Sulfuric acid vapour derivations from negative ion composition data between 25 and 34 km

by

E. ARIJS, D. NEVEJANS, J. INGELS and P. FREDERICK
FOREWORD

The paper: "Sulfuric acid vapour derivations from negative ion composition data between 25 and 34 km" is also published in Geophysical Review Letters.

AVANT-PROPOS

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VOORWOORD

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VORWORT

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Abstract

Negative ion composition measurements obtained during a valve controlled balloon flight were used to derive $\text{H}_2\text{SO}_4 + \text{HSO}_\gamma$ number densities between 25 and 34 km altitude. The data are compared to similar results obtained for other stratospheric temperatures. The implications of the results on our present ideas about aerosol formation are briefly discussed.

Résumé

Les mesures de composition des ions négatifs stratosphériques obtenues durant un vol de ballon à clapet, ont été utilisées pour dériver des concentrations de $(\text{H}_2\text{SO}_4 + \text{HSO})$ entre 25 et 34 km d'altitude. Les données sont comparées à des résultats similaires obtenus pour d'autres températures stratosphériques. Les implications des résultats sur nos idées actuelles sur la formation des aérosols sont discutées brièvement.
Samenvatting

Metingen omtrent het negatieve ionenbestand, bekomen gedurende een vlucht met een ventielgecontroleerde balloon, werden gebruikt om dichtheden van \(\text{H}_2\text{SO}_4 + \text{HSO}_\text{y}\) af te leiden in het hoogtegebied van 25 tot 34 km hoogte. De gegevens worden vergeleken met analoge resultaten, bekomen bij andere stratosferische temperaturen. De weerslag van de resultaten op onze huidige voorstellingen omtrent het aerosolvormingsproces worden beknopt besproken.

Zusammenfassung

Messungen der Zusammensetzung stratosphärischer negativer Ionen, durgeführt dauerend einen Flug mit einem Ventielgesteuerten Ballon, wurden gebraucht um die Teilchenanzahldichte von \(\text{H}_2\text{SO}_4 + \text{HSO}_\text{y}\) ab zu leiten zwischen 25 and 34 km Höhe. Die Resultaten werden mit analogen Ergebnisse, für anderen stratosphärischen Temperaturen bekomen, vergleich. Der Wiederschlag dieser Ergebnissen auf unsere heutige Vorstellung der Aerosolbildung wird kürz besprochen.
INTRODUCTION

Although H$_2$SO$_4$ plays a key role in the formation of aerosols (Turco et al., 1982) measurements of its concentration in the stratosphere have not been possible until very recently.

Stratospheric negative ion composition measurements (Arnold and Henschen, 1978; Arijs et al., 1981) have shown the existence of HSO$_4^-$ cluster ions formed through switching reactions between H$_2$SO$_4$ and NO$_3^-$ cluster ions. Subsequent laboratory studies (Viggiano et al., 1980) revealed the rate constants of these reactions and made it possible to deduce sulfuric acid concentrations in the stratosphere (Arnold and Fabian, 1980). The technique explained briefly hereafter, has been exploited intensively by Arnold and coworkers (Arnold et al., 1981; Viggiano and Arnold, 1981; Viggiano and Arnold, 1983).

Most data published so far were obtained during fall. In the present paper we report some recent measurements performed during summer conditions and briefly discuss the implication on the present ideas about aerosol formation.

EXPERIMENTAL AND MEASUREMENTS

The data shown and discussed hereafter were obtained during a flight performed with a valve controlled balloon on 16 June 1982 from the CNES launching base at Gap-Tallard (S. France).

The balloon borne ion mass spectrometer has been described in detail before (Arijs et al., 1980; Ingels et al., 1978; Nevejans et al., 1982).

The mass range of the spectrometer is limited to 330 amu in the high resolution mode, but use of low resolutions allows detection of ions with masses larger than 330 amu.
For the negative ion measurements two resolution modes have been used. At float altitude spectra were obtained in a moderate resolution mode \((m/Am = 17)\), adequate to resolve the major ions. During descent, series of multiple scans with moderate resolution alternated with single scans in the total ion mode (no DC on the quadrupole rods).

In this way spectra were obtained from 34 km down to 25 km altitude, either at float altitude or during the descending phase of the balloon flight.

RESULTS AND DISCUSSION

Two typical spectra obtained in the moderate resolution mode are shown in figure 1. As observed, the amplitudes of the mass peaks due to \(\text{HSO}_4^-(\text{HNO}_3)_m(\text{H}_2\text{SO}_4)_l\) ions (mass numbers 160, 195 and 293) decrease with decreasing altitude, which illustrates the variation of sulfuric acid number density with altitude.

As pointed out by Arnold and Fabian (1980) the sulfuric acid number density \([\text{H}_2\text{SO}_4]\) can be deduced through the formula:

\[
k[n^-] [\text{H}_2\text{SO}_4] = a[n^+] [n_s^-] \quad (1)
\]

where \([n_+]\) is the total positive ion number density, \(a\) the ion-ion recombination coefficient, \(k\) the reaction rate coefficient for switching reactions between \(\text{NO}_3^-\) cluster ions and \(\text{H}_2\text{SO}_4^\cdot\), \([\text{HSO}_4^-]\) the sum of the number densities of all \(\text{HSO}_4^-\) cluster ions and \([\text{NO}_3^-]\) the total density of all ions with a \(\text{NO}_3^-\) core.

Recently it was discovered (Arnold et al., 1982) that other sulfur bearing gases may contribute to the formation of \(\text{HSO}_4^-\) cluster ions. Therefore formula (1) should be replaced by
**Fig. 1a.** Typical negative ion spectrum in the moderate resolution mode obtained at 32 km after summation of 6 scans of 160 s each.

**Fig. 1b.** Moderate resolution negative ion spectrum obtained at 28 km after summation of 5 scans. Both spectra have been smoothed as described before (Arijs et al., 1982).
\[
\kappa[n^-_N] ([H_2SO_4] + [HSO_y]) = \alpha[n^+[n^-_S] \tag{2}
\]

where HSO_y includes HSO_3 and HSO_5 and any other sulfur compounds that can react with NO_3^- core ions to form HSO_4^- core ions.

The calculation of the ion abundance ratio \([HSO_4^-]/[NO_3^-]\) to be used in formula (2) is straightforward. From spectra as those shown in figure 1, the ion abundances are derived assuming that ion count rates reflect ion number densities. This sounds reasonable in view of the moderate resolution used, which yields small mass discrimination. Due to the limited mass range of the mass filter (330 amu) it may be expected that some ions will not be detected. This is especially true at altitudes above 30 km where mass 391, i.e. \(HSO_4^- (H_2SO_4)_3\), represents a considerable fraction of the total ion signal (Arijs et al., 1981). In order to calculate the signal due to missing ions, the spectra obtained in the total ion mode were used. Thereby it was assumed that all ions with masses larger than 330 amu are due to HSO_4^- core ions, which is confirmed by previous measurements (Arijs et al., 1981; Arnold et al., 1982).

For the ion-ion recombination \(\alpha\), recent data based on in-situ measurements (Rosen and Hofman, 1981), laboratory work (Smith and Adams, 1982) and theoretical calculations (Bates, 1982) are available. In order to compromise between these different data we have used a parametrization of the form:

\[
\alpha = 6 \times 10^{-8} \left( \frac{300}{T} \right)^{0.5} + 1.25 \times 10^{-25} [M](300/T)^4 \text{ cm}^3\text{s}^{-1} \tag{3}
\]

where \(T\) is the temperature in Kelvin and \([M]\) the total gas number density in \(\text{cm}^{-3}\).
This parametrization gives α values, representing an average of the data in the literature referred above, between 30 and 20 km altitude. Above 30 km the value given by formula (3) is in better agreement with the recent data of Bates (1982).

The total positive ion density \([n_+]\) is deduced from the continuity equation \([n_+] = (Q/a)^{1/2}\), where the ion pair production rate \(Q\) is calculated with the parametrization formula of Heaps (1978).

For the reaction rate coefficient \(k\), the recently corrected values, published by Viggiano et al. (1982) were used. A value of \(1.1 \times 10^{-9}\) cm\(^3\) s\(^{-1}\) being reported for the reaction of \(\text{H}_2\text{SO}_4\) with \(\text{NO}_3(\text{HNO}_3)_2\) and the latter being the most abundant \(\text{NO}_3^-\) ion observed in our spectra, only this value will be applied in formula (2).

The \(\text{H}_2\text{SO}_4 + \text{HSO}_3^-\) number densities obtained from the negative ion spectra at different altitudes are shown in figure 2, together with a compilation of data previously obtained by our group and by the group of the Max Planck Institut of Heidelberg (MPIH hereafter) (Viggiano and Arnold, 1982). The data are also compared with different model calculations. The curve marked T is chosen from Turco et al. (1981) whereas the curves labeled H refer to model calculations of Hamill et al. (1982).

The numbered curves are \(\text{H}_2\text{SO}_4\) vapour pressure calculations performed assuming that the water vapour mixing ratio in the stratosphere is constant and that the aerosols consisted of liquid droplets of a homogeneous water/sulfuric acid mixture. Following Hamill et al. (1977) we accepted that the water vapour is in equilibrium with the \(\text{H}_2\text{SO}_4 - \text{H}_2\text{O}\) droplets. Using the vapour pressure data of Gmitro and Vermeulen (1964), which seem to be only reliable for water vapour pressure calculations (Verhoff and Banchero, 1972) the weight percentage of \(\text{H}_2\text{SO}_4\) in the aerosols at different altitudes (or different temperatures)
Fig. 2. - \([\text{H}_2\text{SO}_4 + \text{HSO}_y]\) obtained in-situ by different experiments compared to vapour pressure calculation and recent models. Curve T refers to the work of Turco et al. (1981), H are model calculations of Hamill et al. (1982), R.A. representing the radical agglomeration case.
was calculated. From this and the temperature, the partial vapour pressure of sulfuric acid was derived with the formula reported by Ayers et al. (1980). Curves 1 and 2 represent the results of such a computation for a 3 ppm water mixing ratio and a U.S Standard Atmosphere temperature profile, for spring fall and summer conditions respectively.

As was noted by Viggiano and Arnold (1983) there is good agreement between curve 1 and the MPIH data in the altitude region 35 to 28 km. If a constant water vapour mixing ratio of 1.5 ppm is accepted, this agreement is even excellent (curve 1b). This however is rather fortuituous, regarding the low H₂O mixing ratio (1.5 pp), the large uncertainties on the thermochemical data used to calculate the H₂SO₄ vapour pressure (Verhoff and Bunchero, 1972; Ayers et al., 1980) and the errors on the deduced H₂SO₄ number densities. As can be seen the agreement between our data and curve 2 is rather fair. In order to investigate this, we have plotted the temperature versus time as measured by a tiny bead thermistor during the ascent period of the flight. This measurement took place at night so that errors introduced by solar radiation are non existent. The result is represented by curve 1 in figure 3. For comparison U.S. Standard Atmosphere temperatures at the same altitudes are represented by curve 2 and 3 for summer and spring-fall conditions respectively. Strong fluctuations probably related to balloon motions, occur on the observed temperature, especially at ceiling altitude. Nevertheless it can be concluded that the measured temperature is closer to the spring-fall profile after 00.30 UT, which corresponds to about 30 km. For the derivation of the H₂SO₄ vapour pressure we have therefore adopted two tentative temperature profiles, which fit better to the measured one. Using a simple linear relation \( T = 195.2 + 1.15 z \), where \( z \) is the altitude in km, results in curve 3a of figure 3. Taking the same relationship for \( z \leq 28 \) and \( T = 185 + 1.5 z \) for higher altitudes, results in curve 3b.
Fig. 3.- Temperature versus time during ascent in the balloon flight of 16 June 1982.
Curve 1: temperature as measured; 2: U.S. Standard Atmosphere temperature for summer conditions, derived from the conversion of pressure measurement to altitude; 3: same for spring-fall.
The agreement between experimental data and the $H_2SO_4$ vapour pressures now obtained is within the experimental errors down to about 28 km.

According to model calculations (Turco et al., 1979) the concentration of $H_2SO_4$ is much larger than that of HSO$_3$ above 28 km. It is therefore reasonable to conclude that above this altitude the major compound detected by the present technique is $H_2SO_4$. The agreement of our data with the calculated values of sulfuric acid vapour therefore suggests that the $H_2SO_4$ number density above 28 km is mainly controlled by evaporation from aerosols. This confirms the present ideas about aerosol formation and the measurement of Viggiano and Arnold (1983) who drew the same conclusion.

Below 28 km the data values are much higher than the vapour pressure values, indicating supersaturation which should lead to aerosol growth.

It is tempting to compare the data to the models of Turco and Hamill shown in figure 2. This however is premature since the measured concentrations are referring to $H_2SO_4$ plus all other sulfur compounds which can give rise to HSO$_4^-$ ions. These include HSO$_3^-$, the concentration of which can exceed the $H_2SO_4$ concentration below 28 km (Turco et al., 1981). Therefore a comparison should be made with the total concentration of all sulfur compounds resulting in HSO$_4^-$ clusters. Unfortunately no recent models giving this quantity as a function of altitude and temperature are available yet.

It is evident however from figure 2 that $H_2SO_4 + HSO_4^- y$ concentrations are found which are considerably higher than in the experiments of Viggiano and Arnold (1983). It should be noted that all our data were obtained during descent, so that contamination effects cannot be responsible for the higher values. Furthermore this difference is not
due to the use of a value of the ion-ion recombination coefficient $\alpha$ and the ion production rate $Q$, being different from those used by the referred authors. On the contrary even higher $H_2SO_4 + HSO_y$ concentrations are found with their $\alpha$ and $Q$ values and our ion abundances.

At present however it is difficult to conclude whether this is due to higher stratospheric temperatures, or to recent volcanic eruptions (El Chichon, April 1982). These might have increased considerably the amount of $SO_2$ in the stratosphere and consequently the resulting formation of $H_2SO_4$ and $HSO_y$.

It is clear that more measurements at different stratospheric temperatures are required to elucidate this problem.

CONCLUSIVE REMARKS

A comparison of sulfuric acid vapour pressure calculations with the data set on $H_2SO_4 + HSO_y$ concentrations between 28 and 35 km altitude seems to support the present ideas about the interaction of $H_2SO_4$ with aerosols.

It should be noted however that this interpretation is relying on information, still suffering from some inaccuracies due to different experimental factors which need further investigations. Among those we cite the errors on the sulfuric acid vapour pressure calculations due to the uncertain values of the thermodynamic quantities involved.

As is noticed from figure 2 small changes in temperature and water mixing ratios have a considerable effect on the sulfuric acid partial pressure. A precise measurement of both water mixing ratio and temperature (which is difficult) is therefore needed.
Furthermore the influence of other substances, such as impurities or possibly HNO\textsubscript{3} dissolved in aerosols (Kiang and Hamill, 1974) should be investigated.

Also parameters such as total ion number density, ion-ion recombination coefficient and appropriate ion molecule reaction rate coefficients used to derive trace gas concentrations from ion mass spectra need to be determined with greater accuracy.

Below 28 km the interpretation of the data is hampered by the fact that the present technique cannot distinguish between H\textsubscript{2}SO\textsubscript{4} and other sulfur compounds which may lead to HSO\textsuperscript{4}\textsuperscript{-} cluster ions. Laboratory measurements of the ion-molecule reactions of HSO\textsubscript{3}, HSO\textsubscript{5} and other sulfur compounds with NO\textsuperscript{-} cluster ions may help to assess their role. Furthermore model calculations of these sulfur compounds would be very helpful.

In order to complete our understanding of the problem more in-situ measurements at different altitudes and at different stratospheric temperatures should be performed.

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