial name	Chemical formula	
CFC-11	CCl ₃ F	COCIF
CFC-12	CCl ₂ F ₂	COF ₂
CFC-113	CCl ₂ FCClF ₂	$COClF + COF_2$
CFC-114	CClF ₂ CClF ₂	$2COF_2$
CFC-115	CClF ₂ CF ₃	$2\text{COF}_2 + \text{HF}$
HCFC-22	CHClF2	COF ₂
HCFC-141b	CH ₃ CCl ₂ F	COClF
HCFC-142b	CH ₃ CClF ₂	COF ₂
HFC-23	CHF3	$COF_2 + HF$
HFC-134a	CH ₂ FCF ₃	$COF_2 + 2HF$
HFC-152a	CH ₃ CHF ₂	$2COF_2$
Halon 1211	CBrClF ₂	COF ₂
Halon 1301	CBrF3	$COF_2 + HF$
Halon 2402	CBrF2CBrF2	4HF
COCI	\mathbf{F}^{a}	HF
COF	2^a	2HF

- 935 ^{*a*} These are not source gases, but their degradation products are included for completion.
- 936

937 Table 5: Trends (%/year) derived from the HALOE v19 and ACE-FTS v3.0/v3.5 HF

938 observations.

Dataset	Period	Observed trend	SLIMCAT trend
HALOE	1991-1997	4.97 ± 0.12	4.01
HALOE	1998-2005	1.12 ± 0.08	1.10
ACE-FTS	2004-2012	0.52 ± 0.03	0.48



250 500 750 1000 1250 1500 1750 2000 2250 HF VMR (ppt)

















GOZCARDS

— SLIMCAT

ACE HF trends (January 2004 to December 2012)



SLIMCAT HF trends (January 2004 to December 2012)



SLIMCAT HF trends (January 2004 to December 2012) - fixed to 2000 dynamics





HALOE HF trends (January 1991 to December 1997)



SLIMCAT HF trends (January 1991 to December 1997)









HALOE HF trends (January 1998 to November 2005)



SLIMCAT HF trends (January 1998 to November 2005)







