DETERMINATION OF ACIDITY IN AMBIENT AIR

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ABSTRACT

A laboratory and field study was performed to construct and evaluate an automated, semi-continuous monitor for ambient air concentrations of nitric acid (HNO₃) and ammonia (NH₃) utilizing tungstic acid-coated denuder tubes. In addition, atmospheric concentrations of hydrochloric acid (HCl) were determined. Manual techniques for atmospheric NH₃ measurement were also compared. Interference in HNO₃ measurements was determined with nitrogen dioxide, nitrous acid, and particulate ammonium nitrate. Atmospheric sampling for HNO₃ was done in Riverside, CA during September 1984 with the automated tungstic acid technique (TAT), and by the denuder difference method (DDM), considered the reference procedure for HNO₃. Ammonia and HCl were measured with impregnated filter methods. In addition, NH₃ was measured by a denuder tube collection procedure.

Nitrous acid was the only material studied which exhibited significant interference in nitric acid response with the automated TAT. Atmospheric nitric acid levels with the automated TAT showed 1 hour average values up to 45 µg/m³ (17.5 ppb). These values yielded moderate correlation with O₃ (r = 0.78) and high correlation (r = 0.94) with HNO₃ by the DDM. However, the HNO₃ values averaged about 50% higher than those by the reference procedure. Ammonia measurements with impregnated filters showed high correlation (r = 0.98) with oxalic acid-coated denuder tube collected samples, but the former averaged 12% higher. Ammonia measurements with the automated TAT were not quantitative because of difficulties in calibration. Hydrochloric acid values ranged up to 3 µg/m³ with a pronounced diurnal variation, maxima occurring during the period 0800-1200 or 1200-1600 hr. Particulate chloride levels were similar in magnitude to those for HCl and showed no correlation with HCl.
The statements and conclusions in this report are those of the Contractor and not necessarily those of the State Air Resources Board. The mention of commercial products, their source or their use in connection with material reported herein is not to be construed as either an actual or implied endorsement of such products.
ACKNOWLEDGEMENTS

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I. **SUMMARY AND CONCLUSIONS**

A laboratory and field study was performed to construct and evaluate a semi-real time monitor for ambient air concentrations of nitric acid and ammonia utilizing tungstic acid (WO₄) coated denuder tubes. In addition, atmospheric concentrations of HCl were determined.

The laboratory study focussed on assessing the influence of design parameters to achieve a workable automated sampler for HNO₃ and ammonia. Utilizing a preliminary version of the automated tungstic acid technique (TAT), interference studies were done with NO₂, HONO (nitrous acid), and particulate ammonium nitrate. Of these, only HONO showed significant interference. The apparent response corresponded to at least one third the molar response of nitric acid. The difficulties in preparation and quantitation of HONO made only lower limit measurements of TAT interference possible.

The efficiency of tungstic acid coated tubes at 1 Lpm with 340-1030 ng loadings of HNO₃ at ca. 50% R.H. ranged from 91 to 98%, decreasing with increasing loading. The automated system re-equilibrated from sampling HNO₃ or NH₃ to clean air within one or two cycles if the Teflon ball valve in the inlet was lined with pyrex tubing. Attempts to operate the automated TAT on purified ambient air for desorption and makeup gas flow yielded nonreproducible results and were, therefore, abandoned. Desorption of HNO₃ with helium was shown to provide about 30% higher response compared to desorption with synthetic air.

Alternative methods for NH₃ as well as a method for HCl collection and analysis were assessed with laboratory trials. Oxalic acid-coated denuder tubes yielded NH₃ results agreeing within 20% with those for oxalic acid impregnated quartz fiber filters with better agreement being obtained for 16 hr than for 4 hr trials. HCl collection with triethanolamine-KOH impregnated cellulose filters following a Teflon prefilter was evaluated with several sampling configurations. The procedure adopted was found to provide an efficiency for HCl of at least 92%.

Atmospheric sampling was done on the campus of the University of California (Riverside) with samplers mounted on the roof of a mobile laboratory. In addition
to the automated TAT, HNO$_3$ was measured with 4 or 8 hour resolution by the
denuder difference method (DDM), considered the reference procedure for HNO$_3$.
Ammonia and HCl were measured with 4 or 8 hr resolution with impregnated
filter methods. In addition, NH$_3$ was measured by a denuder tube collection
method with 12 and 24 hr resolution.

Pollution levels ranged from extraordinarily light to moderate photochemical
smog (O$_3$ > 0.30 ppm). Nitric acid levels with the automated TAT showed 1
hour average values up to 45 µg/m$^3$ (17.5 ppb). These values showed moderate
correlation with O$_3$ (r = 0.78) and high correlation (r = 0.94) with HNO$_3$ by the
DDM. However, the HNO$_3$ values averaged about 50% higher than those by the
reference procedure. The difference in results between the automated TAT and
DDM showed positive correlation with NO$_2$ (r = 0.6) perhaps indicative of the
retention on the WO$_x$ tube of a species other than HNO$_3$ whose concentration
correlates with NO$_2$. The difference in results between methods is qualitatively
similar to recent measurements made at a relatively remote site in Colorado
(18).

Ammonia measurements with impregnated filters showed high correlation
(r = 0.98) with oxalic acid-coated denuder tube collected samples but the former
averaged 12% higher. This is consistent with partial volatilization of NH$_4^+$ salts
from the Teflon prefilter. Ammonia measurements with the automated TAT
were not quantitative because of difficulties in calibration. At relatively high
concentrations, TAT response to NH$_3$ atmosphere appeared to be linear, however.
Further modification of the TAT is needed to permit reliable NH$_3$ determinations.

Hydrochloric acid values, estimated by observed levels of gaseous Cl$^-$, ranged
up to 3 µg/m$^3$ with a pronounced diurnal variation, maxima occurring during
the period 0800-1200 or 1200-1600 hr. Particulate chloride levels were similar
in magnitude to those for HCl and showed no correlation with HCl.

The principal conclusions from this study are:

1. An automated WO$_x$ denuder tube HNO$_3$ monitor can be reproducibly
calibrated and used to obtain semi-real time ambient air measurements
which are highly correlated with atmospheric HNO$_3$. 

- 2 -
2. Unidentified atmospheric nitrogen oxides are also retained on WO$_x$ coated denuder tubes which, in the present study, caused a 50\% positive interference. Without modification to improve its selectivity, the automated TAT may have limited applicability as a reference method for nitric acid.

3. NO$_2$ and particulate NH$_4$NO$_3$ do not appear to interfere in the method. However HONO shows a substantial positive interference. It is unlikely that HONO caused a substantial fraction of the observed interference, however.

4. Ammonia is more difficult to sample correctly with the automated TAT. Further work is needed to obtain a reliable, automated NH$_3$ monitor.

5. Atmospheric hydrochloric acid levels in Riverside are low (\(< 3 \mu g/m^3\)) with pronounced daytime maxima.

6. Triethanolamine-KOH impregnated cellulose filters cannot be recommended for HCl collection if ion chromatographic analysis is intended.

7. Phase II of this investigation should include further improvements in the automated TAT to permit NH$_3$ measurements, further interferent studies with the TAT, atmospheric trials at sites with both low and high levels of HCl, NH$_3$ and HNO$_3$, and assessment of the contribution of gaseous acids to atmospheric particulate acidity.
II. INTRODUCTION

A. Gaseous Nitric Acid and Ammonia

Nitric acid is significant as an air pollutant for a variety of reasons; as an inhalation hazard, a precursor of $\text{NH}_4\text{NO}_3$ particles which cause visibility reduction, as a nitrating agent in forming nitro-PAH compounds of increased mutagenicity compared to the parent hydrocarbon, and as a contributor to the acidity of suspended particles, rain water, lakes and vegetation. Ammonia, while not hazardous at atmospheric levels, participates in aerosol formation with $\text{HNO}_3$ and can neutralize the acidity of gaseous and particle-phase acids. Knowledge of $\text{NH}_3$ levels is, therefore, very useful in interpreting visibility data and the spatial variability in atmospheric acidity.

Our preceding studies (1-4) evaluated and compared several procedures for nitric acid measurement. Ammonia was sampled by an impregnated filter procedure. These methods have in common the need for relatively long collection times (~2 hr), water extraction and analyses for nitrate or ammonium ions. In addition, methods relying on dual filters (a prefilter to remove particulate matter with retention of the nitric acid or ammonia on an appropriate impregnated after-filter), are subject to positive errors caused by dissociation of materials such as $\text{NH}_4\text{NO}_3$ on the prefilter.

A recently developed procedure offers promise of overcoming many of these difficulties. The procedure employs a diffusion denuder coated with tungstic acid (5-8). This coating was reported to provide complete retention of both gaseous nitric acid and ammonia. After sample collection, both components are thermally desorbed, the $\text{NH}_3$ trapped by a transfer tube, and the evolved $\text{HNO}_3$ directly measured with a conventional chemiluminescent $\text{NO}_x$ analyzer. Ammonia is subsequently desorbed thermally from the transfer tube, oxidized to NO and measured with the same analyzer.

Prior to the present study, only limited data were available comparing
this technique to the "denuder difference" method (4), currently favored as a reference procedure. Results obtained at Research Triangle Park, N.C. showed agreement within ± 20% (9). However, no atmospheric comparisons had been made under conditions of relatively high nitrate such as found in California's South Coast Air Basin (SCAB).

In addition to this quasi-real time technique, a simple procedure for measuring atmospheric ammonia has been reported by Ferm in which air is drawn through a single 30 cm long oxalic acid-coated denuder tube at a low flow rate (e.g., 1-1.5 Lpm), trapping gas phase NH$_3$ (10). The coating and retained material are extracted by drawing water into the tube and the NH$_3$ determined as NH$_4^+$ with a specific ion electrode. The low flow rate required makes this technique appropriate for relatively long term (e.g., 12-24 hr) sampling. The accuracy and precision of this method are uncertain.

A continuous method for atmospheric NH$_3$ has also been reported relying on formation of a fluorescent derivative. Reproducible measurements of ≥ 0.1 ppb NH$_3$ are claimed, although the collection efficiency of the system for NH$_3$ was only 30% (11).

B. Hydrochloric Acid

Of the acids believed to exist in the polluted troposphere, probably least is known about HCl. Levels up to 8 ppb were observed in Tokyo (12), and it is possible that similar values would be observed in the SCAB. Atmospheric HCl probably results from hydrolysis of lead halides emitted from motor vehicles, from reaction of sea salt with strong acids (e.g. HNO$_3$), by liberation from cloud droplets containing a variety of acids and salts, and by combustion of waste products. The potential significance of HCl as an air pollutant include, an inhalation hazard possibly similar to HNO$_3$, a contributor to the acidification of rain water, lakes, suspended particles and vegetation (by wet and dry deposition), and a source of errors in particulate nitrate values obtained in sampling with inert filters (14).
Consistent with the paucity of HCl data, methodology for HCl sampling remains comparatively primitive. Okita employed a dual filter procedure with the prefilter removing particulate matter and an Na₂CO₃-impregnated after-filter collecting gaseous HCl (12). The resulting values probably represent upper limits to the true HCl levels since chloride can be released from the prefilter by reactions with strong acids.

C. Objectives of the Present Study

The present program has given principal emphasis to the semi-real time method for HNO₃ and NH₃. The variables in design and operation were evaluated together with the influence of potential interferents. Atmospheric measurements were performed in Riverside, California to compare HNO₃ and NH₃ values with those measured by more conventional techniques. The site was chosen to provide what are probably the highest levels of both gases likely to be found in ambient air. In addition gaseous HCl was sampled to obtain what are, to our knowledge, the first urban ambient air HCl measurements in this country.
III. CONSTRUCTION AND INITIAL EVALUATION OF AN AUTOMATED MONITOR FOR HNO₃ AND NH₃

A. Chemistry of the Automated Tungstic Acid Technique (TAT)

The automated tungstic acid technique (TAT) employs Vycor quartz tubes coated initially with blue tungsten oxide (\(W_{18}O_{49}\) plus \(W_{20}O_{58}\)) which is converted by heating in \(O_2\) to the yellow oxide, \(WO_3\). The hydrate of this oxide \(H_2WO_4\) or \(H_2W_2O_7\) (also written \(WO_2(OH)_2\)), is a weak acid suggesting a surface pH of 3-4 (7). Collection of \(NH_3\) is attributed to simple acid-base equilibria such as:

\[
NH_3 + H_2WO_4 \rightleftharpoons NH_4HWO_4 \tag{1}
\]

This equilibrium is reversed by heating to about 350°C. A color change appears to be associated with a change in crystal structure at 320°C. This provides a convenient method for adjusting heating temperature. The mechanism of \(HNO_3\) sorption is unclear but was suggested (7) to involve:

\[
WO_2(OH)_2 + 2 HNO_3 + WO_2(NO_3)_2 + 2H_2O \tag{2}
\]

or

\[
WO_2(OH)_2 + HNO_3 + WO_2(OH)(NO_3) + H_2O \tag{3}
\]

Thermal desorption is suggested (7) to proceed as follows to yield \(NO_2\):

\[
WO_2(NO_3)_2 \xrightarrow{\Delta} 1/2 O_2 + 2NO_2 + WO_3 \tag{4}
\]

More recent studies (21) indicate that NO rather than \(NO_2\) is the species formed following thermal desorption. The formation of NO rather than \(NO_2\) is relatively insignificant, however, since, as shown below, a catalyst downstream of the preconcentrator would convert any \(NO_2\) formed to NO.

B. Design

The automated TAT used by McClenny et al (6) is shown schematically in Figure 1. The automated TAT employed a denuder tube coated with
Figure 1. Schematic of Automated Tungstic Acid Technique
tungstic oxide to retain both gaseous HNO₃ and NH₃ sampling ambient air at 1 Lpm (WO₃ preconcentrator). Following a sampling period of varying length depending on expected concentrations, the collected HNO₃ and NH₃ were desorbed by heating the preconcentrator tube to ca. 350°C (Variac V₁ and nichrome wire heater). The HNO₃ desorbed as NO or NO₂, which passed through the WO₃-coated transfer tube, was converted (if necessary) to NO on a gold catalyst heated at > 600°C, and measured with a chemiluminescent NOₓ analyzer. Ammonia desorbed from the preconcentrator column was retained on the transfer tube. Following analysis of the HNO₃ (as NO), the transfer tube was heated to ca. 350°C desorbing NH₃ which was also converted to NO on the gold catalyst and then analyzed.

McClenney's automated TAT system employed 80/20 v/v He/O₂ as a carrier during desorption. Oxygen was necessary to oxidize NH₃ to NO over the gold catalyst. However, in the automated system of Braman et al, only pure helium was used for desorption (15). Pure oxygen was introduced just ahead of the gold converter to oxidize the NH₃. As a consequence a slow loss of preconcentrator tube activity, apparently by reduction with organic material, was experienced in this system. Consequently, periodic treatment with O₂ was necessary.

The AIHL system began with the McClenney design. The timing for activation of ball valves, solenoid valves, variacs, and integrator was provided by a Chrontrol CT 5 four channel (10 program) timer unit. The unit was initially programmed for a 30 minute cycle time consisting of ten minutes sampling, seven minutes heating of the preconcentrator tube, seven minutes heating of the transfer tube and the remaining six minutes, for cool down before re-cycling. Peaks were integrated by a Hewlett-Packard Model 3380A recorder-integrator. In the prototype AIHL unit built for lab evaluation, flow rates for sampling, carrier and makeup gas (i.e. that needed to satisfy the pumping rate for the chemiluminescent NOₓ analyzer) were regulated with needle valves and flowmeters. These were replaced by mass flow controllers for the atmospheric sampling unit.
C. Theoretical Performance of a Diffusion Denuder

The performance of diffusion denuder tubes having a coating of infinite capacity for the gas to be removed can be approximated by the Gormley-Kennedy equation (23). For viscous, laminar flow, and assuming retention of NH$_3$ and/or HNO$_3$ which contact the walls:

\[
\frac{C}{C_0} = 0.82 \exp(-15 \Delta) + 0.098 \exp(-89 \Delta) + 0.33 \exp(228 \Delta) \tag{5}
\]

Where \( \Delta = \frac{DL\pi}{4Q} \) and

\[D = \text{diffusion coefficient (cm}^2/\text{sec), } 0.25 \text{ for NH}_3, 0.15 \text{ for HNO}_3.\]

\[L = \text{total length of tube (cm), excluding portions uncoated with WO}_x.\]

\[Q = \text{flow rate through tube (cm}^3/\text{sec})\]

\[C_0 = \text{concentration entering tube}\]

\[C = \text{average concentration exiting tube}\]

\[(1 - C/C_0) \times 100 = \text{percent removal}\]

This expression is valid if the Reynolds number is kept below 2000 to prevent turbulent flow. The Reynolds number (\(N_R\)) is given by the equation:

\[N_R = \frac{\rho dv}{n} \tag{6}\]

Where \(\rho\) = density. For air, \(\rho = 1.29 \times 10^{-3} \text{ g/cm}^3\)

\(d = \text{tube internal diameter, cm}\)
Employing 0.4 cm ID tubing, the Reynolds number equals 378 at 1.0 Lpm, the sampling rate used for the preconcentrator tube, and is < 200 for the carrier gas flow rates \( \leq 500 \) mL/min.

Using a 35 cm coated length of 0.4 cm ID tubing for the preconcentrator tube, a sampling rate of 1 Lpm would provide a calculated efficiency of 99.8% for NH\(_3\) and 98.0% for HNO\(_3\). An 8.6 cm coated length transfer tube, and carrier air flow rate of 500 mL/min provide a 96.1% calculated efficiency for NH\(_3\) removal. A 10-fold decrease in carrier flow rate increases the expected removal efficiency to 100%. These calculations assume a 100% efficiency of retention of HNO\(_3\) and NH\(_3\) following collision with the WO\(_x\) surface.

In the automated TAT, the tungstic acid surface can be saturated with NH\(_3\) and HNO\(_3\), making equation (5) potentially inaccurate for predicting the efficiency of this system. Braman et al (7) modified equation (5) by considering that the effective length, \( L \), of the TAT tubes changed exponentially with time. At time \( t \), the effective length is:

\[
L = L_0 \exp(-kt) \tag{7}
\]

Where \( L_0 \) = original effective length (35 cm)

\( k \) = tube depletion rate constant

The constant, \( k \) can be calculated, in turn, from:

\[
k = \frac{\text{feed rate (ng/sec)}}{\text{tube capacity (ng)}} \tag{8}
\]

Combining equation (7) and (5) and neglecting all except the first term
in equation (5), since subsequent terms are normally negligible, leads to the equation:

\[
\frac{C}{C_0} = 0.82 \exp \left( \frac{-15\pi D}{4Q} \right) L_o \exp (-kt) \quad (9)
\]

Braman has observed a capacity for HNO₃ of about 600 ng. Assuming a concentration of 20 µg/m³ HNO₃, at the end of the 10 minute sampling period used in our laboratory studies (at 1 Lpm), the efficiency is calculated as follows:

\[
1 \text{ Lpm} = 1.67 \times 10^{-5} \text{ m}^3/\text{sec}
\]

\[
20 \mu g/m^3 = 2.0 \times 10^4 \text{ ng/m}^3
\]

\[
(1.67 \times 10^{-5} \text{ m}^3/\text{sec}) \times (2.0 \times 10^4 \text{ ng/m}^3) = 0.333 \text{ ng/sec HNO}_3
\]

\[
k = \frac{0.333 \text{ ng/sec}}{600 \text{ ng}} = 5.55 \times 10^{-5} \text{ sec}^{-1}
\]

After 10 min, \(kt = 0.0333\). The flow rate \(Q\) at 1 Lpm is 16.67 cm³/sec. Under these conditions, the calculated percent removal rate after 10 min. sampling is 97.8%. This compares to 98.0% calculated from equation (5). The close agreement in this case is expected since in 10 minutes the HNO₃ loading represents only one third of the expected capacity.

D. Preparation of Tungstic Oxide-Coated Tubes

The coating of tubes with WOₓ by vacuum deposition involves substantially more art than science. As devised by Braman et al. (7), Vycor quartz tubes are coated with the device shown in Figure 2. An 0.02 inch diameter tungsten wire is held under spring tension along the center of the Vycor tube. The tube is mounted within a glass pipe, vacuum sealed at
Figure 2. Apparatus for Coating Hollow Tubes with Tungsten Oxide
both ends and evacuated. A nichrome wire heating coil envelopes the Vycor tube and is used for a preliminary bakeout, supposed to remove carbonaceous material from the tungsten wire and wall of the Vycor tube. With power to the nichrome wire off, current is passed through the tungsten wire depositing a blue-black coating of tungsten oxide (believed to be a tungsten (IV) oxide) on the inner surface of the Vycor tube. After coating is completed, the blue-black coating is oxidized to the yellow tungsten VI oxide by passing $O_2$ through the tube while slowly heating the outside of the Vycor tube with a very short, diffuse natural gas-oxygen flame. The coating technique is detailed in Appendix A. About 20 tungstic acid coated tubes were prepared which exhibited good coating adhesion and uniformity.

Coated tubes were checked for performance with a manual TAT system by briefly sampling relatively high concentrations of HNO$_3$ or NH$_3$, desorption from the heated tube into helium, passage of the products through a gold-coated hollow tube in the presence of oxygen (for conversion of NH$_3$ and NO$_2$ to NO) and then into a NO$_x$ analyzer (TECO Model 14 BE). Acceptable tubes displayed a large NO$_x$ signal whereas poor tubes were expected to show little response. None of the coated tubes tested appeared to be defective by this criterion.

E. Response Time of HNO$_3$ Analyzer

Initial results with the automated TAT suggested that portions of the system upstream from the preconcentrator tube were retaining and then slowly desorbing HNO$_3$. To establish if the motor-driven Teflon ball valve (A in Fig. 1) was a source of significant HNO$_3$ retention, the complete valve unit was removed. With this configuration the atmosphere sampled was changed from dilute HNO$_3$ to purified air only. In contrast to the previously observed slow response, the first or second cycle following the change showed no measurable HNO$_3$. It appeared, therefore, that the all-Teflon (PTFE) ball valve was the source of most, if not all of the HNO$_3$ retention. Careful cleaning of the ball valve had no effect.
To minimize HNO₃ contact with Teflon, the Teflon ball valve was lined with 0.4 mm I.D. Pyrex glass tubing. The change in behavior of the system was dramatic; re-equilibration within one or two cycles was now observed.

F. Efficiency of the Preconcentrator Denuder Tube for HNO₃ Collection during Automated Analysis

To assess the efficiency of the tungstic oxide denuder tube, a NaCl-impregnated Whatman 41 filter was placed downstream of the denuder tube to retain any HNO₃ penetrating the tube during sampling. Since the capacity of the tube is < 1 µg, penetration of the tube during a single sampling cycle would not be measurable by ion chromatography. Accordingly, the system was operated over a total of about 72 hours during which 4.3 m³ of HNO₃ in air were sampled. The observed nitrate on the NaCl impregnated filter corresponded to an HNO₃ concentration of 2.6 µg/m³. Using the mean HNO₃ sampled, 49 µg/m³, the tungstic acid tube is calculated to be 95% efficient under conditions in which ≤ 700 ng of HNO₃ was sampled per cycle. Further trials of denuder tube efficiency in manual operation are given in Section IV B, and yield similar results.

G. Measurement of Ammonia

Ammonia is collected together with HNO₃, at a flow of one Lpm on a 35 cm length WOₓ coating in the preconcentrator tube. The transfer tube can be shorter since NH₃ is collected following desorption from the preconcentrator tube at a flow of only 30-100 ml/min. Trials with 8.6 cm and 14.1 cm coating lengths for transfer tubes, demonstrated that the shorter tube provided adequate capacity and sharper peaks in the subsequent bake-out.

A peak occurring during bake-out of the preconcentrator tube loaded with NH₃ is an artifact here termed "apparent HNO₃". This error typically corresponded to 0.3% of the NH₃ peak area observed in bake-out of the
transfer tube (i.e. the peak ascribed to NH$_3$), and should include NH$_3$ desorbed from the preconcentrator tube but not retained on the transfer tube. Thus, the efficiency of the transfer tube for NH$_3$ retention is inferred to be $\geq 99.7\%$.

The analyzer was observed to respond rapidly to changes in NH$_3$ concentration. The first cycle following changes from, for example, 30 µg/m$^3$ NH$_3$ to synthetic air showed negligible NH$_3$ peak values. Similar findings were made in returning to such relatively high NH$_3$ concentrations.

H. Interference Effects With the TAT

1. Introduction

Assessment of interference from NO$_2$, HNO$_2$ and particulate nitrate was severely hampered by the response behavior of the preliminary system, which used the PTFE Teflon ball valve without glass lining. For example, changing from sampling HNO$_3$ to pure air required about 3 hours to achieve close to background responses. Experiments on interferent effects were designed accordingly.

2. Interference from NO$_2$

Nitrogen dioxide at 30 or 138 ppb was introduced into the system in air or in air containing about 50 µg/m$^3$ (ca. 20 ppb) of nitric acid. No measurable interference was observed. This is consistent with findings by Braman et al (7). Details are given in Appendix B.

3. Interference from Nitrous Acid (HONO)

Appendix C details the procedures used for preparing the acid and assessment of interference. Because only the sum NO$_2$ + HONO could be directly measured, only a lower limit to the true molar response of HONO was obtainable. This indicated that HONO had at least one third the molar response of nitric acid.
Additional trials with a more reliable, automated TAT configuration, and a manual TAT system are needed to confirm this. Such trials will be included in Phase II of the present study. These will also attempt to assess interference less ambiguously by excluding possible co-generated nitric acid and by measuring HONO concentrations more directly. Manual sampling of HONO with single WO\textsubscript{x} coated tubes will also be done to eliminate possible artifacts due to memory or other effects with the automated system.

4. Interference of NH\textsubscript{4}NO\textsubscript{3} Particles in Analysis of HNO\textsubscript{3}

NH\textsubscript{4}NO\textsubscript{3} particles were generated with a nebulizer system from an 0.010\%w solution in water. Particle size was measured with a Royco Model 226 optical particle counter, which indicated that > 50\% of all particles were \( \leq 0.17 \) \( \mu m \) in optical diameter. The measurements were made both at the aerosol generator outlet and immediately downstream of the HNO\textsubscript{3} analyzer inlet. Loss in sample lines was found to be < 10\%. The concentration of NH\textsubscript{4}NO\textsubscript{3} in air was also measured by collection on prefired quartz filters at the generator outlet followed by extraction and NO\textsubscript{3}\textsuperscript{-} determination by ion chromatography.

To insure that the analyzer was responsive, it was operated continuously with NH\textsubscript{3} at 33 \( \mu g/m^3 \) for about 24 hours before switching to NH\textsubscript{4}NO\textsubscript{3}. Based on the peak area for NH\textsubscript{3}, the limit of detection for NH\textsubscript{3} (with 20 min. sampling time) is estimated to be 1 \( \mu g/m^3 \).

Trials at 28 \( \pm 5 \) \( \mu g/m^3 \) and 69 \( \pm 4 \) \( \mu g/m^3 \) NH\textsubscript{4}NO\textsubscript{3} (as NO\textsubscript{3}\textsuperscript{-}) showed no measurable response. Based on the estimated limit of detection for NH\textsubscript{3}, the interference from NH\textsubscript{4}NO\textsubscript{3} is inferred to be < 7\%. This finding is consistent with earlier reports showing negligible loss of NH\textsubscript{4}NO\textsubscript{3} aerosol in MgO-coated denuders (1-3). Phase II will confirm these findings with the improved automated system as well as by manual sampling with WO\textsubscript{x}-coated tubes.
I. The Influence of Operating Variables with the Automated TAT

1. Introduction

During the evolution of a workable system, laboratory trials were conducted on the influence of such variables as carrier gas type and flow rates, temperature for desorption, and location for introducing makeup air and \(O_2\) needed for catalytic oxidation of \(NH_3\). Because many of the variables appear to be interactive, it is unclear that the results and conclusions of some of these trials are relevant to the design subsequently employed for atmospheric sampling. For purpose of the present report, only evaluations clearly relevant to the atmospheric sampling design are mentioned.

Desorption of \(HNO_3\) from the preconcentrator tube must be accompanied by reduction to \(NO_2\) or \(NO\) to avoid capture on the transfer tube and subsequent measurement as apparent \(NH_3\). Such a process should be favored by a high desorption temperature, a relatively slow carrier flow rate, and, possibly, by use of an inert carrier gas. Ammonia desorption must occur as \(NH_3\). Thus chemical transformations should be minimized. The possibility of readsoption of \(NH_3\) in the preconcentrator tube must be minimized, for example, by a high carrier flow rate. However, the carrier flow rate must not be so high as to diminish the efficiency of the transfer tube to trap \(NH_3\). As with \(HNO_3\), an inert carrier might increase the efficiency of transfer of \(NH_3\) to the transfer tube. Since desorption of \(NH_3\) from the transfer tube is followed by oxidation on a gold catalyst at about 625°C, a process which required oxygen, an inert gas cannot be used for the complete analytical cycle.

Substantial effort was devoted to evaluate the automated TAT employing purified ambient air for desorption and to provide the makeup air needed by the \(NO_x\) analyzer. The objective was to eliminate reliance on compressed gas cylinders, a marked disadvantage for prolonged atmospheric sampling.
2. Influence of Carrier Gas Type

To maximize the response of the preliminary automated TAT to both HNO₃ and NH₃ a strategy was adopted in which HNO₃ thermal desorption for initial 3 minutes of the preconcentrator tube heating cycle was done at 60 mL/min helium flow rate*. During this 3 minute period the peak due to HNO₃ is observed, and the NOₓ response returns to baseline. The remaining 6 minutes of the heating period employed a flow of ca. 500 mL/min carrier air (plus the helium flow at 60 mL/min) initially intended to achieve efficient transfer of NH₃ to the transfer tube.

Helium was compared to air as the carrier gas for desorption of HNO₃ during the initial 3 minutes of the preconcentrator tube heating cycle. As shown in Table 1, helium yielded a 30-40% increase in HNO₃ response.

<table>
<thead>
<tr>
<th>TABLE 1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Comparison of Helium and Air as Initial Carrier Gases in Nitric Acid Determination</strong>&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Initial Flowrate</th>
<th>Initial Voltage</th>
<th>Peak Areas X 10⁻⁶</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Carrier Gas</td>
<td>Variac Setting</td>
</tr>
<tr>
<td>3/1/84</td>
<td>0800-1030</td>
<td>5</td>
<td>Helium</td>
</tr>
<tr>
<td></td>
<td>1030-1530</td>
<td>10</td>
<td>air</td>
</tr>
<tr>
<td>3/2/84</td>
<td>1330-1600</td>
<td>6</td>
<td>Helium</td>
</tr>
<tr>
<td></td>
<td>1630-1730</td>
<td>3</td>
<td>air</td>
</tr>
</tbody>
</table>

a. Initial flowrate set by flowmeter = 57 mL/min with air. The same flowmeter setting was used with helium, which subsequently was shown to correspond to about 40 m/min helium.

b. HNO₃ concentration estimated at 50 ± 10 µg/m³.

c. Peak area for "apparent NH₃", possibly an artifact.

*Flow rate measured with a rotameter without correction for use of helium. The flow rate measured with mass flow meter was 40 mL/min. The difference between 40 and 60 mL/min is not considered significant.
3. **Use of Purified Ambient Air for Makeup and Carrier Gas**

Ambient air was purified by passage through a Bendix Model 8833 system utilizing activated charcoal, permanganate-impregnated molecular sieve and an efficient particle filter. Nitric oxide (NO) was not completely removed and yielded a varying baseline. The reproducibility of the automated TAT appeared to be poorer when relying on such air for both makeup and carrier gases compared to the use of bottled gases. Accordingly, this approach was abandoned.
IV. AN IMPROVED, AUTOMATED HNO₃-NH₃ MONITOR FOR SAMPLING IN RIVERSIDE

A. Introduction

This section describes (1) studies with a manual WOₓ technique used to design an improved, automated method, (2) the automated system employed in atmospheric sampling in Riverside, (3) calibration results obtained during atmospheric sampling with the automated TAT, and (4) laboratory studies, principally with NH₃ to elucidate the problems experienced during atmospheric sampling.

B. Preliminary Studies with a Manual Tungstic Oxide Technique

To obtain understanding of the behavior of individual components of an automated TAT system, a manual procedure was employed consisting of a WOₓ preconcentrator tube with and without a transfer tube, a gold NOₓ catalyst converter, and an NOₓ monitor. This was used to evaluate (1) the rate of desorption of NH₃ and HNO₃ from the tubes during heating, (2) the efficiency of a preconcentrator tube for NH₃ and HNO₃, (3) the extent of artifacts due to various components of the automated system, and (4) desirable changes in the automated system to improve reliability and minimize errors. In addition, following atmospheric sampling, problems encountered with NH₃ sampling and calibration were investigated with a manual TAT.

Denuder tubes sampled HNO₃ or NH₃ from diffusion or permeation tubes after dilution with particle free room air. These were thermally desorbed from the tubes with helium at 50 mL/min and mixed with about 700 mL/min O₂ prior to entry in the gold catalyst. The converter consisted of a 4 mm I.D. tube coated on its interior with metallic gold and heated with a nichrome wire heater at dull red heat.

Both NH₃ and HNO₃ yielded very sharp peaks without tailing (retention times 38 sec for HNO₃, about 46 sec for NH₃). When NH₃ and HNO₃
were desorbed together, the peaks were not completely resolved regardless of the desorption temperature. To assess efficiency, two tubes in series were used for sampling and the relative amounts of NO\textsubscript{x} determined on heating each tube. Table 2 summarizes the data obtained and indicates a single tube efficiency of 93 to 98\% at 1 Lpm with 340-1000 ng dosages of HNO\textsubscript{3} and similar values for NH\textsubscript{3}.

**TABLE 2**

Efficiency of Denuder Sampling for HNO\textsubscript{3} and NH\textsubscript{3}\textsuperscript{a,b}

<table>
<thead>
<tr>
<th>Gas</th>
<th>Loading Time (min)</th>
<th>Dosage (ng)</th>
<th>Efficiency, %</th>
<th>Mean Efficiency, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>HNO\textsubscript{3}</td>
<td>1.0</td>
<td>342</td>
<td>99.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>342</td>
<td>97.5</td>
<td>97.7 ± 1.6</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>342</td>
<td>97.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>342</td>
<td>95.9</td>
<td></td>
</tr>
<tr>
<td>HNO\textsubscript{3}</td>
<td>1.5</td>
<td>513</td>
<td>94.3</td>
<td>95.8 ± 2.0</td>
</tr>
<tr>
<td></td>
<td>1.5</td>
<td>513</td>
<td>97.2</td>
<td></td>
</tr>
<tr>
<td>HNO\textsubscript{3}</td>
<td>2.0</td>
<td>684</td>
<td>94.8</td>
<td>93.2 ± 2.8</td>
</tr>
<tr>
<td></td>
<td>2.0</td>
<td>684</td>
<td>95.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.0</td>
<td>684</td>
<td>89.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.0</td>
<td>684</td>
<td>92.3</td>
<td></td>
</tr>
<tr>
<td>HNO\textsubscript{3}</td>
<td>3.0</td>
<td>1026</td>
<td>91.4</td>
<td>91.4</td>
</tr>
<tr>
<td>NH\textsubscript{3}</td>
<td>2.0</td>
<td>-\textsuperscript{c}</td>
<td>97.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.0</td>
<td>-\textsuperscript{c}</td>
<td>99.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.0</td>
<td>-\textsuperscript{c}</td>
<td>96.0</td>
<td>98.0 ± 1.7</td>
</tr>
<tr>
<td></td>
<td>2.0</td>
<td>-\textsuperscript{c}</td>
<td>99.5</td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{a.} Sampling rate 1.0 Lpm.
\textsuperscript{b.} HNO\textsubscript{3} conc. 342 ± 73 \textmu g/m\textsuperscript{3}.
\textsuperscript{c.} Not determined.

To simulate more closely the automated system used for monitoring NH\textsubscript{3} and HNO\textsubscript{3} simultaneously, a transfer tube was attached to the downstream
end of the preconcentrator tube. The extent of NH$_3$ penetration to the detector during heating of the preconcentrator tube (i.e. artifact HNO$_3$) was determined to be $\leq 4\%$. Similarly, with nitric acid, retention and subsequent analysis of HNO$_3$ during heating of the transfer tube (i.e. artifact NH$_3$) was measured and found to be $< 3\%$.

The Teflon 3-way solenoid value used between the preconcentrator tube and transfer tube in the automated system (valve C in Fig. 1) was added to the manual system. The preconcentrator was heated for 2-3 minutes to achieve removal of either NH$_3$ or HNO$_3$. The transfer tube was then heated.

The level of artifact NH$_3$ was $\leq 5\%$ of the HNO$_3$ concentration measured. Similarly, with NH$_3$ the artifact HNO$_3$ was $< 4\%$ of the NH$_3$ measured. Thus the solenoid valve did not appear to introduce substantial error under these conditions.

Finally, the motor driven ball valve for the inlet of the automated system (valve A, fig. 1) was attached ahead of the preconcentrator tube and one or two minute samples drawn in through this valve, venting the flow through the 3-way solenoid valve (i.e. by-passing the transfer tube as in the automated method). Again, no significant sampling artifacts were observed in sampling relatively large dosages of either HNO$_3$ or NH$_3$.

C. The Automated TAT Design Used for Atmospheric Sampling

Based on results with the manual system, as well as studies described in Section III I, the automated system was altered as follows:

1. The carrier gas for desorption and transfer of HNO$_3$ and NH$_3$ was changed to helium only using a flow rate of 50-70 mL/min. This eliminated the introduction of an air carrier following emergence of the HNO$_3$ peak since NH$_3$ could be effectively transferred with helium at $\geq 50$ mL/min.
2. The heating times for the preconcentrator and transfer tubes were each reduced to 3 min. This permitted a change from a 30 min cycle to one as short as 15 minute.

3. The 4 mm I.D. tube coated with gold on its interior replaced the strip of gold foil used previously for catalytic conversion of NO₂ and NH₃ to NO*.3

4. A TECO 14BE was used in place of the Monitor Labs NOₓ analyzer since the TECO yielded much higher resolution.

5. Zero air, required to aid oxidation over the gold catalyst and to make up the difference between the NOₓ analyzer sampling rate and the helium flow rate was altered to be introduced between the transfer tube and gold catalyst rather than ahead of the preconcentrator tube.

An automated TAT unit suitable for field use was constructed in two modules, each in a wooden box for ruggedness and portability. One module contained the Chrontral program module timer, three mass flow controllers and three variable transformers. The second included all solenoid valves, the preconcentrator and transfer tubes, tube heaters, catalytic converter for NH₃, and glass flowmeters to verify the functioning of the mass flow controllers.

The sample inlet consisted of a 6 mm I.D. inverted U tube, total length about 15 cm, followed by Teflon fittings, and the glass-lined, motor driven ball valve. Beyond the valve, the sample passed through about 15 cm of pyrex or quartz tubing before reaching the WOₓ coated section of the quartz tube.

The operating strategy for the automated TAT used in Riverside is summarized in Table 3.

*It was subsequently found that gold in these coated tubes was lost at an excessive rate at the >-600°C temperature required for NH₃ conversion (i.e. the gold foil catalyst was preferable).
TABLE 3
Operating Strategy for the Automated TAT During Automated Sampling

<table>
<thead>
<tr>
<th>Time From Start of Cycle, min</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 to x where x = 5 to 30</td>
<td>Ambient air sampling</td>
</tr>
<tr>
<td>x to (x + 3)</td>
<td>Desorption of HNO₃ and NH₃ from preconcentrator tube with He carrier</td>
</tr>
<tr>
<td>(x + 3) to (x + 6)</td>
<td>Desorption of NH₃ from transfer tube with the carrier at 70 mL/min</td>
</tr>
<tr>
<td>(x + 6) to (x + 10)</td>
<td>Cool down for next cycle</td>
</tr>
</tbody>
</table>

Heater Voltages: Preconcentrator - 95V
Transfer tube - 45V
Au Catalyst tube - 75V

Flow rates: Helium carrier - 60 mL/min
Synthetic air for makeup - 470 mL/min
Sampling rate 0.96 Lpm

D. Strategies for Calibration

Two calibration strategies for HNO₃ and NH₃ were considered: (1) HNO₃ (from a diffusion tube) or NH₃ (from a permeation tube) were separately introduced into a ca. 3 cfm volume of dilution air, and the diluted gases sampled simultaneously with the TAT and impregnated filters. TAT sampling times were held constant and the calibration gas concentration altered by changing the degree of dilution or the oven temperature for the diffusion or permeation tube. This strategy most closely resembled ambient air sampling in that the sample was drawn through the Teflon ball valve inlet, and no alteration of the TAT was required. (2) HNO₃ and NH₃ from constant temperature sources were sampled by manually attaching the WOₓ preconcentrator tube to the source over which 1 Lpm of filtered room air was passed. Dosage was varied by changing the sampling time.
Comparing these two options, it is strongly preferable to calibrate the TAT without the need to dismantle the system to remove the WO\textsubscript{x}-coated preconcentrator tube. However, efforts to employ technique (1) yielded variable and unpredictable results. Memory effects caused by sorption of HNO\textsubscript{3} and NH\textsubscript{3} on glass and Teflon surfaces appeared to contribute to these problems. The alternative of maintaining constant concentration but varying TAT sampling times was not explored but holds promise for work during Phase II. Manual loading of WO\textsubscript{x} tubes was adopted for use in atmospheric sampling in Riverside.

E. Reproducibility of the HNO\textsubscript{3} Calibration During Atmospheric Sampling in Riverside

Nitric acid was supplied by a wafer type permeation tube maintained at a temperature of 83 ± 0.2°C in a Metronics Dynacalibrator oven with a flow rate over the wafer of 1 Lpm. Dilution air was supplied by drawing ambient air through a glass fiber filter followed by a NaCl-impregnated filter to remove atmospheric particles and HNO\textsubscript{3}. The output of HNO\textsubscript{3} from the wafer was monitored by sampling with an NaCl-impregnated filter for 8-24 hr periods. Since the entire output of the wafer passed through the filter or WO\textsubscript{x} tube, accurate flow measurement was unnecessary; the output was calculated in ng/min. For six filter samples collected throughout the period of atmospheric sampling, the mean output was 96 ± 9 ng/min.

The TAT was calibrated each morning between 0530 and 0830 hrs. Results for nine calibrations, based, with one exception, on 4-8 data points spanning the range expected during atmospheric sampling, are shown in Table 4. To assess the reproducibility of the calibration from day to day at a typical daytime HNO\textsubscript{3} dosage, the HNO\textsubscript{3} corresponding to $2 \times 10^6$ area units was calculated by each of the regression equations. From the nine equations the mean HNO\textsubscript{3} was 100.0 ± 10.5 ng for a C.V. of 10.5%.
TABLE 4

Compilation of HNO₃ Calibration Results

<table>
<thead>
<tr>
<th>Date</th>
<th>Time</th>
<th>Intercept</th>
<th>Slope</th>
<th>r</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>9/07/84</td>
<td>evening-2200</td>
<td>-0.0109</td>
<td>0.0186</td>
<td>0.972</td>
<td>7</td>
</tr>
<tr>
<td>9/08/84</td>
<td>0600-0800</td>
<td>-0.0297</td>
<td>0.0178</td>
<td>0.993</td>
<td>5</td>
</tr>
<tr>
<td>9/15/84</td>
<td>0900-1200</td>
<td>-0.306</td>
<td>0.0257</td>
<td>0.937</td>
<td>8</td>
</tr>
<tr>
<td>9/16/84</td>
<td>0600-0900</td>
<td>-0.350</td>
<td>0.0225</td>
<td>0.994</td>
<td>6</td>
</tr>
<tr>
<td>9/17/84</td>
<td>0530-0900</td>
<td>+0.136</td>
<td>0.0166</td>
<td>0.992</td>
<td>5</td>
</tr>
<tr>
<td>9/18/84</td>
<td>0530-0900</td>
<td>+0.074</td>
<td>0.0203</td>
<td>0.999</td>
<td>4</td>
</tr>
<tr>
<td>9/19/84</td>
<td>0530-0700</td>
<td>-0.54</td>
<td>0.0253</td>
<td>0.995</td>
<td>6</td>
</tr>
<tr>
<td>9/20/84</td>
<td>0700-0830</td>
<td>-0.209</td>
<td>0.0237</td>
<td>0.996</td>
<td>5</td>
</tr>
<tr>
<td>9/21/84</td>
<td>0700-0830</td>
<td>-0.394</td>
<td>0.0236</td>
<td>0.999</td>
<td>3</td>
</tr>
</tbody>
</table>

a. \[ y = a + bx \]

\[ y = \text{P}k \text{ Area (Area units } \times 10^{-6}) \]
\[ x = \text{ng HNO}_3 \text{ (as NO}_3^-\text{)} \]
\[ r = \text{Pearson's correlation coefficient} \]
\[ n = \text{number of data points (generally includes two replicates at each level).} \]

F. Calibration for NH₃ During Atmospheric Sampling

During atmospheric sampling, ammonia in air was supplied by a permeation tube maintained at 22.4 ± 0.2°C in a Wallace-Fisher oven inside an air-conditioned mobile laboratory. The flow over the tube was about 1 Lpm of ambient air which had passed through a glass fiber filter followed by a denuder tube coated with phosphorus acid to remove NH₃. The NH₃ emission rate was measured by sampling the total flow through two tandem oxalic acid-impregnated quartz filters. For seven sampling periods, the mean emission rate was 272 ± 33 ng/min.

Substantial difficulties were encountered in calibrating the TAT for NH₃ including lack of reproducibility and non-linearity of response. Atmospheric sampling results frequently exhibited relatively high NH₃ response or negligible values, implying a loss of system conditioning unless high NH₃ levels were encountered. The following section describes supplementary laboratory studies intended to elucidate and overcome these difficulties.
G. Laboratory Evaluation of the Automated TAT Response to NH$_3$ Following Atmospheric Sampling

To elucidate the problems encountered with NH$_3$ calibration and measurements, additional laboratory trials were made. The difference between ambient air temperature (24 hr mean temperature 26-30°C) and laboratory air temperature (estimated 20-24°C) was the only apparent change in condition. No modifications to the automated TAT were made and the system was deliberately not cleaned.

The elevated response observed with low dosages of added NH$_3$ was apparently due, in part, to contamination of the system during atmospheric sampling. Washing of fittings between the transfer tube and gold coated tube NH$_3$ converter as well as the solenoid valve (C in fig. 1) between the preconcentrator and transfer tubes diminished the NH$_3$ response. It is unclear how frequently such cleaning is necessary during atmospheric sampling. Response of the analyzer after such cleaning remained unreproducible, however.

In an effort to improve the reproducibility of the analyzer, the transfer tube was replaced with a fresh one and the gold-coated NH$_3$ converter tube (about 50% devoid of Au coating) replaced with a tube containing gold foil. The linearity of response was substantially improved. The observed ammonia response was now:

$$y = 1.01 + 0.0256x$$

where $y =$ integrator peak area in units $10^6$ area units

$$x = \mu g \text{NH}_3 \text{ (as NH}_4^+\text{)} \text{ in the range 23 to 204 ng NH}_3.$$

The instrument performance remained erratic, however, with dosages $\leq 45$ ng.
A clue to the more fundamental problem with NH₃ sampling was provided by observations of substantial ammonia peaks when helium or other gas scrubbed free of NH₃ was sampled. Furthermore, a clean 50 cm quartz tube (i.e. without WOₓ coating) retained NH₃ effectively when dosages of ca. 100 ng were sampled. It appears, therefore, that substantial amounts of NH₃ are retained on glass tubing walls ahead of the tube and/or on the ca. 8 cm uncoated section* of the quartz tube. When the WOₓ tube was removed for dosing with NH₃ and then replaced, the heating coil was not repositioned in precisely the same location. Thus a variable amount of the NH₃ collected ahead of the WOₓ coating may have desorbed. The ability of the glass and/or quartz tubing ahead of the WOₓ coating to retain NH₃ may account for the difficulties in preconditioning the system for NH₃.

*A clear section ahead of the WOₓ coating is intended to insure laminar flow.
V. LABORATORY EVALUATION OF AMMONIA SAMPLING WITH OXALIC ACID-COATED DENUDER TUBES

A. Experimental

Oxalic acid-glycerol coated denuder tubes were compared to oxalic acid-glycerol impregnated filters for NH$_3$ collection. The latter had previously been shown by us to collect > 99% NH$_3$ at 100 ppb and 50% R.H. Glass tubes, etched for 3 min in 50% HF, were coated with a methanol solution containing 1.5% oxalic acid and 7.2% glycerol. The solvent was removed by passing N$_2$ (99.999%) through each tube for 4-5 min. The mean tube blank was 0.24 ± 0.08 µg NH$_4^+$. The Ferm tubes sampling at 1 Lpm were compared to oxalic acid impregnated, 47 mm quartz fiber (PallFlex QAO) filters sampling at 15 Lpm. Identical lengths of pyrex tubing were used ahead of all samplers to equalize losses to inlet surfaces.

B. Results

Figure 3 compares results for 16-hr (solid dots) and 4 ± 1-hr trials (open dots). The long term samples showed nearly equal results by the two methods, whereas the short term samples differed by about 50%. Including all samples, the regression equation shows about 20% lower values by the Ferm tube method. Since atmospheric sampling for NH$_3$ was to be done for 12-hr and 24-hour periods with the tubes prepared as described, nearly quantitative collection of atmospheric NH$_3$ by the denuder tube as well as the filter method, was expected.
Figure 3. Laboratory Comparison of Denuder Tube and Filter NH$_3$ Determinations

- 16 hr samples
- O 2-4 hr samples

$y = -0.239 + 0.817x$

$r = 0.82$
VI. LABORATORY EVALUATION OF HYDROCHLORIC ACID SAMPLING

A. Introduction

A dual filter technique for gaseous HCl was evaluated in which particulate chloride was removed by a Teflon prefilter and the gaseous HCl captured with a cellulose filter impregnated with a mixture of triethanolamine and KOH. The choice of this technique was based largely on the report of its high efficiency at hi-vol sampler face velocity (14). If confirmed, such sampling would permit relatively short-term (e.g., 4-hr) HCl sampling and, therefore, assessment of diurnal concentration changes.

The objectives of the laboratory experiments were (1) to measure single triethanolamine-KOH filter efficiency, (2) to measure loss of HCl by sorption-desorption on clean Teflon prefilters and on the filter holder, and (3) to evaluate alternative filter configurations for ambient air sampling.

When applied to atmospheric sampling, the chloride measured on the impregnated filter was taken to be the upper limit concentration for atmospheric HCl.

B. Experimental

Gaseous HCl in N₂ was introduced into an airstream purified by passage through charcoal, Purafil, triethanolamine/diatomaceous earth and a HEPA filter. The temperature of the airstream and relative humidity were held constant within 0.5°C and 1% R.H., respectively. After vortex mixing, the diluted HCl was sampled simultaneously with five samplers:

A. A 47 mm diameter, 2 µm pore size Teflon (Zefluor) pre-filter and a 47 mm triethanolamine-KOH impregnated Whatman 41 (T-K) after-filter contained in separate sections of a Nuclepore dual filter holder (Figure 4). The filters were extracted and analyzed separately.
Figure 4. Nuclepore Multiple Holder Adapter for Collection on Several Filters in Series
B. The same filters as in A contained in a single-stage Nuclepore filter holder without a spacer between the filters. The filters were extracted and analyzed together.

C. The same filters and filter holder as in (B) but with a 1000 µm hole size Teflon mesh separating the filters. To achieve an airtight seal it was necessary to use an anodized aluminum filter holder in place of the plastic Nuclepore holder. The filters were extracted and analyzed separately.

D. Two T-K filters contained in the same Nuclepore filter holder without a spacer between filters. The filters were extracted and analyzed together.

E. Two T-K filters each contained in separate sections of a Nuclepore dual filter holder. The filters were extracted and analyzed separately.

All samplers operated at 37.4 Lpm to provide a face velocity approximately equal to that of a hi-vol sampler with 8" x 10" filter at 40 cfm.

Storage of samples prior to extraction was at < 5°C. The filters were extracted in 10 ml of 0.002 M NaHCO₃ by mechanical shaking for 1 hour at room temperature, in plastic screw-capped test tubes. The Teflon filters and Teflon plus T-K filter were agitated for an additional 30-60 min. with a Fisher RotoRack, also at room temperature. Chloride analysis was by ion chromatography. Experiments were conducted at two HCl concentration levels, 4-5 µg/m³ and 20-25 µg/m³ with 3 replications at each level. Samples were collected for four hours at low concentrations and for two hours at the higher concentrations.

C. Results

Chloride recovered from sampler D was used as a measure of the true HCl concentration. Results are expressed as a percent of the sampler D concentration. This strategy is supported by (1) the high single T-K
filter efficiency demonstrated below, and (2) the minimal filter holder surfaces exposed to HCl in this configuration, minimizing possible losses.

**Figures 5a and 5b** show schematically the experimental design as well as mean results for the low and high concentration trials, respectively. Single T-K filter efficiency can be inferred from the results for sampler E and A. Sampler E indicates a mean efficiency of 96% for one T-K filter at both concentrations. The mean Cl⁻ on the T-K after-filter of sampler E, 0.5 and 2.4%, suggests that the T-K prefilter was >97% efficient. Because of possible loss to the filter holder, this is an upper limit value. The sampler A results indicate a loss of about 8% HCl at high concentration and complete recovery at low concentration. Because of HCl retention on Teflon and loss to filter holder surfaces, the value 89.3% for sampler A (Figure 5b) is a lower limit to the single filter efficiency for T-K filters.

Recovery of HCl from atmospheric particle-free Teflon filters reflects the net result of sorption and desorption. Results with sampler A suggest only a small degree of retention, if any (mean 2.4%). Eight percent of the HCl is unaccounted for and may have been lost by desorption from the prefilter prior to extraction in spite of refrigerated storage. Sorption on the prefilter support and walls of the sampler probably contribute to the 8% loss, as well. Lower and upper limit measurements for HCl retention on Teflon filters are 2.4 ± 4.2% and 8.4 ± 7.2%, respectively.

Loss of HCl to the prefilter support and/or walls of the Nuclepore filter holder is best measured by sampler A. Consistent with the preceding discussion, this could range from 0 to 8.4%. The results for sampler B show loss of 24-34% of the HCl sampled. The reason for the greater loss compared to sampler A was not firmly established. Interference by
Figure 5a. Efficiency of HCl Collection on Teflon Plus Triethanolamine-KOH (T-K) Impregnated Filters (% Relative to Sampler D)

Low Concentration Trials

<table>
<thead>
<tr>
<th></th>
<th>Teflon + T-K</th>
<th>Teflon + T-K</th>
<th>2 T-K</th>
<th>T-K</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>93.0 ± 4.4</td>
<td>66.1 ± 9</td>
<td>68.3 ± 12</td>
<td>2.4c</td>
</tr>
<tr>
<td>B</td>
<td>2.4 ± 2.2</td>
<td></td>
<td>48.2 ± 2.4</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E</td>
<td></td>
<td></td>
<td>95.7 ± 8.3</td>
<td></td>
</tr>
</tbody>
</table>

(a) Experimental conditions: 50% R.H.
Flow rates 37.4 LPM
Temp. 20°C
Sampling time 5 hr

b. Concentrations for 3 trials: 3.7, 4.8 and 4.5 µg/m³, respectively.

c. Single trial. Two values discarded as unreliable.
Figure 5b: Efficiency of HCl Collection on Teflon Plus Triethanolamine-KOH (T-K) Impregnated Filters (% Relative to Sampler D)\(^a\)

High Concentration Trials\(^b\)

<table>
<thead>
<tr>
<th>Teflon + T-K</th>
<th>Teflon + T-K</th>
<th>2 T-K</th>
<th>T-K</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.4 ± 4.2</td>
<td>1.1 ± 1.5</td>
<td>100</td>
<td>96.1 ± 12.0</td>
</tr>
<tr>
<td>39.2 ± 5.8</td>
<td>64.8 ± 16.6</td>
<td></td>
<td>0.5 ± 0.7(^c)</td>
</tr>
</tbody>
</table>

\(\text{(no spacer, ext. together)}\) \(\text{(Teflon mesh spacer)}\) \(\text{(no spacer, ext. together)}\)

---

\(\text{a. Experimental conditions: 50\% R.H.}\\
\text{Flow rates 37.4 LPM}\\
\text{Temp. 20} \degree C\\
\text{Sampling time 2 hr}\\
\)

\(\text{b. Concentrations for 3 trials: 19.6, 22.8 and 25.7 µg/m}^3\text{, respectively.}\\
\)

\(\text{c. Two values averaged. Outlier discarded.}\)
the Teflon filter extraction of the T-K filter was not the problem since, in water, the filters pieces separated due to the density difference. However, it was subsequently demonstrated that with a Teflon and T-K filter in the same Nuclepore holder, a vacuum tight seal was not readily achieved. Sampler leakage is the most likely explanations for the greater loss of sampler B.

Regarding possible atmospheric sampling options, samplers A and C, permitting separate analysis of particulate and gaseous chloride (presumed to be mainly HCl in the atmosphere), were the two techniques considered. The consistently low recovery of gaseous chloride with sampler C (65-68%) compared to 89-98% with Sampler A may reflect HCl loss to the anodized aluminum filter holder, and/or to the Teflon mesh, as well as poor sealing of the dual filter plus spacer sandwiched in the holder. The latter is less likely, however, since the loaded filter holder was checked for vacuum tightness before starting HCl collection.

We conclude from this work that for HCl dosages of 36-112 µg sampled at a face velocity of 47 cm/sec:

1. Single T-K filter efficiency for HCl is at least 96%.

2. Retention of gaseous HCl on Teflon filters is at least 2.4%.

3. The configuration shown as Sampler C (Figures 5a and 5b) should not be used for atmospheric sampling because of HCl loss to the analyzed aluminum filter holder and/or the Teflon mesh spacer.

4. Atmospheric sampling should be done with a system analogous to sampler A. By using 88 mm diameter filters* at a flow rate of 4 cfm (face velocity 34.9 cm/sec), relatively short term (e.g., 4-hr) sample collection can be used.

*Two dual filter samplers designed to minimize filter support surface area to minimize losses of reactive gases were furnished by B. Huebert.
VII. ATMOSPHERIC SAMPLING

A. Introduction

Atmospheric sampling was done on the campus of the University of California (Riverside) in collaboration with the Statewide Air Pollution Research Center. The samplers and sampling strategy employed are detailed in Table 5.

The principal objectives of this sampling were:

(1) to evaluate the performance of the automated TAT for HNO$_3$ and NH$_3$ measurement.

(2) to obtain HNO$_3$ measurements by the denuder difference technique (i.e. sampler 3 NO$_3^-$-sampler 2 NO$_3^-$) for comparison with the automated TAT.

(3) to obtain measurements of NH$_3$ by oxalic acid-coated denuder tube and filter collection methods for comparison with one another and with NH$_3$ by the automated TAT.

(4) to measure upper limit values for atmospheric HCl by a dual filter technique.

These measurements were supplemented by those for O$_3$, NO, NO$_2$, dewpoint and temperature to aid in data interpretation.

Simultaneous measurements with these samplers were performed for the period 1600 hr 9/7 to 0800 hr, 9/9 and, again, from 1200 hr 9/15 to 1200 hr 9/21/84. The first period was characterized by exceptionally low pollution levels. Accordingly, sampling was suspended. The second period included days of moderate-to-heavy photochemical smog (O$_3$ maxima > 0.3 ppm).
### TABLE 5.
**Sampling Strategy - Riverside, September 1984**

<table>
<thead>
<tr>
<th>Sampler Number</th>
<th>Sampler</th>
<th>Collection Medium</th>
<th>Samples/day</th>
<th>Sampling Rate, Lpm</th>
<th>Species Measurement</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>TAT</td>
<td>$\text{WO}_x$</td>
<td>Semi-continuous</td>
<td>1.0</td>
<td>$\text{HNO}_3, \text{NH}_3$</td>
</tr>
<tr>
<td>2</td>
<td>total fine $\text{NO}_3^-$</td>
<td>Nylon$^a$ filter (following MgO-coated denuder)</td>
<td>$5^b$</td>
<td>20</td>
<td>fine $\text{NO}_3^-$, $\text{SO}_4^{2-}$</td>
</tr>
<tr>
<td>3</td>
<td>fine particle $\text{NO}_3^-$</td>
<td>Nylon filter</td>
<td>5</td>
<td>20</td>
<td>fine $\text{NO}_3^-$, $\text{SO}_4^{2-} + \text{HNO}_3$</td>
</tr>
<tr>
<td>4</td>
<td>Ferm tube</td>
<td>oxalic acid</td>
<td>2</td>
<td>1.0</td>
<td>$\text{NH}_3$</td>
</tr>
<tr>
<td>5</td>
<td>Ferm tube</td>
<td>oxalic acid</td>
<td>1</td>
<td>1.0</td>
<td>$\text{NH}_3$</td>
</tr>
<tr>
<td>6</td>
<td>dual filter $\text{NH}_3$</td>
<td>Teflon$^c$ prefiltir</td>
<td>5</td>
<td>25</td>
<td>$\text{NH}_4^+$ (particle phase) $\text{NH}_3$ (gas phase)</td>
</tr>
<tr>
<td>7</td>
<td>dual filter $\text{HCl}$</td>
<td>Teflon$^e$ prefiltir</td>
<td>3</td>
<td>4.0 cfm</td>
<td>$\text{Cl}^-$ (particle phase) $\text{Cl}^-$ (gas phase)</td>
</tr>
<tr>
<td>8</td>
<td>Hi-Vol</td>
<td>glass fiber$^f$ filter</td>
<td>1</td>
<td>40 cfm</td>
<td>$\text{NO}_3^-$, $\text{SO}_4^{2-}$</td>
</tr>
</tbody>
</table>

Chemiluminescent $\text{NO}_x$ (Monitor Labs) ---- Continuous 0.75 NO, $\text{NO}_2$

dew point ---- Continuous dew point

dasibi $\text{O}_3$ monitor ---- Continuous $\text{O}_3$

Resistance thermometer device ---- Continuous temperature

---

a. Nylasorb (47 mm dia.) filters, Membrana Corp.
b. 0800-2400 hr, 4 hour samples. 0000-0800 hr, one sample.
c. Two $\mu$m pore size Zeflour (47 mm dia.), Membrana Corp.
d. Pallflex 2500 QAO (47 mm dia.) quartz fiber filters.
e. Three $\mu$m pore size Zefluor (88 mm dia.), Membrana Corp.
f. Schleicher and Schuell, 1HV filters (1981 EPA-Grade).
B. Semi-Real Time Nitric Acid Results

For the period 2200 hours, 9/7 - 0700 9/8, the HNO₃ concentration, measured with the automated TAT, ranged from 0.6 to 0.8 µg/m³. During the daytime period, 0800 - 1700 hr 9/8, the concentration ranged from 0.6 to 1.6 µg/m³ with maxima at 0900 - 0930 and 1330 - 1400 hours. These relatively low values were consistent with ozone concentration < 0.05 ppm.

Nitric acid measurements resumed at noon, 9/15/84 with concentrations ranging from 3 to 6 µg/m³. However equipment failure prevented measurements after 2200 hr. Nitric acid concentrations for five successive 24-hour periods beginning 9/16/84 are shown in Figures 6-10. Concentrations are shown as one hour average values. For periods beginning 9/16 and 9/17, hourly averages are based upon two, 20 minute samples per hour. For the remaining days, hourly averages are based on four, 5 minute samples per hour. Such short sampling times were needed to prevent saturation of the preconcentrator tube.

Short term nitric acid levels, as measured with the TAT, exceeded 40 µg/m³ at their maximum. Maxima were observed during midday. The sharp decrease on 9/16 coincided with rain which persisted in the area from 1400-1800 hours. Figure 11 shows the moderately high (r = 0.78) correlation between HNO₃ by the TAT and hourly average O₃ values.

Figure 12 is a scatter diagram of NO₂ against HNO₃ concentrations and reveals little or no correlation. Thus, although NO₂ may be the principal, relatively stable HNO₃ precursor, other factors (e.g., the concentration of OH radicals and relative humidity) may be more significant in determining HNO₃ concentrations.
Figures 6 and 7. Diurnal Variation of Nitric Acid at Riverside, CA
Figures 8 and 9. Diurnal Variation of Nitric Acid at Riverside, CA
NITRIC ACID, 9/20-9/21/84

CONC. (<ug/m³>)

TIME (PDT)

Figure 10. Diurnal Variation of Nitric Acid at Riverside, CA
Figure 11. Scatter diagram of Atmospheric Ozone vs HNO₃ Concentrations Measured with the Automated TAT
Figure 12. Scatter diagram of Atmospheric NO₂ vs HNO₃ Concentrations Measured With the Automated TAT
C. Fine Particle Nitrate and Sulfate Plus Nitric Acid Results by the Denuder Difference Method

1. Methodology

Sampler 3 nitrate results are taken to represent the sum of fine (i.e. < 2.2 μm) particle nitrate plus nitric acid. Sampler 2, which used a MgO-coated denuder to remove HNO₃, measured fine particulate nitrate, only. The difference (Sampler 3 NO₃⁻) - (Sampler 2 NO₃⁻) measured atmospheric HNO₃.

The accuracy of the procedure requires:

a. The efficient retention of both particulate nitrate and nitric acid on the nylon filters.

b. The efficient removal of HNO₃ by the denuder.

c. The absence of fine particulate nitrate loss in passing through the denuder.

d. The absence of retention on nylon filters of gaseous NOₓ species other than nitric acid.

e. Accurate and precise control of flow rates.

f. Negligible loss of HNO₃ within the teflon-coated cyclone of sampler 3.

Aside from flow rates, only one of these can be readily checked during atmospheric sampling. If fine particulate sulfate is assumed to be similar to fine particle nitrate regarding loss within the denuder, and assuming equivalent sulfate sampling errors, then comparison of sulfate concentrations on sampler 2 and 3 provides an estimate of such loss. Figure 13 compares sulfate on the two
Figure 13. Comparison of Atmospheric Sulfate Collection With and Without a MgO-Coated Denuder
samplers. Results are generally higher on the sampler with denuder (ratio of means 1.17) which might indicate appreciable loss of fine particles. If true, this would result in HNO$_3$ values which are too high using the denuder difference method. Prior studies (19) comparing such sulfate measurements on Teflon filters showed differences of < 1%. We infer that the present sulfate results on nylon filters without the denuder, rather than indicating particle loss in the denuder, are too high because of partial retention of SO$_2$ or other species later analyzed as SO$_4$$^2$ (22).

Flow rates on both samplers were regulated by mass flow controllers and remained constant within about 3%. Factors a-d have been dealt with in prior studies (3,4 and references cited therein) and were not re-investigated here. Factor f will be checked during Phase II of this study.

2. Results

Table 6 lists nitrate and sulfate results together with HNO$_3$ concentrations by the denuder difference method and other measurements. These are discussed in relation to TAT measurements in the following sections.

D. Comparison of HNO$_3$ Measurements by the TAT and Denuder Difference Method

To permit comparison of the TAT and denuder difference method (DDM) results, 1 hour average TAT HNO$_3$ values were averaged for the time periods of filter collection. The results are compared in Figure 14 and indicate high correlation (n = 0.94) between methods for concentrations > 5 $\mu$g/m$^3$, but about 50% higher values by the TAT.

Possible sources of such difference include:
### TABLE 6
Filter Sampling Results Plus Mean Temperatures at Riverside\(^a\)

<table>
<thead>
<tr>
<th>Date</th>
<th>Time (PDT)</th>
<th>Fine (\text{SO}_4)^{2-}</th>
<th>Fine (\text{NO}_3)^{-}</th>
<th>HNO(_3)^{C}</th>
<th>NH(_4^+)^{d}</th>
<th>NH(_3) as NH(_4^+)</th>
<th>T((^\circ)C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>9/7/84</td>
<td>1600-2200</td>
<td>-(^e)</td>
<td>1.29</td>
<td>3.83</td>
<td>-</td>
<td>16.0</td>
<td>26.1</td>
</tr>
<tr>
<td>9/8</td>
<td>2200-0800</td>
<td>-</td>
<td>1.61</td>
<td>0.89</td>
<td>-</td>
<td>7.69</td>
<td>22.2</td>
</tr>
<tr>
<td>0800-1200</td>
<td>-</td>
<td>1.39</td>
<td>4.79</td>
<td>-</td>
<td>9.29</td>
<td>26.5</td>
<td></td>
</tr>
<tr>
<td>1200-1600</td>
<td>-</td>
<td>0.50</td>
<td>3.57</td>
<td>0.52</td>
<td>3.66</td>
<td>30.0</td>
<td></td>
</tr>
<tr>
<td>1600-2000</td>
<td>-</td>
<td>0.40</td>
<td>1.87</td>
<td>0.54</td>
<td>4.12</td>
<td>29.7</td>
<td></td>
</tr>
<tr>
<td>2000-0800</td>
<td>-</td>
<td>2.22</td>
<td>-0.90</td>
<td>0.46</td>
<td>11.8</td>
<td>23.9</td>
<td></td>
</tr>
<tr>
<td>9/15</td>
<td>1200-1700</td>
<td>2.95</td>
<td>12.5</td>
<td>-6.21</td>
<td>-</td>
<td>16.2</td>
<td>-</td>
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<tr>
<td>1700-2000</td>
<td>-</td>
<td>2.12</td>
<td>0.70</td>
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<td>0.92</td>
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<td>2.29</td>
<td>1.07</td>
<td>1.27</td>
<td>1.20</td>
<td>13.4</td>
<td>22.8</td>
</tr>
<tr>
<td>9/16</td>
<td>0000-0800</td>
<td>2.99</td>
<td>4.44</td>
<td>-2.30</td>
<td>3.53</td>
<td>17.0</td>
<td>22.9</td>
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<tr>
<td>0800-1200</td>
<td>-</td>
<td>3.90</td>
<td>3.71</td>
<td>10.7</td>
<td>1.68</td>
<td>18.2</td>
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<tr>
<td>1200-1600</td>
<td>-</td>
<td>3.20</td>
<td>1.37</td>
<td>4.53</td>
<td>1.15</td>
<td>11.8</td>
<td>27.7</td>
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<tr>
<td>1600-2000</td>
<td>-</td>
<td>2.05</td>
<td>1.11</td>
<td>0.60</td>
<td>0.80</td>
<td>15.9</td>
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<td>2000-2400</td>
<td>-</td>
<td>2.67</td>
<td>1.23</td>
<td>-0.56</td>
<td>0.89</td>
<td>20.0</td>
<td>26.2</td>
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<tr>
<td>9/17</td>
<td>0000-0800</td>
<td>2.05</td>
<td>0.69</td>
<td>0.91</td>
<td>1.16</td>
<td>20.0</td>
<td>26.0</td>
</tr>
<tr>
<td>0800-1200</td>
<td>-</td>
<td>3.78</td>
<td>6.79</td>
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<td>-</td>
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<td>5.23</td>
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</tr>
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<td>28.2</td>
</tr>
<tr>
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<td>3.34</td>
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<td>2.99</td>
<td>49.8</td>
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<td>1.05</td>
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<td>17.6</td>
<td>25.9</td>
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<td>0800-1200</td>
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<td>0.20</td>
<td>1.41</td>
<td>3.57</td>
<td>26.2</td>
<td></td>
</tr>
</tbody>
</table>

\(a\) Except as noted, values are in \(\mu\text{g/m}^3\).

\(b\) Mean result for sampler 2 and 3.

\(c\) By the denuder difference method.

\(d\) Particulate NH\(_4^+\) collected without size segregation.

\(e\) Not determined.
Figure 14. Comparison of Atmospheric HNO₃ Concentrations Measured by the Denuder Difference Method and Automated Tungstic Acid Technique

TAT = 1.24 + 1.48DDM
r = 0.94
n = 31
(1) Positive interference in the TAT by nitrogen oxides, including NO₂, nitrous acid (HONO), particulate nitrate (in spite of the absence of measureable interference from NO₂ and particulate nitrate under laboratory conditions), and possibly by organic nitrates. In addition to the interference studies in the present program, PAN, propyl-nitrate, HCN and NO₂ have been shown not to interfere with the TAT (7,20).

(2) Penetration of HNO₃ through the nylon filters employed.

To assess the source of the difference in HNO₃ measured by the TAT and denuder difference method (DDM), Figure 15 plots the difference in results by the two methods against NO₂. The results are positively correlated (r = 0.62) in contrast to the TAT HNO₃ vs. NO₂ (see Figure 12). Since NO₂ is probably not retained on WOX tubes, these data suggest that species whose concentration correlate with NO₂ (e.g., organic nitrates) may be retained. Figure 16 is a scatterdiagram of the TAT-DDM difference against relative humidity over the same averaging periods (4-8 hr). No correlation was observed. Since R.H. was highest at night and since HONO concentrations can build up at night, greatest differences might be expected at high R.H. However, Hoell has observed (21) decreasing collection efficiency for HNO₃ with WOX tubes at increasing R.H., a factor tending to give low TAT results.

Although prior studies (3) have demonstrated the high efficiency of nylon filters for nitric acid, recent unpublished studies (17) suggest that this may not always be the case. It is argued that in the presence of atmospheric moisture, nitric acid forms a gas phase hydrate with a diffusion coefficient substantially below that for HNO₃. The reduced diffusion coefficient would reduce the efficiency of trapping nitric acid during passage through a Nylon filter.

In the present study this was re-investigated using nitric acid diluted with both dry gas (O₂) and filtered room air at 56% R.H. Tungstic oxide-coated tubes sampled ahead and, subsequently, downstream of test filters. The
Figure 15. Scatterdiagram of the Difference in Atmospheric HNO₃ by the TAT and DDM vs NO₂ Concentrations

\[ Y = 3.11 \times X + 39.1 \]

\[ R = 0.622 \]
Figure 16. Scatter diagram of the Difference in Atmospheric HNO₃ by the TAT and DDM vs Relative Humidity
collected acid was then determined by the usual thermal conversion and chemiluminescence detection. Nylon filters sampling at 10 Lpm showed < 6% penetration. With NaCl/Whatman 41 filters at 20 Lpm, the apparent penetration was 3.1%. These results are similar to those we previously observed (3). We conclude that Braman's observations possibly resulted from artifacts in his experimental procedure. The high efficiency of nylon filters for nitric acid was also observed recently by Roberts (18) using a tunable diode laser technique.

The present findings showing about 50% higher HNO₃ by the automated TAT are similar to those reported by Roberts et al. (20) at a site about 50 km west of Boulder, Colorado (Niwot Ridge). Using the nitrate collected on nylon filters by the dual filter technique as a measure of nitric acid, an automated TAT yielded values about a factor of three higher. They concluded that unidentified, probably carbonaceous nitrogen oxides were being retained on WOX tubes and analyzed together with HNO₃. Nitrous acid was not considered a likely source of interference because of the high daytime photolysis rate for this material. The present results contrast with those of McClenny et al. which showed agreement within 20% between the DDM and automated TAT in sampling near Research Triangle Park, N.C. (9).

E. Ammonia Measurements with the TAT, Oxalic Acid-Coated Filters and Denuder Tubes

It was previously noted that problems with the TAT prevented its calibration for NH₃ measurement. The responsiveness of the TAT to atmospheric NH₃ is indicated by Figure 17. This plots NH₃ values obtained with oxalic acid impregnated filters against integrator areas for the peak ascribed to NH₃. Data points enclosed in the box are believed to reflect the loss of conditioning of the TAT when air relatively free of NH₃ was sampled. The remaining points exhibited relatively good linear correlation. At higher NH₃ levels, the TAT appeared to be responding approximately linearly to NH₃. Phase II studies with the TAT will attempt to overcome the problems indicated by these results.
Figure 17. Scatter diagram of Atmospheric NH₃ Concentrations (by Filter Method) vs NH₃ Peak Area with the Automated TAT
In addition to NH$_3$ measurements by the dual filter technique, NH$_3$ was collected with oxalic acid-coated denuder tubes with both 12 hr and 24 hr collection times. Filter results for 4 and 8 hr samples were averaged to permit comparison to denuder NH$_3$ results.

Figure 18 compares filter results to the 12 hr denuder tube NH$_3$ values and indicates excellent correlation, but somewhat higher values by the filter method. The ratio of mean values, (Filter NH$_3$)/(Denuder NH$_3$) was 1.12. Higher NH$_3$ results by the filter method are expected because of dissociation of ammonium salts (especially NH$_4$NO$_3$) on relatively inert Teflon prefilters.

The limited duration of sampling hampers adequate comparison of 12 hr and 24 hr NH$_3$ denuder results. For five, 24 hour periods, the ratio of means, (24-hr NH$_3$ calculated from 12 hr samples)/(observed 24-hr NH$_3$) was 1.23 indicating higher average results for 12 hr samples. These findings are consistent with diminished collection efficiency after prolonged sampling at the relatively high NH$_3$ levels existing in Riverside. The capacity of the tubes for NH$_3$ was not established by laboratory trials.

F. Particulate NH$_4^+$ Results

Particulate NH$_4^+$ concentrations were measured without size segregation using a 47 mm Teflon (Zefluor) filter ahead of the oxalic acid filters intended to retain NH$_3$. The results given in Table 6 range from 0.5 to 17 µg/m$^3$. These values may be compared to those calculated from fine particle SO$_4^{2-}$ and NO$_3^-$ with the assumptions that (1) all NH$_4^+$ exists on particles < 2.5 µm, and (2) SO$_4^{2-}$ and NO$_3^-$ are present as (NH$_4$)$_2$SO$_4$ and NH$_4$NO$_3$, respectively. The comparison is given in Figure 19 and indicates moderate correlation ($r = 0.87$) with somewhat higher calculated values compared to those observed. Such findings are consistent with the contributions of cations other than NH$_4^+$.

The fraction of NH$_4^+$ in the particle phase may reflect, in part, the dissociation of NH$_4$NO$_3$. Figure 20 is a scatter diagram of the fraction (partic. NH$_4^+$)/(partic. NH$_4^+$ + NH$_3$ as NH$_4^+$) against temperature. Clearly, the proportion of particle phase NH$_4^+$ is not principally governed by temperature.
Figure 18. Scatter diagram of atmospheric NH$_3$ by Oxalic Acid-Coated Denuder Collection vs Oxalic Acid-Impregnated Filter Collection
Figure 19. Comparison of Observed and Calculated Atmospheric Particulate NH$_4^+$ Concentrations
Figure 20. Scatterdiagram of Mean Ambient Temperature vs the Fraction of NH$_4^+$ plus NH$_3$ in the Particle State
G. HCl and Particulate Chloride

Table 7 summarizes results for gaseous chloride determination taken as upper limit estimates of the concentrations of HCl at the site. Levels ranged from below detection to $3.1 \, \mu g/m^3$ as Cl$^-$ (or 2.0 ppb). A pronounced diurnal pattern was observed with maxima usually in the daytime periods 0800-1200 or 1200-1600 hr.

In the next phase of this study HCl measurements will be made in the western part of the Los Angeles basin where much higher levels are expected. Correlations between the concentrations of gaseous acids (i.e. HCl and HNO$_3$) and atmospheric particulate strong acidity will be sought.

**TABLE 7**

Concentration of Gaseous Chloride as a Measure of HCl at Riverside ($\mu g/m^3$ as Cl$^-$)

<table>
<thead>
<tr>
<th>Date</th>
<th>0000-0800 hr</th>
<th>0800-1200 hr</th>
<th>1200-1600 hr</th>
<th>1600-2000 hr</th>
<th>2000-2400 hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>9/8/84</td>
<td>0.33</td>
<td>0.93</td>
<td>0.40</td>
<td>&lt;0.2</td>
<td>N.D.$^a$</td>
</tr>
<tr>
<td>9/15/84</td>
<td>N.D.</td>
<td>N.D.</td>
<td>0.46</td>
<td>&lt;0.3</td>
<td>&lt;0.3</td>
</tr>
<tr>
<td>9/16/84</td>
<td>0.25</td>
<td>N.D.</td>
<td>0.17</td>
<td>0.25</td>
<td>&lt;0.26</td>
</tr>
<tr>
<td>9/17/84</td>
<td>0.38</td>
<td>1.76</td>
<td>2.77</td>
<td>1.50</td>
<td>0.28</td>
</tr>
<tr>
<td>9/18/84</td>
<td>0.48</td>
<td>3.08</td>
<td>1.46</td>
<td>0.47</td>
<td>0.43</td>
</tr>
<tr>
<td>9/19/84</td>
<td>0.93</td>
<td>1.91</td>
<td>1.13</td>
<td>0.94</td>
<td>0.3</td>
</tr>
<tr>
<td>9/20/84</td>
<td>0.12</td>
<td>1.12</td>
<td>0.76</td>
<td>1.32</td>
<td>0.3</td>
</tr>
<tr>
<td>9/21/84</td>
<td>0.83</td>
<td>1.09</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
</tr>
</tbody>
</table>

a. Not determined.

Sea salt is probably the most significant source of particulate chloride in California samples. Because relatively low levels of particulate chloride were anticipated in Riverside, Teflon prefilters (from which such chloride was determined) were run consistently for 8-hr periods. Table 8 summarize the data obtained. Throughout the period 9/8 - 9/19/84, particulate Cl$^-$ concentrations remained $< 0.8 \, \mu g/m^3$. Substantially higher values were found on 9/20 - 9/21. No linear correlation was observed between particulate chloride and gaseous chloride ($r = -0.16$).
### Table 8
Concentration of Particulate Chloride at Riverside (μg/m³)

<table>
<thead>
<tr>
<th>Date</th>
<th>0000-0800 hr</th>
<th>0800-1600 hr</th>
<th>1600-2400 hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>9/15/84</td>
<td>N.D&lt;sup&gt;a&lt;/sup&gt;</td>
<td>N.D.</td>
<td>0.11</td>
</tr>
<tr>
<td>9/16/84</td>
<td>0.76</td>
<td>0.13</td>
<td>0.21</td>
</tr>
<tr>
<td>9/17/84</td>
<td>0.10</td>
<td>0.16</td>
<td>0.21</td>
</tr>
<tr>
<td>9/18/84</td>
<td>0.71</td>
<td>0.21</td>
<td>0.15</td>
</tr>
<tr>
<td>9/19/84</td>
<td>0.39</td>
<td>&lt;0.15</td>
<td>0.27</td>
</tr>
<tr>
<td>9/20/84</td>
<td>0.47</td>
<td>N.D&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.22</td>
</tr>
<tr>
<td>9/21/84</td>
<td>4.67</td>
<td>1.38</td>
<td>N.D.</td>
</tr>
</tbody>
</table>

<sup>a</sup> Not determined.

<sup>b</sup> For 0800-1200 hr only.

### H. Hi-Vol (Glass Fiber) Filter Results and Comparison of Nitrate Values

Twenty four-hour Hi-Vol filter samples were collected employing glass fiber filters. The filter batch (Schleicher and Schuell 1HV, 1981 EPA Grade) had been previously shown to retain efficiently both particle phase nitrate and nitric acid at TIN levels up to 32 μg/cm² of filter area (19). Hi-Vol SO₄<sup>2-</sup> and NO₃<sup>-</sup> concentrations are shown in Table 9 together with fine total inorganic nitrate (FTIN) values for four 24 hr periods. Equipment malfunction prevented obtaining two additional Hi-Vol samples. FTIN results shown were 24 hr averages calculated from short term samples obtained with Sampler 3.

Hi-Vol NO₃<sup>-</sup> and FTIN are strongly correlated with consistently higher values being obtained with the Hi-Vol. The differences may reflect retention of coarse NO₃<sup>-</sup> as well as more efficient retention of such materials as organic nitrates on the glass fiber filters.

Hi-Vol NO₃<sup>-</sup> as well as FTIN are notable for the substantial day-to-day variability in contrast to SO₄<sup>2-</sup>; the range in 24-h average SO₄<sup>2-</sup> concentration was < 25%.
TABLE 9
Hi-Vol Sampler Results and Comparison of Hi-Vol NO$_3^-$ and FTIN (µg/m$^3$)

<table>
<thead>
<tr>
<th>Date$^a$</th>
<th>Hi-Vol SO$_4^{=}$</th>
<th>Hi-Vol NO$_3^-$</th>
<th>FTIN$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>9/8-9/9</td>
<td>11.1</td>
<td>3.58</td>
<td>2.73</td>
</tr>
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<td>9/17-9/18</td>
<td>11.0</td>
<td>23.0</td>
<td>19.8</td>
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<td>9/18-9/19</td>
<td>11.7</td>
<td>28.3</td>
<td>26.8</td>
</tr>
<tr>
<td>9/20-9/21</td>
<td>9.09</td>
<td>16.6</td>
<td>9.62</td>
</tr>
</tbody>
</table>

Hi-Vol NO$_3^-$ = 3.63 + 0.966 (FTIN)

$r = 0.967$

$n = 4$

---

a. Hi-Vol sampler employed Schleicher & Schuell 1HV glass fiber filters (1981 EPA Grade) and sampled from 0800-0800 hr.

b. Fine total inorganic nitrate as measured by sampler 3.
VIII. REFERENCES


APPENDIX A

PREPARATION OF TUNGSTIC ACID DENUDER TUBES

A. INTRODUCTION

Hollow tubes are coated by vacuum deposition of polymeric blue tungsten oxide then thermally oxidized to tungstic acid.

B. MATERIALS

1. HF - 50% solution.
2. NaOH - 50% solution.
3. Vycor (quartz) tubing, 4 mm I.D., 45 cm length.
4. Metal inserts (short sections of tubing < 4 mm O.D. inserted at each end of tube to be coated to prevent coating the ends of the Vycor tubes).

C. TUBE PREPARATION - New Tubes

1. Supporting the tube vertically, apply suction through a short section of plastic tubing and draw HF solution to above the level of the quartz tube and hold for about two min.
2. Allow HF to drain and rinse thoroughly with distilled water.
3. Similarly treat the tube with 50% NaOH.
4. Rinse well with distilled water.
5. Allow tube to air dry. Do not oven dry tube to avoid contamination of HF cleaned surface.

- A1 -
D. TUBE PREPARATION - Previously Coated Tubes

1. Follow steps 1 and 2 above.

2. Repeat process two or three times. Most of the tungstic acid should begin to flake off quickly. Although not obvious, portions still adhering will also be loosened.

3. Remove tube from HF solution and rinse both inside and outside of tube with running tap water. Flush for 2-3 min. until all traces of tungstic acid are gone. Scrape stubborn spots out with a piece of stiff "bare" copper wire.

4. Finish rinsing with distilled water.

E. COATING TUBES

1. Install tube inserts and tungsten wire in coating apparatus (Figure 2 in text). Make sure the inserts are properly in place and the tungsten wire is centered and straight.

2. Support the apparatus vertically, start the pump and vacuum dry the tube. Increase voltage of external heating coil until it is a barely visible dull red heat (about 110 V with AIHL system).

3. Continue to pump until vacuum is < 100 Torr.

4. Turn heater off and increase tungsten wire voltage setting, while monitoring current flow, until wire is a very bright orange heat (9.2 amps, AC, with AIHL system). For a new wire, note the powerstat setting at the bright orange heat and rapidly decrease then increase the current to that setting several times. Check that the wire is straight and centered in tube.
5. A blue-black coating will be visible in about 5 min., with a heavier coating on the ends. Check the system about every five min. to track coating progress. After 15 to 20 min. the tungsten filament will no longer be visible on the ends and barely visible in the middle.

6. When the glow of the wire is no longer visible in the center, turn off the tungsten powerstat (too thick a coat will result in a granular appearance which may flake when oxidized). When coating chamber is cool, disassemble coating apparatus and remove tube.

F. OXIDATION OF COATING

1. Turn gas-oxygen torch on using burner tip No. 5. Adjust the fuel-oxygen mixture to provide a cool flame (yellow-orange center).

2. Connect coated tube to oxygen feed through swivel, rotate the tube, and begin heating one end of the blue-black coating to a dull red glow.

3. Continually rotate the tube to avoid local overheating. Overheating will vaporize and create holes in the coating. Following oxidation, properly heated sections on cooling will be bright, translucent, uniform yellow in appearance. Advance the burner tip slowly along the entire coated section of the tube. Transparency (always most apparent in the center section) is indicative of a thin coat but does not necessarily mean poor collection efficiency.

4. When all the coating has been satisfactorily oxidized, increase the oxygen flow to the torch to provide a hotter, blue flame and heat the uncoated ends of the tube to red heat to burn off all traces of tungsten from these sections.

5. When cool, disconnect tube and immediately seal tube ends with Parafilm to avoid contamination of coating.
APPENDIX B

INTERFERENCE OF NO₂ IN HNO₃ MEASUREMENTS WITH THE TAT

Efforts to detect interference effects were hampered by the response behavior of the preliminary version of the TAT system. Since changing from HNO₃ to purified air required ca. 3 hours to achieve close to background HNO₃ response, the apparent instrument response to NO₂ was expected to vary with time and the prior history of the analyzer. The first trials with NO₂ employed the analyzer well conditioned for HNO₃, changing test atmospheres from HNO₃ in air to NO₂ in air. Table B-1 gives the results. The initial cycles following the change to NO₂ showed a large increase (ca. 50%) in apparent HNO₃ peak area. Over a 3 hour period this decreased to about half the initial value with NO₂ added, but the peak area remained high (7.6 ± 0.6 X 10⁵). Changing from NO₂ to synthetic air or 80/20 He-O₂ caused no measurable decrease in response (mean 6.6 ± 0.9 X 10⁵). Thus NO₂ appears to have no long-term interference capability. It remains unclear why the blank was so high on this day.

To minimize the uncertainty in interpreting instrument response to interferents, NO₂ was added in mixture with dilute HNO₃ following about 3 hours of conditioning with HNO₃. The addition was done in such a way as to keep the HNO₃ concentration unchanged (within 2%), at about 50 μg/m³ or 20 ppb (est. from prior calibration). The final NO₂ level was 57 μg/m³ (30 ppb). Following addition of NO₂ the apparent HNO₃ peak remained unchanged in area [(8.66 ± 0.54) X 10⁵ vs. (8.69 ± 0.27) X 10⁵], indicating no measurable interference. The change from NO₂-HNO₃ to synthetic air only caused an immediate sharp drop in measured HNO₃.
TABLE B-1

INTERFERENCE OF NITROGEN DIOXIDE IN THE AUTOMATED HNO₃ ANALYZER

<table>
<thead>
<tr>
<th>Atmosphere Sampled</th>
<th>Cycle</th>
<th>TAT Response (Area \times 10^{-5})</th>
</tr>
</thead>
<tbody>
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<td>15.2</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>12.1</td>
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<td>Synthetic air</td>
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a. Each cycle is 30 minutes including a 10 minute sample collection. Cycles 1-14 are consecutive cycles on the same day.

b. An NaCl-impregnated filter was placed in the sampling line just ahead of the inlet to the TAT to remove HNO₃ possibly present as a contaminant from tubing walls.
APPENDIX C

INTERFERENCE OF NITROUS ACID IN HNO₃ MEASUREMENT WITH THE TAT

A. EXPERIMENTAL

Nitrous acid was prepared following a procedure (1) developed at the University of California (Riverside) Statewide Air Pollution Research Center (SAPRC). In this procedure 0.012 M NaNO₃ was added dropwise to a slowly stirred, 5% w H₂SO₄ solution at room temperature, while flowing N₂ through the all glass and Teflon system. Further dilution with N₂ yielded HONO concentrations < 150 ppb. Nitrous acid preparation is made difficult by the co-generation of NO₂ and NO. The studies at SAPRC found molar ratios of HONO/(NO + NO₂ + HONO) of up to 0.5.

In the present study, an NOₓ analyzer provided measurement of the NO formed and of the sum NO₂ + HONO. With this sum and the reported maximum mole fraction of HONO, an upper limit to the HONO concentration could be calculated.

The procedure used for HONO trials included:

1. Conditioning of the TAT system with HNO₃.

2. Calibration of the NOₓ monitor for NO and NOₓ using a known concentration of NO in air.

3. Measurement with the NOₓ monitor the NO and NO + NO₂ + HONO (NOₓ) in the diluted nitrous acid in N₂ immediately before and after the 10 minute sampling by the TAT system. Measurements before and after sampling were needed because the NOₓ from the HONO generator decreased with time. The averages of the NOₓ and NO measurements together with the maximum mole fraction of HONO, were used to determine the upper limit HONO concentration being sampled by the TAT.
4. Measuring the TAT response to HONO as apparent HNO₃.

5. Measuring the TAT response to HNO₃ while simultaneously measuring the HNO₃ concentration by filter collection using NaCl-impregnated Whatman 41 filters.

B. RESULTS

Table C-1 summarizes results for trials with nitrous acid. Earlier work (Appendix B) demonstrated that NO₂ does not interfere. Since interference from NO is much less probable, the present results clearly suggest that HONO does interfere significantly. The results appear to show that HONO is up to 1/3 as responsive as HNO₃ in the TAT. However, determining the efficiency of HONO for interference is made difficult by the use of generally higher levels of HONO compared to HNO₃. Relatively high HONO concentrations were needed to permit real-time measurement of the HONO with the NOₓ analyzer. During some of these trials the WOₓ preconcentrator column may have been saturated by HONO. As a result peak area responses would have been too low for HONO making the calculated response per ppb too low. In addition, since HONO values are upper limit concentrations, response factors should be lower limit values only. Thus it cannot be concluded from the current data that HONO is substantially less responsive than HNO₃ in the TAT.

As an alternative hypothesis to direct interference by HONO, nitrous acid can decompose via the reaction:

\[
3 \text{HNO}_2 + 2 \text{NO} + \text{HNO}_3 + \text{H}_2\text{O}
\]

The generation of nitric acid by decomposition via this route would, of course, produce an interference with a response about 1/3 of that for HNO₃. If such decomposition occurs within the TAT it would represent a source of interference to be expected in atmospheric sampling. If, however, it occurred within the HONO generation system or the Teflon lines transporting the diluted HONO to the TAT, this would be an experimental artifact. The present data do not discriminate between the possible pathways to interference by HONO. Use of
a nylon filter to remove HNO$_3$ ahead of the TAT will eliminate this ambiguity in trials to be done in Phase II.

Quantitative retention of nitrous acid on nylon wool has recently been demonstrated (R. Atkinson, personal communication, 1984). Thus HONO can be measured by a difference method.

\[(\text{HONO}) = (\text{NO}_x)_\text{total} - (\text{NO}_x)_\text{Nylon}\]

where: \((\text{NO}_x)_\text{Nylon}\) is the total NO$_x$ (with HNO$_3$ previously removed by nylon filter) penetrating a tube pack with Nylon wool.

Reference

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a. Estimated assuming mole fraction of HONO is ≤ 0.5.
b. Since HONO concentrations are upper limit values, response factors for HONO are lower limit values.
c. Corrected for blank with synthetic air.
APPENDIX D

ANALYSIS OF HCl AND PARTICULATE CHLORIDE IN ATMOSPHERIC SAMPLES

Teflon prefILTER samples were extracted in 0.0018 NaHCO₃ solution and analyzed by ion chromatography for chloride. Filter blanks for 88 mm diameter filters were low (3.2 ± 4.2 μg/filter). The triethanolamine-KOH impregnated Whatman 41 (T-K) filters were extracted as above. However, ion chromatographic analysis proved unfeasible because of interfering peaks. While such peaks had been observed in laboratory studies as well, their relative magnitude had been much lower, permitting chloride measurement. T-K filter extracts were successfully analyzed by potentiometric titration with AgNO₃.

Potentiometric titration of chloride was done in 15 ml 0.5 M HNO₃, to make the sample extract acidic. This HNO₃ solution was saturated with AgCl by adding 2 μg Cl⁻ and titrating to the potentiometric endpoint with AgNO₃ using a Model ABU 12 autoburette (Radiometer). An 0.8 ml aliquot of the sample extract was then added and the solution again titrated to the endpoint (i.e. the point of maximum rate of change of potential).

The recovery of chloride was assessed with one and two μg samples and ranged from 65-80% with these samples. Precision was assessed with standards and ranged from about 5% with 10 μg samples to about 20% with 1 μg samples. No corrections were applied to the resulting atmospheric data. Further work is needed to assess more completely the precision and accuracy of this method with the atmospheric samples.

During Phase II of this study, alternative sampling media (e.g. Na₂CO₃ impregnated filters) will be evaluated to permit improved quantitation of HCL (as Cl⁻) by ion chromatography.