

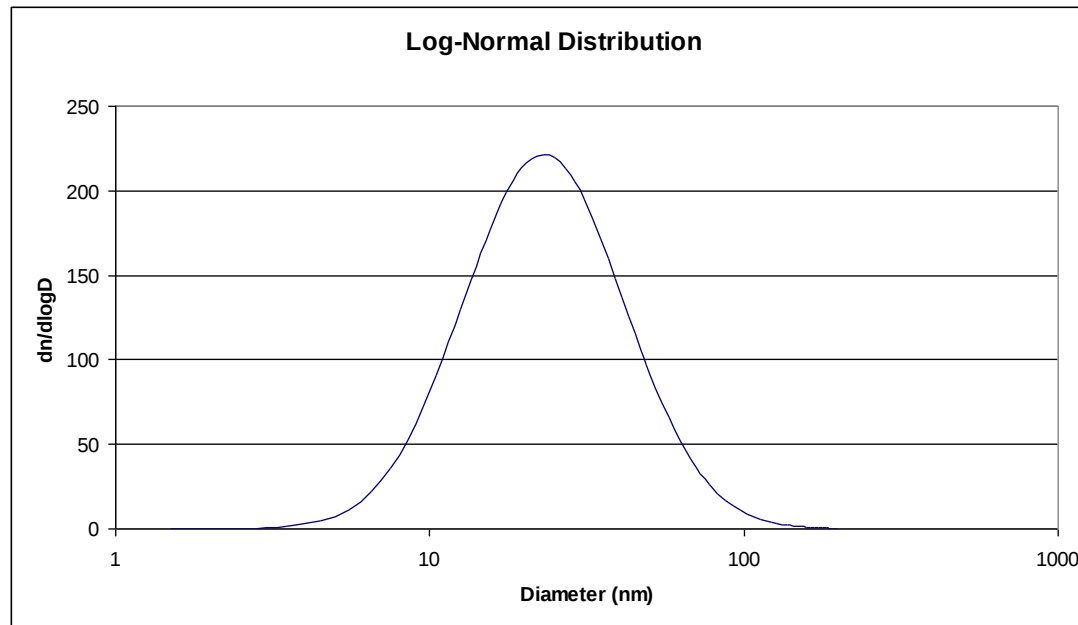
Aerosols and Cloud microphysics

Lecture 4 - ENV 407

Aerosol lognormal distributions

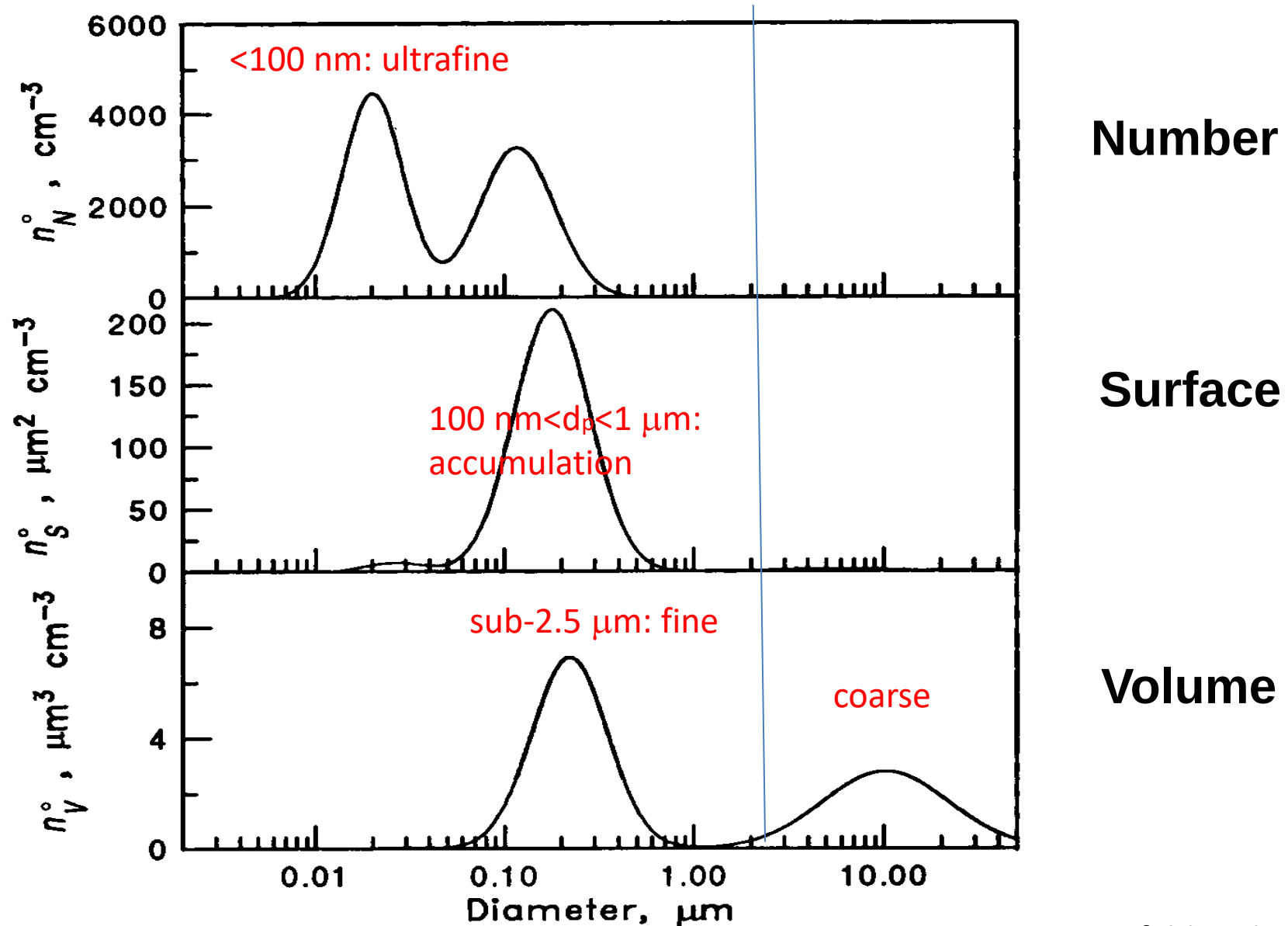
- Appears as a normal distribution when x-axis is plotted on log scale

$$n(D) = \frac{N}{(2\pi)^{1/2} \ln \sigma_D} \exp \left[-\frac{(\ln D - \ln D_g)^2}{2 \ln^2 \sigma_D} \right]$$

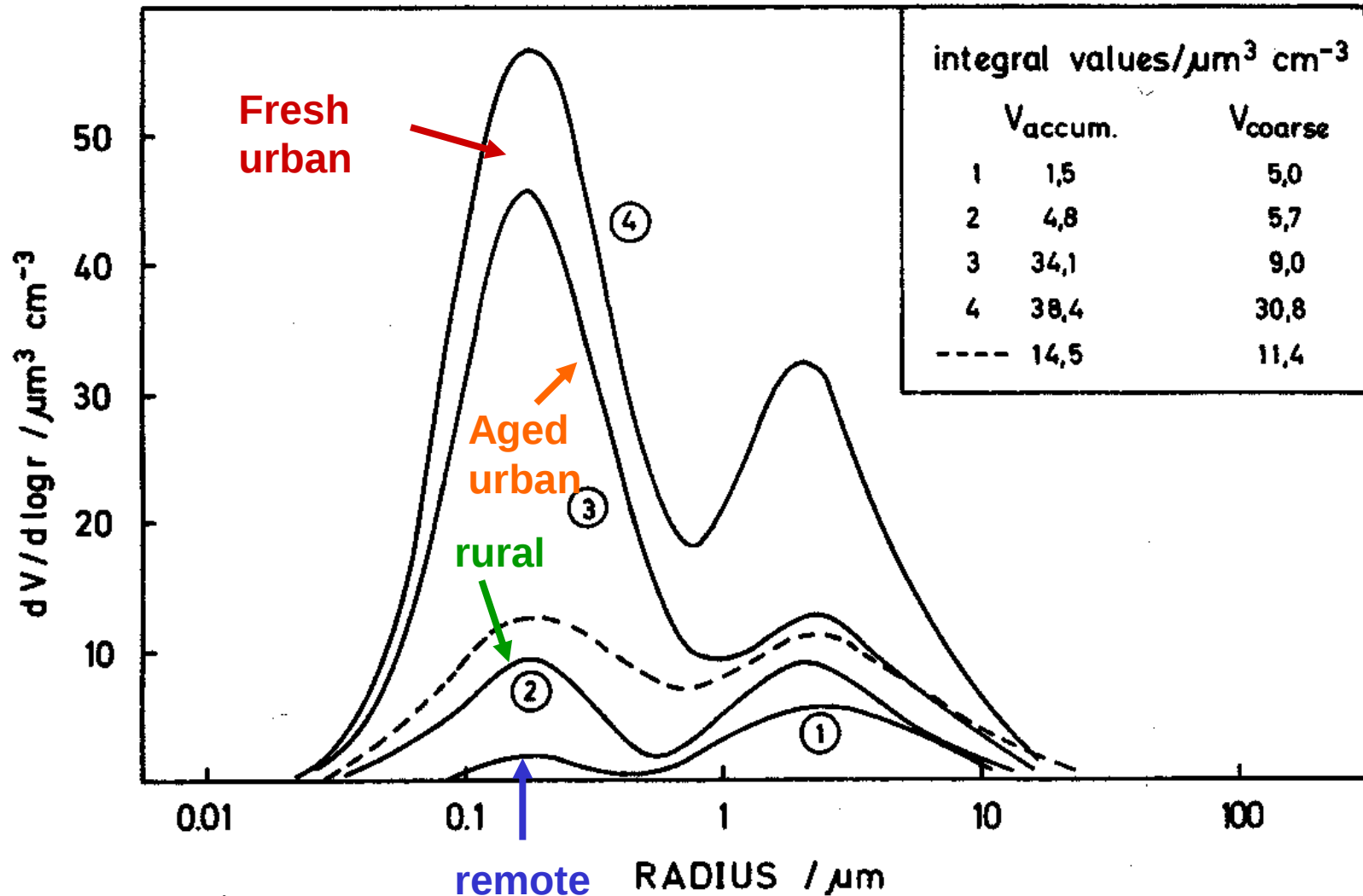


Geometric Mean
Diameter = 23 nm;
Geometric Standard
Deviation (σ) = 1.8

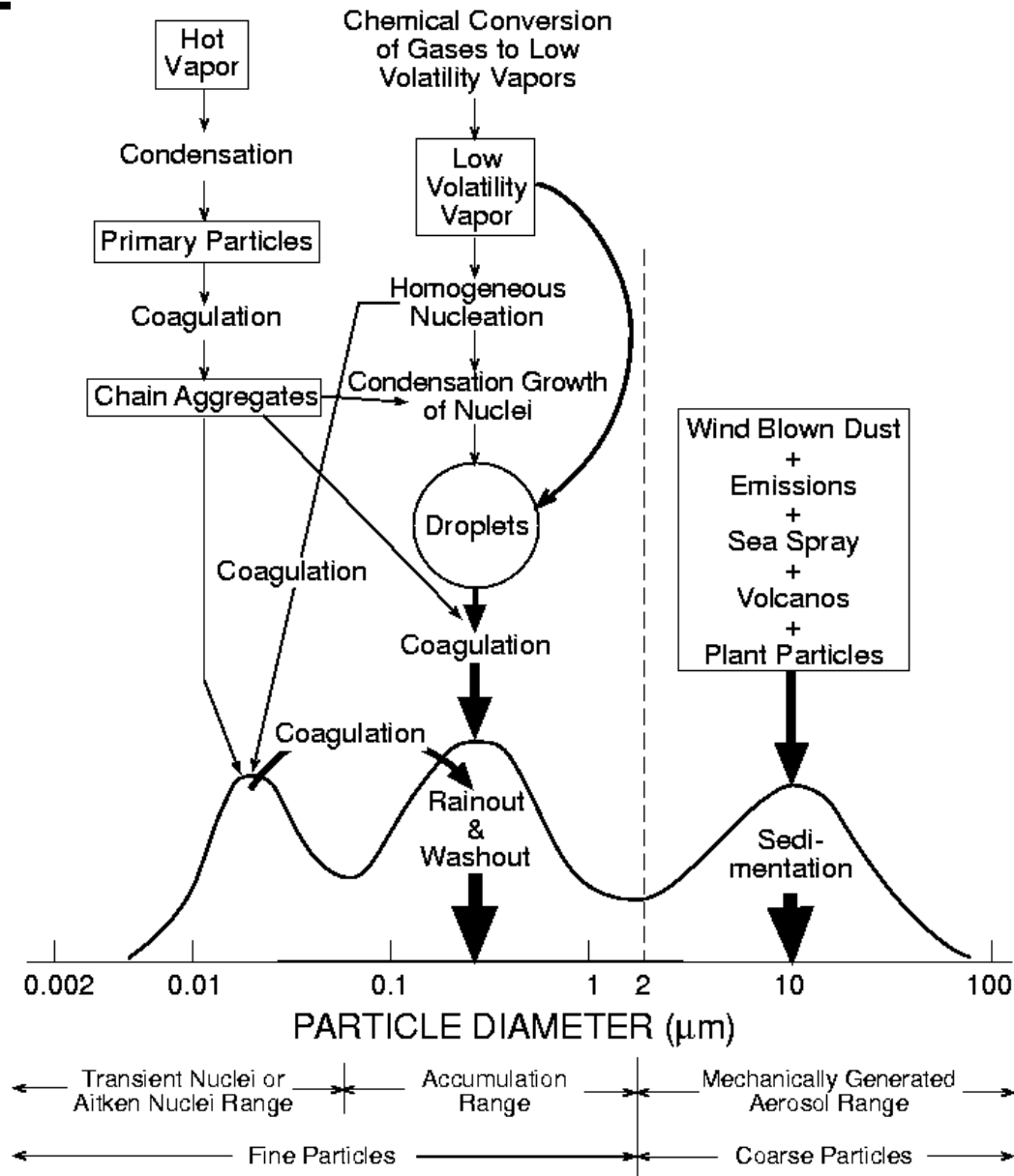
Size Distribution: Remote continental air



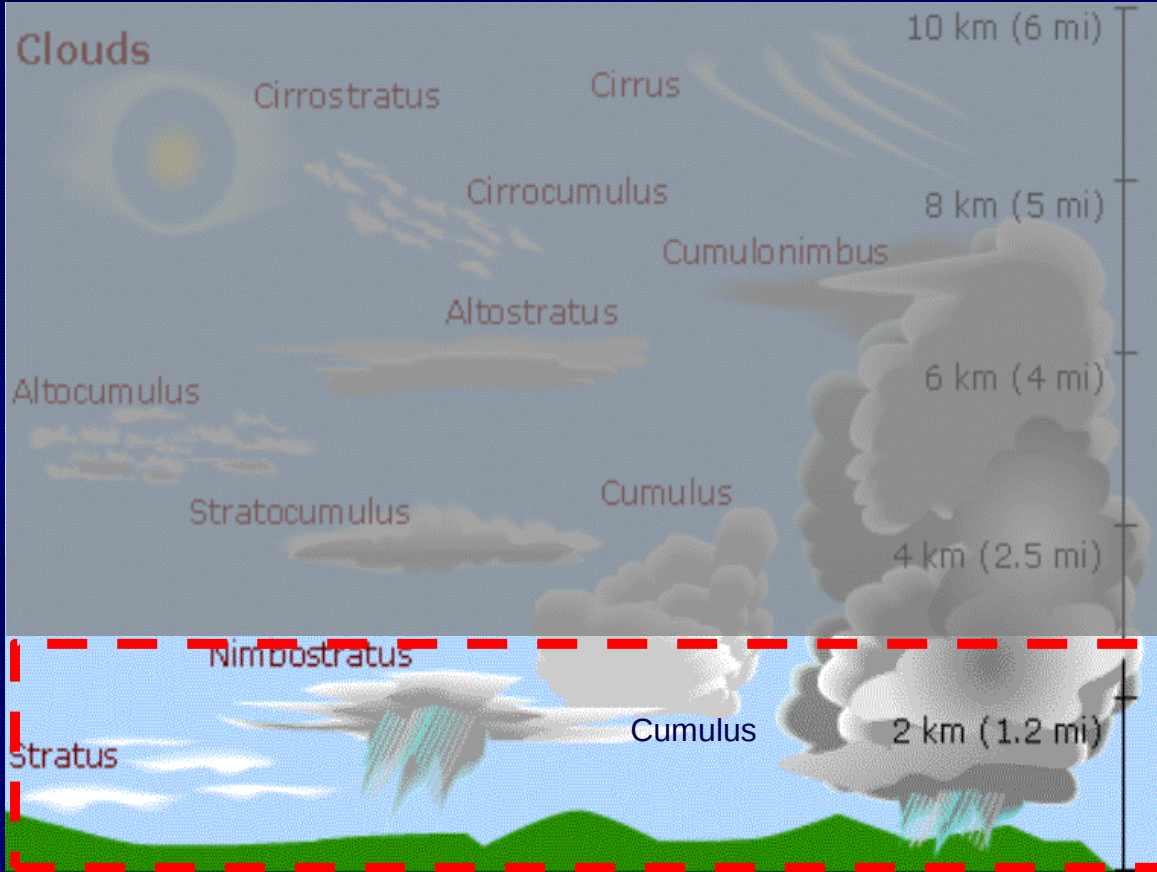
Size distributions vary a lot



Size distribution is shaped by the processes present



LIQUID cloud microphysics



- **Ice (cold) clouds:**
Made of ice crystals at $T < 235 \text{ K}$.
- **Mixed Phase clouds:**
Mixture of liquid droplets and ice for T between 235 and 273K
- **Liquid (warm) clouds:**
Made of liquid droplets at $T > 273 \text{ K}$

Cloud particles are not created directly from the vapor phase but from **suspended aerosol particles**

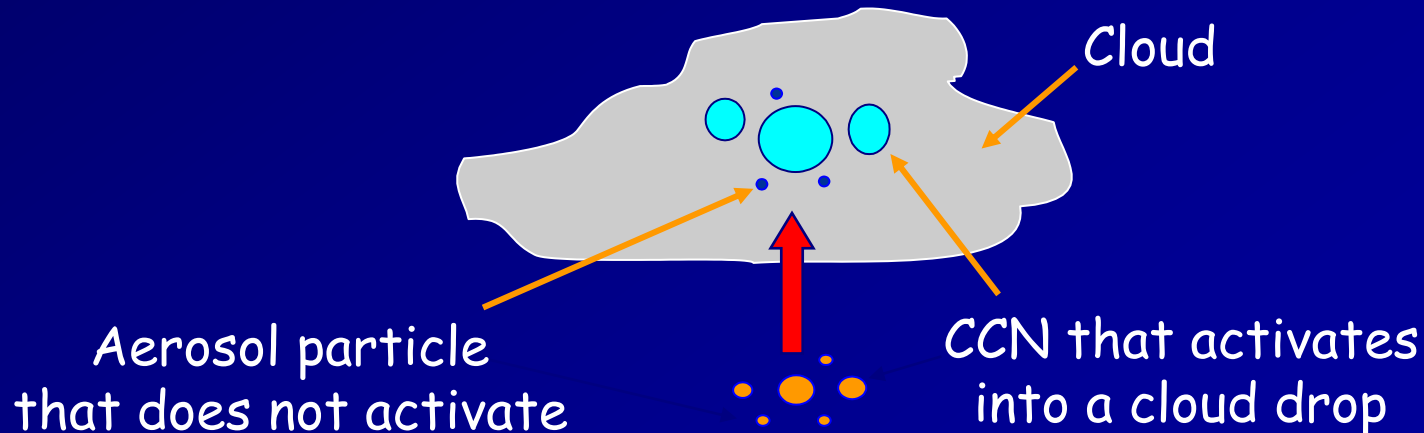
How do (liquid water) clouds form?

Clouds form in regions of the atmosphere where there is too much water vapor (it is "supersaturated").

This happens when air is cooled (primarily through expansion in updraft regions and radiative cooling).

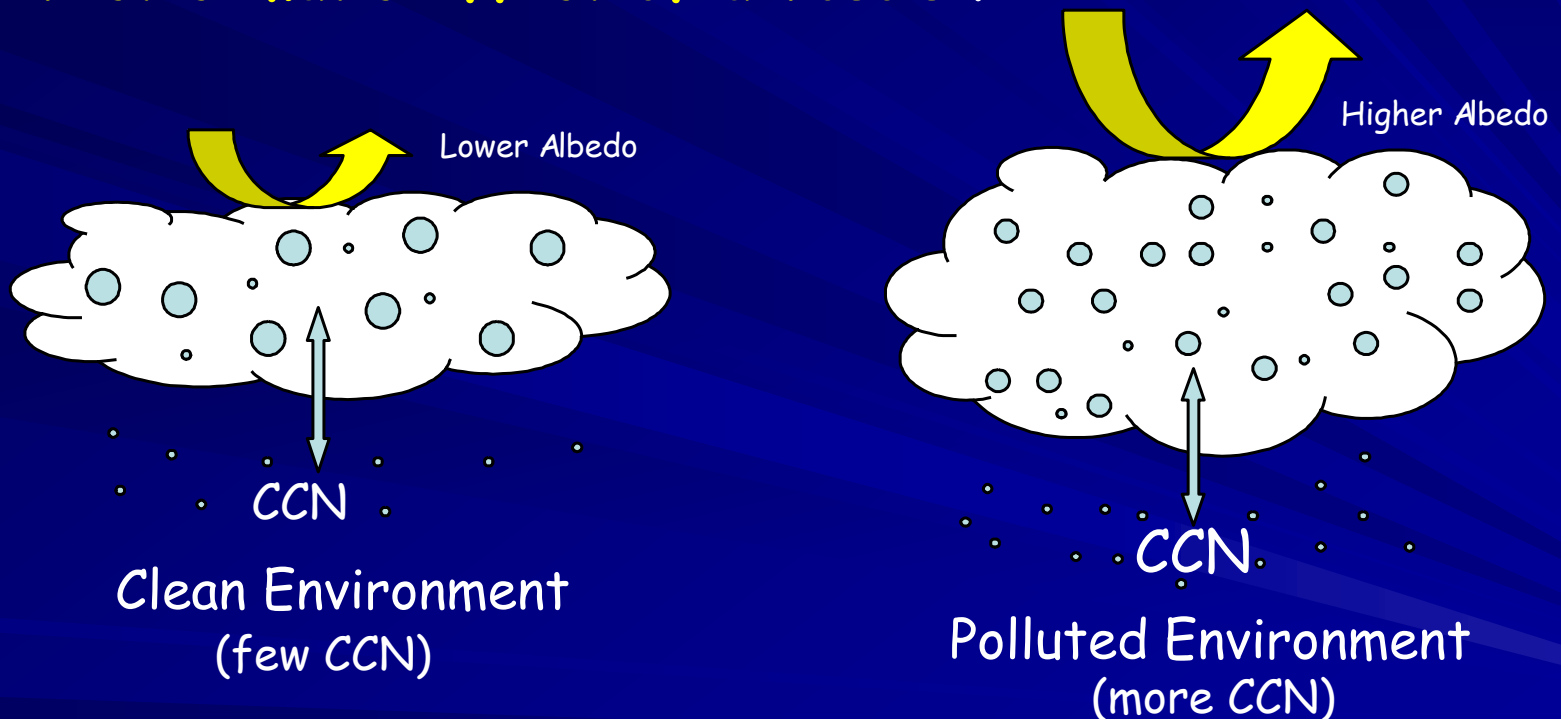
Cloud droplets nucleate on pre-existing particles found in the atmosphere (aerosols) with $\sim 0.1\mu\text{m}$ diameter.

Aerosols that can become droplets are called cloud condensation nuclei (CCN).



Increases in aerosol affects warm clouds

You make clouds that are "whiter", precipitate less (persist longer) and potentially cover larger areas of the globe. This is thought to yield a net cooling on climate and is termed as the **"indirect climatic effect of aerosols"**.

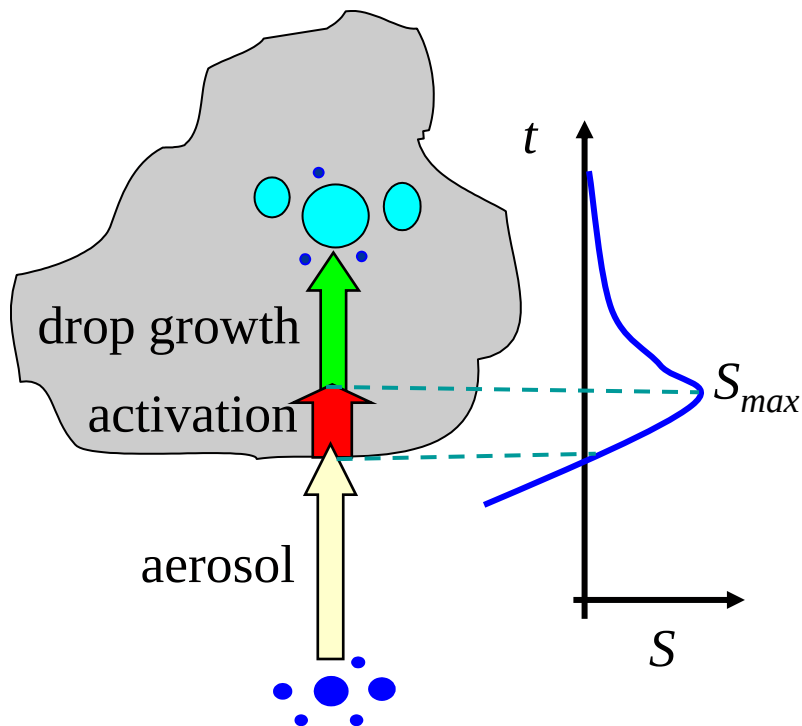


Increasing particles tends to cool climate (potentially alot).
Quantitative assessments done with climate models.

Droplet formation: The essence

Goal: Link cloud droplet concentration with precursor aerosol

Approach: Use the “simple story of cloud formation”.

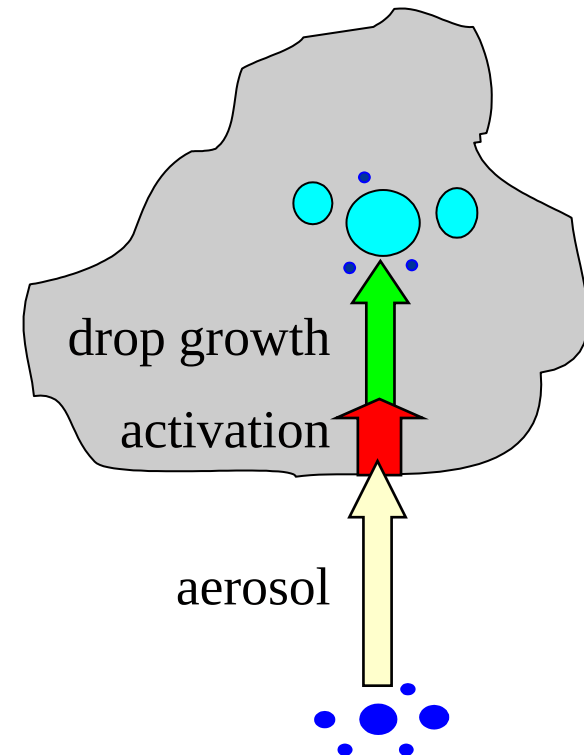
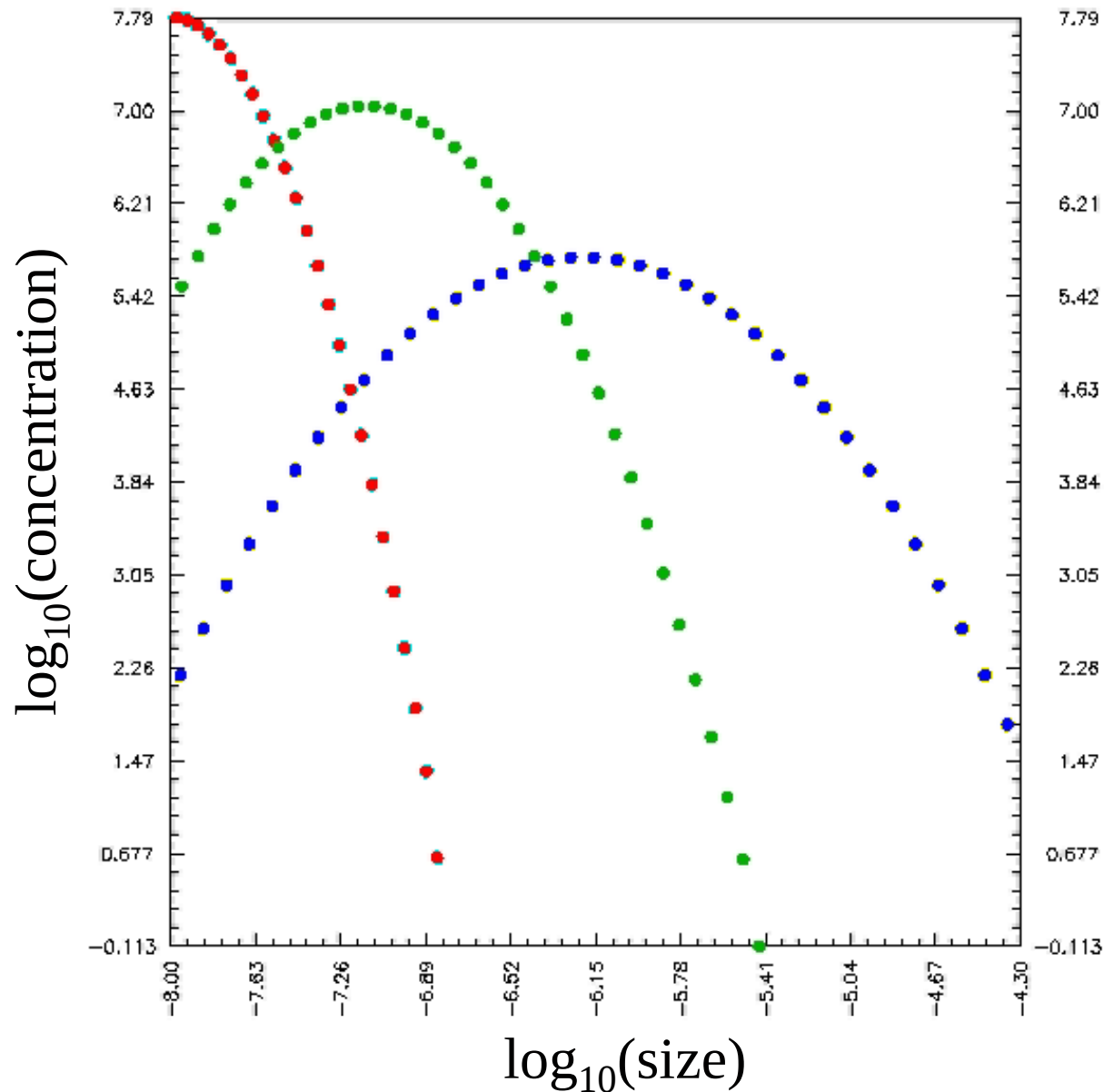


Conceptual steps are:

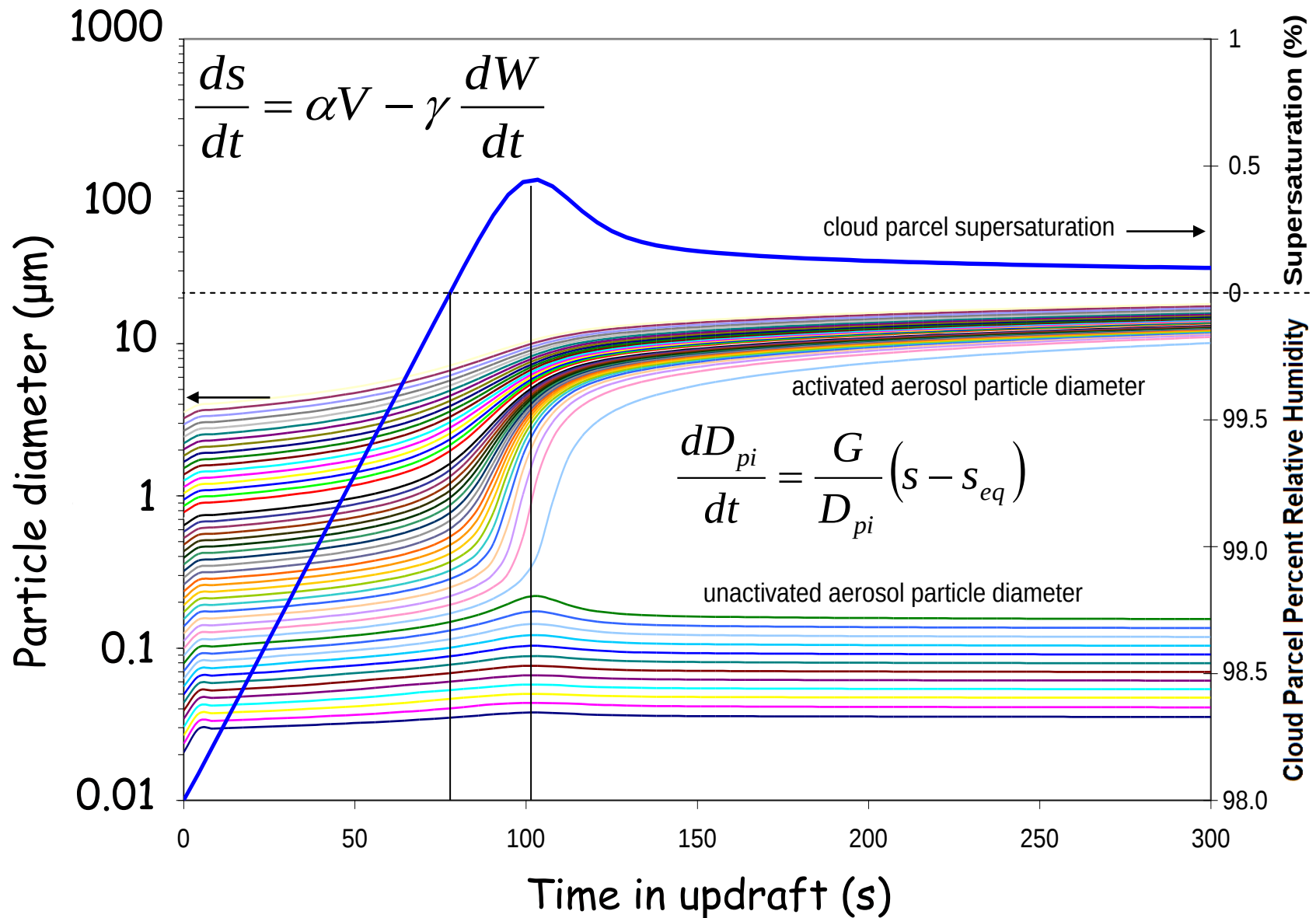
- Air parcel cools, exceeds dew point
- Water vapor is **supersaturated**
- Droplets start forming on existing CCN.
- Condensation of water on droplets becomes intense.
- Humidity reaches a **maximum**
- No more additional drops form

A “classical” nucleation/growth problem

Simulation of cloud droplet formation



Simulation of cloud droplet formation



Drop formation: coupled nonlinear system

Supersaturation (and s_{max}) depends strongly on the expansion rate (updraft velocity V) and the condensation rate of water:

$$\frac{ds}{dt} = \alpha V - \gamma \left(\frac{dW}{dt} \right) \quad \leftarrow \text{Water condensation rate on droplets}$$

Water condensation rate on droplets determined from the contribution of each aerosol "size class" i

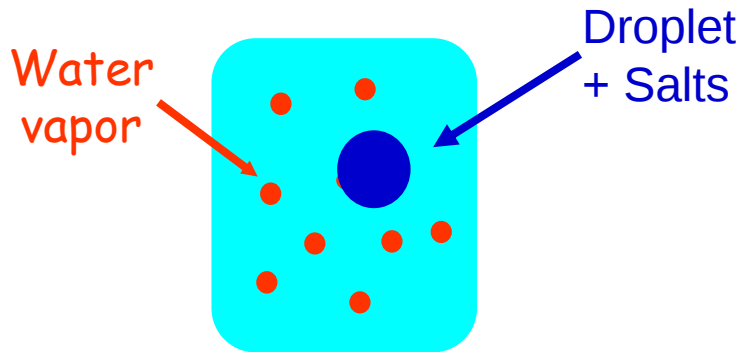
$$\frac{dD_{p,i}}{dt} = \frac{G_i}{D_{p,i}} \left(s - s_{eq,i} \right) \quad \begin{array}{l} \leftarrow \text{Mass transfer coefficient of water from the gas phase} \\ \leftarrow \text{Equilibrium saturation (relative humidity) of droplet} \end{array}$$

We need to know:

1. $S_{eq,i}$ for aerosol and droplets (**thermodynamics**)
2. G_i depends on the water vapor diffusivity (P, T) and the water vapor mass transfer coefficient (**kinetics**).

Thermodynamics 101: essentials

- Equilibrium vapor pressure of water over pure water



$$P_{H_2O(g)} = P_{sat}(T)$$

- Effects of dissolved solutes

Dissolved salts decrease the energy of your system (why?).

This reduces the equilibrium $P_{H_2O(g)} < P_{sat}(T)$

Mol fraction of water in solution $\frac{n_w}{n_w + n_{salts}}$

$$P_{H_2O(g)} = P_{sat}(T) x_{H_2O(l)}$$

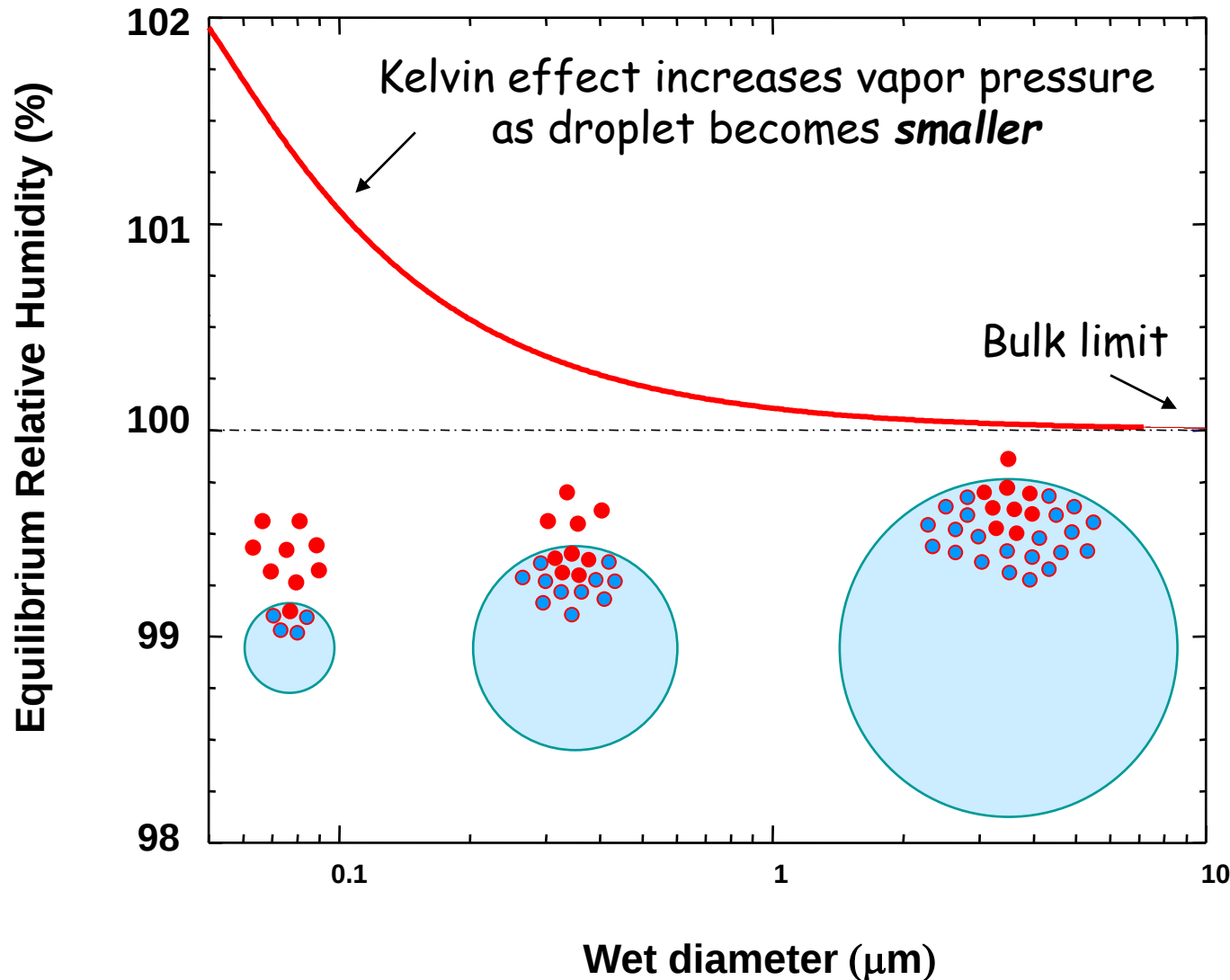
Raoult's Law (for ideal solutions)

Droplet thermo: special considerations

- Aerosol thermo mentioned before considered aerosol as a “bulk” system, where there is an infinite amount of each phase for interaction.
- “Bulk” thermodynamics thus assume that interfaces are “flat”.
- Sometimes this is not a good approximation.
- “Curvature” effects may need to be included in the thermodynamic expressions.
- Main parameters expression curvature effects:
 - Interfacial tension (“surface tension”)
 - Radius of curvature (most often, aerosol/drop radius)

Including curvature: Thermodynamics of droplets

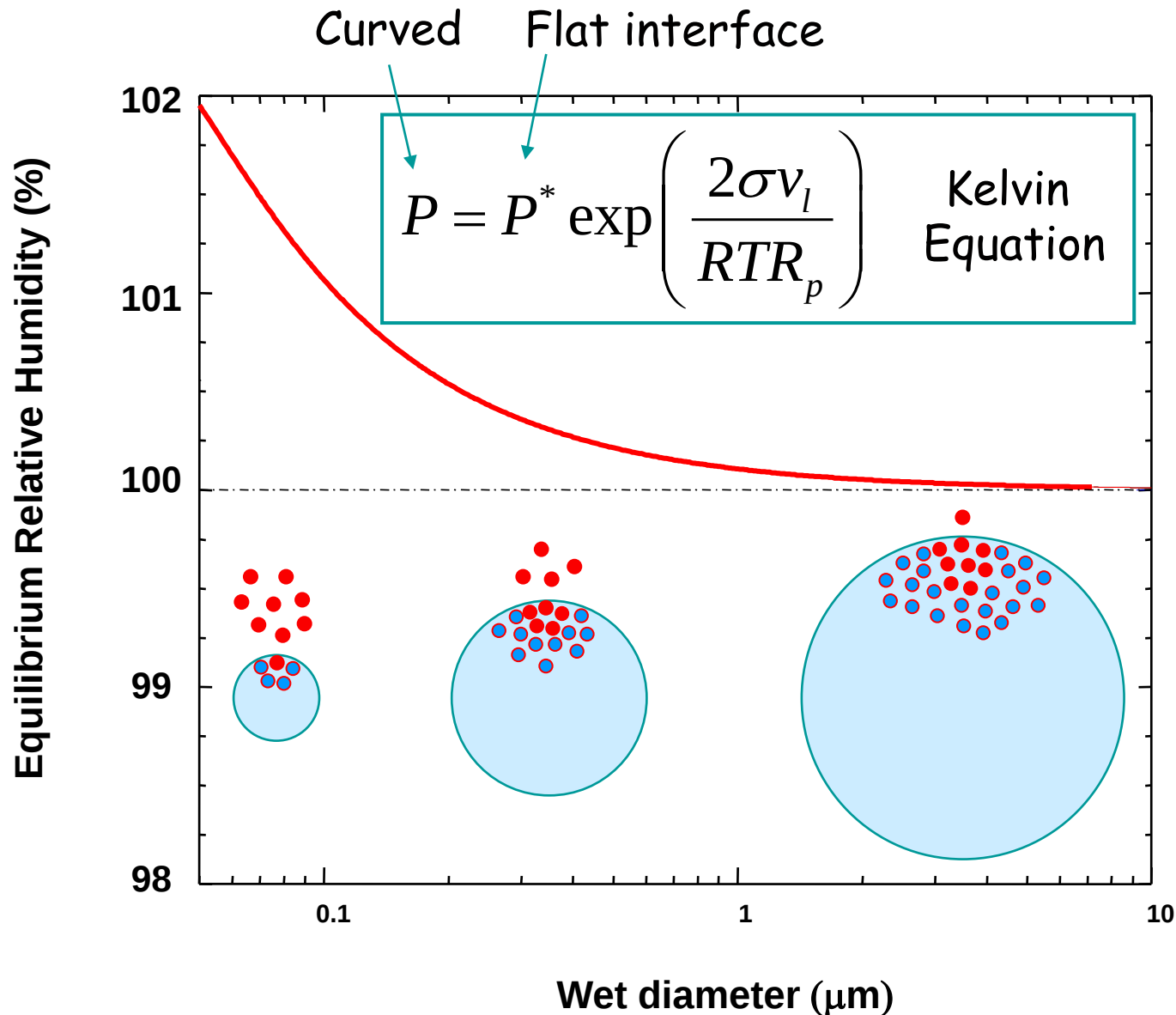
Impact of curvature: “the physical explanation”



As the droplet size decreases, its equilibrium vapor pressure increases (“Kelvin effect”)

Less molecules around in small drops to “keep” H_2O in the droplet phase

Including curvature: Thermodynamics of droplets



As the droplet size decreases, its equilibrium vapor pressure increases ("Kelvin effect")

Hugely important equation.

Needed for any nucleation process

Thermodynamics of droplets: Köhler equation

Apply Kelvin equation to a pure water droplet (i.e., σ_w and $v_l = \frac{M_w}{\rho_w}$)

$$P = P^* \exp\left(\frac{4M_w\sigma}{RT\rho_w D_p}\right)$$

Dissolved substances in the drop depress water vapor pressure. If $\sigma_w, v_l \sim \text{const.}$ then only P^* changes (given by Raoult's law: $P^* = P^{sat} \gamma_w x_w$)

$$\frac{P}{P^{sat}} = x_w \gamma_w \exp\left(\frac{4M_w\sigma}{RT\rho_w D_p}\right) \quad \text{Köhler Equation}$$

Equilibrium relative humidity of a particle when it has absorbed water and acquired a wet diameter, D_p

Thermodynamics of droplets: Köhler equation

One can then invoke simplifying assumptions:

$$x_w = \frac{n_w}{n_w + in_s} = 1 - \frac{in_s}{n_w + in_s} \sim 1 - \frac{in_s}{n_w} = 1 - \frac{in_s}{\frac{\pi}{6} D_p^3 \frac{\rho_w}{M_w}} = 1 - \frac{6 \frac{M_w}{\pi \rho_w} in_s}{D_p^3}$$

$$= 1 - \frac{B}{D_p^3} \quad \text{where} \quad B = \frac{6 M_w}{\pi \rho_w} in_s$$

Moles of solute in droplet
van't Hoff factor of solute in droplet

$$\gamma_w \sim 1 \quad \text{and} \quad \exp\left(\frac{4M_w\sigma}{RT\rho_w D_p}\right) \sim 1 + \frac{A}{D_p} \quad \text{where} \quad A = \frac{4M_w\sigma}{RT\rho_w}$$

Substitution into full Köhler equation, and considering leading terms:

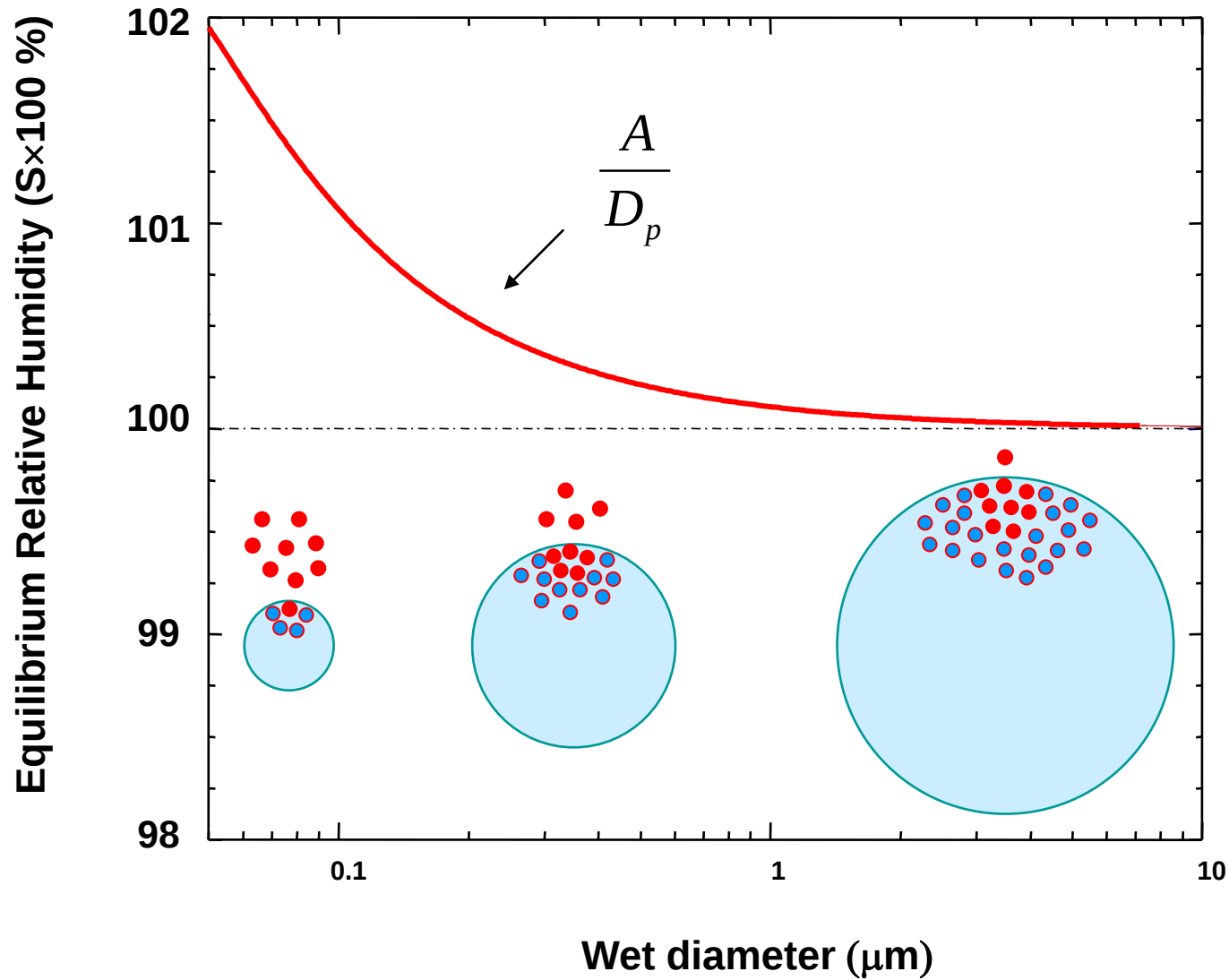
$$S = \frac{P}{P^{sat}} = 1 + \frac{A}{D_p} - \frac{B}{D_p^3}$$

Simplified Köhler equation

↑ Saturation ratio
↑ "Kelvin" term
↑ "Raoult" term

Thermodynamics of droplets: Köhler equation

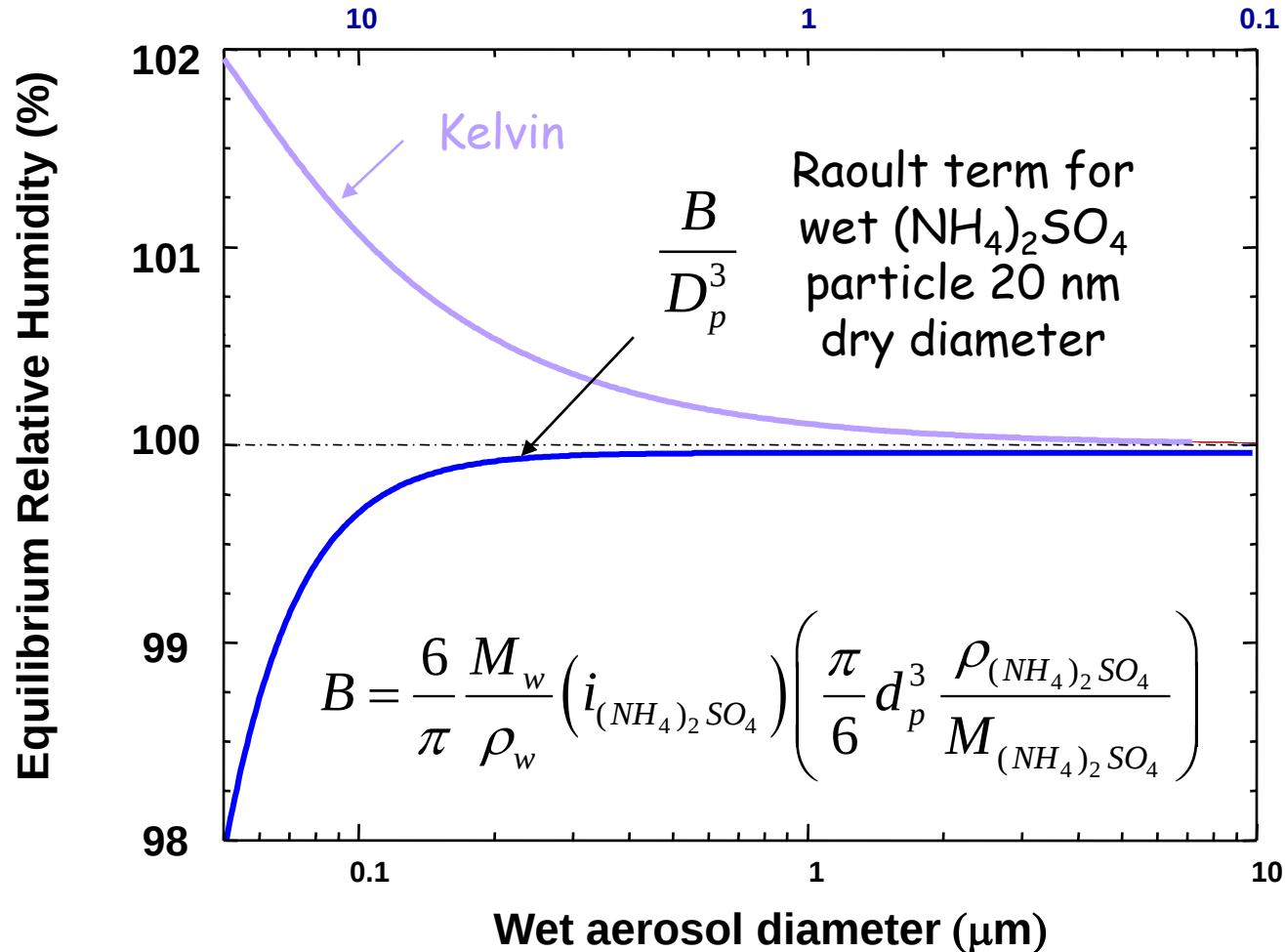
First plot the Kelvin term



Thermodynamics of droplets: Köhler equation

...then plot the Raoult term

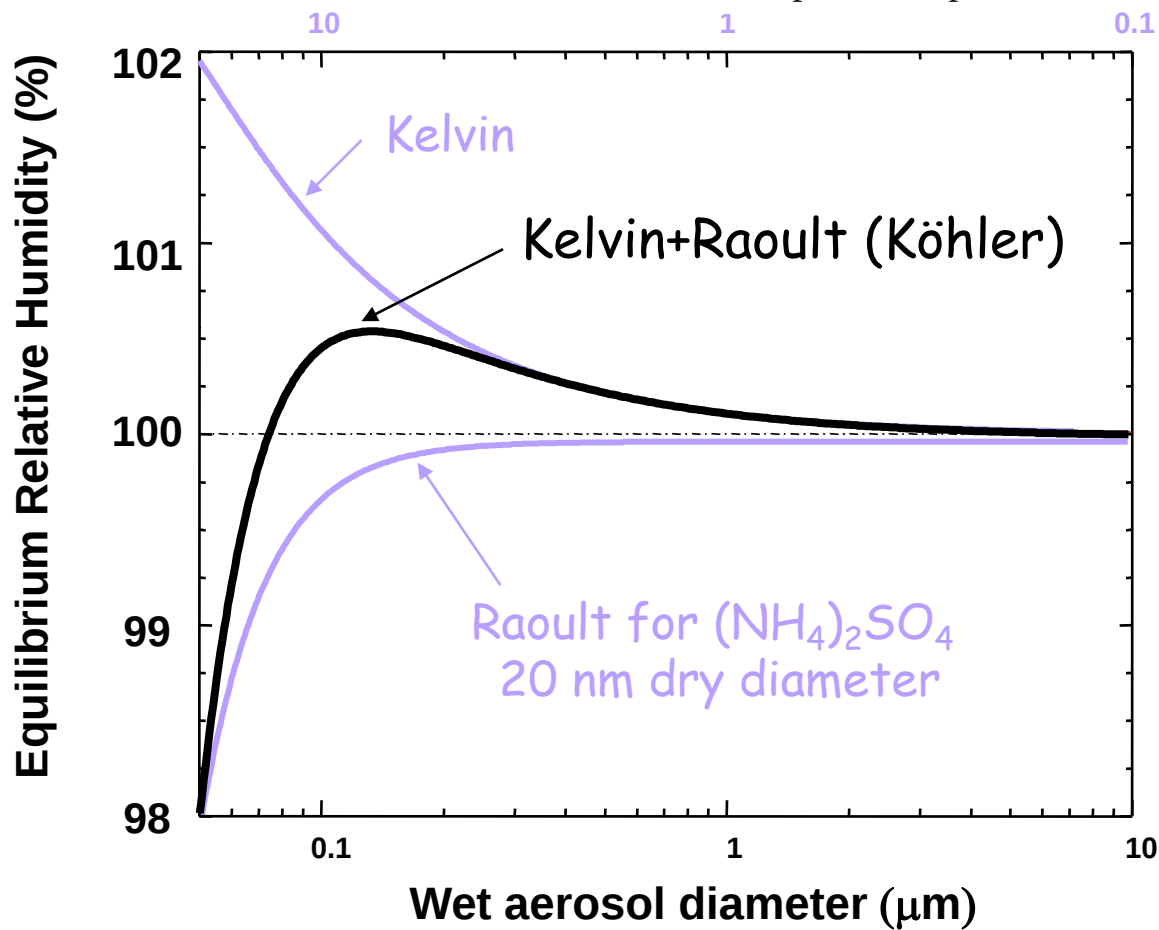
Solute Concentration (M)



Thermodynamics of droplets: Köhler equation

Both effects together: equilibrium vapor pressure of a wet aerosol.

$$S = \frac{P}{P^{sat}} = 1 + \frac{A}{D_p} - \frac{B}{D_p^3}$$



The combined Kelvin and Raoult effects is the simplified **Köhler equation**.

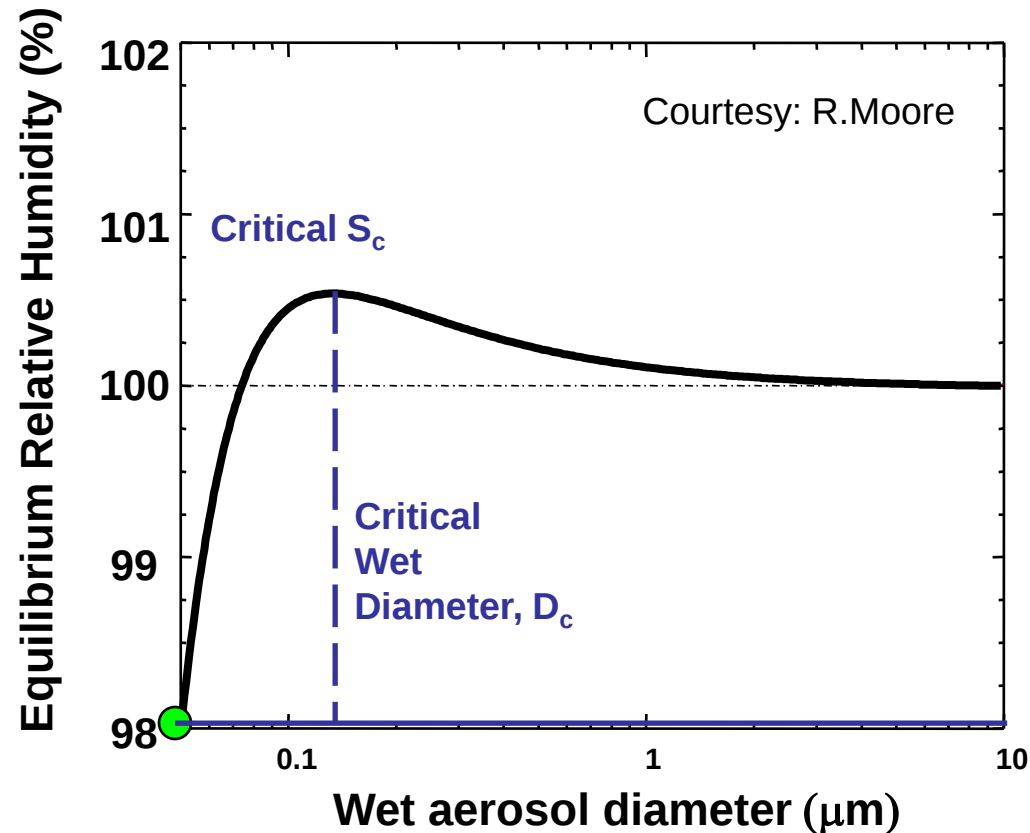
You can be in equilibrium **even if** you are above saturation.

Regions of stability/instability of ambient droplets

Dynamical behavior of an aerosol particle in a variable RH environment.

Wet Aerosol
(Haze)

Cloud Droplet

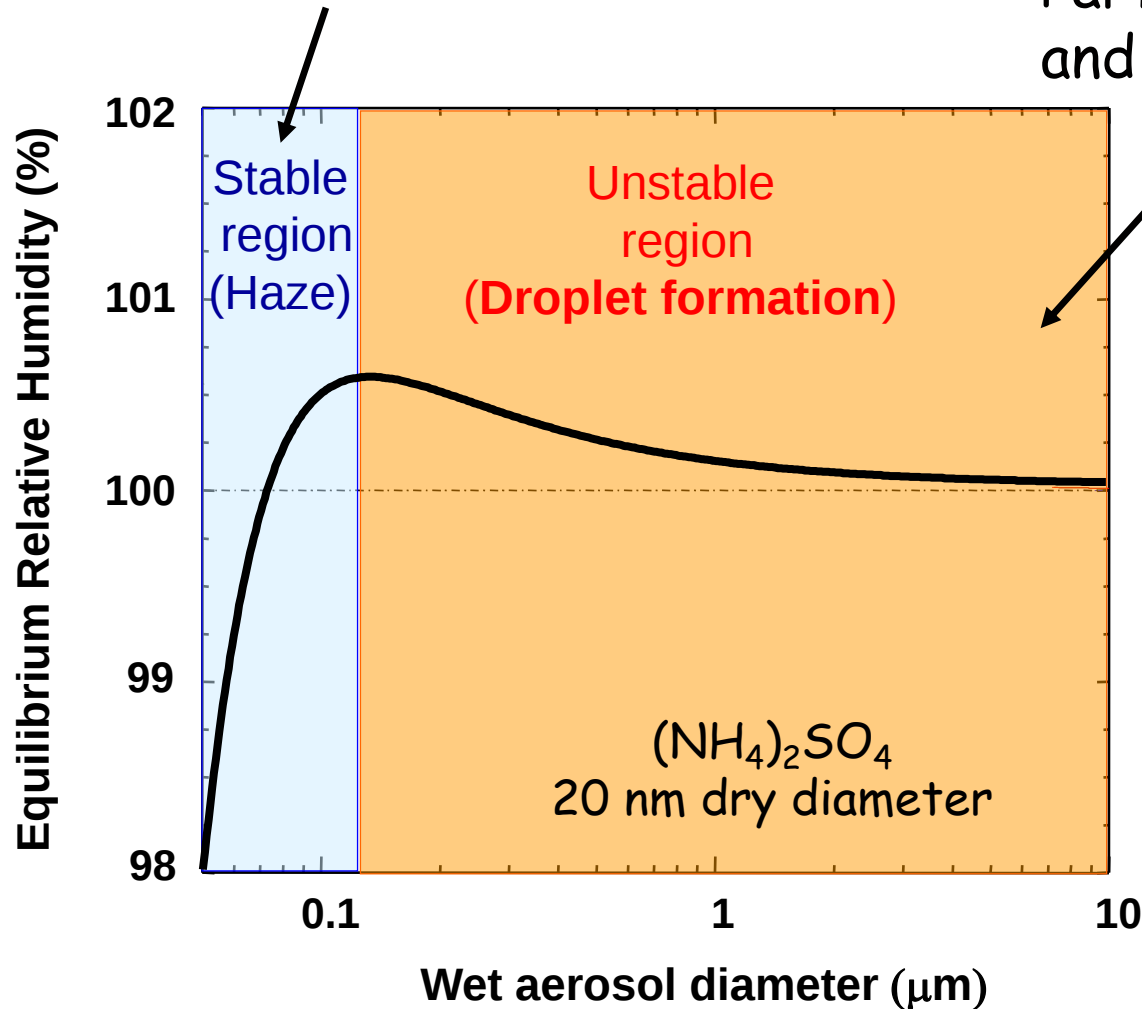


If ambient S exceeds the maximum, particles grow uncontrollably.
They are said then to act as Cloud Condensation Nuclei (CCN)

When does an aerosol particle act as a CCN ?

Ambient RH **less** than S_c ->
stable equilibrium.

Ambient RH **above** S_c ->
unstable equilibrium.
Particles act as CCN
and make droplets



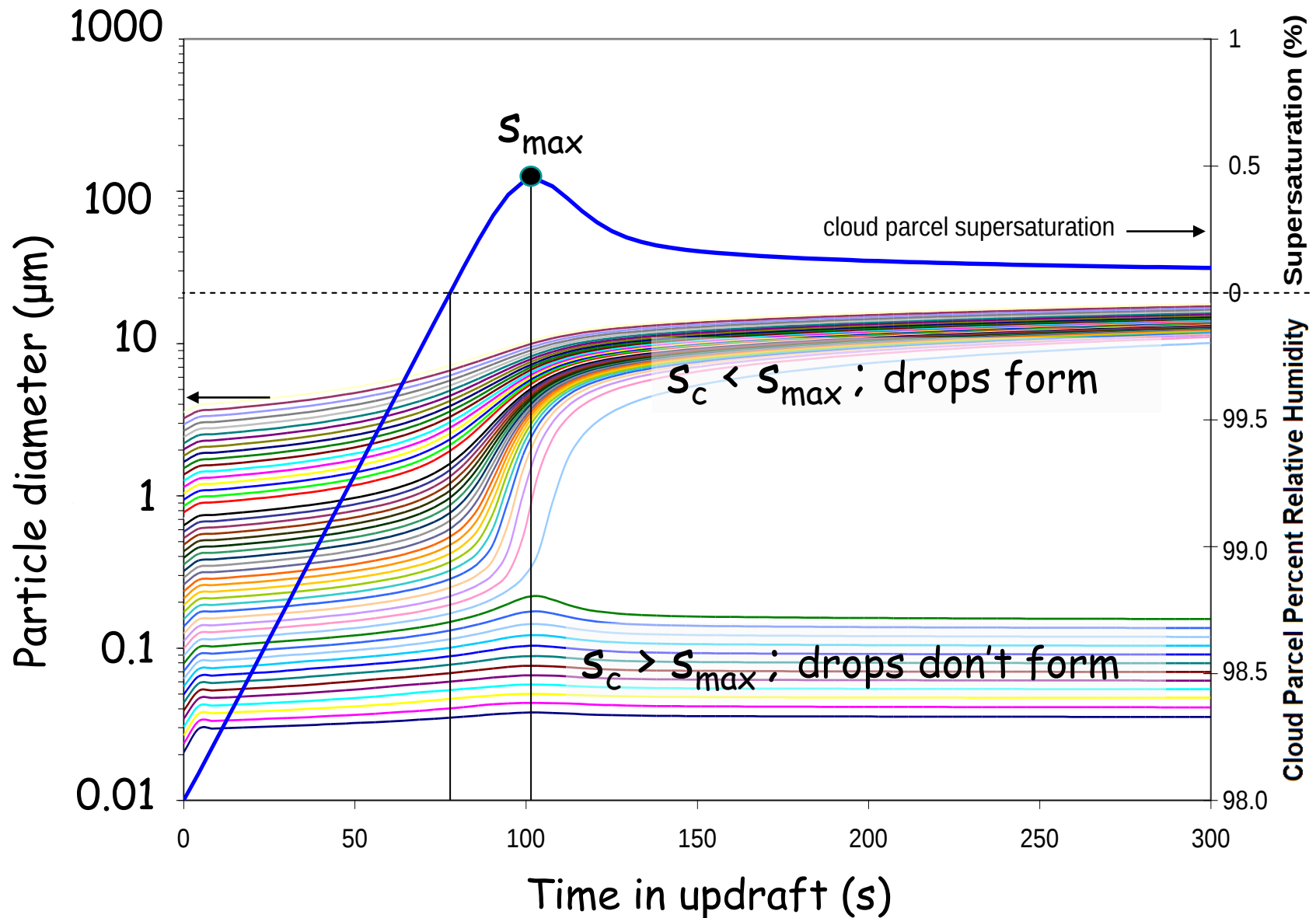
Köhler theory:

$$S_c = \left(\frac{4A^3}{27B} \right)^{1/2}$$

$$S_c \sim d_{\text{dry}}^{-3/2}, \epsilon_{\text{soluble}}^{-1/2}$$

Size is more
important than
composition

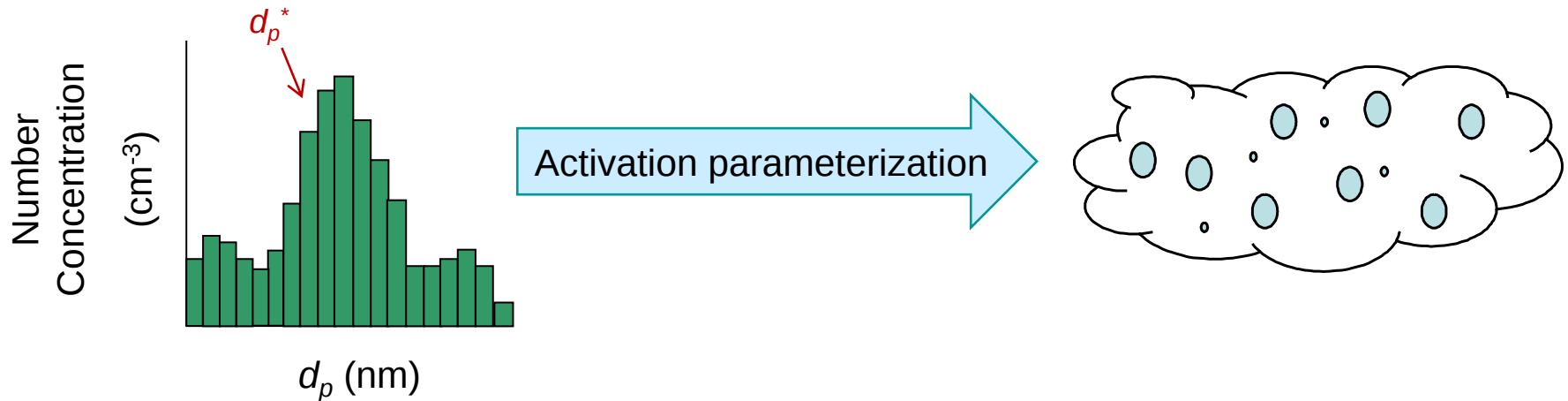
Now we understand droplet formation



Describing droplet formation in models...

Droplet calculation in models then is:

Calculated size distribution + κ + vertical velocity

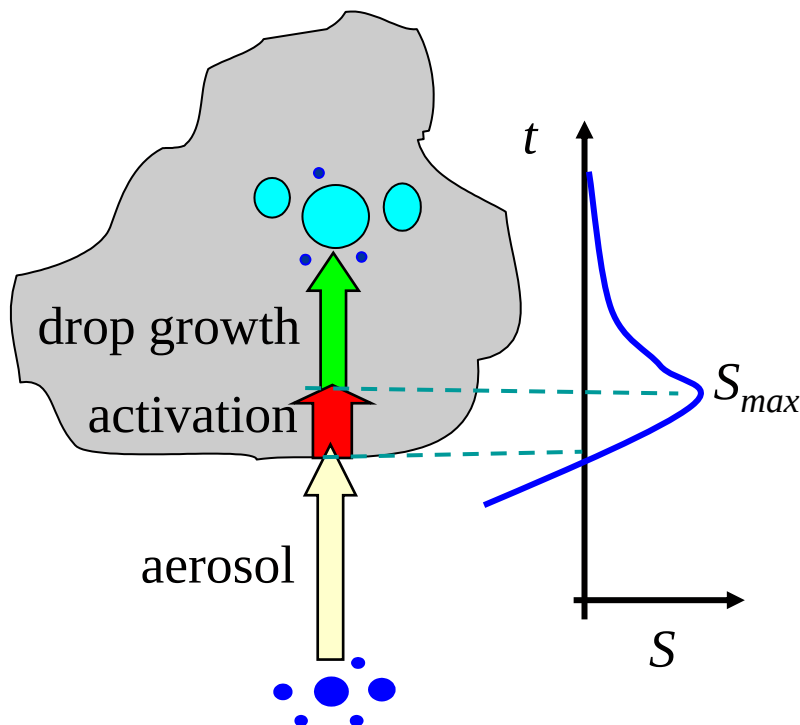


Activation parameterization is either a correlation or a solution to the parcel model equations that describe the activation process in clouds.

Droplet number needs CCN and max.cloud RH...

Algorithm for calculating N_d :
(Mechanistic parameterization)

1. Calculate S_{max} (approach-dependent)
2. N_d is equal to the CCN with $s_c \leq S_{max}$



Mechanistic Parameterizations:

Twomey (1959); Abdul-Razzak et al., (1998); Nenes and Seinfeld, (2003); Fountoukis and Nenes, (2005); Kumar et al. (2009), Morales and Nenes (2014), and others.

Input: P, T, vertical wind, particle size distribution, composition.

Output: Cloud properties (droplet number, size distribution).

Comprehensive review & intercomparison:

Ghan, et al., JAMES (2011); Morales and Nenes (2014)

Is this description of droplet formation real

Evaluate with in-situ data from airborne platforms



Observed Aerosol size
distribution & composition

Observed Cloud updraft
Velocity (PDF)

Predicted Drop Number
(Parameterization)

Compare

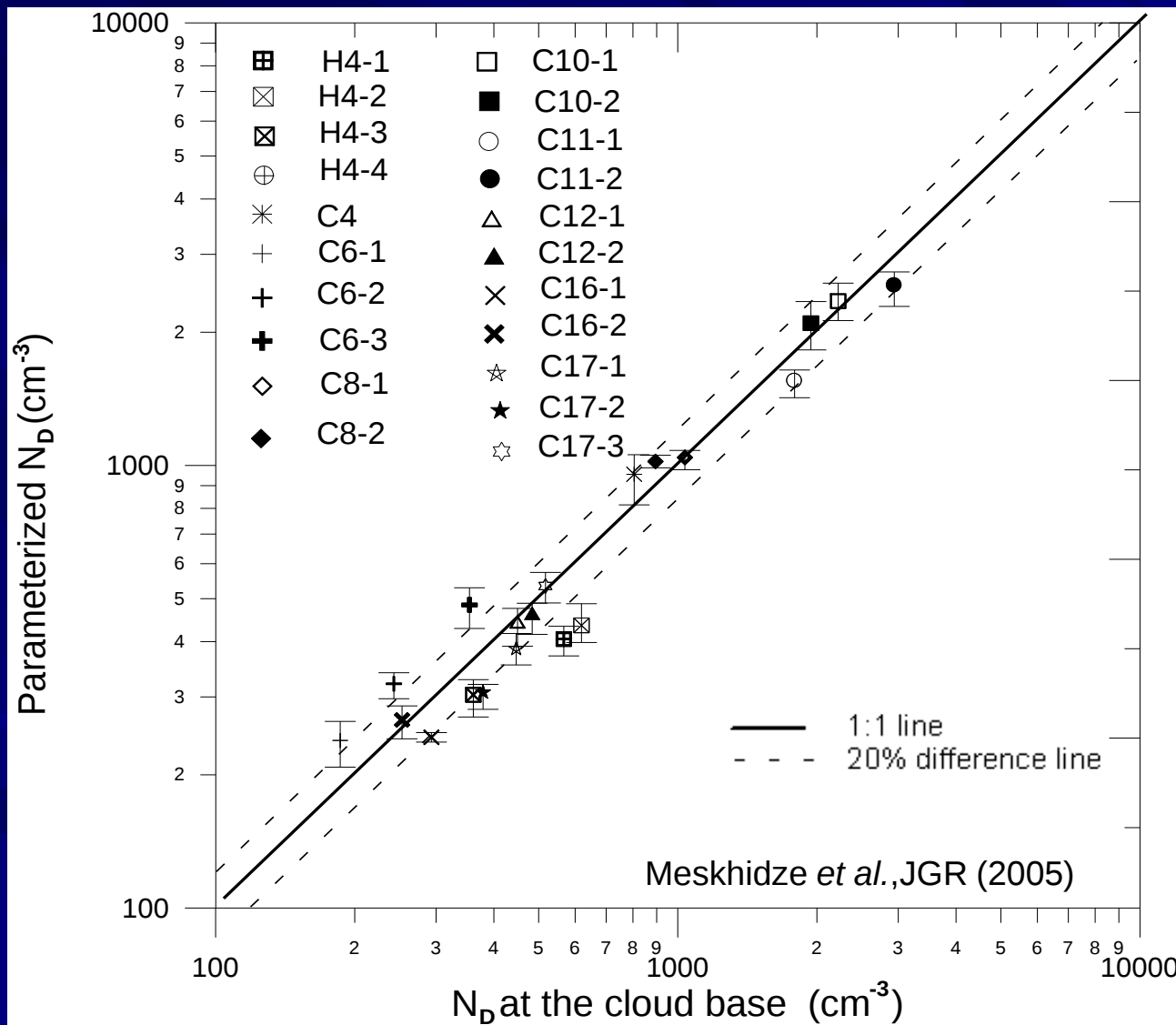
Observed Drop Number
Concentration



CRYSTAL-FACE (2002) Cumulus clouds



CIRPAS Twin Otter



Parameterized
agrees with
observed cloud
droplet number

Agreement to
within a few %
(on average)!

...when you know
everything
about aerosol
composition and
size

Issue: aerosols are complex



Primary emissions

Automobiles, industry, domestic, vegetation, forest fires, seasalt, ...

Secondary transformations

Oxidation of precursors (by O_3 , H_2O_2 , OH , NO_3 , etc.) generates organic compounds.

Reaction of volatile bases (NH_3) with acids, dust and seasalt form salts like $(NH_4)_2SO_4$.



Parameterizing “characteristic” CCN activity...

Petters and Kreidenweis (2007) expressed the Kohler theory parameter B in terms of a “hygroscopicity parameter”, κ

$$S_c = \left(\frac{4A^3}{27B} \right)^{1/2} = \left(\frac{4A^3}{27\kappa d^3} \right)^{1/2} \Rightarrow S_c = \left(\frac{4A^3}{27\kappa} \right)^{1/2} d^{-3/2}$$

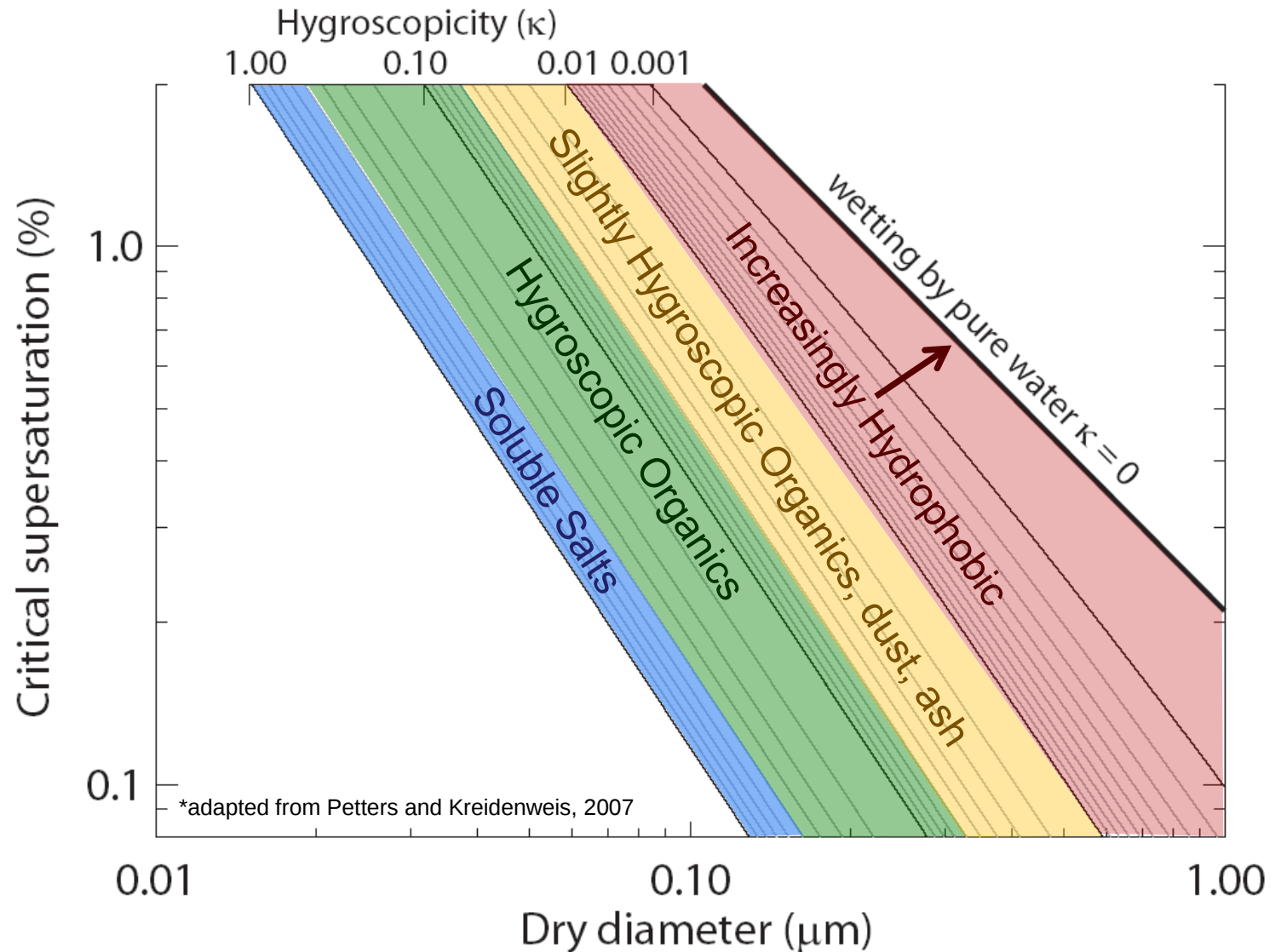
$\kappa \sim 1$ for seasalt, ~ 0.6 for $(\text{NH}_4)_2\text{SO}_4$, $\sim 0-0.3$ for organics

κ rarely exceeds 1 in atmospheric aerosol

Simple way to think of κ : the “equivalent” volume fraction of seasalt in the aerosol (the rest being insoluble).

$\kappa \sim 0.6 \Rightarrow$ particle behaves as 60% seasalt, 40% insoluble

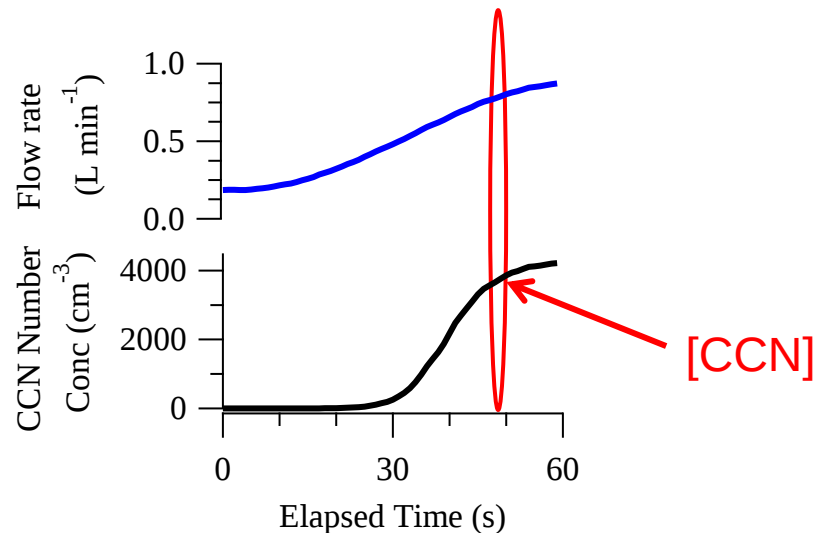
Hygroscopicity Space



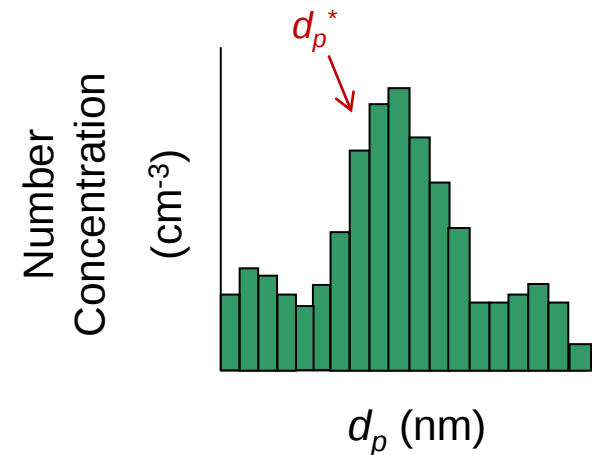
- κ is used to parameterize the activation of particles in the atmosphere

Getting κ from CCN Measurements

1. Measure CCN concentration, $[CCN]$, at a given s^*



2. Find where backwards integrated size distribution = $[CCN]$ to obtain the critical diameter, d_p^*



3. Calculate κ

$$\kappa \approx \frac{4A^3}{27d_p^3 s^{*2}} = \frac{M_w \rho_s}{\rho_w M_s} v \varepsilon_s$$

Measuring CCN: Basic Principles

Operation: Expose particle sample to a known water vapor supersaturation, and measure those that become droplets.

Desired range: 0.01% - 1.0% supersaturation (S)

Main challenges:

Generating a highly controlled level of supersaturation

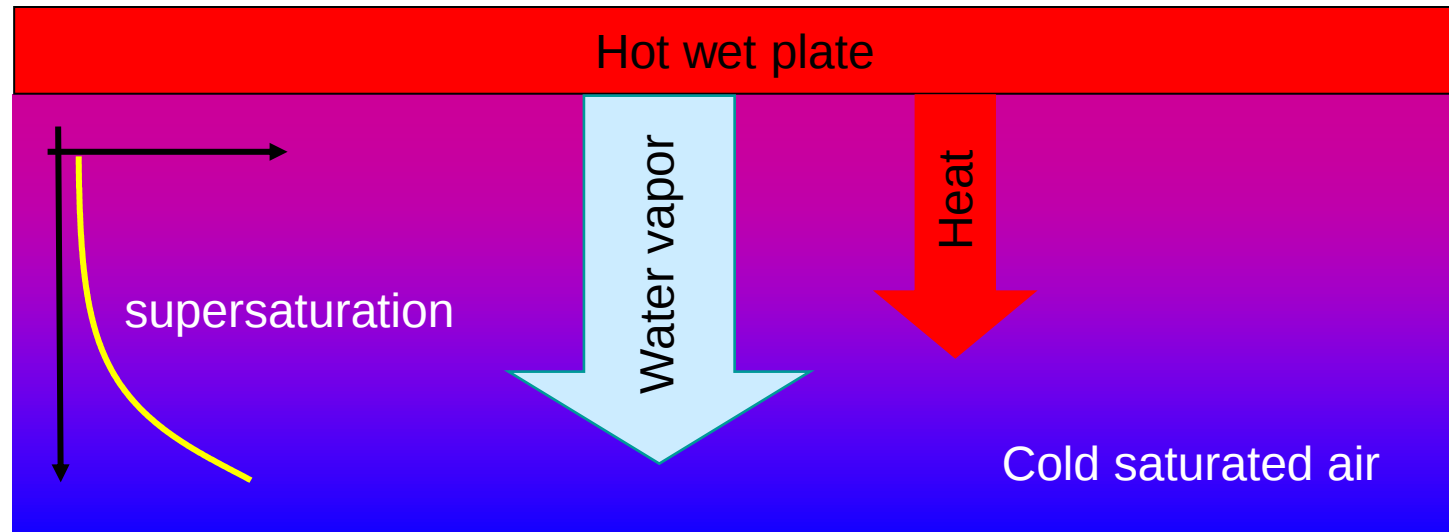
- *Fluctuations in supersaturation are always a problem*
- *This is more difficult for low supersaturations*

Droplet detection

- *Need to differentiate from particles that do not become droplets*
- *Generally more difficult for low supersaturations*

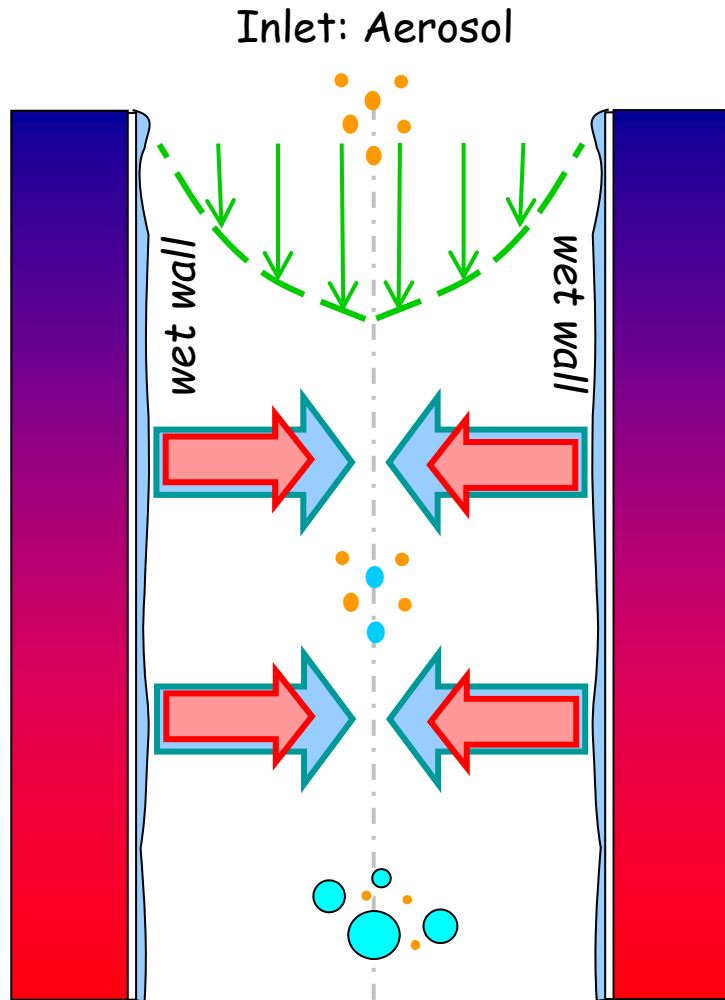
CCN Instruments: Generating Supersaturation

Relative Diffusion Instruments: Exploiting the difference in diffusivity between heat and water vapor.



- Generate a condition where saturated water vapor diffuses into a (saturated) colder region.
- *Supersaturation* develops in the "cold" region (why? Hint: H_2O molar mass 18, air molar mass ~ 29).

Continuous-Flow Streamwise Thermal Gradient Chamber



Outlet: [Droplets] = [CCN]

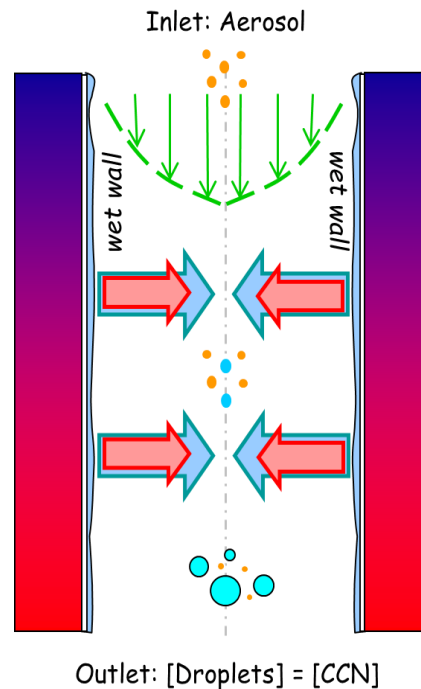
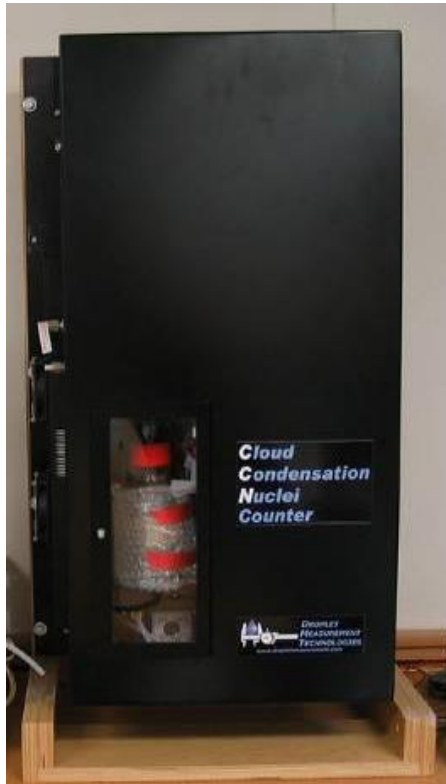
- Standard CCN measurement (>120 instruments in operation).
- Metal cylinder with wetted walls
- Streamwise Temp. Gradient
- Water diffuses faster than heat
- Supersaturation, S , generated at the centerline = f (Flowrate, Pressure, and Temp. Gradient)
- Operated as a *spectrometer* using Scanning Flow CCN Analysis (Moore and Nenes, 2009)

Roberts and Nenes (2005), US Patent 7,656,510

Lance et al., (2006), Lathem and Nenes (2011),
Raatikainen et al. (2012, 2013)

In-situ (and remote sensing data) key for studying the aerosol-CCN (microphysical) link

The CFSTGC or "DMT CCN Counter"



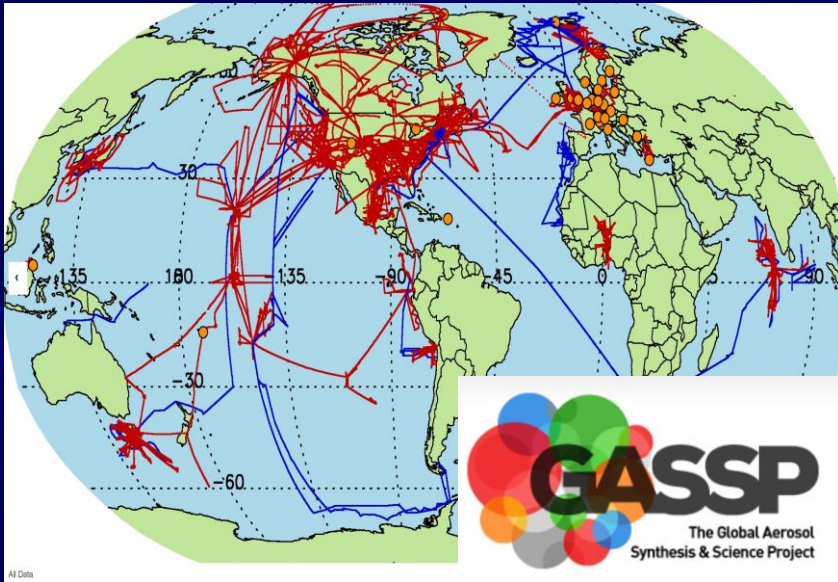
- Metal cylinder with wetted walls
- Streamwise Temp. Gradient
- Water diffuses faster than heat to centerline of flowtube
- Supersaturation, S , generated at the centerline = f (Flowrate, Pressure, and Temp. Gradient)

Roberts and Nenes (2005), US Patent 7,656,510

Used to test theory and develop a climatology over 15 years...

The community has obtained a CCN climatology over the 15 years

Some locations sampled ...



Measured:

CCN, Aerosol concentrations and size distributions, and aerosol chemistry

Cloud hydrometeor distributions (liquid/ice) and dynamics.

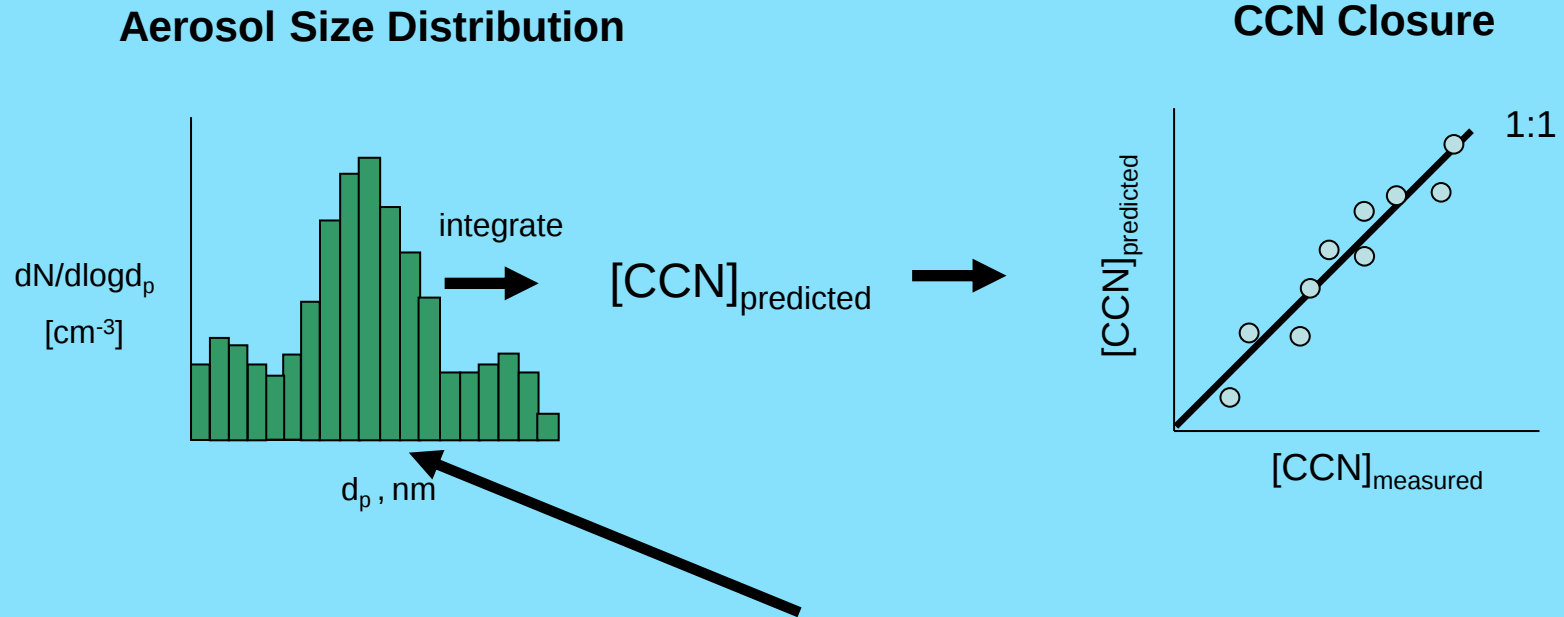
Environments:

Arctic, urban pollution, biomass burning, marine aerosol, hurricanes, oil spills, the tropics....



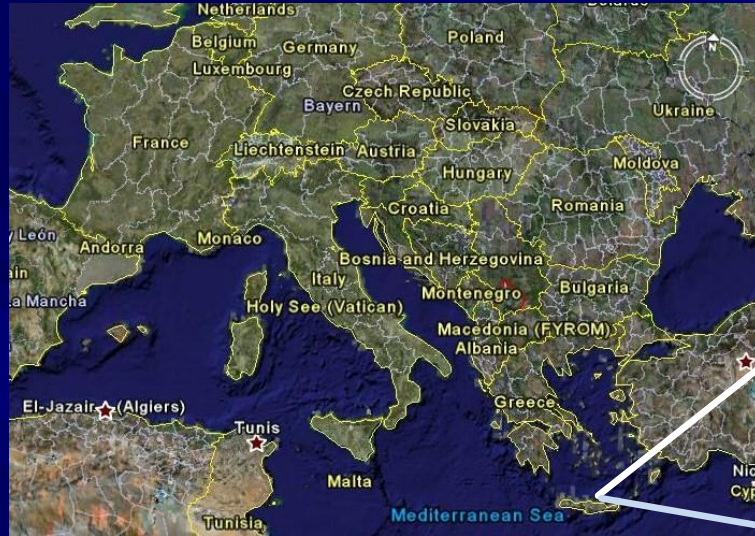
Testing CCN activation theory: CCN "Closure" studies

Compare measurements of CCN to predictions using Köhler activation theory and κ description



Use theory to predict the particles that can act as CCN based on measured chemical composition and CCN instrument supersaturation.

(The first) Finokalia Aerosol Measurement Campaign (FAME-07) - Summer 2007



DMT CCN counter
*Supersaturation
range: 0.2-1.0%*

TSI 3080 SMPS
Size: 20-460 nm

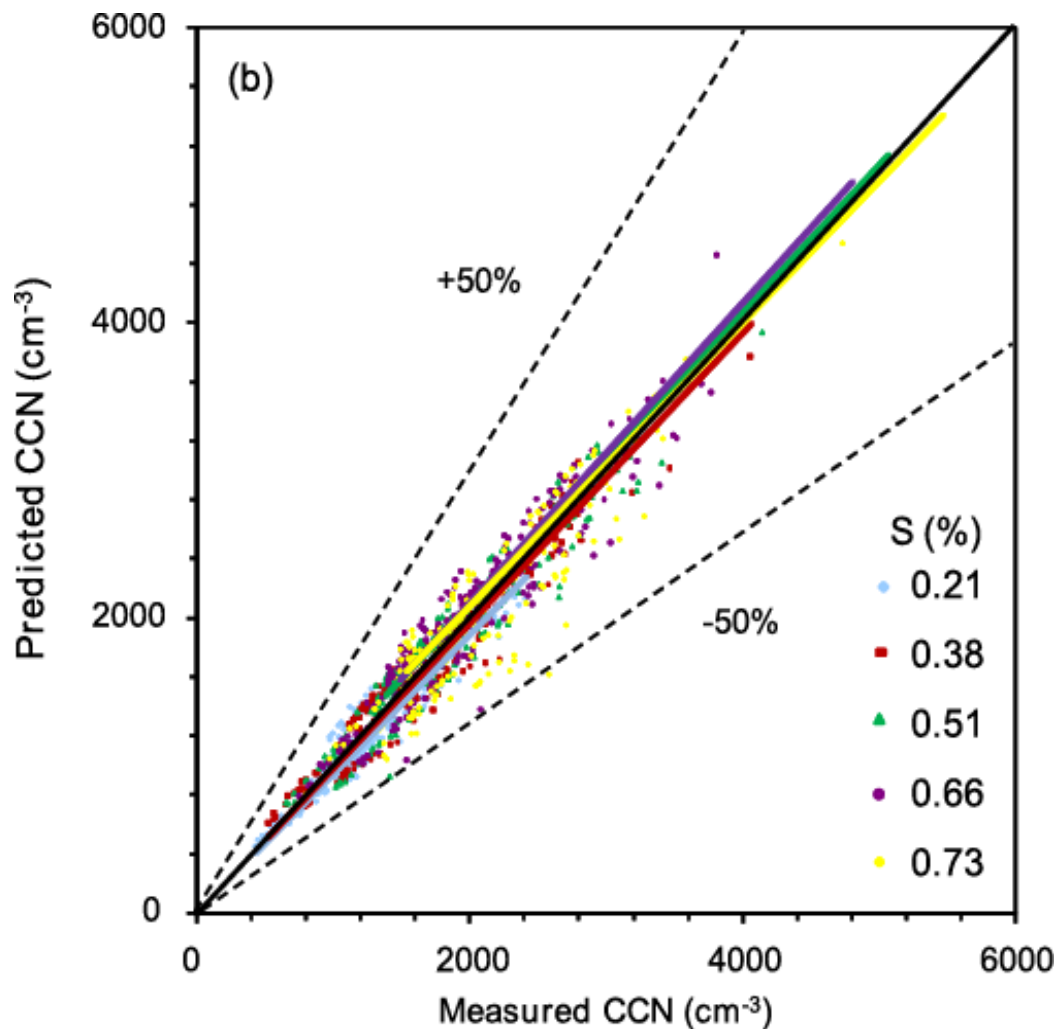
Low-vol impactor
*Ionic composition
measured via IC*

*WSOC/EC/OC also
measured*



(Bougiatioti et al., ACP, 2009)

Finokalia Aerosol Measurement Campaign (FAME-07) - CCN closure



Organics: very oxidized (MO-OOA) $\kappa \sim 0.15$

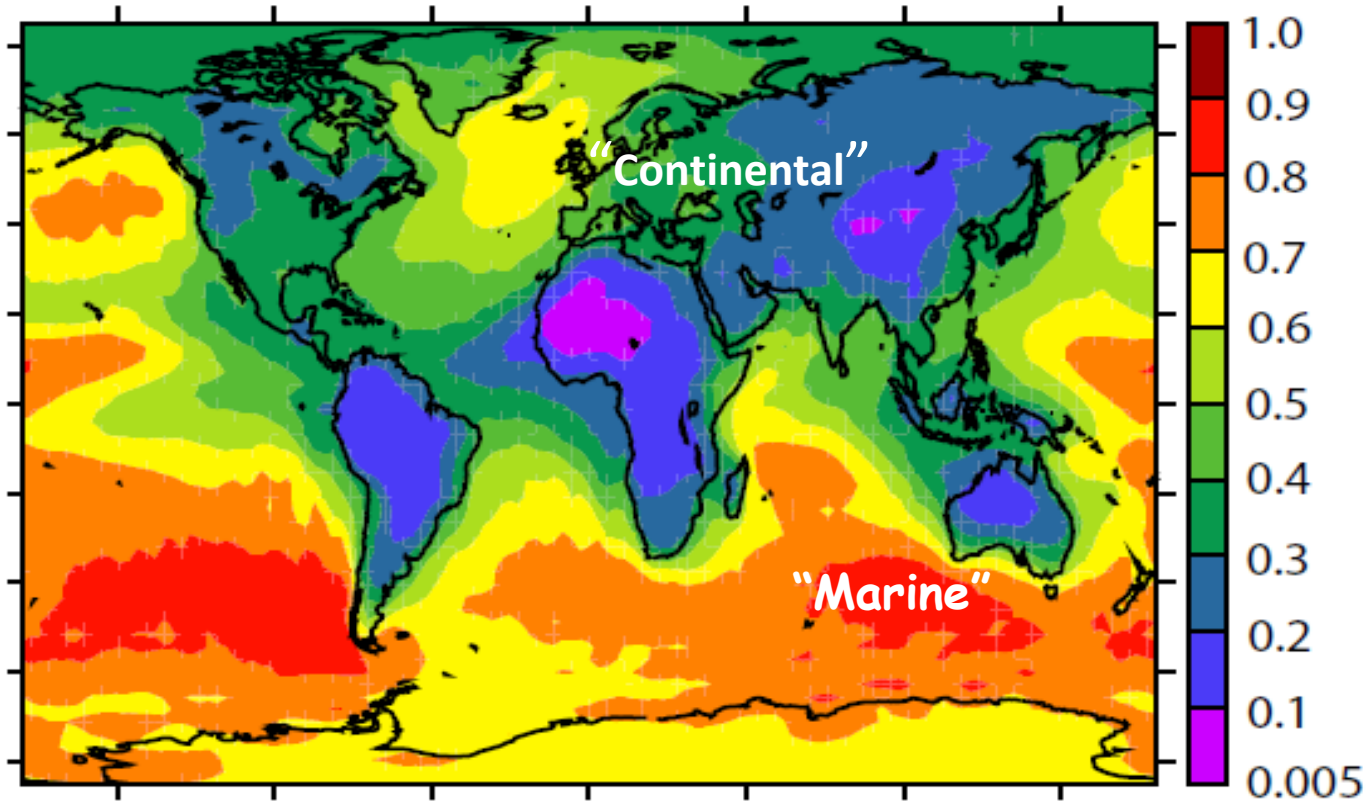
Excellent CCN closure (error $\sim 2\%$).

Köhler (CCN) theory *really* works.

Simple treatments of hygroscopicity in general work "well enough" for CCN/CDNC predictions

(Bougiatioti et al., ACP, 2009)

Global “average” distribution for κ



Pringle et al., ACP, (2010)

Fig. 2. Annual mean distribution of κ at the altitude of the planetary boundary layer.