Origin of condensation nuclei in the springtime polar stratosphere

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Abstract. An enhanced sulfate aerosol layer has been observed near 25 km accompanying springtime ozone depletion in the Antarctic stratosphere. We use a one-dimensional aerosol model that includes photochemistry, particle nucleation, condensation growth, coagulation, and sedimentation to study the origin of the layer. Annual cycles of sunlight, temperature, and ozone are incorporated into the model. Our results indicate that binary homogeneous nucleation leads to the formation of very small droplets of sulfuric acid and water under conditions of low temperature and production of H$_2$SO$_4$ following polar sunrise. Photodissociation of carbonyl sulfide (OCS) alone, however, cannot provide sufficient SO$_2$ to create the observed condensation nuclei (CN) layer. When subsidence of SO$_2$ from very high altitudes in the polar night vortex is incorporated into the model, the CN layer is reasonably reproduced. The model predictions, based on the subsidence in polar vortex, agree with in situ measurements of particle concentration, vertical distribution, and persistence during polar spring.

Introduction

Balloon-borne counters have been able to measure aerosols with radii greater than 0.01 $\mu$m in the stratosphere since 1973 [Hofmann and Rosen, 1977; Rosen and Hofmann, 1977]. These fine particles are generally referred to as condensation nuclei (CN) because of their small size and their role as sites for condensation of sulfuric acid vapor [Hofmann and Rosen, 1985; Hofmann, 1990]. Measurements over Antarctica have revealed an enhanced CN layer between 20 and 30 km with a maximum particle concentration exceeding 100 cm$^{-3}$ at about 25 km [Hofmann and Rosen, 1985; Hofmann, 1988; Wilson et al., 1989]. The layer has been observed to appear repeatedly in each spring from September to November. Hofmann et al. [1989] found that the CN layer and the severe ozone depletion (ozone hole) in the south polar region are concurrent, and both CN concentrations and ozone abundance are sensitive to local temperature. Moreover, coincident with the CN layer and the ozone hole is an observed decrease in concentration of "optical-active aerosols" (OA) with particle radii greater than 0.15 $\mu$m. Recent measurements in the Arctic during winter show a similar layering effect [Hofmann, 1990]. Stratospheric CN, which are most likely liquid sulfuric acid and water solutions, are believed to serve as sites for heterogeneous chemical reactions. Thus it is of great interest to investigate the mechanisms which lead to the formation of these CN layers.

Several mechanisms for the formation of the CN layer have been proposed. Rosen and Hofmann [1983] suggested that sulfur-bearing gases (carbonyl sulfide (OCS)) could accumulate in the polar stratosphere during winter due to the lack of sunlight and produce high concentrations of H$_2$SO$_4$ vapor in spring after sunrise. However, Oppenheimer [1987] argued that photodecomposition of OCS is a slow process because of the large zenith angle in the polar region; the production of H$_2$SO$_4$ through OCS photolysis is not sufficient for CN formation. Thus he proposed that photooxidation of OCS or reaction of OCS with elevated levels of OH might account for an enhancement in sulfuric acid vapor leading to the formation of the CN layer.

Because CN layers are always observed to be accompanied by low temperatures, Hofmann [1988] suggested that low temperatures in the winter polar stratosphere support homogeneous nucleation and rapid growth of fine particles due to excessive supersaturation of H$_2$SO$_4$ vapor. Hamill et al. [1990] studied the homogeneous nucleation of ambient sulfuric acid at polar winter low temperatures and concluded that even with modest acid concentrations the formation of H$_2$SO$_4$/H$_2$O droplets would be efficient. However, such acid concentrations may not exist in polar winter because production of H$_2$SO$_4$ through photochemical processes ceases after polar sunset, while the acid vapor still condenses to preexisting particles as temperatures are decreasing. Hamill et al. did not include the loss of H$_2$SO$_4$ to preexisting particles. Their model did not reproduce the observed sea-
Table 1. Photochemical Reactions of Sulfur Components in the Stratosphere

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R_1$ $S + O_2 + \text{hv} \rightarrow \text{SO} + O$</td>
<td>$k_1 = 2.3 \times 10^{-12}$</td>
</tr>
<tr>
<td>$R_2$ $\text{SO} + O_2 + \text{hv} \rightarrow \text{SO}_2 + O$</td>
<td>$k_2 = 2.6 \times 10^{-13} \ e^{-2400/T}$</td>
</tr>
<tr>
<td>$R_3$ $\text{SO}_2 + \text{OH} + \text{M} \rightarrow \text{HSO}_3 + \text{M}$</td>
<td>$k_3 = 3.0 \times 10^{-31}, n = 3.3^*$</td>
</tr>
<tr>
<td>$R_4$ $\text{HSO}_3 + \text{O}_2 \rightarrow \text{SO}_3 + \text{HO}_2$</td>
<td>$k_4 = 1.5 \times 10^{-12}, m = 0^*$</td>
</tr>
<tr>
<td>$R_5$ $\text{SO}_3 + \text{H}_2\text{O} \rightarrow \text{H}_2\text{SO}_4$</td>
<td>$k_5 = 1.3 \times 10^{-12} \ e^{-330/T}$</td>
</tr>
<tr>
<td>$R_6$ $\text{OCS} + \text{O} \rightarrow \text{SO} + \text{CO}$</td>
<td>$k_6 = 2.1 \times 10^{-11} \ e^{-2200/T}$</td>
</tr>
<tr>
<td>$J_1$ $\text{OCS} + \text{hv} \rightarrow \text{S} + \text{CO} (\lambda &lt; 228 \text{ nm})$</td>
<td></td>
</tr>
<tr>
<td>$J_2$ $\text{SO}_2 + \text{hv} \rightarrow \text{SO} + \text{O} (\lambda &lt; 220 \text{ nm})$</td>
<td></td>
</tr>
</tbody>
</table>

*Three-body reactions with $K_1 = (K_0[M]/[1 + K_0[M]/K_0]) \times 0.6^{1.5 \log_{10}(K_0[M]/K_0)^{1/2}}$, where $K_0 = K_0^{300}(T/300)^{-m}$ and $K_3 = K_3^{300}(T/300)^{-n}$.  
$^*$This reaction may require a surface or a small ion cluster to balance energy and momentum.  
Nevertheless, the reaction is apparently very rapid under atmospheric conditions.

seasonal nor altitudinal characteristics of the CN layers. Rosen and Hofmann [1983] proposed that evaporation of sulfate aerosol in a stratospheric warming event followed by sufficiently rapid cooling could be responsible for the CN formation. However, the evaporation of aerosols may not release sufficient acid vapor for CN formation because the concentrations of preexisting aerosol near 30 km are very low.

Measurements of long-lived tracers suggest organized downward motion within the polar night vortex [Toon et al., 1989; Russell et al., 1993]. For example, data from the Halogen Occultation Experiment (HALOE) instrument on the Upper Atmosphere Research Satellite (UARS) show that methane (CH₄) mixing ratios in polar stratosphere are as low as those in the mesosphere. Moreover, the data suggest that air with lower CH₄ mixing ratios in the polar vortex does not mix with the stratospheric air surrounding the vortex until the vortex breaks down in spring [Russell et al., 1993]. It can be inferred that the air at the top of the springtime polar stratosphere originates from the mesosphere by descending motion in the polar vortex associated with radiative cooling during the winter. SO₂ may achieve a high mixing ratio in the mesosphere through photolysis of H₂SO₄. Although accurate measurements of the H₂SO₄ ultraviolet absorption spectrum are not available, photodissociation at wavelength below ~250 nm is likely. SO₂ generated in the mesosphere may then be transported into the stratospheric polar vortex by subsidence. There, SO₂ has a relatively long lifetime because of the absence of OH during polar winter. In this situation the mesosphere can be considered as a sulfur source for CN formation in spring after sunrise. Hofmann et al. [1988] also suggested that a possible source of sulfuric acid may be transport from sunlight regions in lower latitude with entrance into the vortex above 30 km. Then subsidence and radiative cooling in the lower stratosphere may result in particle formation.

Dust particles and micrometeorites from the ablation of micrometeors in the upper atmosphere may contribute to the CN concentrations above 20 km [Hunten et al., 1980; Turco et al., 1981]. These particles can affect the properties of stratospheric aerosols by serving as condensation nuclei for sulfuric acid droplets, enhancing the sulfate aerosol concentrations and reducing average particle sizes when sulfuric acid becomes supersaturated.

Polar stratospheric clouds (PSCs) may also modulate CN populations. PSCs can scavenge preexisting sulfur particles and thus decrease the total aerosol surface area in spring. The decreased aerosol surface area limits H₂SO₄ vapor loss onto the preexisting particle surfaces and maintains high H₂SO₄ vapor concentrations for homogeneous nucleation and CN formation.

In this study we elaborate on these various hypotheses by conducting a series of numerical experiments that include in a consistent way chemistry, aerosol microphysics, annual variations of temperature and sunlight, micrometeorite fluxes, and polar vortex dynamics.

Model Description

To simulate the behavior of CN in the springtime polar stratosphere, we incorporated sulfur photochemistry and microphysics in a one-dimensional version of the tracer model developed by Toon et al. [1988]. Table 1 gives the sulfur photochemical reactions which are important in the lower stratosphere. Under volcanically quiescent stratospheric conditions, the sulfur oxides originate primarily from OCS transported through the tropopause, as first suggested by Crutzen [1976]. After reaching the stratosphere, photodecomposition of OCS ($J_1$) releases sulfur atoms (S), which are quickly converted into sulfur dioxide (SO₂) through $R_1$ and $R_2$. The hydroxyl (OH) radical is critical in oxidizing SO₂ to sulfuric acid (H₂SO₄) through reactions $R_3$ to $R_5$. Because $R_4$ and $R_5$ are relatively fast, sulfuric acid production is limited mainly by reactions $J_1$ and $R_3$. In this study, the baseline profiles of OH, O, and H₂O are assumed to be fixed with concentrations taken from the photochemical model of Brasseur and Solomon [1986]. All the chemical reaction rates correspond to standard compilations [DeMore et al., 1990].

Three types of aerosols are considered in this model: involatile (or solid) particles which originate from the troposphere (we also refer these "tropospheric" CN as Aitken particles), pure H₂SO₄/H₂O particles, and mixed aerosols...
consisting of Aitken particles in H₂SO₄/H₂O solutions. Aitken particles are transported from the troposphere into the lower stratosphere and provide the core for mixed aerosols either through heterogeneous nucleation or coagulation between Aitken particles and pure H₂SO₄-H₂O particles. Mixed aerosols interact with H₂SO₄ vapor by condensation and evaporation processes. If a mixed aerosol is completely evaporated, then its core is released and returned to the "Aitken" category. Pure H₂SO₄/H₂O aerosols can be produced only by homogeneous nucleation, when the supersaturation of H₂SO₄/H₂O vapor exceeds a critical value. An efficient algorithm to calculate homogeneous and heterogeneous nucleation rates [Zhao, 1993] is used here. Details concerning the other microphysical processes and corresponding numerical algorithms can be found in the works of Turco et al. [1979], Toon et al. [1988], and Zhao [1993]. In the calculations that follow, each type of particle (Aitken nuclei, pure, and mixed particles) is subdivided by size. The smallest size treated in the model is about 0.001 μm. The total number of particles in our simulations corresponds to the sum over the whole size bins. The observed CN correspond to the total number of particles with sizes greater than about 0.01 μm. To compare with observations, the concentration of CN in our simulations is by summing all the appropriate size bins in the model.

**Simulation Parameters**

**Benchmark Simulation**

There are two purposes for investigating the background aerosols before simulating transient CN layers. First, we need to verify the capability of the model to generate realistic vertical concentration profiles of the background aerosols and key chemical species. Second, the polar stratospheric model is initialized using the background aerosol profiles, which approximate the state of the aerosol layer in summer. To achieve a well-characterized background state, we first integrate the model to a steady state using 10-year mean temperatures between 70° and 80°S and a monthly mean ozone profile. The tropopause was set at 9 km and was kept constant through all of the numerical experiments described here. The baseline vertical profiles of SO₂ (Figure 1) and aerosol concentration (Figure 2) are reasonable comparing with observations. The computed size distributions (Figure 3) are similar to those calculated by Turco et al. [1979]. The simulated CN concentration is about 5 cm⁻³ at 20 km and less than 1 cm⁻³ at 30 km; these values are close to the annual average aerosol concentrations measured near the south pole at the same heights.

**Annual Cycle of Temperature**

The observed high correlation between CN concentrations and low stratospheric temperatures suggests the importance of temperature variation in forcing the anomalous CN. In our "polar" simulations the annual cycle of temperature is deduced from observed zonal mean temperatures between 70° and 80°S for 1984 (Figure 4). These data present a typical picture of temperature variations in the southern polar region; similar temperatures hold in other years. In general, temperatures decrease from fall through winter and recover in the spring. The lowest temperature in Figure 4 is less than 190°K, near 18 km in later winter and early spring. An approximation to the period of twilight in the polar region is superimposed on this temperature field. Near latitude of 75°S, sunset occurs in early May and sunrise in late August. Figure 4 shows clearly that in spring after sunrise, stratospheric temperatures can remain low for a period of several weeks. This cycle of temperature and sunlight implies that the photochemical production of H₂SO₄ vapor recovers after polar sunrise, while the equilibrium vapor pressure of H₂SO₄ under low-stratospheric temperature is still very small. Therefore the acid vapor concentrations can easily
photodecomposition of OCS has been suggested as one cause of the CN layer [Rosen and Hofmann, 1983]. OCS photolysis is controlled by solar ultraviolet radiation, which is particularly sensitive to ozone abundances. Therefore to calculate the photodissociation rate of OCS in the southern polar region more accurately, monthly average ozone profiles derived from balloon soundings at Syowa (69°S) in 1990 were adopted. Stratospheric ozone concentrations decline in September, reach minimum values in October, and gradually recover in November (Figure 5). These spring ozone "hole" profiles have reduced ozone abundances that favor the photodecomposition of OCS.

**Baseline CN Simulation**

Using the annual cycles of zonal mean temperature, solar illumination, and ozone concentrations derived from measurements in the southern polar region and the benchmark stratosphere as an initial state, a "baseline" simulation of CN formation was performed to investigate CN layer formation. This baseline model represents the best first-guess parameterization for stratospheric CN formation, based on existing observations and theories. In the baseline run, integration begins at day 80, when the initial state is very close to the summer climatology and runs to day 365, when the solution essentially returns to the background state.

The simulated \( \text{H}_2\text{SO}_4 \) mixing ratio drops dramatically as polar sunset advances at around day 120 and remains near zero until sunrise the following spring (Figure 6). The loss of \( \text{H}_2\text{SO}_4 \) vapor occurs because the normal balance between \( \text{H}_2\text{SO}_4 \) vapor production and condensation on preexisting aerosols cannot be maintained when solar illumination vanishes, and the OH radicals that generate \( \text{H}_2\text{SO}_4 \) disappear. In addition, decreasing temperatures favor \( \text{H}_2\text{SO}_4 \) loss onto the preexisting droplets. Below 35 km, \( \text{H}_2\text{SO}_4 \) vapor reappears rapidly after sunrise on day 230; and photochemical reactions driven by sunlight reestablish the production of \( \text{H}_2\text{SO}_4 \). Subsequently, the \( \text{H}_2\text{SO}_4 \) mixing ratio fluctuates in accord with temperature changes. Above about 40 km the \( \text{H}_2\text{SO}_4 \) reduction in midwinter does not occur because of higher temperatures there that prevent condensation. Figure 7 provides several snapshots of the evolution of the vertical distribution of \( \text{H}_2\text{SO}_4 \) concentrations. During the winter period the evaporation level for sulfuric acid (i.e., where sulfuric acid vapor becomes undersaturated and evaporation can occur) rises from 37 to 45 km owing to the wintertime trend of temperature decrease in this region.

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**Figure 3.** Calculated particle size distributions of the background aerosols at altitudes of 17, 21, 25, and 29 km. These size distributions compare favorably with measurements summarized by Turco et al. [1982].

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**Figure 4.** Isotherms of observed zonal belt-mean temperatures between 70° and 80°S latitudes for 1984. The data are derived from the National Meteorological Center analysis. The shaded area indicates the period of darkness during polar winter at a latitude of 75°S.

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**Figure 5.** Monthly mean profiles of observed ozone concentrations with unit of \( 10^{12} \text{ cm}^{-3} \) at Syowa Station (69°S, 40°E) in 1990 [Nankyoku and Kansokutai, 1990].
Along with the rapid recovery of H$_2$SO$_4$ just after polar sunrise, temperatures in the lower stratosphere are near their minimum values (Figure 4). This creates an ideal situation for particle formation by nucleation, which depends on H$_2$SO$_4$ concentration and temperature. Indeed, the simulated total aerosol concentrations (all particles larger than 0.001 $\mu$m) between the tropopause and 35 km increase dramatically due to the new particle formation by homogeneous nucleation shortly after sunrise (Figure 8). The aerosol concentrations produced by homogeneous nucleation can initially exceed 10$^3$ cm$^{-3}$ near 25 km. These newly formed sulfate embryos, however, cannot grow to measurable “CN” sizes. Only a few particles near 30 km are found reaching measurable sizes (greater than 0.01 $\mu$m) and the springtime perturbation of CN concentration is hardly noticeable (Figure 9). The simulated baseline CN profiles are not comparable to the observed CN layer (Figure 10). Moreover, the aerosol size distributions in the baseline experiment show that although a large number of fine H$_2$SO$_4$/H$_2$O particles are initially produced by homogeneous nucleation right after sunrise, they do not grow to CN sizes because of a shortage of H$_2$SO$_4$ vapor and quick coagulation with ambient particles (Figure 11).

The baseline experiment fails to produce a substantial CN layer. The photodissociation rate of OCS is too low to produce much H$_2$SO$_4$ vapor in early spring ($<10^{-9}$ s$^{-1}$) owing to the weak solar illumination. Hence the conversion of ambient OCS into H$_2$SO$_4$ cannot be the dominant mechanism in the formation of the CN layer in polar spring. This is true even if the OCS concentration is allowed to accumulate through vertical diffusion over the winter. In fact, the polar vortex is highly stable against upward vertical mixing, and the vertical profiles of most long-lived tracers suggest sustained subsidence in the winter polar vortex, which should reduce OCS concentrations at 25-30 km. It follows from the baseline simulation that certain mechanisms not treated in the baseline experiment must play an important role in providing extra sulfur for the formation of new CN.
Role of Subsiding Motion in the Polar Vortex

A Lagrangian visualization using a three-dimensional model simulation provides a detailed picture of the airflow in the middle atmosphere [Fisher et al., 1993]. Air from the mesosphere executes an inward and downward spiral into the polar vortex. The numerical simulation offers an explanation of HALOE observations of low mixing ratios of methane in the polar stratosphere [Russell et al., 1993]. The subsiding motion in the polar vortex transports mesospheric air with low mixing ratios of methane down into the stratosphere. Similarly, the subsiding motion would transport a higher mixing ratio of sulfur dioxide into the polar stratosphere from the mesosphere. A high mixing ratio of SO₂ in the mesosphere is likely from the photolysis of H₂SO₄ at those altitudes [Turco et al., 1979]. The SO₂ transported downward would not be destroyed in the polar night because of the absence of OH radicals, hence the descending air can provide a critical sulfur reservoir for CN formation in the spring. The weak CN layer in the baseline experiment is thus compensated by a mesospheric sulfur source.

There are no quantitative observations of the amplitude, structure, or duration of the descending motion in the polar vortex, but three-dimensional model simulations [Fisher et al., 1993] provide useful guidelines. The model results show that the mean poleward velocity in the mesosphere is about 40° per month, whereas the subsiding velocity in the upper stratosphere of the polar vortex is about 12 km per month in July. The descending motion decreases to about 3 km per month in the middle stratosphere and the poleward drift is also much slower than in the mesosphere. The vortex circulation can maintain this structure for up to 6 months.

On the basis of these model results we postulate a subsiding velocity field as

$$w = w_0 \left( \frac{\rho}{\rho_T} \right)^\alpha \cos \left( \frac{\pi}{180} \left( t - 220 \right) \right), \quad 120 < t < 310. \quad (1)$$

Here $t$ is time (day), $\rho_T$ is the air density at the top layer of the model, $w_0 = 0.5$ cm/s, and $\alpha = 0.4$. This idealized formulation of vertical descent qualitatively agrees with the results of the three-dimensional simulation.

The effect of the corresponding poleward airflow is treated consistently by including an additional term in the one-dimensional model to ensure mass continuity. First, note that the horizontal advection is related to the vertical velocity divergence through the mass continuity equation. Moreover, the continuity equation can be written for any tracer in the form

$$\frac{\partial C}{\partial t} + \nabla \cdot (\vec{v} \rho C) + \frac{\partial (w C)}{\partial z} = \nabla \cdot \rho \vec{K} \nabla \left( \frac{C}{\rho} \right) + P - L, \quad (2)$$

Figure 10. (a) and (b) Vertical distributions of CN concentrations at different times in the baseline model simulation, compared to observations [Hofmann and Rosen, 1985].

Figure 11. Aerosol size distributions at different times in the baseline model simulation.
If zonal symmetry is assumed, the horizontal advection term can be approximated by differencing from the pole over the radius $l$ of the vortex as

$$\nabla_H \cdot (\vec{v}_H C) = -\frac{\mu_H C_a}{l}. \tag{3}$$

Here the subscript $a$ indicates values in ambient air outside the vortex at the same altitude, and $\mu_H$ is the average horizontal poleward velocity. Similarly, the mass continuity equation can be written as

$$-\frac{\mu_H}{l} \frac{\partial (w \rho)}{\partial z} = 0. \tag{4}$$

Then, in a one-dimensional tracer representation we have

$$\frac{\partial C}{\partial t} + \frac{\partial (wC)}{\partial z} = C_a \frac{\partial (w \rho)}{\partial z} + \frac{\partial}{\partial z} \rho K \frac{\partial}{\partial z} \left( \frac{C}{\rho} \right) + P - L. \tag{5}$$

The first term on the right-hand side of (5) can be viewed as the effect of horizontal mass convergence associated with net vertical subsidence. Whereas the vertical velocity used here increases with altitude, the downward mass flux actually decreases with altitude, and the horizontal convergence effectively brings air into the vortex from outside. Both the vertical motion and the horizontal convergence decrease in the lower stratosphere, and there is little accumulation of mass in the stratospheric column.

As a variation of a baseline experiment, a SO$_2$ mixing ratio of 0.1 parts per billion by volume (ppbv) is assumed at the upper boundary (60 km). Similarly, the mixing ratio of H$_2$SO$_4$ in air descending from mesosphere is set to 0.1 ppbv, and the concentrations of preexisting aerosols are set to zero. These concentrations for SO$_2$ and H$_2$SO$_4$ are taken from photochemical simulations of the stratospheric sulfur cycle [e.g., Turco et al., 1979, 1982]; however, the actual concentrations are uncertain. As will be shown, the results are not sensitive to the H$_2$SO$_4$ concentration but are sensitive to SO$_2$.

![Figure 12](image-url)

**Figure 12.** Vertical distributions of CN concentrations at different times in the experiment including a subsiding motion from the mesosphere. Compare with data shown in Figure 10.

![Figure 13](image-url)

**Figure 13.** The vertical distribution of large aerosol particles with $r > 0.15 \mu m$, versus time in the simulation with subsidence.

After including such a descending motion and associated fluxes at the upper boundary, the model generates dense CN layers during spring after sunrise that agree with observational data (Figure 12). The layer appears at higher altitudes in early spring and shifts downward as time progresses. The maximum concentration of CN particles is situated near 29 km in September and 24 km in October. The background particles in this case also have lower concentrations during winter and spring because of the downward flow, which brings particle-free air into the upper and middle stratosphere. The number of large particles is reduced significantly (Figure 13). Hence subsidence is also consistent with the aerosol hole observed by balloon-borne and satellite soundings [Hofmann and Rosen, 1983; McCormick et al., 1983].

The vertical flux of SO$_2$ at upper boundary is crucial to forming the CN layer. During fall the photolysis of OCS decreases and the SO$_2$ mixing ratio above 20 km decreases through reaction with OH. The photochemical source of SO$_2$ remains insignificant throughout the winter. Moreover, the production of SO$_2$ from OCS is too slow to generate a CN layer in the springtime. Once the downward transport of SO$_2$ from the mesosphere into the stratospheric vortex is included, however, the SO$_2$ mixing ratio can be maintained at an elevated level, as shown in Figure 14.

Variations in the H$_2$SO$_4$ mixing ratios for the case with subsidence are similar to those in Figure 6, except the mixing ratio of H$_2$SO$_4$ has a higher value near 30 km in winter and early spring due to the subsidence motion; however, the total aerosol and CN concentrations (Figures 15 and 16, respectively) are considerably different from those in Figures 8 and 9. The perturbations in the aerosol concentrations following sunrise are far more intense and persistent. Note particularly the strong enhancement in CN concentrations from 18 to 30 km during springtime in Figure 16. By comparing Figures 15 and 16, it becomes evident that the enhancement in CN concentrations occurs mainly above 18 km, although many small droplets are generated below this level (these very small particles are not counted as CN). At the lower altitudes the ambient aerosols capture all of the small droplets by coagulation before they grow to CN sizes by condensation. The vertical distribution of the CN con-
Figure 15. Variation of total aerosol concentration (number per cubic centimeter) with time and altitude in the experiment with a subsiding motion. The concentration contours are shown with the logarithm of the concentrations. The contour interval is 0.5 log units.

Figure 16. Variation of CN concentration (number per cubic centimeter) with time and altitude in the experiment with a subsiding motion. The CN concentrations are shown as isopleths of constant concentration to logarithm. The contour interval is 0.5 log units.

Figure 17. Aerosol size distributions at different times in the experiment with a subsiding motion.

Concentration near the evaporation level at 35 km (Figure 15) is quite sensitive to local temperature variation. A clear correlation between fluctuations in the temperature and simulated CN is seen during springtime.

The simulated aerosol size distributions for the case with subsidence reveal the detailed evolution of these aerosols (Figure 17). The aerosol size distribution in winter is similar to the ambient size distribution, as might be expected, although the size spectrum shifts to smaller-sized concentrations as some of the bigger particles fall out. Ten days after sunrise, however, a large number of very small $\text{H}_2\text{SO}_4/\text{H}_2\text{O}$ particles are generated by homogeneous nucleation, and the size distribution is bimodal. Unlike the baseline case, the aerosol size spectra for days 50 and 80 (after sunrise) indicate that the small particles have grown to larger sizes, and the typical particle radius on day 80 is at 0.01 $\mu$m.

Figure 18 shows the complete evolution of the CN distribution from sunset through the end of the year at altitudes from 10 to 60 km. Clearly, a significant enhancement in the CN concentration occurs during spring (September, October, and November) after sunrise. The peak in the CN layer moves from the highest altitude early in spring down to 23 km in October. The maximum concentration of CN particles reaches 40 cm$^{-3}$, with the main layer located between 18 and 37 km. Below 15 km, CN-sized particles are lost by coagulating with the larger number of ambient aerosols. Above 37 km, higher temperatures cause the volatile CN droplets to evaporate. The simulated CN layer is consistent with measurements of the intensity, altitude range, and time of occurrence.

In the simulations, subsiding air carries higher mixing ratios of $\text{SO}_2$ from the mesosphere into the stratospheric polar vortex thereby creating a CN layer, which seems to provide a reasonable explanation for CN layer formation and the springtime aerosol hole. The case with subsidence also reproduces the observed major features and evolution of the polar CN layer. We conducted a series of sensitivity experiments with different subsidence velocities and mesospheric $\text{SO}_2$ mixing ratios. The intensity of the CN layer increases with higher $\text{SO}_2$ mixing ratios and rates of descending air. It
Modified Photochemistry

To test the importance of oxidation of OCS by OH radicals originally proposed by Oppenheimer [1987] and other related chemical processes, the following photochemical reactions involving OCS were considered.

\[
\begin{align*}
(J_1) & \quad \text{OCS} + h\nu \rightarrow \text{S} + \text{CO} \\
(R_6) & \quad \text{O} + \text{OCS} \rightarrow \text{SO} + \text{CO} \\
(R_7) & \quad \text{OH} + \text{OCS} \rightarrow \text{SH} + \text{CO}_2 \\
(R_8) & \quad \text{ClO} + \text{OCS} \rightarrow \text{products} \\
(R_9) & \quad \text{NO}_3 + \text{OCS} \rightarrow \text{products} \\
(R_{10}) & \quad \text{Cl} + \text{OCS} \rightarrow \text{SCI} + \text{CO} \\
\end{align*}
\]

The first two reactions are included in the baseline model. As we noted earlier, the photolysis rate of OCS \((J_1)\) is too small in polar spring to provide an adequate sulfur source for the observed CN layer. An OCS photodecomposition rate, \(I_{\text{OCS}}\), of roughly \(10^{-7}\) s\(^{-1}\) is necessary to create a realistic CN layer. At 25 km, the reaction rate with atom oxygen, \(I_6 = R_6[O]\) is about \(10^{-8}\) [Brasseur and Solomon, 1986]. This is obviously too slow. The reaction of OH with OCS \((R_7)\) suggested by Oppenheimer [1987], with an upper limit concentration for OH of \(10^{-7}\) cm\(^{-3}\), is again too small \((I_{\text{OH}} = 10^{-8}\) s\(^{-1}\)\). Measurements show that ClO mixing ratios in the lower polar stratosphere during spring can exceed 1 ppbv, more than 100 times greater than normal background stratospheric concentrations [de Zafra et al., 1987; Solomon et al., 1987]. Thus the OCS lifetime against decomposition by reaction \(R_9\) could become as large as about \(10^{-7}\) s\(^{-1}\), which is an order of magnitude greater than reaction \(I_6\). This chemical reaction may contribute to the conversion of OCS to \(\text{H}_2\text{SO}_4\) vapor. \(R_9\) and \(R_{10}\) would be more than an order of magnitude less important than \(R_8\), because the concentrations of \(\text{NO}_3\) and Cl are so low.

A ClO concentration field is constructed according to observational data from de Zafra et al. [1987] and Solomon et al. [1987] as

\[
[\text{ClO}] = c(z) \cos \left(\frac{\pi (t - 240)}{60}\right), \quad 230 < t < 270 \quad (7)
\]

where

\[
c(z) = \begin{cases} 
\exp \left(\frac{-(z - 20)^4}{2}\right), & z < 20 \text{ km} \\
\exp \left(\frac{-(z - 20)^2}{2}\right), & z > 20 \text{ km} 
\end{cases} \quad (8)
\]

Here \(t\) is time (day) and \([\text{ClO}]\) is ClO mixing ratio (ppbv).

When we incorporate reaction \(R_9\) in the baseline model with the ClO concentrations shown in Figure 19 and the upper limit (room temperature) rate coefficient of \(10^{-16}\) cm\(^3\) s\(^{-1}\), CN layers are formed between 15 and 30 km with peak concentrations of more than 100 cm\(^{-3}\) (Figure 20). This surplus of CN reflects particle formation, condensation, and coagulation in the presence of abundant \(\text{H}_2\text{SO}_4\) vapor generated by the decomposition of OCS.

Among the photochemical processes listed above, reaction \(R_9\) is the most likely to provide sufficient \(\text{H}_2\text{SO}_4\) vapor for CN formation in polar spring. The reaction rate coefficient, however, is probably much smaller than the room temperature upper limit [DeMore et al., 1990] used in the simulation. At much lower stratospheric temperatures than the room temperature the rate coefficient may be 1 to 2 orders of magnitude smaller. Further experimental studies of this mechanism are needed to eliminate this possibility.

Effect of Micrometeorites

To study the impact on CN layer formation of an additional particle source associated with micrometeorites, we also did some experiments which include a constant flux of small particles at the upper boundary according to the results from Hunten et al. [1980] and Turco et al. [1980]. These fine meteoric aerosols slowly migrate downward by diffusion in
Figure 19. (a)-(c) CIO mixing ratios used in the numerical experiment with reaction \( R_8 \) set to its upper limit value. The enhanced CIO concentrations were constructed using equation (7) referring to the measurements from de Zafra et al. [1992].

The one-dimensional simulation. Figure 21 shows the steady state aerosol number size distributions obtained by integrating the model 300 days under conditions corresponding to the background state with a meteoric aerosol source at the upper boundary. Even though there are a larger number of meteoric dust particles throughout the upper stratosphere, these do not influence CN concentrations. The aerosol size spectra (Figure 21) show a micrometeorite mode below a 0.01-\( \mu \)m radius, with a typical radius of 0.003 \( \mu \)m. The initial size of the meteoric dust particles, although quite uncertain, are very unlikely to be larger than 0.01 \( \mu \)m [Hunten et al., 1980]. Accordingly, micrometeoritic particles themselves are unlikely to change the CN (\( r > 0.01 \) \( \mu \)m) concentration, or to form a distinct CN layer.

The micrometeoritic particles were incorporated in the CN simulation as an initial distribution (Figure 21) in equilibrium with a fixed flux at the upper boundary. Other conditions were similar to those in the baseline experiment. The resultant CN concentrations, including the vertical distributions and temporal evolution, are almost the same as in the baseline model. When micrometeorites are coupled with either subsiding motion or reaction \( R_8 \) in the model, minor variations in CN concentrations occurred when heterogeneous nucleation was assumed to have a contact angle of 50° (moderate nuclei). The overall features of the CN layers remained unchanged. Clearly, the influence of micrometeorites is a secondary factor in CN layer formation.

We also considered an extreme case by assuming (1) all the micrometeorites serve as very efficient condensation nuclei for sulfate aerosols and are nucleated in a very short time once \( \text{H}_2\text{SO}_4 \) vapor is supersaturated and (2) homogeneous nucleation is switched off so that no new particles are
generated. When we incorporated conditions 1 and 2 into the model with otherwise the same conditions as in baseline experiment, the model results are almost the same as those in baseline experiment. A CN layer still does not appear. Even if the micrometeorites are assumed to be excellent nuclei for sulfuric acid and water solutions, they did not enhance CN formation in this experiment.

When conditions 1 and 2 are combined with vortex subsidence and sulfur flux at the upper boundary of the model, a CN layer is produced similar to that shown in Figure 12. However, the scenario of CN formation is distinctly different from that in the similar case without meteorites. In the case illustrated in Figure 12 the new sulfuric acid/water droplets are produced by homogeneous nucleation. In contrast, in this case, it is the heterogeneous nucleation of sulfuric acid/water solutions on these preexisting micrometeorites that is the key mechanism to initialize the CN formation. As we can see in Figure 21, the concentrations of micrometeorites can be more than 100 cm\(^{-3}\) above 30 km. These micrometeorites are heterogeneously nucleated and grow to CN sizes (>0.01 \(\mu m\)) by condensation as they are transported to the lower stratosphere by subsidence.

According to the results from numerous numerical experiments we found that if sufficient concentrations of wettable micrometeorites exist, we can not rule out the possibility of CN layer formation through heterogeneous nucleation of sulfuric acid and water solutions on these meteorites. However, the presence of sufficient \(H_2SO_4\) vapor concentrations is the most crucial factor controlling the occurrence of CN events. With the subsidence supplying the extra sulfur from the mesosphere, a CN layer can be generated either through homogeneous or heterogeneous nucleation, or by both.

**Influence of Polar Stratospheric Clouds (PSCs)**

It has been proposed that the largest sulfate particles in the stratosphere may freeze and serve as condensation nuclei for PSCs. The removal of aerosols from the polar vortex may be related to PSC formation in winter and early spring when temperatures drop below about 193°-195°K (refer to the temperature variation in Figure 4). The larger PSC particles fall to lower altitudes more quickly than sulfate aerosols and thereby serve as a sink. With the largest sulfate aerosols removed, \(H_2SO_4\) vapor loss by condensation on particle surfaces is reduced, which may favor new particle formation by nucleation.

To test the influence of PSCs on CN formation, it was assumed that whenever the temperature falls below 195°K, large sulfate particles \((r > 2.5 \mu m)\) are scavenged by PSCs. The results suggest that the loss of aerosol surface area due to PSC sedimentation during winter is not a critical mechanism for CN formation. That is, the baseline experiment does not generate a CN layer even if this process is included; however, when it is combined with vortex subsidence or the CIO reaction, the predicted CN layers are slightly intensified.

**Summary and Discussion**

The major conclusions derived from numerous model experiments regarding polar CN layer formation are summarized in Table 2. We found that the variations in photochemical reactions corresponding to the conditions of polar twilight and low temperature control the first appearance of new particle formation. Specifically, OH radicals play an important role in producing sulfuric acid vapor from \(SO_2\), and low temperatures are crucial for homogeneous nucleation, which generates large numbers of small \(H_2SO_4/H_2O\) particles. However, if OCS photolysis is the only mechanism considered, the newly formed particles cannot grow to observable CN sizes because of the shortage of \(H_2SO_4\) vapor. Not only is the photodissociation rate of OCS too small under the weak solar illumination at large zenith angles in early spring
but also the OCS concentrations would be suppressed by subsidence in the polar vortex over the winter season.

Air descending from the mesosphere in the polar vortex carries higher SO\textsubscript{2} mixing ratios into the stratosphere and provides an extra sulfur source for the formation of a CN layer. In a one-dimensional model, realistic CN layers are produced when the vortex dynamical process is included. The major features of the CN layer observed by balloonborne instruments are accurately depicted in the “subsidence” experiment. For instance, the layer appears at higher altitudes in early austral spring and shifts downward to near 25 km by October. The layers are confined to altitudes between 18 and 37 km. Below 18 km, the ambient particles absorb sulfuric acid vapor fast enough to prevent new CN production. Above 37 km the local temperatures are high enough to prevent H\textsubscript{2}SO\textsubscript{4} vapor from nucleating. The subsidence experiment also explains the small concentrations of optically active aerosols during winter and spring resulting from the intrusion of aerosol-free air from high altitudes. Polar subsidence and photochemistry of SO\textsubscript{2} seems to be the most plausible explanation for CN layer formation in polar spring. Our calculations are based on the parameters such as temperature, subsidence, and solar illumination for the Antarctic stratosphere, but these conditions are similar over Arctic. We believe CN events in both hemispheres have similar formation mechanisms.

By analyzing all of the reactions involving stratospheric OCS that might initiate sulfate aerosol production in spring, we have found that none of these reactions are likely to provide sufficient H\textsubscript{2}SO\textsubscript{4} vapor for CN layer formation and persistence. The photochemical production of SO\textsubscript{2} by the reaction between OCS and ClO seems to be a secondary process at best. The OCS + ClO reaction, although quite uncertain in the absence of sufficient laboratory data, would, at its upper limit room temperature value, be barely adequate to generate a weak CN layer.

Other sensitivity tests were also conducted. We conclude, for example, that micrometeorites may cause minor variations in CN concentrations. Heterogeneous nucleation on micrometeorites could be important for CN formation if the micrometeorites are good condensation nuclei for sulfuric acid and water solutions and have sufficient concentrations in the stratosphere. Homogeneous nucleation in most cases appears to be necessary to explain the increase in CN concentrations. Indeed, the total sulfate mass in the CN layer implies such a large H\textsubscript{2}SO\textsubscript{4} vapor production rate that homogeneous nucleation at polar temperature seems unavoidable. Reductions in the background sulfate aerosol surface area caused by PSC condensation and sedimentation can help CN layer formation according to our model simulations. However, PSC effects cause minor variations in CN layer properties and cannot explain the appearance of the layer or its overall characteristics.

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References


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