Models of Clouds, Precipitation and Storms
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INTRODUCTION

Clouds play an important role for life on earth. Apart from influencing, for example, the radiative balance of the atmosphere and the lifetime of atmospheric trace constituents, they are the essential element in the hydrological cycle. Clouds transport the evaporated water of the oceans to the continents where the precipitation releases the water load. This release of water can be more or less vigorous depending on the energy stored in the cloud in the form of condensed hydrometeors (liquid or solid). This energy depends on the amount of available moisture and the way the vertical lifting necessary for cloud formation proceeds. We distinguish here mainly two different forms with varying extensions on the horizontal scale: the gentle uplift associated to the large-scale lifting, for example at a frontal zone, and the vigorous small-scale ascent associated with convection, knowing that also mixed forms of the lifting occur.

Consequently, when deciding to model clouds, precipitation, and storms, we immediately encounter a problem of scale: in order to model correctly the large-scale dynamics, we need to consider a horizontal domain of several thousand kilometers. And in order to consider correctly the form and the size of the condensed hydrometeors, we need also to take into account processes that take place on the micrometer scale.

Below, we quickly review the type and scale of microphysical processes that need to be considered (Flossmann and Laj, 1998) before presenting the different types of approaches used in cloud modeling in order to account for the scale problem.
MICROPHYSICAL PROCESSES

For a complete description see, for example, Pruppacher and Klett (1997), Rogers and Yau (1989), Cotton and Anthes (1989), and Houze (1993).

Nucleation of Drops

Our atmosphere does not allow the formation of drops by agglomerating water vapor molecules (homogeneous nucleation) alone. In order to form a tiny drop of $2 \times 10^{-3} \mu m$ radius, assembled of 800 molecules, it would require a relative humidity of 200%. Consequently, in the atmosphere droplets form on already existing nuclei. A subset of the aerosol particles present in every air mass provides these necessary nuclei.

Aerosol particles typically encountered in the atmosphere have a size range between $10^{-3}$ and $10 \mu m$. Their total number concentration varies between 100 and 100 000 cm$^{-3}$ and their mass can reach up to several hundreds of $\mu g \text{m}^{-3}$. It has been proposed to distinguish three different size ranges for the particles corresponding to different formation and destruction mechanisms:

- Table 1 points out that Aitken and large aerosol particles are formed by condensation processes from the gas phase and aggregation. The gases can be of natural or anthropogenic origin. However, giant aerosol particles have a mechanical origin, which is mostly natural.

- Depending on their size, chemical composition and the ambient relative humidity aerosol particles take up a certain amount of water (Köhler curve, see Pruppacher and Klett, 1997). So even in the absence of a cloud, the atmosphere is full of tiny droplets representing swollen aerosol particles, which can become visible, for example, in an atmospheric haze.

- At a given size, particles carrying more hygroscopic material can take up more water before achieving a critical size at a critical relative humidity (supersaturation below 5%). When a given aerosol particle has passed its critical size, we call it activated. It serves as a cloud condensation nucleus (CCN). Then, it does not require a further increase in relative humidity to make it grow and it is considered a cloud droplet as long as supersaturation prevails. At a given ambient supersaturation, all aerosol particles whose critical supersaturation is below the ambient one will be activated and transformed into drops.

Condensation and Evaporation

Once moist aerosol particles have been activated to form drops, they grow according to the droplet growth equation

$$\frac{d \text{mass}}{dt} = \left( \frac{s_{sw} - 2M_w \sigma_{s,h} - \frac{v \rho_w \phi_v M_w \rho_i r^3}{M_s \rho_w (r^3 - r_N^3)}}{\rho_w RT} \right) f_s$$

Equation (1) describes the change of the drop mass $m$ (radius $r$) as a function of the supersaturation $s_{sw} = e_a/\varepsilon_{sat,w}$ (ratio of the vapor pressure and the saturation vapor pressure). Herein, $T$: temperature, $l_v$: latent heat of evaporation, $D'$: modified diffusion coefficient, $K'$: modified thermal conductivity, $\sigma_{s,h}$: surface tension of the solution droplet, $\rho_w$, $\rho_i$: density of water and salt respectively, $M_w$, $M_s$: molecular weight of water and salt, $v$: number of ions, $\phi_v$: osmotic coefficient, $K$: universal gas constant, $e_a$: mass fraction of soluble material, $r_N$: radius of the insoluble aerosol nucleus, $f_s$: is the ventilation coefficient which is due to the falling motion of drops in the atmosphere (for more information see Pruppacher and Klett, 1997).

The growth rates given by this formula are rapid for small droplets. However, they decrease with an increasing radius and reduce considerably for drops larger than $10 \mu m$ radius (see Pruppacher and Klett, 1997). Consequently, the larger the drop, the longer it will take for it to grow further, leading to a halt in growth by condensation at around $30 \mu m$ in radius. Consequently, in order to form precipitation-sized drops other growth mechanisms interfere as detailed below.

Fall Speed of Drops

The terminal velocity of a drop is determined by a balance between buoyancy-corrected gravitation and the drag force.

Table 1 Classification of aerosols after Junge and Whitby and the different formation and removal processes most important for the various size classes of aerosol particles

<table>
<thead>
<tr>
<th>Size classes ($\mu m$)</th>
<th>Junge (1963)</th>
<th>Whitby (1978)</th>
<th>Formation</th>
<th>Removal</th>
</tr>
</thead>
<tbody>
<tr>
<td>$r &lt; 0.1$</td>
<td>Aitken particles</td>
<td>Nuclei or transient mode</td>
<td>Condensation from the gas phase</td>
<td>Aggregation</td>
</tr>
<tr>
<td>$0.1 &lt; r &lt; 1$</td>
<td>Large particles</td>
<td>Accumulation mode</td>
<td>Condensation, aggregation</td>
<td>Washout and rainout</td>
</tr>
<tr>
<td>$r &gt; 1$</td>
<td>Giant particles</td>
<td>Coarse mode</td>
<td>Sea spray, wind-blown dust, volcanoes, plant debris, diesel engines</td>
<td>Sedimentation</td>
</tr>
</tbody>
</table>

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Fall Speed of Drops

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acting on the drop. In Table 2, some values for the terminal velocity as a function of drop radius are given.

These values pertain to normal surface conditions. At higher altitudes the fall speed increases as a result of a decrease in air density. These fall speeds are determined in wind tunnel measurements because an analytical calculation using the balance between the acting forces is not possible. This is due to the fact that large drops are no longer spherical. They get deformed by the airstream and develop an indentation on the upwind side. Furthermore, they start to oscillate because of internal and external vortices that develop and because of electrical charges in the atmosphere and the drop. These features do not allow a correct analytical formulation of the drag.

**Collision and Coalescence of Drops**

Because of the difference in their size, drops have different terminal velocities. Thus, they can overtake each other while falling and collide to form a larger drop with the sum of the original masses. The probability for a small drop \( r_2 \) that is located inside the geometrical sweep-out volume of the larger drop \( r_1 \) to collide with this drop is given by the collision efficiency \( E_{\text{coll}} \) (Figure 1). This collision efficiency is strongly dependent on the size of the two drops. If the large drop is smaller than 20 \( \mu \text{m} \), a collision is essentially impossible, as the terminal velocity is too small and the small drop is just carried around the large drop by the airstream. In a wide-size range, the collision efficiency takes values of unity. When the two drops are roughly of equal size, \( E_{\text{coll}} \) can even exceed unity because of the wake capture of the trailing drop (see Pruppacher and Klett, 1997, for details).

\[
E_{\text{coll}} = \frac{\pi y_c^2}{\pi (r_1 + r_2)^2}
\]

**Figure 1** Schematical display of the collision efficiency

The process of collision is not always accompanied by coalescence, a fact that is described by the coalescence efficiency. Colliding drops may bounce apart, coalesce temporarily, and then separate again later, or coalesce temporarily and then shatter.

The collection efficiency of the process of drop–drop collision can be written as the product of the collision and the coalescence efficiency:

\[
E(m_1, m_2) = E_{\text{coll}} E_{\text{coal}}
\]

and this efficiency will determine the rate of precipitation formation in clouds that do not develop an ice phase.

It is important to note that collection is enhanced by a broad droplet size spectrum that depends essentially on the dynamics of the clouds but also on the nature and sizes of the aerosol particles, as well as turbulence in clouds, and even radiative cooling effects. The broadness of the distribution is influenced not only by the number of CCN but also by the presence of giant or ultragiant particles, which can serve as coalescence embryos.

**Breakup of Drops**

Drops in the atmosphere do not grow indefinitely. The upper size limit is around 6 mm in diameter. Drops arriving at this size range develop a tendency to break up. We can distinguish two main mechanisms:

- **Spontaneous breakup:** the deformation of large drops and the induced oscillation becomes so important that the drop becomes hydrodynamically unstable and breaks up
- **Collisional break up:** during the process of collision/coalescence, the newly formed drop is too unstable and breaks up forming two large drops, slightly smaller than the original drops, and a number of small satellite droplets (see discussion on coalescence above)

**Nucleation of Ice Particles**

In the atmosphere, liquid drops can exist even at temperatures well below 0°C, that is, in a supercooled state. In fact, significant numbers of ice particles start to form only below –5°C coexisting mostly still with liquid drops. Homogeneous freezing of liquid droplets depends on the size; large droplets can freeze homogeneously at temperatures of around –33°C, whereas by –40°C even the smallest droplets freeze homogeneously. Thus, at –40°C the last liquid has disappeared. In the temperature range between

<table>
<thead>
<tr>
<th>Radius ( r ) in ( \mu \text{m} )</th>
<th>50</th>
<th>100</th>
<th>200</th>
<th>500</th>
<th>1000</th>
<th>2000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terminal velocity ( V_{\infty} ) in ms(^{-1} )</td>
<td>~0.25</td>
<td>~0.8</td>
<td>~1.5</td>
<td>~4</td>
<td>6.5</td>
<td>8.5</td>
</tr>
</tbody>
</table>
–5 and –40°C, the presence of ice-forming nuclei is necessary to initiate the formation of an ice crystal. These ice nuclei (IN) are aerosol particles that can act in four main ways:

- **Deposition mode:** water is adsorbed directly from the vapor phase onto the surface of an IN where it is transformed into ice.
- **Condensation–freezing mode:** this is a hybrid process that requires supersaturation with respect to water. Here, the CCN that has formed the drop acts now as an IN. This process seems far more effective than the deposition mode.
- **Freezing mode:** the IN, scavenged by the drop, initiates the ice phase from within a supercooled water droplet.
- **Contact mode:** the IN initiates the ice phase at the moment of contact with the supercooled drop.

The number of IN depends on the chemical properties of the aerosol particles. It has been found that there exists a dependency on supersaturation (Meyers et al., 1992) and also on temperature (Fletcher, 1962). In contrast to CCN, a good IN should be insoluble and dispose already of a crystalline-type structure to facilitate the formation of the ice lattice (e.g. silicate). Some bacteria have also been identified as excellent IN. For a comprehensive review of the biogenic versus anthropogenic sources of IN, see Szyrmer and Zawadzki (1997).

**Deposition and Sublimation**

Water droplets in the atmosphere have a spherical appearance facilitating their analytical treatment. Ice crystals present more problems since they appear in a number of different shapes depending on temperature and the grade of water supply. Here we find densely packed structures like needles, columns, and plates, as well as light dendritic structures (for details see e.g. Pruppacher and Klett, 1997). The actual growth rate of these crystals by water vapor deposition depends on the crystal form as well as temperature and humidity.

This deposition growth can take place at the same time as the condensation growth of droplets, if the ambient relative humidity is high enough to maintain supersaturation over liquid-water and over ice. However, the equilibrium curves of H₂O allow also the case that the air is supersaturated with respect to ice and subsaturated with respect to liquid water. Then, the drops evaporate and the vapor condenses onto the ice crystals (Bergeron–Findeisen effect; Pruppacher and Klett, 1997).

**Fall Speed of Ice Particles**

Ice particles like droplets have a terminal velocity that depends heavily on the size of the particle. Additionally, they depend on the shape of the particle and its density, which can take values between 0.98 g cm\(^{-3}\) and 0.2 g cm\(^{-3}\). A light dendritic structured ice particle will, thus, have a lower terminal velocity than a dense plate-like crystal even if both have the same diameter. For aggregated and rimed structures, the same applies.

**Aggregation and Rimming**

For droplets we noted that condensation alone cannot develop precipitation, and we need the process of collision/coalescence. The same applies for ice crystals. Their growth by deposition of water vapor alone is also limited. Thus, a process of collision is necessary to produce larger aggregates. Two different mechanisms are possible:

- **The collision of ice crystals among themselves:** this process is called aggregation. It is responsible for the formation of snow crystals. This process is efficient in two different temperature regimes, around –10 to –15°C and around 0°C. Around –10 to –15°C, the crystals develop a dendritic structure and, thus, upon the collision of two crystals they easily succeed in interlinking with their branches. Around 0°C, the crystals develop a “pseudo liquid layer” (a micro layer of melting ice) at their surface which enables them to create a link with the other crystal via freezing. Outside these specific temperature ranges, the collision of two ice crystals is rarely successful, thus, not resulting in a snow flake.

- **The collision of ice crystals with liquid drops:** this process is called riming. Here, the drop is frozen upon collision with the crystal and several of these collisions transform the ice crystal into a graupel and then into a hail particle. This is a very efficient way of precipitation production. Sometimes the latent heat involved in this riming process is so important that not all the total captured liquid can be frozen. Normally, the remaining liquid will be incorporated into the solid structure forming a “spongy” ice. In a current meso-scale model (MSL), this is not treated. Thus, the excess liquid is shed and reattributed to the liquid water, a process called wet growth.

**Secondary Ice Particle Formation**

During the above-discussed processes of aggregation and riming, an ice multiplication mechanism can occur. If the crystals involved in the collision have a dendritic structure, parts can easily break off and form new ice crystals. Hallett and Mossop (1974) also proposed a process leading to secondary ice formation during the process of riming. If the drop involved freezes from the outside-in, then first a solid shell will form with a liquid core. When the liquid core starts to freeze, the volume inside the ice shell is too small to receive the forming ice. This fact will explode the shell and eject liquid material in the air, which will...
immediately freeze and form new ice crystals. This process is called **splintering**.

**Melting**

Even though a precipitating particle might have formed through riming of ice crystals, it can arrive at the ground as a liquid raindrop. This depends on the altitude of the 0 °C level in the atmosphere. Below this level, ice aggregates will start to melt and, depending on the fall distance, will arrive at the ground as partly or completely melted hydrometeors. A 3-mm ice particle can fall at a distance between 1 and 3 km before complete melting, depending on its initial density and its warm environment (see Pruppacher and Klett, 1997, for details).

**THE SCALE PROBLEM**

Models are an assembly of equations that describe the phenomenon to be studied. For a cloud model, this includes equations describing the change in time of temperature, humidity, pressure, density, and three-dimensional (3-D) wind field, and some variables that represent the cloud, such as drop or crystal numbers or liquid water content. These equations emerge from the concepts of conservation of mass and energy, Newton’s law, and the perfect gas law and can be written in a general form:

\[
\begin{align*}
\frac{\partial \rho}{\partial t} + \text{div}(\rho \mathbf{v}) &= 0 \\
\frac{\partial T}{\partial t} + \text{div}(\rho \mathbf{v} T) &= \frac{1}{c_p} \text{div} \mathbf{F}_T + \frac{1}{\rho c_p} \frac{dp}{dt} + \frac{L}{c_p} C_{\text{con}} \\
&+ \frac{L_{\text{sub}}}{c_p} C_{\text{sub}} + \frac{L_{\text{melt}}}{c_p} C_{\text{melt}} \\
\frac{\partial}{\partial t}(\rho \mathbf{v}) + \text{div}(\rho \mathbf{v} \mathbf{v}) &= \text{grad} p - \mathbf{g} + \text{div} \mathbf{F} \\
\frac{\partial}{\partial t} \rho q_v + \text{div}(\rho \mathbf{v} q_v) &= \text{div} \mathbf{F}_v - C_{\text{con}} - C_{\text{sub}} - C_{\text{melt}} \\
\frac{\partial R}{\partial t} + \text{div}(\mathbf{v} R) + \frac{\partial}{\partial z}(V_{\infty, \text{drop}} \mathbf{N}_{\text{drop}}) &= \text{div} \mathbf{F}_{\text{drop}} + S_{\text{drop}} \\
\frac{\partial R}{\partial t} + \text{div}(\mathbf{v} R) + \frac{\partial}{\partial z}(V_{\infty, \text{crystal}} \mathbf{N}_{\text{crystal}}) &= \text{div} \mathbf{F}_{\text{crystal}} + S_{\text{crystal}} \\
p/\rho &= nRT
\end{align*}
\]

with \( \rho \): density of air, \( \mathbf{v} \): 3-D wind field, \( T \): temperature, \( q_v \): water vapor mixing ratio, \( c_p \): specific heat at constant pressure, \( \mathbf{F}_T \): sensible heat-flux, \( p \): pressure, \( L \), \( L_{\text{sub}} \), \( L_{\text{melt}} \): latent heat of condensation, sublimation, and melting, \( C_{\text{con}}, C_{\text{sub}}, C_{\text{melt}} \): rate of phase change of condensation, sublimation, and melting, \( \mathbf{g} \): acceleration of gravity, \( \tau \): friction tensor, \( N_{\text{drop}}, N_{\text{crystal}} \): number of drops and crystals of a given size per unit volume, \( \mathbf{F}_{\text{drop}}, \mathbf{F}_{\text{crystal}} \): diffusion fluxes of drops and crystals, \( S_{\text{drop}}, S_{\text{crystal}} \): source, sinks, and transfer terms of drops and crystals, \( n \): number of moles, \( R \): universal gas constant. (Instead of temperature, models often calculate potential temperature \( \theta = T(p_0/p)^{R/c_p} \).)

These equations are, then, solved numerically at a number of points in space and time, while the points represent a certain interval in time or space. As the equations result from continuity principles, these intervals should be infinitely small. In practice, the grid boxes have a finite size. Thus, to be correct, we need to average the equations over finite increments in space and time:

\[
\begin{align*}
\bar{\psi} &= \frac{1}{\Delta t \Delta x \Delta y \Delta z} \int_{t}^{t+\Delta t} \int_{x}^{x+\Delta x} \int_{y}^{y+\Delta y} \int_{z}^{z+\Delta z} \psi \, dz \, dy \, dx \, dt \\
\psi &= \bar{\psi} + \psi'(5)
\end{align*}
\]

This modifies the equations in such a way that we now calculate average values over a grid box that modifies the equations so that a new term appears in every prognostic equation taking into account the subgrid changes of space and time inside the box:

\[
\begin{align*}
\frac{\partial \bar{\rho}}{\partial t} + \text{div}(\bar{\rho} \bar{\mathbf{v}}) &= 0 \\
\frac{\partial \bar{T}}{\partial t} + \text{div}(\bar{\rho} \bar{\mathbf{v}} \bar{T}) &= \frac{1}{c_p} \text{div} \bar{\mathbf{F}}_T + \frac{1}{\bar{\rho} c_p} \frac{d\bar{p}}{dt} + \frac{\bar{L}}{c_p} \bar{C}_{\text{con}} \\
&+ \frac{\bar{L}_{\text{sub}}}{c_p} \bar{C}_{\text{sub}} + \frac{\bar{L}_{\text{melt}}}{c_p} \bar{C}_{\text{melt}} \\
\frac{\partial \bar{\rho} \bar{\mathbf{v}}}{\partial t} + \text{div}(\bar{\rho} \bar{\mathbf{v}} \bar{\mathbf{v}}) &= \text{grad} \bar{p} - \bar{\mathbf{g}} + \text{div} \bar{\mathbf{F}} \\
\frac{\partial \bar{\rho} q_v}{\partial t} + \text{div}(\bar{\rho} \bar{\mathbf{v}} q_v) &= \text{div} \bar{\mathbf{F}}_v - \bar{C}_{\text{con}} - \bar{C}_{\text{sub}} - \bar{C}_{\text{melt}} \\
\frac{\partial \bar{R}}{\partial t} + \text{div}(\bar{\mathbf{v}} \bar{R}) + \frac{\partial}{\partial z}(\bar{V}_{\infty, \text{drop}} \bar{\mathbf{N}}_{\text{drop}}) &= \text{div} \bar{\mathbf{F}}_{\text{drop}} + \bar{S}_{\text{drop}} \\
\frac{\partial \bar{R}}{\partial t} + \text{div}(\bar{\mathbf{v}} \bar{R}) + \frac{\partial}{\partial z}(\bar{V}_{\infty, \text{crystal}} \bar{\mathbf{N}}_{\text{crystal}}) &= \text{div} \bar{\mathbf{F}}_{\text{crystal}} + \bar{S}_{\text{crystal}} \\
\bar{p}/\bar{\rho} &= nRT
\end{align*}
\]

The new terms have the form:

\[
\begin{align*}
\bar{\mathbf{F}}_T &= -\bar{\rho} \bar{\mathbf{v}}' \bar{\mathbf{v}}, \quad \tau_{\text{turb}} = -\bar{\rho} \bar{\mathbf{v}}' \bar{\mathbf{v}}'
\end{align*}
\]

\[
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\]
and depend on the subgrid fluctuations $\psi'$ in space and time of the considered variables. These terms need to be parameterized as a function of the variables $\bar{\psi}$ of the resolved scale. This approach is known under the term closure, whereby different theories of closure (first, second order, etc.) can be found in the literature (Ogura and Phillips, 1962). In the following, for simplicity of writing, the overbar will be dropped; however, all equations are averaged equations.

Considering, that an appropriate closure approach has been selected, the foregoing equations can be solved numerically in a type of grid such as that displayed in Figure 2, covering the entire cloud area and the affected environment. The averaging done in equations (6) and (7) now allows finite grid increments. However, the microphysical processes discussed above make it evident that essential processes in clouds operate at the scale of micrometers, requiring very small intervals in space. But, on the other hand, clouds sometimes form systems of several thousand kilometers, requiring a large spatial domain to be covered. One is, thus, confronted with the problem that actual computer power does not allow us to cover domains of 1000 km$^2$ with grid increments of 1 m$^2$ (this would require 10$^{12}$ grid boxes).

A similar argumentation applies to the time increment. Time and space increments are linked in an explicit numerical treatment of the system of equations by the Courant–Friedrich–Levy (CFL) criterium (Jacobson, 1999):

$$\left( \frac{\Delta t}{\Delta x} \right) U \leq 1$$

with $U$ being the maximum transport velocity considered. This criterium ensures that the fasted signal does not jump a grid box in one time step. The fasted transport velocities in the atmosphere are those of the sound. They are not physically relevant in cloud models and are, thus, filtered by suppressing local density fluctuations in the equations (an elastic or soundproof assumption) (Ogura and Phillips, 1962).

However, the numerical effort for a high resolution of the processes stays considerable. Consequently, strategies to reduce the simulation time need to be developed. These consist in compromises concerning the dynamics, the microphysics, and the resolution.

SIMPLIFICATION IN THE DYNAMICS

3-D Models

The appropriate way to simulate the dynamics of the atmosphere is a 3-D representation of space. Here, a classical coordinate system would be a Cartesian one with $x$, $y$, and $z$ in the directions of space (see Figure 2). In order to take into account the topography, often the vertical coordinate $z$ is replaced by one that follows the terrain. Phillips (1957) designed such a terrain-following coordinate system for use in hydrostatic, numerical prediction models by defining:

$$\sigma = \frac{p}{p_s} \tag{9}$$

where $p_s$ represents surface pressure and $\sigma$ varies from a value $\sigma = 1$ at the ground to $\sigma = 0$ at the top of the atmosphere. Nowadays, however, most of the 3-D models (Clark, 1977; Dudhia, 1993; Pielke et al., 1992) devoted to cloud simulations use the vertical coordinate:

$$z^* = H \frac{z - z_s}{H - z_s} \tag{10}$$

The surface height above some reference level is given by $z_s$ (e.g. sea level) and the height of the model top is given by $H$.

Three-dimensional models that cover an entire continent equally need to take into account in the horizontal grid the deformation of the coordinate system by the curvature of Earth and the large-scale Coriolis force. These models use stereographic coordinate systems. Some of the most widely used 3-D models are currently RAMS (Pielke et al., 1992; Cotton et al., 2003), MM5 (Dudhia, 1993), and Clark (1977), among others.

2-D Models

For special case studies concerning phenomena that are mainly two-dimensional (2-D; flow over a mountain range, e.g.) or in order to reduce computer calculation times, 3-D dynamics can be reduced to two dimensions. This eliminates one coordinate (e.g. $y$ in a Cartesian coordinate system) (e.g. Orville and Kopp, 1977; Soong and Ogura, 1980)

Another configuration, especially adapted in simulating isolated cumulus clouds consists of a cylindrical model with a symmetry with respect to the angular coordinate. Such
a model configuration has been used, for example in the models of Shino (1983), Murray and Koenig (1975), and Reisin et al. (1996).

1-D and 0-D Models

The cylindrical model philosophy has also been applied in the 1.5-D model of Asai and Kasahara (1967) where in fact two one-dimensional models are connected to represent the updraft and the downdraft region of a convective cloud.

Nowadays, one-dimensional models are mainly used to test microphysical schemes. In this case, there exists just a vertical axis $z$, subdivided into several layers, representing the updraft region of the cloud.

Finally, the least sophisticated dynamics is that of an air parcel. In this concept, we consider a volume of air separated from the environment by an immaterial surface. For the entraining air parcels, they exchange mass and heat with the environment; otherwise, they are considered as adiabatic. Air parcels can represent an interesting concept if they are driven by a 3-D flow field, because they permit highly resolved microphysics to be followed in a larger-scale dynamics that does not consider microphysics (Feingold et al., 1998a). Here, care needs to be taken to ensure that the parcel bulk properties are adequately constrained to those from the driving 3-D model.

MODELS OF CLOUD MICROPHYSICS

Drops and ice crystals exist in a cloud in varying number concentrations and with different sizes. We distinguish cloud drops of radii between 1 $\mu$m and 30 $\mu$m, drizzle drops of up to 200 $\mu$m, and rain drops up to 6 mm. Ice crystals have diameters of up to 100 $\mu$m, snow flakes, graupel, and hail particles have no clear upper size but can become larger than raindrops. The number of hydrometeors per unit volume of a given size is fundamental for the evolution of the cloud and, thus, should be considered in a cloud model.

Explicit or Bin-resolving Cloud Models

The most logical approach is to introduce an explicit number density distribution function $f(m) \, dm$, which follows the number per unit volume of a given type of hydrometers in a given mass class between $m$ and $m + dm$ (see Figure 3). Inside a bin, class most models assume a uniform value. There exist, however, also models that assume inside a drop bin class a distribution of aerosol particles or chemical compounds (e.g. Wobrock et al., 2002).

Warm Clouds

Most ground fogs or low-level clouds contain only liquid drops, and no ice crystals. If they develop precipitation, this will take place by the collision and coalescence of liquid drops. These clouds are called warm clouds. In order to model the size distribution of droplets in warm clouds, a drop number density distribution function $f_d(m)$ is defined. It allows the continuous distribution of drops over the entire spectrum to be captured in a discrete function.

Since the liquid hydrometers need to cover a range from 1 $\mu$m drops up to 6 mm raindrops, which covers at least three orders of magnitude, an equidistant grid in size or mass is not appropriate. Consequently, Berry and Reinhardt (1974a–c) have introduced a logarithmic size grid which allows a doubling of mass every JRS category:

$$r(j) = r(1)2^{(j-1)/3JRS}$$

$r(1)$ gives the minimum radius (e.g. 1 $\mu$m) and JRS can be 1, 2, 3, or more. In most models JRS = 2 is chosen, as a compromise between the necessary resolution of drop spectrum (Silverman and Glass, 1973) and the computer times of the simulation. For each of the resulting drop-size classes (on the order to 70), the following prognostic equation has to be solved:

$$\frac{\partial f_d(m, \tilde{r}, t)}{\partial t} = -\text{div}(\vec{v} f_d(m, \tilde{r}, t)) + \frac{\partial}{\partial z}(V_{\infty} f_d(m, \tilde{r}, t)) + \text{div}(\vec{\nu}_{\text{mix}} f_d)$$

$$+ \left[ \frac{\partial}{\partial t} f_d(m, \tilde{r}, t) \right]_{\text{nuc}} + \left[ \frac{\partial}{\partial t} f_d(m, \tilde{r}, t) \right]_{\text{cond/evaporation}}$$

$$+ \left[ \frac{\partial}{\partial t} f_d(m, \tilde{r}, t) \right]_{\text{coll}} + \left[ \frac{\partial}{\partial t} f_d(m, \tilde{r}, t) \right]_{\text{breakup}}$$

The term on the left-hand side calculates the change of the number of drops at a certain location $\tilde{r}$ per unit volume between a mass $m$ and $m + dm$ per time interval $dt$ because
of the physical effects represented by the terms on the right-hand side which are: transport with the scale velocity $\vec{v}$, the sedimentation with the terminal velocity $V_\infty$, transport through turbulence, the change in number concentration due to nucleation of drops, due to condensation or evaporation, due to collision and coalescence among the drops, and due to breakup of drops. The mathematical expressions for these terms follow the physics detailed above.

A simple way to calculate the nucleation of drops is to prescribe the number of cloud condensation nuclei as a function of supersaturation:

$$N_{CCN} = c s^{k}_{vw} \quad (13)$$

with $c, k$ constants which are adapted to the air mass considered, as, for example, in Twomey (1959). In addition, an assumption is necessary on the size of the freshly nucleated drops. More sophisticated models (Flossmann et al., 1985) follow the number of aerosol particles and their chemical composition in order to calculate as a function of supersaturation (Köhler equation, Pruppacher and Klett, 1997) the number and the size of activated aerosol particles.

Once formed, the drops grow or shrink by condensation or evaporation respectively:

$$\left[ \frac{\partial}{\partial t} f_d(m) \right]_{\text{cond/eva}} = -\frac{\partial}{\partial m} \left( \frac{d m_{\text{cond/eva}}}{dt} f_d(m) \right) \quad (14)$$

where the individual droplet growth rate is given from equation (1). The process of collision and coalescence of drops is calculated following Berry and Reinhardt (1974a–c):

$$\left[ \frac{\partial}{\partial t} f_d(m) \right]_{\text{coll}} = \int_{0}^{m/2} f_d(m-m') f_d(m') K(m-m', m') \, dm' - \int_{0}^{\infty} f_d(m) f_d(m') K(m, m') \, dm' \quad (15)$$

Here, the first term on the right-hand side describes the gain of $m$-sized drops due to collision between drops of sizes $(m-m')$ and $m'$. The second term describes the loss of $m$-sized drops due to all types of collision. $K$ represents the coalescence kernels that give the probability of such a collision to happen. $E$ is the collection efficiency.

$$K(m, m') = \pi(r + r') |V_{\infty, \text{drop}}(m) - V_{\infty, \text{drop}}(m')| E(m, m') \quad (16)$$

The breakup term can be parameterized either as a spontaneous breakup (see e.g. Hall, 1980; Srivastava, 1971) or as a collisional breakup (Low and List, 1982a,b).

In solving these equations coupled with the 1, 2, or 3–D dynamics as detailed above, the calculation of the evolution of the drop-size distribution as a function of time and space inside and below the cloud, as well as on the ground is allowed.

However, because of the large number of variables to be updated at every time step, only a few models available can calculate explicit cloud microphysics in a 3-D framework (e.g. Feingold et al., 1994 Kogan, 1991 Khvorostyanov, 1995). Most models are restricted to two dimensions (Hall, 1980) or – one dimension (Ogura and Takahashi, 1973). The parcel model discussed above is an exception, since it does not allow precipitation to be calculated. It can, thus, only be used to simulate the evolution of a nonprecipitating cloud (Feingold et al., 1998), or at the most estimate the amount of precipitable water.

### Cold Clouds

Most clouds in midlatitudes form ice particles and the formation of precipitation involves the ice phase. The ice phase can also be modeled by a bin approach if a certain number of additional size distributions are considered – one distribution function to follow the pristine (non or little rimed) crystals: $f_i$, one distribution function for the heavily rimed particles like graupel or hail $f_h$, and one distribution function for snow flakes, that is, aggregates of crystals $f_a$. As ice particles generally can become larger than liquid drops, more categories need to be considered. Also, because of the different possible forms of ice crystals (columns, needles, plates, etc.) additional distribution functions can be added. However, most models follow just one species of crystal shape (plate-like; e.g. Alheit et al., 1990). Only a few models calculate the explicit cold microphysics in three-dimensions (Ovtchinnikov and Kogan, 2000). Most models use reduced dynamics, for example, 2-D (Respondek et al., 1995), 1 – D, or parcel models (Alheit et al., 1990). Takahashi had the first paper on bin-resolved microphysics of mixed-phase clouds in the 1970s. Reisin et al. (1996) is another review on mixed-phase bin models of considerable sophistication. It has been used in 2-D models of cumuli and also Arctic stratus clouds (Harrington et al., 2000).

### Bulk Parameterizations

Between 1965 and 1975, when cloud modeling started, due to the limitations of early computers, bin modeling in a reasonable dynamic framework was completely out of reach.

### Warm Clouds

Consequently, Kessler (1969) proposed a simple parameterization of the microphysics of a warm cloud (Figure 4).
MODELS OF CLOUDS, PRECIPITATION AND STORMS

It aimed to simulate a convective cloud and just calculated the amount of liquid water attached to the small, non-precipitating drops (\( r \) smaller than 50 \( \mu \text{m} \)) and the amount of liquid water attached to precipitation-sized drops (\( r \) larger than 50 \( \mu \text{m} \)). However, due to its simplicity this scheme was quickly adapted to all types of clouds, and still has its place in many weather prediction models.

The basic equations of this model allow calculation of the evolution with time of the water vapor mixing ratio \( q_v \), the cloud water mixing ratio \( q_c \), and the rain water mixing ratio \( q_R \).

\[
\frac{\partial q_v}{\partial t} = -\text{div}(\vec{v}q_v) + \text{div}(\vec{F}_{\text{turb}}) \\
- P_{\text{auto}} - P_{\text{acc}} + C_{\text{con,c}} \\
\frac{\partial q_c}{\partial t} = -\text{div}(\vec{v}q_c) + \frac{\partial}{\partial z} V_R q_c \\
+ \text{div}(\vec{F}_{\text{turb}}) + P_{\text{auto}} + P_{\text{acc}} - C_{\text{eva,R}}
\]

Here, Kessler (1969) proposes for the condensation rate of cloud drops \( C_{\text{con,c}} \) a saturation adjustment. This means that all dynamically generated supersaturation is immediately converted to cloud water. And in cases of subsaturation, an amount of cloud water is evaporated that restores saturation, if possible. Obviously, this is a rather crude treatment that does not consider the role of the CCN population and the supersaturation.

\( P_{\text{auto}} \) is the autoconversion rate, that is, the collision of cloud water that forms rainwater. It is assumed to be a linear function of the cloud water content, only influenced by threshold values \( q_{\text{crit}} \) determining the onset of the autoconversion process. In the literature, numerous different values for \( q_{\text{crit}} \) have been proposed.

\[
P_{\text{auto}} = k(q_{\text{crit}} - q_c) \\
k = 10^{-3} \text{s}^{-1} \text{ if } q_c > q_{\text{crit}}, \\
\text{otherwise } k = 0
\]

The accretion rate \( P_{\text{acc}} \) details the amount of rainwater created by further collision with cloud water. Kessler (1969) calculates the accretion rate by:

\[
P_{\text{acc}} = \int_{0}^{\infty} \frac{dn_{\text{acc}}}{dr} N_R(r) \, dr
\]

by calculating the increase of mass of one raindrop \( m_{\text{acc}} \) of radius \( r \) due to the collision with a cloud water population.
presumed monodisperse represented by \( q_c \) with a collision efficiency of \( E \).

\[
\frac{dm_{\text{acc}}}{dr} = \pi r^2 EV_k(r) \rho q_c \tag{20}
\]

In assuming that the raindrops obey a Marshall–Palmer distribution, (Marshall and Palmer, 1948):

\[
N_k(r) = N_{k,0} e^{-2\lambda r} \tag{21}
\]

where \( \lambda \): shape parameter, and assuming a simple analytical function for the drop terminal velocity:

\[
V_k(r) = 130\sqrt{r} \tag{22}
\]

yielding

\[
V_R = \frac{\int_0^\infty V_k(r) N_k(r) dr}{\int_0^\infty N_k(r) dr} \tag{23}
\]

in m s\(^{-1}\).

Autocollection of cloud or raindrops are not considered in this approach. The evaporation rate of raindrops below cloud base can be calculated by

\[
C_{\text{eva},R} = -\int \frac{dm_{\text{con/eva}}}{dr} N_k(r) dr \tag{24}
\]

Numerous authors have since modified the terms for the parameterizations of the different processes, keeping, however, the basic concept (Berry, 1965; Orville and Kopp, 1977, among others).

**Cold Clouds**

The basic concept of the warm cloud bulk parameterization was soon extended to cold clouds, distinguishing two or three categories of ice. One, corresponding to nonprecipitating cloud drops is called cloud ice. For the precipitating cloud ice, one can, for example, distinguish graupel from snow.

An example for cloud ice and graupel can be found in Orville and Kopp (1977); an example for cloud ice, graupel, and snow can be found in Lin et al. (1983), and an example using pristine, such as non- or little-rimed crystals (ice A) and rimed crystals (ice B) can be found in Koenig and Murray (1976). A new approach to couple bulk and explicit microphysics can be found in Farley and Orville (1986), which was recently applied to a 3-D model (Wobrock et al., 2003). It should be mentioned that the approach of Farley and Orville is a hybrid approach in which hail is simulated in bins, although continuous accretion approximations are used.

Examples for pure bulk ice physics in a 3-D model can be found in Reisner et al. (1998) and for a 2-D model in Grabowski et al. (1996).

**The Semispectral Microphysics Parameterizations**

As a compromise between the bin schemes that follow the spectra of hydrometeors in discrete size classes or bins and the bulk parameterizations that follow just the water mass associated to 5 or 6 classes of hydrometeors, recently, semispectral parameterizations have been developed.

These parameterizations, similar to the bulk ones, follow a limited number of categories. Within one category, however, they assume a size distribution of a log-normal or gamma type, and then calculate the time evolution of the first (total number) and third (mass) moment of these distribution functions. Example of these approaches can be found in Ferrier (1994), Meyers et al. (1997), Cohard and Pinty (2000), Seifert and Beheng (2003), and Caro et al. (2003). Actually, Meyers et al. was already extended to the ice phase. Moreover, their approach used approximate solutions or look-up table solutions to the stochastic collection equations as opposed to the continuous accretion approximations to collection. Subsequently, Feingold et al. (1998b) extended the emulation of a bin model in a bulk scheme to autoconversion and sedimentation.

More recently Saleeby and Cotton (2004) added a second mode to the cloud droplet spectrum which permitted a more accurate representation of collection (autoconversion) and sedimentation, and permitted the explicit activation of CCN (including a concept for giant CCN) and IN. Others such as Milbrandt and Yau (2004) extended this approach to a three-moment scheme in which all degrees of freedom in the specified basis functions are predicted. This approach offers an interesting compromise between the detailed bin approach and bulk schemes. The reduction in computational effort will probably mean that even operational forecast models will move towards these schemes in the future.

**CUMULUS PARAMETERIZATIONS**

If the size of the grid increments becomes larger than 10 km, it is no longer possible to resolve the convective clouds explicitly (stratiform clouds with large horizontal extension can still be resolved). But since the energetic processes associated with convective clouds are important, their net impact on the scale values needs to be parameterized, in addition to the turbulence parameterization already discussed. This gives rise to cumulus parameterizations which can be traced back to the development of numerical prediction models. Smagorinsky (1956) was the first to introduce a cumulus parameterization when he adjusted the vertical derivative of an “effective static stability” which included...
the heat released during condensation. This gave rise to the first approaches of cumulus parameterizations called convec-
tive adjustment (e.g. Krishnamurti and Moxim, 1971; Manabe et al., 1965; Kurilhara, 1973). As these approaches
turned out too crude, a second generation of schemes
attempted to link the convection to the large-scale mass,
motion, or energy convergence, while using some form of
“cloud model” (mostly rising plumes and compensating
downdrafts) to vertically distribute mass, moisture, and
energy by a flux scheme (e.g. Kuo, 1965, 1974; Anthes,
large-scale control was introduced by Arakawa and Schub-
bert (1974). They assumed that the rate of stabilization by
an ensemble of cumulus clouds balances the rate at which
the large scale makes buoyant energy available for convec-
tion. For a complete review of the approaches, see Emanuel
and Raymond (1993).

All cumulus parameterization schemes depend severely
on the size of the grid box, which they need to stabilize
completely or only partially. Consequently, they will only
apply to the case for which they were developed and cannot
easily be generalized for other grid sizes.

A recent approach, for example, Grabowski et al. (2001)
and Khairoutdinov and Randall (2001) concerns the de-
velopment of so-called “super-parameterizations” that have
now been implemented in several global circulation mod-
els. This approach consists basically in running a 2-D
cloud-resolving model at each grid point and average the
cloud-resolving model data to determine heating, moisten-
ing, and precipitation rates.

NUMERICS

The foregoing physical treatments of the different processes
involved in clouds enter the balance equations. These equa-
tions, then, need to be solved numerically on a computer.
However, before coding, the differential expressions in the
terms need to be discretized in order to be linked to the
scale variables. The real art of cloud models these days lies
here. Numerous studies have been performed on the best
numerical scheme to solve the transport equation (positive-
definite schemes; e.g. Smolarkiewicz, 1984; Bott, 1989)
and the integrals in the drop–drop collision/coalescence
process (e.g. Berry and Reinhardt, 1974a-c; Bott, 2000),
and it is the best way to distribute the calculated val-
ues in a grid, among others. Another serious numerical
problem lies in the calculation of the supersaturation due
to dynamical and microphysical tendencies. These ten-
dencies should be solved at the same time and not one
after the other as in the models. As these tendencies are
very sensible, they require extremely small time steps,
which in connection with the timescales of the other pro-
cesses introduce a numerical stiffness problem. Numerous
approaches concerning this problem have been proposed
in the literature. Next to the explicit method, implicit
methods were also developed. Equally, the state-of-the-
art cloud models now dispose of an interactive two-way
grid nesting. This allows zooming with finer resolution
into a domain of special interest and transferring this
information to the larger grid outside afterwards. These
techniques create a new way to address the scale prob-
lem, in following the large-scale dynamics in a coarse
grid and then, to zoom with a finer grid into the cloud
region. Another technique that advances cloud model-
ing is the assimilation of remote sensing data (e.g. from
satellite) into a model in order to improve model ini-
tialization. Furthermore, the calculated evolution can be
constrained by observation or outputs of other models.
For a summary of current modeling techniques, see Jacob-
son (1999).

CONCLUSION

This article attempts to provide an introduction to the
complex subject of modeling clouds, the production of
precipitation, and the development of cloud and storm
systems. The elements intervening in cloud modeling
have been exposed, starting from a description of the
physical phenomena. On the basis of the occurring scale
problem, a number of approaches for simplification were
presented. These parameterizations are generic and can
be combined as a function of the addressed problem. In
addition, other modules that have not been discussed in
the article need to be added, such as a radiation model, a
surface model, a chemistry model, and so on. The main
problem is achieving a balance between the degree of
parameterization, the number of physical and chemical
processes relevant to a particular problem, and the available
computing resources. Owing to the increasing capacity of
computers, however, less and less parameterizations are
necessary and the resulting models are more and more
complete. Consequently, the current state-of-the-art MSMs
(e.g. RAMS, MM5, Clark, etc.) have large capacities.
Essentially the same models can be used to study problems
as diverse as rainfall over large areas and over many hours,
resulting from frontal cloud systems, the development of
local storms with damaging hail and flooding, transport of
pollutants by small convective clouds, and photochemical
reactions in fog. Next to these models a variety of different
models exist which are more or less restricted to a
specific question.

FURTHER READING

REFERENCES


Keywords: microphysics processes; modeling techniques; bulk microphysics; bin microphysics; cumulus parameterization; numerical techniques