An Investigation of South Pole HOx Chemistry: Comparison of Model Results with ISCAT Observations

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Abstract. Unexpected high levels of OH and NO were recorded at the South Pole (SP) Atmospheric Research Observatory during the 1998-99 ISCAT field study. Model simulations suggest a major photochemical linkage between observed OH and NO. A detailed comparison of the observations with model predictions revealed good agreement for OH at NO levels between 120 and 380 pptv. However, the model tended to overestimate OH for NO levels < 120 pptv, while it underestimated OH at levels > 380 pptv. The reasons for these deviations appear not to involve NO directly but rather HOx radical scavenging for the low NO conditions and additional HOx sources for the high NO conditions. Because of the elevated levels of NO and highly activated HOx photochemistry, the SP was found to be a strong net source of surface ozone. It is quite likely that the strong oxidizing environment found at the South Pole extends over the entire polar plateau.

1.0. Introduction

The photochemistry of near surface air at the South Pole (SP) has received minimal attention from the modeling community in past years. This most likely reflects the fact that other than O3 and CO, little data existed to justify detailed model studies. The recent field study ISCAT (Investigation of Sulfur Chemistry in the Antarctic Troposphere) provides extensive photochemical observations so that a comprehensive modeling analysis can be carried out to investigate SP summertime photochemical processes. The ISCAT program represents an extension of the earlier SCATE program (Sulfur Chemistry in the Antarctic Troposphere Experiment) that examined sulfur chemistry at Palmer Station located on the Antarctic coast. This extension to the South Pole recognizes that the different chemical forms of sulfur found in Antarctic ice cores can serve as useful climate proxies (e.g., Legrand, 1997 and references therein). It also highlights the continuing need to improve our quantitative understanding of the dynamical and chemical factors influencing sulfur speciation and deposition at the SP.

Among the important chemical factors is the role played by local photochemistry in altering species before their deposition.

Figure 1. Simplified HOx-NOx-CH4 photochemical scheme reflecting South Pole chemistry.
rate coefficients were based on in-situ spectrally resolved actinic flux measurements. The division of data subsets L1, L2, M, and H was based on observed NO levels.

photochemical observations included: NO, OH, and photolysis rate coefficients for J(O'D), J(NO2), J(HNO3), J(H2O2), J(CH3O), J(HONO), J(CH3OOH), J(CH3ONO2), and J(PAN).

The NO data was reported at a time resolution of 1 min. from Nov. 30, '98 to Jan. 4, '99. The modified chemiluminescence NO sensor had a 2σ detection limit (LOD) of 6 pptv. The OH data was also taken at 1 min. intervals for Dec. 12–30, '98. The OH observations were recorded using the SICIMS (select-ion chemical-ionization mass spectrometer) technique which had a nominal LOD of ~5x10^5 molecule cm^(-3) based on a 5 min. integration period. The estimated total error (i.e., precision plus accuracy) was ±75% (2σ). At S/N ratios of 5/1 or greater, the systematic error dominated, leading to an overall uncertainty of ~±60%. More detailed discussions of these measurements can be found in Davis et al., Mauldin et al., and Lefer et al. [all this issue]. In addition, there were also very limited atmospheric HNO3 measurements (Richard Arimoto, unpublished results).

All data collected during the ISCAT study involved sampling from either the second floor of the NOAA operated Atmospheric Research Observatory (ARO) building or from the roof of the ARO building. The OH and NO sampling inlets at the second floor extended out 1~2 m from the building, at a height of ~10 m above the snow surface. All sampling inlets were mounted on the side of the ARO building facing the prevailing wind, e.g., out of the "clean air sector" (0°~120°).

2.2. Model Description

The photochemical box model used in this study was similar to that described previously by [Crawford et al., 1999]. The model assumes that all short lived species to be in photochemical stationary state. The longer lived species (e.g. H2O2, CH3OOH, HNO3, etc) were assumed to be at steady state. Model runs were constrained by observational values of O3, NO, CO, H2O, CH4, M, and H was based on observed NO levels.

3.0. Results and Discussion

3.1. OH: Observations vs. Model predictions

In the text that follows, only when both OH and NO values were recorded simultaneously was an "observation" versus "model" comparison considered. In addition, comparisons were only made when temporal variations in NO over a 20 min period were ≤ 25%. This filtering procedure resulted in a total of 316 independent 10 min. OH values, or ~80% of the total OH data.

As shown in Figure 2, in general, the agreement between model and observations was quite good, i.e., ~80% of the results were well within the combined uncertainties of the model and measurements. The median value for the ratio of observation to model (O/M) for all data was 1.01. This result, however, is misleading in that an inspection of the O/M ratio shows that the value for this ratio varies significantly as a function of the NO mixing ratio. As illustrated in Figure 2, O/M values are significantly less than 1 for NO levels <120 pptv while significantly higher than unity for NO levels >380 pptv.

Figure 2. Ratio of observed OH to model calculated values as a function of observed NO. The division of data subsets L1, L2, M, and H was based on observed NO levels.

Figure 3. Observed (a) and model calculated (b) OH as a function of observed NO and O3 levels. The three O3 levels correspond to the 5th and 95th percentile and median values.
For mid-range values (120 to 380 pptv), the model predictions calculated for this subset is 0.8. We have labeled this data subset as group "M".

For NO values > 380 pptv, the median O/M value is 1.6. These data have been labeled as group "H". The lowest O/M subset, labeled here as "L", has been further subdivided into groups L1 and L2. The lowest of these, L1, is made up of data primarily collected on 12/17 and reflect NO values that are shown as typically ≤ 50 pptv. The median O/M ratio for this data group is 0.21. The second "L" grouping (L2), recorded on 12/27, resulted in an estimated O/M ratio of 0.45 and corresponds to a median NO level of 100 pptv. Data groups L and H have been examined in sections 3.1.1 and 3.1.2 in greater detail, since no evidence was found for an instrument malfunction during the times of ISCAT OH sampling. Thus, data groups "H" and "L" are viewed as providing a strong indication that other factors likely influenced the SP OH which were not included in our standard model runs.

As shown in Figure 1, elevated NO promotes OH recycling through the HO2 + NO reaction. However, because of the follow-on reaction of OH + NO, steady increases in the level of NO do not lead to monotonic increases in OH. In fact, one typically finds that the OH concentration peaks at intermediate levels of NO as shown in Figure 3a. For the typical conditions found during ISCAT, Figure 3b shows that model predicted OH values peak at 126 pptv of NO2 whereas, the experimental OH maximum, as shown in Figure 3a, indicates near max values of OH as occurring over a rather broad range of NO (i.e., 100 to 200 pptv). However, this can be largely understood in terms of the influence of varying levels of O3. As seen in Figure 3b, where the top and bottom curves define the 5th and 95th percentile for observed O3, the optimum NO value is seen shifting from 89 to 158 pptv, respectively. Variations of other parameters (e.g., H2O) can also cause small shifts (i.e., ≤ 15%).

### Table 1. South Pole OH budget

| OH Source | OH Sinks | Percent Contribution |
|-----------|----------|----------------------|
| O(D) + H2O | OH + CO | 6% 51% |
| HO2 + NO | OH + CH4 | 92% 20% |
| HO2 + O3 | OH + NO2 | 2% 9% |
|           | OH + O2 |               |
|           | OH + H2 |               |
|           | OH + CH2O | 6% 4% |
|           | Others |               |

Total 100% 100%

### 3.1.1. Detailed Investigation of Data groups L1 and L2.

An important atmospheric characteristic revealed from examining data subsets L1 and L2 was that very high dew points were recorded for both sampling days (i.e., -25 °C for 12/17 and -29 °C for 12/27). In fact, these were the only cloudy/foggy days during ISCAT for which both model runs and observations were available. As suggested by [Mauldin et al., this issue], the environmental conditions on these two days points strongly toward the possibility that there were additional losses of HOx radicals due to droplet scavenging. Previous studies have reported evidence of substantial HO2 loss in clouds [Mauldin et al., 1998; Cantrell et al., 1996]. To simulate this loss in the current study, we introduced a first order "k" value for HO2 with an assigned sticking coefficient, y, of unity. This scavenging process was further assumed to be irreversible. Given the temperature and dew point on 12/17 and 12/27, we then took the supercooled water droplet size distribution as falling within the range of 5 - 10 μm and as having an estimated number density of 5 to 15/cm3 [Austin Hogan, unpublished results]. Using the midpoint of these ranges, the resulting first order scavenging rate was estimated at 9.0 × 10^-3 s^-1 [Fuchs and Sutugin, 1970].

The required "k" values needed to bring the model and observations into a high level of agreement were determined to be 8.0 × 10^-3 s^-1 and 3.5 × 10^-3 s^-1 for L1 and L2, respectively. Thus, within the range of values cited for droplet parameters, a modification of the standard model was able to bring both days' predictions into reasonable agreement with the observations. The corresponding lifetimes for HO2 are 2.1 and 4.8 minutes.

### 3.1.2. Detailed Investigation of Data Group H.

As discussed in the above text, the group "H" appear unique in that the observed OH is substantially higher than the standard model prediction. In this context, recent polar observations indicating snow emissions of CH2O, H2O2, and HONO as a possible source of HOX radicals would appear to be quite relevant [McConnell et al., 1997; Sumner and Shepson, 1999; Hutterli et al., 1999; and Dibb et al., 1999]. To simulate the effect of the new sources, we have carried out sensitivity runs to estimate the level of each of these species required to remove the model underestimation. Then, the estimated levels have been compared with available data from various polar sites. In the case of CH2O, additions of only 80 pptv were found to be sufficient. Observed CH2O levels at Summit, Greenland have ranged from 100 - 450 pptv and those at Alert, Canada from 52 - 690 pptv.

Similarly, for H2O2 and HONO we estimate that 190 pptv and 4 pptv, respectively, would be required. These values can be compared to summertime SP H2O2 measurements that have ranged from 80 to 280 pptv [McConnell et al., 1997] and Summit, Greenland observations of HONO that have been found to be as high as 10 pptv [Dibb et al., 1999]. Collectively, these findings would suggest that modifications to the standard model, involving the inclusion of other HOx sources, can readily account for observed OH values exceeding those predicted using the standard model.

On the other hand, if similar levels of these new HOx sources are applied to our group M data, the model predicted OH is increased by a factor of 1.3 to 2.5, which is significantly higher than the observed OH. By contrast, for data groups L1 and L2, the same addition of CH2O and H2O2 shifts the predicted OH up by less than 20%. HONO, however, increases predicted OH by more than a factor of 2. To reconcile these discrepancies, one has to assume that the actual levels of these additional HOx sources would to a large extent parallel NO levels, which as noted by Davis et al. [this issue] is strongly controlled by the atmospheric mixing depth at SP. If true, then the general trend would be one in which the levels of all four species would be significantly modulated by shifts in the mixing depth. Thus, the OH results for data groups "M" and "L" would not be expected to be as strongly influenced by the proposed additional HOx sources as group "H". Suffice it to say, any comprehensive

### Table 2. South Pole HOx budget

| HOx Source | Percent Contribution | HOx Sink | Percent Contribution |
|------------|----------------------|----------|----------------------|
| O(D) + H2O | 37%                  | HNO3 + DD | 52%                  |
| CH4        | 63%                  | HNO3 + OH | 10%                  |
| Chemistry  |                      | HONO2 + DD | 22%                  |
|            |                      | HONO2 + OH | 6%                   |
|            |                      | OH + HO2 | 8%                   |
|            |                      | Others | 2%                   |

Total 100% 100%
understanding of this SP HOx chemistry will require measurements of all four species.

3.2 SP OH and HOx Photochemical Budget

As shown in Table 1, primary production of OH from the reaction O('D)/H2O accounts for only 6% of the total. The major source is from the recycling reaction, NO + HO2. Major OH sinks involve reaction with CO, CH4, and NO2, e.g., 51%, 20%, and 9%, respectively. The remainder is due to the reaction of OH with O3 (7%), H2 (6%) and CH4 (4%).

For HOx (shown in Table 2), the reaction O('D)/H2O defines only approximately 1/3 of the total production. The largest contributor is from CH4 oxidation. Quite interestingly, at NO levels of 225 pptv, the CH2O yield from CH4 oxidation is nearly 100%. Thus, the net HOx production from CH4 is ~50% of the initiating reaction rate. Major loss channels for HOx involve the long-lived nitrogen species, i.e., HNO3 and HO2NO2. Approximately 90% of the HOx lost can be accounted for by a combination of dry deposition reaction with OH.

3.3 Consequences of Intense SP HOx Chemistry

One of the interesting consequences of the intense surface layer photochemistry at the SP is the prediction that significant net O3 should be a result. By extension, the same statement should also apply to the entire Antarctic plateau region. This finding is quite unique relative to what is typically found at a remote surface site where NO levels are usually quite low. In the latter case one normally finds net photochemical O3 destruction. Based on the SP observations, the predicated net photochemical production of O3, P(O3), is estimated to range from 1 to 6 ppbv/day (e.g., see Crawford et al. [this issue]). The major pathway for this formation involves the reaction HOx/NO (see Figure 1). SP photochemical O3 destruction ranges from 0.3 to 0.8 ppbv/day, the two most important destruction processes being OH/O3 and OH/NO2.

The results from the first ISCAT field study have once again demonstrated that atmospheric surprises are still be found. Quite remarkable is the finding that the near surface atmospheric layer at SP, with greatly enhanced NO levels, produces a 24 hour average oxidizing level (i.e. OH = 1.7 x 106 molec./cm3) which rivals that of equatorial marine environments. It is also 20 times higher than the 24 hour average value estimated for Palmer Station, Antarctica [Jefferson et al., 1998].

Since for most chemical species deposition to the snow surface occurs during the summer months [Bergin et al., 1998], these results raise some interesting new questions about the degree to which some species might be modified before being deposited at the surface. In conjunction with growing evidence that extensive oxidative processes are also occurring within the snowpack (i.e., firm) [e.g., Sumner and Shepson et al., 1999], the interpretation of the concentration levels of some climate proxy species in ice cores may need to be reexamined. These results also point to the need for new research to explore the impact from surface emissions of NO and other trace gases on near surface OH levels for other snow covered regions.

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