PHYSICAL-CHEMICAL PROCESSES OF CLOUD ACTIVATION STUDIED WITH A DESKTOP CLOUD MODEL

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Abstract

A model of the activation process of a zero-dimensional cloud is described. The model runs in Mathematica®, a platform-independent environment for numerical computation. Key input parameters are updraft velocity and the dry distribution of number and composition of aerosol particles; 25 bins in the dry radius range 9 to 280 nm provide an accurate representation of microphysical dynamics. As the parcel is cooled water vapor is taken up by aerosol particles, governed by the difference between the ambient water vapor partial pressure and the equilibrium partial pressure above each drop, as given by the Köhler expression. As environmental supersaturation increases, initially larger particles, and then successively smaller particles may activate. The environmental supersaturation reaches a maximum and then decreases, effectively limiting the radius range of activated particles. This "desktop cloud" can be used to explore a variety of phenomena: the dependence of maximum supersaturation on environmental variables such as updraft velocity, aerosol properties and the presence of soluble and/or reactive gases; to examine the locus of uptake and aqueous-phase reaction of gases and the resultant influence on the particle size distribution; and to test parametrizations of aerosol influences on cloud microphysics.
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Work in Progress
Background

Clouds form when air containing water vapor and aerosol particles is cooled below the dew point. Water vapor condenses on aerosol particles.

The equilibrium vapor pressure of water $p(a)$ above a liquid water drop of radius $a$ is controlled by Raoult's law (vapor pressure lowering by solute) and the Kelvin equation (free energy of surface tension) and is strongly a function of drop radius, according to the Köhler expression (1923, 1936):

$$p(a) = p_w(T) \exp \left[ \frac{2M_w \sigma}{R_g T \rho a} - \frac{v m_s / M_s}{(\frac{4}{3} \pi a^3 \rho - m_s) / M_w} \right]$$

where

- $p_w(T)$ is the saturation vapor pressure of pure water at the ambient temperature
- $M_w, M_s$ are the molecular weights of water and solute
- $\sigma$ is the solution-air surface tension
- $R_g$ is the gas constant; $T$ is the temperature
- $\rho$ is the solution density
- $v$ is the van't Hoff factor (equiv / mol) including nonideality corrections
- $m_s$ is the solute mass
**Köhler curves** give saturation ratio vs. radius for fixed solute mass per particle:

![Diagram showing Köhler curves](image)

**Critical saturation** is maximum in curve; **critical radius** is corresponding radius value. Critical saturation decreases with increasing solute mass.
Cloud Droplet Activation

Activation of an aerosol particle to form a cloud droplet occurs when the environmental water vapor pressure exceeds Köhler maximum, with resultant condensation of water vapor, sufficiently long for the drop radius to exceed the corresponding critical radius.

- Once activated the drop grows under kinetic control.
- Activation is inherently kinetic, being controlled mainly by vapor-phase diffusion and heat transfer.

During cloud formation the environmental supersaturation in a given air parcel typically exhibits a short transient maximum triggered by initial adiabatic cooling and quenched by condensation of water vapor onto the newly available surface area of the activated cloud droplets, which serves as a runaway sink for water vapor.

The time history of the saturation ratio depends intrinsically on the interaction of water vapor with aerosol and gaseous solutes.
Importance

Cloud droplet activation determines the droplet number concentration and size distribution of the resultant cloud and the distribution of soluble gases taken up in the cloudwater.

The efficiency of activation of aerosol particles is of interest in cloud chemistry and microphysics.

- Major influence on *deposition* of aerosol materials in rain.
- An enhanced concentration of cloud droplets can *suppress precipitation*.
- The radiative properties of clouds: A greater concentration of aerosol particles generally results in a greater concentration of cloud droplets and *brighter clouds*.

These phenomena are important in considerations of *anthropogenic climate change*.

This Study. . .

Examines the dependence of cloud droplet activation on controlling processes by examining its *kinetics*.

Examines the time constants of droplet growth relative to duration of maximum supersaturation, gaseous diffusion, and the like.

Provides a differentiated picture of growth of size classes of drops and their uptake of water vapor and soluble gases.

Provides dependence of cloud droplet activation on controlling parameters that can be used to test approximations suitable for large scale atmospheric models.
Approach

**Zero-Dimensional box model** describing increase in water vapor saturation ratio and resultant mass-transfer processes pertinent to cloud formation. Cloud updraft velocity and initial aerosol populations are treated as independent variables. Updraft velocity is treated as constant in results reported here, but any time-dependent updraft velocity can be employed. Temperature decrease due to updraft velocity is manifested in the difference between ambient partial pressure of water and equilibrium partial pressure as given by the Clausius-Clapeyron equation. Coagulation and precipitation development are not represented in the model. Soluble gases (e.g. HNO₃) can be added and their fate and influence as solute can be determined. Kinetics of mass transport processes are examined by solution of coupled differential equations describing the radii of classes of aerosol particles using Mathematica®.

Key Independent Variables

- Number and size of aerosol particles
- Updraft velocity
- Presence or absence of soluble gas (HNO₃)
Thermodynamic Driving Force

*Temperature change* as a function of time $\Delta T(t)$ is given by *updraft velocity* $w$ and *lapse rate* $g / c_{p,a}$, taking into account *latent heat release* due to condensed water:

$$
\Delta T(t) = -\frac{g w t}{c_{p,a}} + \frac{C_w L_w}{\rho_a c_{p,a}}
$$

where $g$ is the acceleration of gravity

$c_{p,a}$ is the heat capacity of air at constant pressure

$C_w$ is liquid water content, kg m$^{-3}$

$L_w$ is the latent heat of condensation of water vapor, J kg$^{-1}$

$\rho_a$ is the density of air

Change in *equilibrium vapor pressure of water* (driving force for water condensation) is given by the Clausius-Calpeyron equation as

$$
\Delta p_w(t) = \exp\left[-\frac{L_w M_w \Delta T(t)}{R_g T^2}\right]
$$
Droplet Growth Equations

The rate of accretion of water mass is governed by molecular diffusion:

\[
\frac{dm_{w,i}}{dt} = M_w (4\pi a_i^2) \cdot \frac{D_{\text{eff},w,i}}{a_i} \cdot \left[ p_w(\infty) - p_w(a_i) \right] / R_g T
\]

where

- \( M_w \) is the molecular weight of water
- \( m_{w,i} \) is the mass of water in a single drop of class \( i \)
- \( a_i \) is the radius of drop of class \( i \)
- \( p_w(a_i) \), \( p_w(\infty) \) are the gas-phase partial pressure of water at the drop surface and in the bulk, respectively, and
- \( D_{\text{eff},w,i} \) is the effective diffusion coefficient of water vapor for drop of class \( i \)

The corresponding radial growth equation is

\[
\frac{da_i}{dt} = \frac{1}{(4\pi a_i^2) \rho} \frac{dm_{w,i}}{dt} = \frac{1}{a_i \rho} M_w \frac{D_{\text{eff},w,i}}{a_i} \cdot \left[ p_w(\infty) - p_w(a_i) \right] / R_g T
\]

Radial growth rate is inversely proportional to the radius. Small drops “catch up” with large drops, narrowing the radius spectrum.
The **effective diffusion coefficient** $D_{\text{eff, w}}$ of water vapor takes into account inhibition of mass transfer due to latent heat release on condensation and mass and thermal accommodation at the air-water interface:

$$
D_{\text{eff, w}} = \frac{1}{\frac{1}{D_w} + \frac{1}{\frac{1}{4}v_w a \alpha} + \frac{L_w^2 M_w^2 p_w(\infty)}{R_g T^3} \left( \frac{1}{k_a} + \frac{1}{\frac{1}{4}v_a \rho_a c_{p,a} a \beta} \right)}
$$

where $D_w$ is the diffusion coefficient of water vapor in air

- $k_a$ is the thermal conductivity of air
- $v_w, v_a$ are the mean molecular velocities of water vapor and air
- $\alpha, \beta$ are mass and thermal accommodation coefficients (0.5, 1)
- $L_w$ is the latent heat of condensation of water vapor
- $\rho_a$ is the density of air
- $c_{p,a}$ is the heat capacity of air at constant pressure
Size Distribution that served as Basis of Calculations

Seasalt, though low in number, dominates volume concentration.

Drop Classes used in the Calculations

**Critical radius vs. Dry radius**

**Critical supersaturation**

**Color code** allows history of individual drop classes to be readily followed. Particles are treated as (NH4)2SO4. Largest four classes (sea salt) are omitted in most calculations.
Example Results

Updraft velocity $w$ taken as 0.5 m s$^{-1}$, typical of moderately strong updraft (stratocumulus). Calculation starts at relative humidity 90%, with dry adiabatic lapse rate. ($\text{(NH}_4\text{)}_2\text{SO}_4$ mass loading 0.275 µg m$^{-3}$; mixing ratio 0.048 nmol/mol(air) (ppb)).

Break at ~320 s is due to latent heat release at onset of cloud formation.
Note *sharp maximum in saturation ratio* followed by decrease as water vapor condenses on newly activated cloud drops.

Note abrupt change of drop radii upon activation. Note compression of cloud drop spectrum.

Note “knife-edge” separation of activated and unactivated particles. Unactivated particles grow and shrink as saturation ratio increases and then decreases as water is taken up on activated droplets.
Unactivated droplets exhibit saturation ratio equal to that of the environment, also plotted but hidden under the nest of curves at the top.

Colors denote last two classes of unactivated drops; black curve denotes environmental saturation ratio.

Note slight lag in equilibrium saturation ratio due to mass-transport kinetics.
Droplet Growth Kinetics

Fractional rate of change of radius of particles during activation \( \frac{1}{a} \frac{da}{dt} \)
Fractional growth evaluated as \( \frac{1}{a} \frac{da}{dt} \)

Evaluated as \( \frac{1}{a_{eq}} \frac{da_{eq}}{dt} = \frac{1}{a_{eq}} \frac{da_{eq}}{dp_w} \frac{dp_w}{dt} \)

The near equality establishes that **mass transport kinetics are not limiting activation** of the drops under these conditions; contrast Chuang, Charlson & Seinfeld (1997).
Sea salt droplets get a “head start” and the rest of the cloud droplets never catch up. The three largest classes of droplets never reach supersaturation but are effectively cloud droplets anyway, as pointed out by Hänel (1987).
Results with 1 ppb HNO$_3$ included

HNO$_3$ partial pressure vs. time

Note abrupt decrease as drops activate.

HNO$_3$ mass in activated drops vs. time

After activation mass is constant because low HNO$_3$ vapor pressure precludes exchange among drops.
HNO₃ aqueous concentration exerts a complex behavior that reflects mass transport of water vapor and HNO₃ and droplet growth (dilution).

As drops activate HNO₃ concentration rapidly decreases because of dilution.

HNO₃ concentration in last two unactivated classes increases as drop radii decrease as water transfers to newly activated drops.

The nonuniform HNO₃ concentrations would be expected to influence strongly pH-dependent reactions such as aqueous oxidation of SO₂ by O₃.
## Summary of Results

<table>
<thead>
<tr>
<th>(NH$_4$)$_2$SO$_4$ mass loading</th>
<th>(NH$_4$)$_2$SO$_4$ mixing ratio</th>
<th>Updraft velocity</th>
<th>Maximum Super-saturation</th>
<th>Droplet Number Conc.</th>
<th>Fraction NSS mass scavenged</th>
</tr>
</thead>
<tbody>
<tr>
<td>µg m$^{-3}$</td>
<td>nmol/mol(air) (ppb)</td>
<td>m s$^{-1}$</td>
<td>%</td>
<td>cm$^{-3}$</td>
<td></td>
</tr>
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<td>2.75</td>
<td>0.48</td>
<td>0.5</td>
<td>0.26</td>
<td>997</td>
<td>0.935</td>
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<td>0.275</td>
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<td>0.5</td>
<td>0.50</td>
<td>140</td>
<td>0.955</td>
</tr>
<tr>
<td>0.275 + seasalt</td>
<td>0.048 + 0.64</td>
<td>0.5</td>
<td>0.47</td>
<td>140</td>
<td>0.955</td>
</tr>
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<td>2.0</td>
<td>0.51</td>
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<td>0.955</td>
</tr>
<tr>
<td>2.75 + HNO$_3$</td>
<td>0.48 + 1</td>
<td>0.5</td>
<td>0.22</td>
<td>997</td>
<td>0.935</td>
</tr>
</tbody>
</table>
Conclusions

A *highly flexible model* (“desktop cloud”) has been presented that describes the mass-transport kinetics of water vapor condensation and cloud droplet activation.

The input parameters to the model can readily be changed to study cloud droplet activation for any situation of interest.

This model readily displays drop-size dependent processes occurring during cloud formation.

For the cases studied the *mass transport of water vapor is sufficiently fast that Köhler equilibrium can be assumed for activating drops*.

Large drops may not activate but become effective cloud drops anyway.

The reversible uptake of even a highly soluble gas such as HNO₃ exerts a complex dependence on drop size and time.

This model can be used to develop parameters for large scale models or to test alternative formulations such as moment methods.

Acknowledgment

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This presentation is available on the web at [http://www.ecd.bnl.gov/steve/pubs.html](http://www.ecd.bnl.gov/steve/pubs.html)