Airborne measurements of cirrus-activated C\textsubscript{2}Cl\textsubscript{4} depletion in the upper troposphere with evidence against Cl reactions

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[1] Airborne whole air samples collected over the western Pacific in spring, 2001 showed depletion of tetrachloroethylene (C\textsubscript{2}Cl\textsubscript{4}) in every upper tropospheric (UT) air parcel that had interacted with large areas of cirrus less than three days upwind. The amount of C\textsubscript{2}Cl\textsubscript{4} depletion showed a negative correlation with time since the interaction, consistent with the C\textsubscript{2}Cl\textsubscript{4}-depleted air parcels mixing with the ambient air as they moved downwind of the cirrus. Ethane and C\textsubscript{2}Cl\textsubscript{4} both react relatively quickly with atomic chlorine (Cl) but ethane was not significantly depleted in these same air parcels, indicating that the C\textsubscript{2}Cl\textsubscript{4} depletion cannot be attributed to Cl chemistry alone. Based on the minimum ethane depletion that can be detected by our measurements, a daytime upper limit of roughly 3 \times 10\textsuperscript{9} atom cm\textsuperscript{-3} is indirectly estimated for heterogeneous chlorine activation by cirrus clouds in the UT in tropical- and mid-latitudes during spring. INDEX TERMS: 0317 Atmospheric Composition and Structure: Chemical kinetic and photochemical properties; 0320 Atmospheric Composition and Structure: Cloud physics and chemistry; 0365 Atmospheric Composition and Structure: Troposphere—composition and chemistry; 0368 Atmospheric Composition and Structure: Troposphere—constituent transport and chemistry; KEYWORDS: Tetrachloroethene, chlorine activation, heterogeneous chemistry, TRACE-P, cirrus. Citation: Simpson, I. J., O. W. Wingenter, D. J. Westberg, H. E. Fuelberg, C. M. Kiley, J. H. Crawford, S. Meinardi, D. R. Blake, and F. S. Rowland, Airborne measurements of cirrus-activated C\textsubscript{2}Cl\textsubscript{4} depletion in the upper troposphere with evidence against Cl reactions, Geophys. Res. Lett., 30(20), 2025, doi:10.1029/2003GL017598, 2003.

1. Introduction

[2] Heterogeneous reactions on the ice surfaces of polar stratospheric clouds are known to enhance the conversion of HCl and Cl\textsubscript{2}NO\textsubscript{2} to more reactive species such as Cl\textsubscript{2}, thereby playing a critical role in activating Cl and augmenting stratospheric ozone depletion [see Solomon et al., 1997 and references therein]. Several modeling studies suggest that cirrus clouds are likely to similarly activate chlorine in the tropopause region, with the magnitude and duration of resulting ClO enhancement depending upon season, latitude and altitude [Borrmann et al., 1996; Solomon et al., 1997; Bregman et al., 2002]. In support of this, ice particles in cirrus (T = 235 K) and aircraft condensation trails (T = 220 K) are reported to take up large amounts of atmospheric HCl [Thibert and Dominic, 1997], with the uptake coefficient increasing with decreasing temperature [Hynes et al., 2001].

[3] Airborne studies have investigated chlorine activation by cirrus near the tropopause and in the lowermost stratosphere for various latitudes and seasons. Lelieveld et al. [1999] observed enhanced CO/ethane ratios over Sweden, from which they inferred mean diurnal Cl values of up to 1.0 \times 10\textsuperscript{9} atom cm\textsuperscript{-3} based on heterogeneous activation. Enhanced ClO has been observed at polar latitudes during winter (averaging 15–20 pptv; Thornton et al., 2003), and at midlatitudes during spring (>90 pptv; Keim et al., 1996) and winter (2.7 pptv in cirrus and up to 70 pptv near the tropopause; Borrmann et al., 1997). The Thornton et al. and Borrmann et al. studies refrained from or were not able to link the ClO enhancements to chlorine activation by cirrus, and the results by Keim et al. were consistent with heterogeneous reactions. By contrast, Smith et al. [2001] suggest that midlatitude ozone loss is not controlled by chlorine activation by cirrus near the tropopause, based on a consistent absence of ClO enhancements in summer. However, this disparity among the midlatitude studies may reflect different conditions (water subsaturation in the case of Smith et al.) rather than conflicting results [Thornton et al., 2003]. Here we use aircraft measurements of hydrocarbons and halocarbons at midlatitudes during spring, together with backward trajectories and satellite images, to indirectly investigate chemical processing by cirrus in the UT between 10–12 km. This is the first study of its kind in the UT.

2. Experimental

[4] The measurements were made during the Transport and Chemical Evolution over the Pacific (TRACE-P) experiment (February–April, 2001) [Jacob et al., 2003], using airborne whole air sampling and laboratory gas chromatographic analysis with flame ionization-, electron capture-, and mass spectrometer detection [Colman et al., 2001]. During 17 science flights of the NASA DC-8 aircraft, 2725 whole air samples were collected between 0–12 km, including 231 between 10–12 km. The samples were collected into individual conditioned, evacuated 2-L stain-
influence. Secondly, because the ratios become less reliable.

Selected Compounds with OH (O3 > 100 ppbv and CO < 50 ppbv) were removed from UT. First, air parcels that showed stratospheric influence any ratio increases were consistent with Cl chemistry in the

four gases have similar \( k_{\text{OH}} \) values whereas \( C_2\text{Cl}_4 \) and ethane have much larger \( k_{\text{Cl}} \) values (Table 1), and an increase in these ratios may indicate recent Cl chemistry. Ethane and \( C_2\text{Cl}_4 \) have previously been used to evaluate tropospheric Cl levels [Rudolph et al., 1996; Singh et al., 1996; Wingenter et al., 1999].

The possible influence of Cl chemistry in the UT was investigated using ratios of compounds with similar rate constants for reaction with OH (\( k_{\text{OH}} \)) and quite different rate constants for reaction with Cl (\( k_{\text{Cl}} \)), specifically \( C_2\text{Cl}_2/ C_2\text{Cl}_4, \) CHCl3/CH2Cl2, ethane/CHCl3, and CHCl3/ethane. All four gases have similar \( k_{\text{OH}} \) values whereas \( C_2\text{Cl}_4 \) and ethane have much larger \( k_{\text{Cl}} \) values (Table 1), and an increase in these ratios may indicate recent Cl chemistry. Ethane and \( C_2\text{Cl}_4 \) have previously been used to evaluate tropospheric Cl levels [Rudolph et al., 1996; Singh et al., 1996; Wingenter et al., 1999].

The following data editing procedures ensured that any ratio increases were consistent with Cl chemistry in the UT. First, air parcels that showed stratospheric influence (\( O_3 > 100 \) ppbv and \( CO < 50 \) ppbv) were removed from further analysis. Of 14 air parcels sampled for at least 20 minutes at altitudes \( \geq 10.7 \) km, two showed stratospheric influence. Secondly, because the ratios become less reliable at very low mixing ratios, only mixing ratios \( > 5 \times \text{LOD} \) were used. Though this cutoff prevented us from using other compound pairs such as benzene/ethyne, the mixing ratios of \( C_2\text{Cl}_4 \) and ethane were always \( > 5 \times \text{LOD} \) in the UT air samples (mean \( \pm 1\sigma \) of 1.9 \( \pm 1.3 \) pptv for \( C_2\text{Cl}_4 \) and 584 \( \pm 205 \) pptv for ethane). Thirdly, the ratios were inspected to ensure that any enhancements were caused by depleted \( C_2\text{Cl}_4 \) or ethane (consistent with possible Cl chemistry) rather than enhanced \( C_2\text{Cl}_2 \) or CHCl3. In our experience \( C_2\text{Cl}_4 \) and \( C_2\text{Cl}_2 \) emissions are often co-located, and here CHCl3 correlated more poorly with \( C_2\text{Cl}_4 \) than did \( C_2\text{Cl}_2 \). Fourthly, because Cl chemistry has been previously detected in the marine boundary layer (MBL), the air parcels were tested for possible deep convective influence from the MBL. Five-day backward trajectories were calculated every 5 minutes for each flight leg using a kinematic model described in Fuelberg et al. [1996, 2003]. Of the 12 air parcels considered here, seven had 5-day backward trajectories that remained in the UT, with no evidence for MBL influence.

| Compound  | \( k_{\text{OH}} \) (cm\(^3\) molec\(^{-1}\) s\(^{-1}\)) | \( k_{\text{Cl}} \) (cm\(^3\) molec\(^{-1}\) s\(^{-1}\)) |
|-----------|-----------------|-----------------|
| CHCl3     | 3.3 \( \times 10^{-14} \) | 2.0 \( \times 10^{-14} \) |
| CH2Cl2    | 3.2 \( \times 10^{-14} \) | 6.7 \( \times 10^{-14} \) |
| Ethane    | 7.6 \( \times 10^{-14} \) | 5.3 \( \times 10^{-14} \) |
| C2Cl4     | 4.0 \( \times 10^{-14} \) | 5.8 \( \times 10^{-14} \) |

\( ^*\text{DeMore et al. [1997].} \)

\( ^6\text{Atkinson [1997].} \)

[5] After each flight, the canisters were shipped to our UCI laboratory and analyzed for 78 compounds within 10 days of sampling. Ethane, \( C_2\text{Cl}_4 \), methylene chloride (\( \text{CH}_2\text{Cl}_2 \)) and chloroform (\( \text{CHCl}_3 \)) were used in this study. Their limits of detection (LOD) are: ethane (3 pptv); \( C_2\text{Cl}_4 \) (0.05 pptv); \( \text{CH}_2\text{Cl}_2 \) (1 pptv); \( \text{CHCl}_3 \) (0.1 pptv). The precisions are: ethane (0.5%); \( C_2\text{Cl}_4 \) and \( \text{CHCl}_3 \) (1%); \( \text{CH}_2\text{Cl}_2 \) (5%). The accuracy is 5% for ethane and \( \text{CHCl}_3 \), and 10% for \( C_2\text{Cl}_4 \) and \( \text{CH}_2\text{Cl}_2 \).

[6] The possible influence of Cl chemistry in the UT was investigated using ratios of compounds with similar rate constants for reaction with OH (\( k_{\text{OH}} \)) and quite different rate constants for reaction with Cl (\( k_{\text{Cl}} \)), specifically \( C_2\text{Cl}_2/ C_2\text{Cl}_4, \) CHCl3/ \( C_2\text{Cl}_4 \), \( \text{CH}_2\text{Cl}_2/ \text{ethane} \) and CHCl3/ethane. All four gases have similar \( k_{\text{OH}} \) values whereas \( C_2\text{Cl}_4 \) and ethane have much larger \( k_{\text{Cl}} \) values (Table 1), and an increase in these ratios may indicate recent Cl chemistry. Ethane and \( C_2\text{Cl}_4 \) have previously been used to evaluate tropospheric Cl levels [Rudolph et al., 1996; Singh et al., 1996; Wingenter et al., 1999].

[7] The following data editing procedures ensured that any ratio increases were consistent with Cl chemistry in the UT. First, air parcels that showed stratospheric influence (\( O_3 > 100 \) ppbv and \( CO < 50 \) ppbv) were removed from further analysis. Of 14 air parcels sampled for at least 20 minutes at altitudes \( \geq 10.7 \) km, two showed stratospheric influence. Secondly, because the ratios become less reliable at very low mixing ratios, only mixing ratios \( > 5 \times \text{LOD} \) were used. Though this cutoff prevented us from using other compound pairs such as benzene/ethyne, the mixing ratios of \( C_2\text{Cl}_4 \) and ethane were always \( > 5 \times \text{LOD} \) in the UT air samples (mean \( \pm 1\sigma \) of 1.9 \( \pm 1.3 \) pptv for \( C_2\text{Cl}_4 \) and 584 \( \pm 205 \) pptv for ethane). Thirdly, the ratios were inspected to ensure that any enhancements were caused by depleted \( C_2\text{Cl}_4 \) or ethane (consistent with possible Cl chemistry) rather than enhanced \( C_2\text{Cl}_2 \) or CHCl3. In our experience \( C_2\text{Cl}_4 \) and \( C_2\text{Cl}_2 \) emissions are often co-located, and here CHCl3 correlated more poorly with \( C_2\text{Cl}_4 \) than did \( C_2\text{Cl}_2 \). Fourthly, because Cl chemistry has been previously detected in the marine boundary layer (MBL), the air parcels were tested for possible deep convective influence from the MBL. Five-day backward trajectories were calculated every 5 minutes for each flight leg using a kinematic model described in Fuelberg et al. [1996, 2003]. Of the 12 air parcels considered here, seven had 5-day backward trajectories that remained in the UT, with no evidence for MBL influence.

3. Results and Discussion

[9] Using the edited data set, \( \text{CH}_2\text{Cl}_2/ C_2\text{Cl}_4 \) and CHCl3/ \( C_2\text{Cl}_4 \) were notably enhanced in the UT, particularly between 11–12 km (Figure 1 shows \( \text{CH}_2\text{Cl}_2/ C_2\text{Cl}_4 \)). The enhanced ratios (depleted \( C_2\text{Cl}_4 \)) occurred during encounters with four separate air parcels sampled in the region 19–31°N and 129°E–160°W (Table 2). The average (\( \pm 1\sigma \)) \( \text{CH}_2\text{Cl}_2/ C_2\text{Cl}_4 \) and CHCl3/ \( C_2\text{Cl}_4 \) ratios from the four encounters (24 \( \pm 8 \) and 9 \( \pm 3 \) pptv/ppbv, respectively) were ~3 times larger than those measured for 25 background UT air samples at comparable altitudes (11–12 km; Table 2). By contrast, the average ratios of \( \text{CH}_2\text{Cl}_2/ \text{ethane} \) and CHCl3/ethane in the same four air parcels (25 \( \pm 4 \) and 10 \( \pm 2 \) pptv/ppbv, respectively) were not significantly different from the background ratios at 11–12 km, though we note that values for individual “depleted \( C_2\text{Cl}_4 \)” and “undepleted \( C_2\text{Cl}_4 \)” flights sometimes differed. Because \( C_2\text{Cl}_4 \) and ethane have similar \( k_{\text{OH}} \) and \( k_{\text{Cl}} \) values at UT

![Figure 1](image-url) Figure 1. Altitude profile of \( \text{CH}_2\text{Cl}_2/ C_2\text{Cl}_4 \) for air samples that passed the data editing procedures outlined in the text. Open circles are from the four air parcels that showed depleted \( C_2\text{Cl}_4 \).
temperatures and pressures (Table 1), the failure of ethane to corroborate the C$_2$Cl$_4$ findings indicates that Cl chemistry alone does not explain the observed C$_2$Cl$_4$ depletion.

[10] To understand the cause of the C$_2$Cl$_4$ depletion, all seven air parcels were tested for upwind interactions with cirrus. Interestingly, all four air parcels that showed elevated CH$_2$Cl$_2$/C$_2$Cl$_4$ and CHCl$_3$/C$_2$Cl$_4$ also had major interactions with cirrus (lasting 15–36 hours) 1–3 days upwind (Table 2). The cirrus were located over eastern Asia and the western Pacific ($5^\circ$–$40^\circ$N and $90^\circ$E–$160^\circ$W; see website) and were associated with frontal activity for the D4 and D18 air parcels and with tropical convection for D9 and D16. Using D4 as an example, the backward trajectory heights (8–13 km or 150–300 mb) overlap the estimated cirrus height (10–13 km or 225–240 K). At these temperatures the clouds are expected to be mostly ice with little supercooled water. Hygristor measurements from a Shanghai radiosonde sounding the day before sampling are near saturation from 5–11 km (200–500 mb), providing consistent evidence that the upper level air was humid and is expected to contain cirrus. All four air parcels showed similar evidence for interaction between the trajectories and cirrus.

[11] Of the three UT air parcels that do not show depleted C$_2$Cl$_4$, two do not interact with cirrus for at least 5 days upwind (Table 2). By contrast, the D13 air parcel shows an 11-hour interaction with cirrus between 2.7–3.1 days upwind, even though C$_2$Cl$_4$ was not depleted. This behavior was investigated by plotting the CH$_2$Cl$_2$/C$_2$Cl$_4$ ratio (normalized to the change of CH$_2$Cl$_2$/C$_2$Cl$_4$ with altitude in background air) against time since the upwind cirrus interaction (Figure 2a). Bearing in mind that only four data points are available in Figure 2a, the magnitude of C$_2$Cl$_4$ depletion shows a negative correlation with time ($r^2 = 0.86$ for an exponential fit), which is consistent with the C$_2$Cl$_4$-depleted air parcels mixing with the ambient air as they move downwind of the cirrus. Taking the ratio uncertainty into account, the normalized CH$_2$Cl$_2$/C$_2$Cl$_4$ ratio crosses the background value of 1 within the characteristic 3–5 day mixing time for an air parcel in the UT [R. Newell, pers. comm., 2002]. That is, the D13 air parcel is likely to have almost completely mixed with the ambient air by the time it was sampled, consistent with the observed lack of C$_2$Cl$_4$ depletion.

[12] Unlike CH$_2$Cl$_2$/C$_2$Cl$_4$, the ratio of CH$_2$Cl$_2$/ethane shows no correlation with time since the upwind cirrus interaction (Figure 2b). The depletion of C$_2$Cl$_4$ but not ethane following encounters with cirrus indicates a primary depletion mechanism other than reaction with Cl. We rule out reaction with bromine (Br) based on a very slow C$_2$Cl$_4$ + Br rate constant ($5 \times 10^{-15}$ at 220 K; Ramacher et al., 2001) and an estimated upper limit Br range of 0.5–2 pptv. Tetrachloroethene has a much lower vapour pressure than ethane, and in snow doping experiments at Summit, Greenland we have found that addition of equal amounts of C$_2$Cl$_4$ and ethane leads to a 10–100 fold greater amount of ethane downstream as a result of preferential C$_2$Cl$_4$ adsorption onto

### Table 2. Details of Seven UT Air Parcels That Were Inspected for Possible Influence From Cl Activation by Cirrus

| DC-8 Flight | Sampling Date (UTC) | Sampling Interval (UTC) | Main Sampling Alt. (km) | n | CH$_2$Cl$_2$/C$_2$Cl$_4$ (pptv/ppmv) | CHCl$_3$/C$_2$Cl$_4$ (pptv/ppmv) | CH$_2$Cl$_2$/Ethane (pptv/ppbv) | CHCl$_3$/Ethane (pptv/ppbv) | Time since cirrus (days) | Duration of interaction (hours) |
|-------------|---------------------|--------------------------|-------------------------|---|-----------------------------------|----------------------------------|---------------------------------|-------------------------------|-----------------------------|-------------------------------|
| D4          | Feb.26–27           | 23:40–0:49               | 11.9                    | 9 | 25 ± 6                           | 10 ± 4                           | 27 ± 3                          | 12 ± 2                        | 1.2–1.9                     | 18                            |
| D9          | Mar.11              | 6:30–6:51                | 11.9                    | 4 | 18 ± 5                           | 6 ± 2                            | 24 ± 3                          | 8 ± 1                         | 1.8–3.2                     | 36                            |
| D16         | Mar.30              | 1:08–1:53                | 11.0–11.3               | 9 | 22 ± 8                           | 8 ± 3                            | 25 ± 5                          | 9 ± 2                         | 1.4–2.3                     | 21                            |
| D18         | Apr.4               | 6:28–7:00                | 11.3                    | 5 | 32 ± 9                           | 11 ± 3                           | 24 ± 2                          | 9 ± 1                         | 0.8–1.4                     | 15                            |
| D3–20       | Feb.28–Apr.9        | n/a                      | 11–12                   | 25 | 9 ± 3                           | 3 ± 1                            | 30 ± 6                          | 11 ± 4                        | n/a                         | n/a                           |
| D5          | Feb.27–28           | 23:41–0:05               | 10.4–10.7               | 4 | 10 ± 1                           | 4 ± 1                            | 35 ± 1                          | 15 ± 1                        | none                        | none                          |
| D6          | Mar.4               | 5:44–6:10                | 10.7                    | 5 | 11 ± 1                           | 6 ± 1                            | 22 ± 2                          | 12 ± 2                        | none                        | none                          |
| D13         | Mar.21              | 1:37–2:24                | 10.7                    | 9 | 8 ± 1                            | 3 ± 1                            | 36 ± 2                          | 14 ± 1                        | 2.7–3.1                     | 11                            |

Uncertainty estimates represent ±1σ; n = number of samples; none = no cirrus interaction for at least 5 days upwind.

Figure 2. Time since cirrus interaction versus the normalized ratio of (a) CH$_2$Cl$_2$/C$_2$Cl$_4$ and (b) CH$_2$Cl$_2$/ethane for the four UT air parcels that showed depleted C$_2$Cl$_4$. The horizontal range shows the duration of the cirrus interaction, and the vertical error bars show propagated uncertainty based on the standard deviation of the mean.
the snow. While recognizing that the snow results may not extrapolate to ice surfaces, it is possible that C$_2$Cl$_4$ is preferentially adsorbed onto the cirrus particles, then becomes depleted via an addition reaction across its double bond, surface photolysis, or sedimentation. However the exact cause of the C$_2$Cl$_4$ depletion is unknown.

[13] The ClO values presented in Section 1 can be used to estimate roughly the extent that ethane would be depleted under such conditions. (Of the dozens of compounds that we measure, ethane is one of our most precise measurements: precision = 3 pptv at UT mixing ratios.) In the UT under TRACE-P conditions (265 mb; 223 K; O$_3$ = 50 ppbv; CO = 120 ppbv; SZA = 30°), we modeled a ClO/Cl ratio of roughly 330 for NO = 50 pptv and 170 for NO = 100 pptv. During TRACE-P, NO$_x$ levels over the western Pacific were generally <100 pptv, and background values at all altitudes were ~25 pptv [Talbot et al., 2003]. Choosing a ClO/Cl ratio of 300 (NO ~ 50 pptv) yields roughly 0.01 and 0.03 pptv Cl, respectively, for 2.7 and 10 pptv Cl. By comparison, background Cl levels in the northern hemisphere free troposphere are estimated to be <10$^{-3}$ atom cm$^{-3}$ (<0.0002 pptv Cl at 250 mb) [Rudolph et al., 1996]. Using ln([ethane]/[ethane]$_0$) = -k$_{Cl}$ [Cl] t (where t is the duration of interaction with cirrus, and the background UT value [ethane]$_0$ was ~610 pptv during TRACE-P), 0.01 pptv Cl would cause ethane to be depleted by 65 pptv for 12 hours. Such an ethane depletion is well within our measurement capability but was not observed during TRACE-P, suggesting that Cl levels of 0.01 pptv (ClO levels of 2.7 pptv) were not present.

[14] An upper limit for Cl can be indirectly calculated based on the minimum detectable ethane depletion. The natural variability of ethane in a typical 20-minute UT flight leg is 5 pptv (1σ), and we estimate that 40 pptv (which would be diluted to roughly 30 pptv after 1 day of downwind travel; see Figure 2a) is the minimum ethane depletion that we could confidently detect in the UT. A 40 pptv ethane depletion requires 3 × 10$^4$ atom Cl cm$^{-3}$ (0.005 pptv Cl) for 12 hours of photochemical processing in the UT, and 1.5 × 10$^4$ atom Cl cm$^{-3}$ (0.002 pptv Cl) for 1 day. Therefore, if it occurs, the maximum Cl activation by UT cirrus in tropical- and mid-latitudes during spring is likely within an order of magnitude of background tropospheric Cl levels, and is not expected to exceed 3 × 10$^3$ atom Cl cm$^{-3}$. The limited Cl activation by UT cirrus is consistent with laboratory studies by Hynes et al. [2002], who found that HCl uptake onto ice is inhibited by HNO$_3$ at $T >$ 210 K.

4. Conclusions

[15] Airborne C$_2$Cl$_4$ and ethane measurements were used to indirectly investigate the possibility of heterogeneous chlorine activation by UT cirrus. Of seven UT air parcels that had not been recently influenced by the stratosphere or lower troposphere, four had 15–36 hour interactions with cirrus between 1–3 days upwind. Tetrachloroethene was depleted in each of these air parcels, and the magnitude of the depletion was negatively correlated with time since the interaction. A lack of significant ethane depletion in these air parcels rules out the possibility that Cl was the primary cause of the C$_2$Cl$_4$ depletion. We tentatively speculate that C$_2$Cl$_4$ was preferentially adsorbed onto the clouds, followed by chemical processing at the ice surface and/or sedimentation.

[16] Ethane is abundant in the UT and it is one of our most precise measurements. We used a minimum detectable ethane depletion of 40 pptv to indirectly estimate an upper-limit Cl abundance of 3 × 10$^4$ atom Cl cm$^{-3}$ (0.005 pptv Cl) in the vicinity of UT cirrus in tropical- and mid-latitudes during spring. Assuming a ClO/Cl ratio of 300 in the springtime UT, this corresponds to an upper limit of 1.5 pptv ClO. This value lies in the low range of modeled ClO enhancements of 1–100 pptv for chlorine activation by cirrus in cold conditions (near and above the tropopause), and suggests that cirrus in relatively warmer conditions (in the UT) play a much more limited role in midlatitude chlorine activation during spring.

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