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5	Theoretical analysis of mixing in liquid clouds. Part IV: DSD evolution
6	and mixing diagrams
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8	Mark Pinsky, and Alexander Khain
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10	Department of Atmospheric Sciences, The Hebrew University of Jerusalem, Israel
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23	Communicating author: Alexander Khain, The Hebrew University of Jerusalem,
24	khain@vms.huji.ac.il
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28 Abstract

Evolution of droplet size distribution (DSD) due to mixing between cloudy and dry 29 volumes is investigated for different values of the cloud fraction and for different initial DSD 30 shapes. The analysis is performed using a diffusion-evaporation model which describes time-31 dependent processes of turbulent diffusion and droplet evaporation within a mixing volume. 32 Time evolution of the DSD characteristics such as droplet concentration, LWC and mean 33 volume radii is analyzed. The mixing diagrams are plotted for the final mixing stages. It is 34 shown that the difference between the mixing diagrams for homogeneous and inhomogeneous 35 36 mixing is insignificant and decreases with an increase in the DSD width. The dependencies of normalized cube of the mean volume radius on the cloud fraction were compared with those 37 on normalized droplet concentration and found to be quite different. In case the normalized 38 droplet concentration is used, mixing diagrams do not show any significant dependence on 39 relative humidity in the dry volume. 40

The main conclusion of the study is that traditional mixing diagrams cannot serve as areliable tool for analysis of mixing type.

43

44 **Keywords:** turbulent mixing, droplet evaporation, DSD evolution, mixing diagram

46 **1. Introduction**

The effects of mixing of cloudy air with surrounding dry air on cloud microphysics are still the focus of many studies (see overview by Devenish et al., 2012). Processes of mixing are investigated in observations (Yum et al., 2015; Bera at al., 2016a,b), Large Eddy Simulations (Andrejczuk et al., 2009; Khain et al., 2017) and Direct Numerical Simulations (Kumar et al., 2014, 2017). Processes of mixing and their effects on droplet size distributions were recently investigated in a set of theoretical studies (Yang et al., 2016; Korolev et al., 2016 (hereafter, Pt1); Pinsky et al., 2016 a,b).

The Pt1 presented analysis of conventional (classical) concept of mixing and introduced the 54 55 main parameters characterizing homogeneous and extremely inhomogeneous mixing. In the 56 classical concept two volumes, cloudy and droplet free one, mix within an unmovable adiabatic mixing volume. At a monodisperse initial droplet size distribution (DSD), homogeneous mixing 57 leads to a decrease in droplet size and droplet mass content, while the number of droplets 58 remains unchanged. Extremely inhomogeneous mixing is characterized by decreasing the 59 number of droplets due to full evaporation of some fraction of droplets penetrating the initially 60 dry air volume while the DSD shape in the cloud volume remains unchanged. As a result of 61 extremely inhomogeneous mixing, droplet number decreases while the mean volume radii 62 63 remain unchanged. At a polydisperse DSDs, the extreme homogeneous mixing is characterized by proportional changes in DSD for all droplet radii (Pt1). Since widely used mixing diagrams 64 describe the final equilibrium stage of mixing within the mixing volume they do not contain 65 information about changes in microphysical quantities in the course of mixing. 66

Pinsky et al. (2016a, hereafter Pt2) analyzed the time evolution of initially monodisperse and polydisperse DSD during homogeneous mixing. It was shown that result of mixing strongly depends on the shape of the initial DSD. At a wide DSD, evaporation of droplets (first of all, of the smallest ones) is not accompanied by a decrease in the mean volume or effective radius. Moreover, the values of the radii may even increase over time. This result indicates that the widely used criterion of separation of mixing types based on the behavior of
the mean volume radius during mixing is not generally relevant and may be wrong in
application to real clouds.

Pinsky et.al. (2016b, hereafter Pt3) introduced a diffusion-evaporation model which 75 describes evolution DSDs and all the microphysical variables due to two simultaneously 76 occurring processes: turbulent diffusion and droplet evaporation. Mixing between two equal 77 volumes of subsaturated and cloudy air was analyzed, i.e. it was assumed that the cloud 78 volume fraction $\mu = 1/2$. The initial DSD in the cloudy volume was assumed monodisperse. 79 These simplified assumptions allowed to reduce the turbulent mixing equations to two-80 parametric ones. The first parameter is the Damkölher number, Da, which is the ratio of the 81 characteristic mixing time to the characteristic phase relaxation time. The second parameter is 82 the potential evaporation parameter R characterizing the ratio between the amount of water 83 vapor needed to saturate the initially dry volume and the amount of available liquid water in 84 the cloudy volume. 85

Within the Da - R space, in addition to the two extreme mixing types defined in the 86 classical concept, two more mixing regimes were distinguished, namely, intermediate and 87 inhomogeneous mixing. It was shown that any type of mixing leads to formation of a tail of 88 89 small droplets, i.e. to DSD broadening. It was also shown that the relative humidity in the initially dry volume rapidly increases due to both water vapor diffusion and evaporation of 90 penetrating droplets. As a result, the mean volume and effectice radii in the initially dry 91 volume rapidly approach the values typical of cloudy volume. At the same time, the liquid 92 93 water content (LWC) remains significantly lower than that in the cloudy volume during much longer time than required for the effective droplet radius to grow. 94

In the present study (Pt4) we continue investigating the turbulent mixing between an initially droplet free volume (referred to as dry volume) and a cloudy volume. The focus of the study is investigation of DSD temporal evolution and analysis of the final equilibrium 98 DSD. In comparison to Pt3, the problem analyzed in this study is more sophisticated in99 several aspects:

The dependences of different mixing characteristics on cloud volume fraction 0≤ µ≤1
are analyzed. In this case the equations of turbulent mixing cannot be reduced to the twoparametric problem as it was done in Pt3.

103 • The initial DSDs in cloud volume are polydisperse. We use both narrow and wide 104 initial DSD described by Gamma distributions with different sets of parameters. The DSD are the same as those used in Pt2. Mechanisms of formation of wide DSDs in clouds including 105 DSDs in undiluted cloud cores were investigated in several studies [e.g., Khain et al., 2000; 106 Pinky and Khain, 2002; Segal et al., 2004; Prabha et al., 2011]. These studies show the DSD 107 broadening is caused by in-cloud nucleation of droplets within clouds as well as by collisions 108 109 between cloud droplets. It was shown that DSDs in adiabatic volumes can be wide and first raindrops or drizzle drop arise namely in non-diluted adiabatic cloud parcels [Khain et al., 110 2013; Magaritz-Ronen et al., 2016]. We use both narrow and wide DSDs in the form of 111 112 Gamma distribution with typical parameters used in different cloud resolving models. The 113 DSDs that are used as initial ones in cloudy volumes could be formed also under influence of mixing during their previous history. The mechanisms of the formation of initial DSD are not 114 115 of interest in the study since that do not affect the analysis.

The equation for supersaturation, used in this study, is valid at low humidity in the
initially dry volume and is more general and compared with that used in Pt3, which makes the
DSD calculations more accurate.

At the same time, some simplifications used in Pt3 are retained in this study. The vertical movement of the entire mixing volume is neglected; collisions between droplets and droplet sedimentation are not allowed. Also, we consider a 1D diffusion-evaporation problem. We neglect the changes of temperature in the course of mixing, which is possibly a less significant simplification. All these simplifications allow to reveal the effects of turbulent mixing andevaporation on DSD evolution.

125

126 **2.** Formulation of the problem and model design

In this study, the process of mixing is investigated basing on the solution of 1D diffusionevaporation equation (see also Pt3). According to this equation, evaporation of droplets due to negative supersaturation in the mixing volume takes place simultaneously with turbulent mixing. Since droplets within the volume are under different negative supersaturation values until the final equilibrium is reached, the modeled mixing is inhomogeneous. The droplets can evaporate either partially or totally. The evaporation leads to a decrease in droplet sizes and in droplet concentration.

Like in Pt3, the process of turbulent diffusion is described by a 1D equation of turbulent diffusion. The equation does not describe formation of separate turbulent filaments. Instead, it describes averaged effects of turbulent vortices of different scales by modeling of turbulent diffusion, characterized by a typical value of turbulent diffusion coefficient K. The mixing is assumed to be driven by isotropic turbulence at scales within the inertial sub-range where Richardson's law is valid. In this case, turbulent coefficient is evaluated as in Monin and Yaglom (1975):

141
$$K(L) = C\varepsilon^{1/3}L^{4/3}$$
 (1)

In Eq. (1) ε is the turbulent kinetic energy dissipation rate and C = 0.2 is a constant (Monin and Yaglom, 1975), Boffetta and Sokolov (2002). Eq. (1) means that we consider the effects of turbulent diffusion at scales much larger than the Kolmogorov microscale, i.e. the effects of molecular diffusion are neglected. In the simulations, we use L = 40 m and $\varepsilon = 20 cm^2 s^{-3}$. It means that in the present study mixing is performed by vortices smaller than several tens of meters which agrees with measurements in warm Cu (Gerber et al. 2008). The value of turbulent kinetic energy dissipation rate chosen is also typical for small Cu (e.g. Gerber et al.

149 2008). These parameters correspond to the values of Da of several hundred. The model allows 150 utilization of other values of L and ε typical of other cloud type (say, deep convective 151 clouds) which can change results quantitatively, but not qualitatively.

152

153 Geometry of mixing and the initial conditions

154 The conceptual scheme presenting mixing geometry and the initial conditions used in the 155 following analysis are shown in **Figure 1**.

156

157 **Fig 1 here**

158

At t = 0 the mixing volume of length L is divided into two volumes: the cloud volume of 159 length μL (Fig.1, left) and the dry volume of length $(1-\mu)L$ (Fig.1, right), where $0 \le \mu \le 1$ 160 is the cloud volume fraction. The entire volume is assumed closed, i.e. adiabatic. At t = 0 the 161 cloud volume is assumed saturated, so the supersaturation $S_1 = 0$. This volume is also 162 characterized by the initial distribution of the square of the droplet radii $g_1(\sigma)$, where $\sigma = r^2$. 163 164 The initial liquid water mixing ratio in the cloudy volume is equal to $q_{w1} = \frac{4\pi\rho_w}{3\rho_a} \int_{0}^{\infty} \sigma^{3/2} g_1(\sigma) d\sigma$. The integral of $g_1(\sigma)$ over σ is equal to the initial droplet 165

166 concentration in the cloud volume $N_1 = \int_{0}^{\infty} g_1(\sigma) d\sigma$. The initial droplet concentration in the

167 dry volume is $N_2 = 0$, the initial negative supersaturation in this volume is $S_2 < 0$ and the 168 initial liquid water mixing ratio $q_{w2} = 0$. Therefore, the initial profiles of these quantities 169 along the x-axis are step functions:

171
$$N(x,0) = \begin{cases} N_1 & \text{if } 0 \le x < \mu L \\ 0 & \text{if } \mu L \le x < L \end{cases}$$
 (2a)

172
$$S(x,0) = \begin{cases} 0 & \text{if } 0 \le x < \mu L \\ S_2 & \text{if } \mu L \le x < L \end{cases}$$
(2b)

173
$$q_w(x,0) = \begin{cases} q_{w1} & \text{if } 0 \le x < \mu L \\ 0 & \text{if } \mu L \le x < L \end{cases}$$
 (2c)

The initial profile of droplet concentration is shown in Fig. 1b. This is the simplest inhomogeneous mixing scheme, wherein mixing takes place only in the x-direction, and the vertical velocity is neglected.

Since the total volume is adiabatic, the fluxes of different quantities through the left andright boundaries at any time instance are equal to zero, i.e.

180

181
$$\frac{\partial N(0,t)}{\partial x} = \frac{\partial N(L,t)}{\partial x} = 0; \quad \frac{\partial q_w(0,t)}{\partial x} = \frac{\partial q_w(L,t)}{\partial x} = 0; \quad \frac{\partial q_v(0,t)}{\partial x} = \frac{\partial q_v(L,t)}{\partial x} = 0 \quad (3)$$

182 where q_v is the water vapor mixing ratio.

To investigate of mixing process for different initial DSD, we assume that DSD in the cloudvolume can be represented by a Gamma distribution:

185
$$f(r,t=0) = \frac{N_0}{\Gamma(\alpha)\beta} \left(\frac{r}{\beta}\right)^{\alpha-1} \exp\left(-\frac{r}{\beta}\right)$$
(4)

where N_0 is an intercept parameter, α is a shape parameter and β is a slope parameter of distribution. The DSD f(r) relates to distribution $g_1(\sigma)$ as $f(r) = 2rg_1(\sigma)$. We performed simulations with both initially wide and narrow DSDs. The width of DSD is determined by a set of parameters. The parameters of the initial Gamma distributions used in this study are presented in **Table 1**. Parameters of the distributions are chosen in such a way that the modal radii of DSD and the values of LWC are the same for both distributions. These distributions were used in Pt2 for analysis of homogeneous mixing.

193

194 Table 1 here

Conservative quantity $\Gamma(x,t)$

198 The supersaturation equation for an adiabatic immovable volume can be written in the

199 form
$$\frac{1}{S+1}\frac{dS}{dt} = -A_2 \frac{dq_w}{dt}$$
, where S is supersaturation over water, and the coefficient

 $A_2 = \frac{1}{q_v} + \frac{L_w^2}{c_p R_v T^2}$ is slightly dependent on temperature (Korolev and Mazin, 2003) (notations

201 of other variables are presented in **Appendix**). In our analysis we consider A_2 to be a 202 constant. As follows from the supersaturation equation, the quantity

204
$$\Gamma(x,t) = \ln[S(x,t)+1] + A_2 q_w(x,t)$$
 (5)

is a conservative quantity, i.e. it is invariant with respect to phase transitions. In Eq. (5), |S(x,t)| can be comparable with unity by the order of magnitude. The conservative quantity $\Gamma(x,t)$ obeys the following equation for turbulent diffusion

210
$$\frac{\partial \Gamma(x,t)}{\partial t} = K \frac{\partial^2 \Gamma(x,t)}{\partial x^2}$$
(6)

212 with the adiabatic (no flux) condition at the left and right boundaries $\frac{\partial \Gamma(0,t)}{\partial x} = \frac{\partial \Gamma(L,t)}{\partial x} = 0$

and the initial profile at
$$t = 0$$

215
$$\Gamma(x,0) = \begin{cases} A_2 q_{w1} & \text{if } 0 \le x < \mu L \\ \ln[S_2 + 1] & \text{if } \mu L \le x < L \end{cases}$$
(7)

From Eq. (7) it follows that $\Gamma(x,0)$ is positive in the cloud volume and negative in the initially dry volume. The mean value of function $\Gamma(x,0)$ can be written as follows:

220
$$\overline{\Gamma} = \frac{1}{L} \int_{0}^{L} \Gamma(x,0) dx = \frac{A_2 q_{w1}}{L} \int_{0}^{\mu L} dx + \frac{\ln[S_2 + 1]}{L} \int_{\mu L}^{L} dx = \mu A_2 q_{w1} + (1 - \mu) \ln[S_2 + 1] \quad (8)$$

221

222 $\overline{\Gamma}$ can be either positive or negative. In the latter case a complete evaporation of droplets in the 223 course of mixing takes place.

The solution of Eq. (6) with the initial condition (7) is (Polyanin et al., 2004):

$$\Gamma(x,t) = \sum_{n=0}^{\infty} a_n \exp\left(-\frac{Kn^2 \pi^2 t}{L^2}\right) \cos\left(\frac{n\pi x}{L}\right) =$$
225 $\mu A_2 q_{w1} + (1-\mu) \ln[S_2+1] -$
2($\ln[S_2+1] - A_2 q_{w1}$) $\sum_{n=1}^{\infty} \frac{\sin(n\pi\mu)}{n\pi} \exp\left(-\frac{Kn^2 \pi^2 t}{L^2}\right) \cos\left(\frac{n\pi x}{L}\right)$
(9)

226 One can see that function $\Gamma(x,t)$ depends on three independent parameters A_2q_{w1} , S_2 and μ . 227 This function does not depend on the shape of the initial DSD in the cloud volume. In the final 228 state when $t \to \infty$, $\Gamma(x,t)$ is :

229
$$\Gamma(t=\infty) = \mu A_2 q_{w1} + (1-\mu) \ln[S_2 + 1]$$
 (10)

230 Therefore, $\Gamma(t = \infty)$ depends on the cloud fraction and the initial values of liquid water 231 mixing ratio in the cloud volume and the relative humidity in initially dry volume.

The final equilibrium values of supersaturation $S(x,\infty)$ and liquid water mixing ratio $q_w(x,\infty)$ can be calculated using Eq. (5). The case $\Gamma(t=\infty) > 0$ corresponds to the equilibrium state with $S(x,\infty) = 0$ and $q_w(x,\infty) = \mu q_{w1} + (1-\mu) \frac{\ln[S_2+1]}{A_2}$, when droplets

remain, but do not evaporate any longer.

The case $\Gamma(t=\infty) < 0$ corresponds to the equilibrium state with $q_w(x,\infty) = 0$ and $S(x,\infty) = (1+S_2)^{1-\mu} \exp(\mu A_2 q_{w1}) - 1$. In this equilibrium state droplets are totally evaporated, and volume remains subsaturated $S(x,\infty) < 0$. At given q_{w1} and S_2 , there is a critical value of the cloud fraction μ_{cr} which separates these two possible final equilibrium states. This critical value corresponds to $\Gamma(t=\infty) = 0$ and can be calculated from Eq. (10) as:

241

242
$$\mu_{cr} = \frac{\ln[S_2 + 1]}{\ln[S_2 + 1] - A_2 q_{wl}}$$
(11)

243

244 Another expression for μ_{cr} was formulated in Pt1.

The examples of spatial-temporal variations of function $\Gamma(x,t)$ for different cloud fractions and initial RH=80% are shown in **Figure 2.**

247

248 Fig 2 here

249

Upper panels $\mu = 0.1$ correspond to the case of final total droplet evaporation and negative 250 final function Γ , whereas the middle and bottom rows $\mu = 0.5$ and $\mu = 0.9$ illustrate partial 251 evaporation cases when the total mixing volume reaches saturation. It is interesting that the 252 253 time required for the final equilibrium state to be reached practically does not depend on the cloud fraction, being ~180 seconds for the illustrated cases. The cases $\mu = 0.1$ and $\mu = 0.9$ 254 demonstrate a strong non-symmetric spatial variability of $\Gamma(x)$ function during the first 50 255 seconds. At $\mu = 0.5$, a nearly full compensation between saturation deficit in the dry volume 256 and available liquid water in the cloud volume takes place if at the equilibrium state 257 $S(x,\infty) = q_w(x,\infty) = \Gamma(x,\infty) = 0$. However, the compensation at $\mu = 0.5$ is not full because of 258 the nonlinearity of Γ in Eq. (5). 259

261 Diffusion-evaporation equation for DSD

To formulate the diffusion-evaporation equation we use a simplified equation for droplet evaporation (Pruppacher and Klett, 1997), in which the curvature term and the chemical composition term are omitted

$$265 \quad \frac{d\sigma}{dt} = \frac{2S}{F} \tag{12}$$

266 where $F = \frac{\rho_w L_w^2}{k_a R_v T^2} + \frac{\rho_w R_v T}{e_w (T)D} = const$ (Notations of other variables are presented in Appendix.)

267 The solution of Eq. (12) is

268
$$\sigma(t) = \frac{2}{F} \int_{0}^{t} S(t') dt' + \sigma_0$$
(13)

Eq. (13) means that in the course of evaporation, distribution $g(\sigma)$ shifts to the left without changing its shape. The diffusion-evaporation equation for function $g(x,t,\sigma)$ can be written in the form

272

273
$$\frac{\partial g}{\partial t} = K \frac{\partial^2 g}{\partial x^2} + \frac{\partial}{\partial \sigma} \left(\frac{d\sigma}{dt} g \right)$$
(14)

274 Combining Eqs. (12) and (14) yields

275
$$\frac{\partial g(x,t,\sigma)}{\partial t} = K \frac{\partial^2 g(x,t,\sigma)}{\partial x^2} + \frac{2S}{F} \frac{\partial g(x,t,\sigma)}{\partial \sigma}$$
(15)

276

Eq. (15) is similar to the diffusion-evaporation equation for size distribution function used in Pt 3. The first term on the right hand side of Eq. (15) describes the effect of turbulent diffusion, while the second term describes the changes of size distribution due to droplet evaporation. To close this equation, one can use Eq. (5) written as

282
$$S(x,t) = \exp[\Gamma(x,t) - A_2 q_w(x,t)] - 1,$$
 (16)

and the equation for liquid water mixing ratio

285

286
$$q_w(x,t) = \frac{4\pi\rho_w}{3\rho_a} \int_0^\infty \sigma^{3/2} g(x,t,\sigma) d\sigma$$
(17)

287 The equation system (15-17) for distribution $g(x,t,\sigma)$ should be solved under the following

288 initial condition

289
$$g(x,0,\sigma) = \begin{cases} g_1(\sigma) & \text{if } 0 \le x < \mu L \\ 0 & \text{if } \mu L \le x < L \end{cases}$$
(18)

and using the Neumann boundary conditions

291

292
$$\frac{\partial g(0,t,\sigma)}{\partial x} = \frac{\partial g(L,t,\sigma)}{\partial x} = 0$$
(19)

293

These equations were solved numerically on a linear grid of droplet radii r_j being within the range 0-50 µm, where j = 1...50 are the bin numbers. The number of grid points along the x-axis was set equal to 81. In numerical calculations, the "evaporation term" in Eq. (15) was approximated as

298
$$\frac{2S}{F} \frac{\partial g(x,t,\sigma)}{\partial \sigma} \approx \frac{g\left(x,t,\sigma + \frac{2S}{F}\Delta t\right) - g\left(x,t,\sigma\right)}{\Delta t}.$$
 (20)

299

A shift and subsequent remapping of DSD using the method proposed by Kovetz and Olund's (1969) were implemented to solve Eq. (20) with the help of MATLAB solver PDEPE. After calculation of $g(x,t,\sigma_j)$ function, DSD $f(x,t,r_j)$ was calculated using the relationship $f(x,t,r_j) = 2r_jg(x,t,\sigma_j)$.

305 3. Spatial-temporal variations of DSD and of DSD parameters

Mixing may take a significant time. Cloud microphysical parameters measured in *in-situ* observations correspond to different stages of this transient mixing process. During mixing, DSDs and its parameters change substantially, which makes it reasonable to analyze these time changes.

310 Figure 3 shows time evolution of initially narrow DSD in the centers of the cloudy volume and of the initially dry volume. The values of DSD in the initially cloudy volume decrease 311 while there are no significant changes in the DSD shape. At $\mu = 0.7$, the droplet radius 312 corresponding to the DSD maximum remains unchanged during mixing staying equal to 10 313 314 μm . At $\mu = 0.3$ the effect of droplet diffusion on DSD is stronger, and mixing leads not only to a decrease in the DSD values, but also to a decrease in the peak droplet radius in the 315 cloudy volume. Both at $\mu = 0.3$ and $\mu = 0.7$, mixing leads to broadening of the initial DSD 316 due to the appearance of the tail of small droplets. The tail of small droplets is especially 317 pronounced in the initially dry volume since maximum evaporation of penetrated droplets. 318

The rate of the DSD growth in the initially dry volume, depends on the value of the cloud 319 fraction. At a low cloud fraction, DSD maximum (i.e. drop concentration and drop mass) 320 remains substantially lower for the most period of mixing process than that in the cloudy 321 volume. At the same time, the radius corresponding to the DSD maximum increases reaching 322 80% of its maximum value already within the first 5 s. This is due to the fast increase in the 323 324 relative humidity during mixing, so large droplets penetrating the initially dry volume do not decrease in size anyhow significantly determining the values of modal, mean volume and 325 effective radii. Thus, we see two stages of DSD evolution within in the initially dry volumes: 326 327 at the first stage penetrated droplets evaporate totally or partially forming the tail of small droplets. The formation of the tail of smallest droplets does not lead to a significant changes 328 of the size of the largest droplets. Note that according to equation of diffusion 329

growth/evaporation in of sub-saturation conditions, the rate of droplet radii decreases inverse 330 proportionally to the droplet radius. It means that if, say, radius of a 2 μm droplet decreases 331 twice during a certain time instance, the radius of 20 μm droplet will decrease by less than 332 0.1 μm , i.e. remains approximately unchanged. At this stage diffusion of water vapor from 333 334 cloudy volume and evaporation of penetrating droplets lead to a rapid growth of relative humidity RH. This growth of RH decreases evaporation rate of droplets penetrating initially 335 dry volume later. At the second stage mixing leads to the increase in the droplet number due 336 337 to droplet diffusion from cloudy volume. Since, RH is high, this diffusion is not accompanied by significant change droplet sizes, so DSD grows similarly at all radii. 338

339

340 Figure 3 here

341

At the initially wide DSD (**Figure 4**), the radii of the DSD maximum do not change. It means that at the initial RH= 80%, mixing and evaporation lead to a fast saturation of the initially dry volume, after which the peak radius remains unchanged in this volume. In the initially cloud volume RH remains close to 100% so the DSD decrease is related to dilution by the air from initially dry volume.

347

348 Figure 4 here

349

It is interesting that at $\mu = 0.3$ in the initially dry volume, DSD reaches its maximum during the transition period (Fig.4, at t=80s), and then decreases toward the equilibrium state. This behavior is caused by the competition between the diffusion and droplet evaporation.

Figure 5 shows spatial dependences of droplet concentration, LWC and the mean volume radius within the mixing volume at different time instances at narrow initial DSD. At small values of the cloud fraction, diffusion of water vapor and droplets, as well as droplet

256	
356	evaporation lead to a fast decrease in droplet concentration and in LWC in the initially cloud
357	volume. The mean volume radius in this volume decreases by about 15% in the course of
358	mixing. It is natural that at large cloud fraction, droplet concentration and LWC in the initially
359	cloudy volume decrease slowly, while these quantities in the initially dry volume increase
360	rapidly. At both small and large cloud fractions, the mean volume radius in the initially dry
361	volume grows rapidly during the mixing toward its values in the initially cloudy volumes,
362	even if droplet concentration and LWC remain much lower than in the adjacent cloud volume.
363	
364	Figure 5 here
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366	Figure 6 shows the spatial dependences of droplet concentration, LWC and the mean
367	volume radius within the mixing volume at different time instances at wide initial DSD.
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370	Figure 6 here
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372 373	A specific feature of mixing at a wide DSD is the increase in the mean volume radius, so the
	A specific feature of mixing at a wide DSD is the increase in the mean volume radius, so the ratio $\frac{r_v}{r_{v0}} > 1$. In the course of mixing, the mean volume radius maximum is reached in the
373	ratio $\frac{r_{\nu}}{r_{\nu}} > 1$. In the course of mixing, the mean volume radius maximum is reached in the
373 374	ratio $\frac{r_v}{r_{v0}} > 1$. In the course of mixing, the mean volume radius maximum is reached in the
373 374 375	ratio $\frac{r_v}{r_{v0}} > 1$. In the course of mixing, the mean volume radius maximum is reached in the initially dry volumes. This result can be attributed to the fact that in this volume smaller
373 374 375 376	ratio $\frac{r_v}{r_{v0}} > 1$. In the course of mixing, the mean volume radius maximum is reached in the initially dry volumes. This result can be attributed to the fact that in this volume smaller droplets fully evaporate, so the concentration of large droplets increases with respect to
373 374 375 376 377	ratio $\frac{r_v}{r_{v0}} > 1$. In the course of mixing, the mean volume radius maximum is reached in the initially dry volumes. This result can be attributed to the fact that in this volume smaller droplets fully evaporate, so the concentration of large droplets increases with respect to concentration of smaller droplets (Fig. 4, right column). Scattering diagrams plotted using <i>in</i> -

study shows that the behavior of $\frac{r_v}{r_{v0}}$ with time in the course of mixing may depend of the DSD shape in the initially cloud volume that determines relationship between concentrations of small and large droplets in course of mixing. Of course, the DSD shape is only one possible reason of appearance of points with $\frac{r_v}{r_{v0}} > 1$ on the scattering diagram.

385 We see that the transition to the final equilibrium state within the volume with the spatial 386 scale of 40 m is about 5 min (Fig. 8), which is a comparatively long period of time compared to the characteristic times of other microphysical processes, including droplet evaporation. 387 388 During this time the DSD changes substantially, especially at small cloud fraction. The mean volume radius in the initially dry volume increases much faster than LWC. As a result, mean 389 volume radius in such volume rapidly reaches the values typical of cloudy air, while LWC 390 still remains substantially lower than in the cloudy volume. Despite some DSD broadening, 391 the final DSDs in the mixing volume resemble those in the initially cloud volumes. The main 392 393 effect of mixing is lowering the DSD values as the cloud fraction decreases.

- 394
- 395

4. Equilibrium state and mixing diagram

This study reconsiders the classical theory of mixing diagrams. In the classical theory 396 two volumes (cloudy and droplet free) mix with each other within a given unmovable mixing 397 volume (see review by Korolev et al., 2016). Mixing diagrams are typically plotted for times 398 399 when all variables become uniform within the mixing volume, i.e when the equilibrium state 400 is reached. We plot the mixing diagram using the same simplifications used in the plotting 401 classical mixing diagrams, namely: no vertical motions and no collisions are assumed. These 402 assumptions allow to reveal better the microphysical effects of turbulent mixing. It is widely 403 assumed that the mixing type is determined by the Damkohler number that depends only on 404 drop relaxation time and mixing time. No averaged vertical velocity and no collision rate are 405 included into this criterion.

We extend the theory, however, in several important aspects concerning microphysical effects: a) we consider time dependent process of mixing and b) initial droplet size distributions are assumed polydisperse.

Mixing considered in the present study always leads to the equilibrium state. As was explained above, two equilibrium states are possible. The first one is characterized by the total evaporation of cloud droplets $q_w(x,\infty) = 0$, whereas the second one occurs if the air in the mixing volume becomes saturated, i.e. when $S(x,\infty) = 0$. At the given initial values of q_{w1} in the cloud volume and of S_2 in the initially dry volume, there always exists the cloud fraction μ_{cr} (Eq. 11) separating these two states.

415

416 **4.1.** The process of achieving the equilibrium state

Figure 7 shows the dependences of the time required to reach the equilibrium on the cloud fraction, at different initial relative humidity values in the dry volume and two initial DSDs (the parameters are presented in Tab.1). The characteristic time is defined here as the time from the beginning of mixing to the time instance when inequality $\delta = \frac{\overline{N}(t) - \overline{N}(\infty)}{\overline{N}(0) - \overline{N}(\infty)} < 0.01$ becomes valid. The mean droplet concentration is calculated by averaging along *x*-axes $(\overline{N}(t) = \frac{1}{L} \int_{0}^{L} N(x,t) dx)$. In case of a total evaporation, $\overline{N}(\infty) = 0$.

423

424 Figure 7 here

425

Each curve in Fig. 7 consists of two branches. The left branches correspond to the total evaporation regime, while the right branches correspond to the partial evaporation at equilibrium. The maximum time corresponds to the situation when the available amount of liquid water is approximately equal to the saturation deficit. A similar result was obtained in 430 Pt1 and Pt2 for homogeneous mixing. The maximum values of the characteristic time are 431 about 4 min for a mixing volume of 40 m in length. The right branches show that the 432 characteristic time decreases with increasing cloud fraction. Despite some differences in the 433 curve slopes, the characteristic times for wide and narrow DSD are quite similar.

Figure 8 shows dependences of normalized cube of the mean volume radius on the cloud 434 fraction at different time instances for two values of x: x=0 (solid lines) corresponds to the 435 initially cloudy volume, and x = L (dashed line) corresponds to the initially dry volume. The 436 figure is plotted for the narrow DSD for two values of RH_2 : 60% and 95%. Despite the fact 437 that the diffusion-evaporation equation allows simulating using any initial RH, we do not 438 consider in our examples the cases of very low RH of dry volume. It is because at very low 439 RH, say, RH=20%, the cloud fraction should exceed 0.8 to prevent total droplet evaporation 440 in the equilibrium state (at LWC=1 g/kg). At the same time, we are interested in the 441 equilibrium state at which droplets exist. Note that at the lateral edges of warm Cu a shell of 442 humid air arises around cloud, so RH of the entrained air should be high enough (e.g. Gerber 443 et al., 2008). 444

445

446 **Figure 8 here**

447

The curve plotted for the time instance of 300 s corresponds to the equilibrium state (hereafter the equilibrium curve). The curves above the equilibrium curve correspond to the initially cloudy volume, and the curves below the equilibrium curve correspond to the initially dry volume. One can see how curves of both types approach the same final state. During the

452 mixing the curves move over the $\left(\frac{r_{\nu}}{r_{\nu 0}}\right)^3 - \mu$ plane toward the equilibrium curve. As a result,

the curves plotted in Fig.8, corresponding to different time instances of the mixing, togethercover the entire area of the panels.

455 During this movement the distance from the curves to the horizontal line
$$\left(\frac{r_{\nu}}{r_{\nu 0}}\right)^3 = 1$$
 changes,

and the curves slopes increase. In our case of L = 40 m, the mixing remains inhomogeneous

20

457 the during entire mixing process, so the change in the distance from the curves to the 458 horizontal line $\left(\frac{r_{\nu}}{r_{\nu 0}}\right)^3 = 1$ characterizes the temporal changes over the mixing process, but not

459 a change in mixing type.

456

It is noteworthy in this relation that scattering diagrams plotted using *in-situ* observations reflect mixing between different multiple volumes at different stages of the mixing process. Accordingly, points in the scattering diagrams can be far from the equilibrium location. Fig. 8 indicates, therefore, that scattering diagrams show snapshots of transient mixing process when the distance from points in the diagrams to line $\left(\frac{r_v}{r_{v0}}\right)^3 = 1$ characterize the stage of the

465 mixing process, but not the mixing type.

466 The dependences of normalized cube of the mean volume radius on the cloud fraction at 467 different time instances at wide DSD also indicate approaching to the equilibrium curve,

468 while all the curves correspond to
$$\left(\frac{r_{\nu}}{r_{\nu 0}}\right)^3 > 1$$
 (not shown).

469 Note that in several studies normalized effective radius is used for plotting scattering and 470 mixing diagrams, but not mean volume radius (Gerber et al. 2008; Freud et al., 2011). 471 Comparison of scattering and mixing diagrams in the study plotted using mean volume and 472 effective radii did not reveal any significant differences (not shown).

473

474 **4.2. Mixing diagrams**

Using the diffusion-evaporation equations (15-17) we calculated the equilibrium DSD for
different initial relative humidity values and different cloud fractions. Each calculation was

performed for both narrow and wide initial DSD (parameters shown in Tab.1). These
equilibrium DSD were used to calculate mixing diagrams showing dependences of normalized
cube of the effective radius on the cloud fraction.

The corresponding mixing diagrams for homogeneous mixing case were also calculated for comparison. To this effect, the supersaturation and DSD in both the cloud and the dry volumes were aligned, taking into account the cloud fraction value μ . The alignment led to the following initial values of supersaturation and DSD within the mixing volume:

484

485
$$S_0 = (1 - \mu)S_2; \ g_0(\sigma) = \mu g_1(\sigma)$$
 (21)

486

Upon the alignment, time evolution values of DSD under homogeneous evaporation in an adiabatic immovable parcel were calculated until the equilibrium state was reached. These equilibrium DSD were used to calculate mixing diagrams for homogeneous mixing. To do this, we used the parcel model proposed by Korolev (1995) that describes evaporation by means of equations with temperature-dependent parameters. **Figure 9** shows the mixing diagrams plotted for initial narrow and wide DSD cases.

493

```
494 Figure 9 here
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495

496

497 While all the curves in the mixing diagram for narrow DSD are below the straight line 498 $\left(\frac{r_v}{r_{v0}}\right)^3 = 1$, the curves for wide DSD are above this line. The explanation of this effect is given 499 in Section 3 (Fig. 6). The curves plotted for homogeneous and inhomogeneous mixing 500 demonstrate an important feature. Namely, at given values of RH and q_{w1} in the initially dry 501 volume, the values μ_{cr} of the cloud fraction at which all the droplets evaporate are 502 approximately the same for any type of mixing. This condition is the consequence of the mass 503 conservation law determined by Eq. (11) and does not depend of the initial DSD shape. In 504 standard mixing diagrams (e.g. Lehmann et al., 2009; Gerber et al., 2008; Freud et al., 2011),

505 the horizontal straight line
$$\left(\frac{r_v}{r_{v0}}\right)^3 = 1$$
 (or $\left(\frac{r_e}{r_{e0}}\right)^3 = 1$) is typically plotted for the entire range of

the cloud fraction [0...1], while the curves corresponding to homogeneous mixing are plotted for different RH within the range $[\mu_{cr}(RH_2)...1]$. As a result, the high difference between extremely inhomogeneous and homogeneous mixing types is clearly seen at low RH and at small cloud fractions. The condition that μ_{cr} is the same for different mixing types indicates that the mixing diagrams may look nearly similar for $\mu > \mu_{cr}$. It means that the range of the cloud fractions required for comparison of diagrams aimed at determination of a mixing type shortens as RH_2 values in the surrounding air decrease.

513 The comparison of the left and the right panels in Fig. 9 shows that the differences between the diagrams for homogeneous and inhomogeneous mixing types are more 514 515 pronounced for initially narrow DSD. The maximum difference should take place for monodisperse DSD considered in in Pt1, Pt2 and Pt3. Within the range of $\mu > \mu_{cr}$, the 516 517 distance between the curves corresponding to different mixing regimes is small even for narrow DSD and low RH_2 . The lower difference is related to the fact that at high RH_2 the 518 curves in the mixing diagrams are close to the horizontal straight line in both regimes, while at 519 low RH_2 , μ_{cr} is small and both curves should drop to zero in the vicinity of $\mu = \mu_{cr}$. 520

521 As regards the wide DSD case, the difference between the curves corresponding to 522 different mixing type is negligible (Fig. 9, right)

523

524 **4.3. Effect of the relative humidity**

In measurements carried out at cloud boundaries and in cloud simulations, the cloud 525 fraction is not known, therefore it is widely accepted to use normalized droplet concentration 526 instead of the cloud fraction (Burnet and Brenguier, 2007; Gerber et al., 2008: Lehmann et al., 527 2009). Droplet concentration is normalized by the maximum value along the airplane traverse. 528 The difference between the cloud fraction and normalized droplet concentration is obvious: 529 the cloud fraction is a parameter given as the initial condition. At the same time, normalized 530 droplet concentration changes with time and space due to complete evaporation of some 531 droplet fraction. Figure 10 shows dependencies of normalized droplet concentration on the 532 cloud fraction at the equilibrium final state of mixing. One can see a substantial deviation 533 534 from 1:1 linear dependence, especially at low RH. As we know, droplet concentration decreases in the course of both homogeneous and inhomogeneous mixing if the initial DSD 535 are polydisperse. The fraction of totally evaporating droplets increases with decreasing RH_2 . 536 As expected, droplet concentration in homogeneous mixing is higher than that in 537 inhomogeneous mixing. The difference between droplet concentrations at wide DSD is lower 538 539 than at narrow DSD.

540

541 Fig. 10 here

542

543 Figure 11 shows the dependencies $\left(\frac{r_v}{r_{v0}}\right)^3$ on normalized droplet concentration for narrow

and wide DSD in inhomogeneous mixing. The normalization by droplet concentration in the initially cloud volume at t = 0 was used. Taking into account the dependences of normalized droplet concentration on the cloud fraction μ (Fig. 10), one can get the curves shown in Fig. 11 which actually coincide at different RH_2 . The lack of the sensitivity to RH_2 can be attributed to the fact that a decrease in RH leads to a decrease in normalized droplet concentration, so the curves corresponding to low RH in Fig. 9 shift to the left when the normalized droplet concentration is used instead of μ . The shape of the dependences in Fig 11 (right) is explained by an increase in the mean volume radius with decreasing droplet concentration.

553

- 554 **Fig 11 here**
- 555

Thus, the mixing diagrams plotted in the plane $\left(\frac{r_v}{r_{v0}}\right)^3$ vs normalized droplet concentration do not depend on the relative humidity of the surrounding dry air. This result indicates an additional difficulty in distinguishing between mixing types based on scattering diagrams plotted using *in-situ* data in these axes. The concentration of observed points in $\left(\frac{r}{r}\right)^3$

560 these scattering diagrams close to the line $\left(\frac{r_{\nu}}{r_{\nu 0}}\right)^3 = 1$ is often interpreted as an indication of

homogeneous mixing, but at high RH in the surrounding air (Gerber et al., 2008; Lehmann et
al., 2009). High values of RH in the penetrating air volumes are usually explained by
formation of a layer of most air around the cloud boundary (Gerber et al., 2008, Knight and
Miller, 1998).

The reference values of droplet concentration and the effective radius used for normalization in the present study are taken as the initial values in the cloud volume before it mixes with the neighbouring dry volume. In real *in-situ* measurements the reference values of these quantities are typically chosen in a less diluted cloud volume along the airplane traverse. This reference volume may be quite remote from the particular mixing volume. It can lead to

570 a shift of the mixing diagram with respect to the $\left(\frac{r_v}{r_{v0}}\right)^3 = 1$ line, as well as to a large variation

in mixing diagram shapes, unrelated, however, to the mixing type (e.g., Lehmann et al.,2009).

5. Discussion and conclusion

575 This study extends the analysis of mixing performed in Pt3 where the diffusionevaporation equation served as the basis, the initial DSD were assumed monodisperse and 576 the cloud fraction was chosen as $\mu = 1/2$. In the present study, the analysis focuses on the 577 temporal and spatial evolution of initially polidisperse DSD and investigates mixing diagrams 578 obtained for narrow and wide initial DSD within a wide range of the cloud fraction values (0.1 579 580 - 0.95). It is shown that results of mixing and the structure of mixing diagrams depend on the initial DSD shape. This finding indicates that mixing is a multi-parametrical problem that 581 582 cannot be determined by a single parameter (e.g. the Damkölher number as often assumed) or 583 even by two parameters (the Damkölher number and the potential evaporation parameters as assumed in Pt3). The temporal changes of DSD and their moments during mixing are 584 calculated. Although DSD broaden, they tend to remain similar to the original DSD. The main 585 changes come from the cloud air dilution by the dry air, which leads to a decrease in droplet 586 concentration for all droplet sizes. The changes of DSD and its shape are minimum in the 587 588 initially cloud volumes, especially at significant cloud fractions. The droplet radii corresponding to the DSD peak do not change anyhow significantly. In the initially dry 589 volumes, mixing and evaporation of penetrated droplets leads to a rapid increase in RH. 590 591 Consequently, large droplets penetrating these volumes do not change their sizes significantly. As a result, the mean volume radius in these volumes rapidly increases and reaches the values 592 typical of cloud volumes, while LWC remains lower than in the cloud volume for most of the 593 mixing time. At narrow DSD, the mean volume (and effective) radius remains smaller than 594 that in the initially cloud volume. At wide DSD, the mean volume (and effective) radius may 595 become larger than that in the initial DSD. This increase in the effective radius is attributed to 596 the fact that evaporation of smaller droplets leads to the increase in the fraction of larger 597 598 droplets in the DSD. In this study, and in Pt3 it is shown that mixing leads to DSD

599 broadening. This contrasts with the classical theory, when initially monodisperse DSDs 600 remain monodisperse in course of mixing. This problem is analyzed in detail in Pt 3. Note 601 that in real clouds DSD there are many mechanism leading to DSDs broadening (e.g. Pinsky 602 and Khain 2002).

Dependences of normalized cube of the mean volume radius on the cloud fraction $(r_v / r_{v0})^3$ as a function of μ at different time instances form the set of curves filling the entire $(r_v / r_{v0})^3 - \mu$ plane. Therefore, both the slope and the distance of these curves in respect to the horizontal line $(r_v / r_{v0})^3 = 1$ change with time. It means that this distance characterizes the temporal changes in the course of mixing, but not the mixing type (which remains inhomogeneous during the entire mixing time). The mixing process is comparatively long (several minutes), so the final equilibrium stage is hardly achievable in real clouds.

It is highly significant that the critical values of the cloud fraction μ_{cr} corresponding to total droplet evaporation are the same for any mixing type. It means that the curves in a mixing diagram corresponding to homogeneous and inhomogeneous mixing types should be compared only within the range of $\mu > \mu_{cr}$. The range width of $\mu > \mu_{cr}$ decreases with decreasing relative humidity in the initially dry volume. Taking into account significant scattering of observed points, this condition greatly hampers the problem of how to distinguish between mixing types,

Another important result of the study is that mixing diagrams for homogeneous and inhomogeneous mixing plotted for polydisperse DSD do not differ much. The largest difference takes place for initially narrow DSD (the maximum difference takes place for initially monodisperse DSD), but even in this case the difference is not large enough to reliably distinguish mixing type, owing the significant scatter of observed data. At wide DSD, this difference between mixing diagrams for homogeneous and inhomogeneous becomes negligibly small.

The cloud fraction μ is a predefined parameter and is not determined from observations. 624 Consequently, in the analysis of *in-situ* measurements the normalized droplet concentration is 625 typically used instead of the cloud fraction. However, there is a significant difference 626 between the cloud fraction prescribed a priori and the normalized droplet concentration that 627 628 changes due to total evaporation of some fraction of droplets. We have shown that the utilization of normalized droplet concentration in mixing diagrams is not equivalent to the 629 630 utilization of the cloud fraction. The important conclusion is that when mixing diagrams are plotted using the normalized concentration, the sensitivity to RH disappears. This conclusion 631 is valid even when RH in the initially dry volume is as low as 60%. This conclusion clearly 632 contradicts the wide-spread assumption that mixing types can be easily distinguished in 633 634 mixing diagrams in case of low relative humidity of the surrounding air.

In the present study as well as in Pt3 and LES performed by Andrejczuk et al. (2006, 2009), Khain et al. (2017) it is shown that time needed to establishing of equilibrium either quite long or even never reached. It means that the scattering diagrams observed in situ are just snapshots of the transient mixing process. However, since the classic mixing diagrams are plotted namely for equilibrium states, we investigate the transition to such equilibrium assuming that the mixing volume remains adiabatic (i.e. isolated) during the entire period of mixing. This is, of course, a serious simplification made to compare the results with those predicted by classical concept.

To sum up, our general conclusion is that the simplifications underlying the classical concept of mixing are too crude, making it impossible to use mixing diagrams for comprehensive analysis of mixing and especially for determination of mixing types. At the same time, mixing diagrams may contain useful information concerning the DSD width.

646

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Appendix. List of symbols

Symbol	Description	Units	
A ₂	$\frac{1}{q_v} + \frac{L_w^2}{c_p R_v T^2} \text{, coefficient}$	-	
a_n	Fourier series coefficients	-	
С	Richardson's law constant	-	
C _p	specific heat capacity of moist air at constant pressure	J kg ⁻¹ K ⁻¹	
D	coefficient of water vapor diffusion in air	$m^2 s^{-1}$	
Da	Damkölher number	-	
е	water vapor pressure	N m ⁻²	
e_w	saturation vapor pressure above flat surface of water	N m ⁻²	
F	$\left(\frac{\rho_w L_w^2}{k_a R_v T^2} + \frac{\rho_w R_v T}{e_w (T) D}\right), \text{ coefficient}$	m ⁻² s	
f(r)	droplet size distribution	m ⁻⁴	
g(r)	droplet size distribution	m ⁻⁵	
$g_0(\sigma)$	initial distribution of square radius in homogeneous mixing	m ⁻⁵	
$g_1(\sigma)$	initial distribution of square radius	m ⁻⁵	
ka	coefficient of air heat conductivity	J m ⁻¹ s ⁻¹ K ⁻¹	

K	turbulent diffusion coefficient	m^2s^{-1}
L	characteristic spatial scale of mixing	m
$L_{_{\!W}}$	latent heat for liquid water	J kg ⁻¹
Ν	droplet concentration	m ⁻³
N_0	Parameter of Gamma distribution	m ⁻³
\overline{N}	mean droplet concentration	m ⁻³
N_1	initial droplet concentration in cloud volume	m ⁻³
р	pressure of moist air	N m ⁻²
q_v	water vapor mixing ratio (mass of water vapor per 1 kg of dry	-
ai	ir)	
$q_{_W}$	liquid water mixing ratio (mass of liquid water per 1 kg of dry	-
ai	ir)	
$q_{\scriptscriptstyle w1}$	liquid water mixing ratio in cloud volume	-
R	$\frac{S_2}{A_2 q_{w1}}$, non-dimensional parameter	-
R_a	specific gas constant of moist air	J kg ⁻¹ K ⁻¹
R_{ν}	specific gas constant of water vapor	J kg ⁻¹ K ⁻¹
r	droplet radius	m
<i>r</i> ₁	initial droplet radius	m
r_e	effective radius	m
r_{e0}	initial effective radius	m
S	$e/e_w - 1$, supersaturation over water	-
S_2	initial supersaturation in the dry volume	-
S_0	initial supersaturation in homogeneous mixing	-

	Т	temperature	Κ
	t	time	S
	x	distance	m
	α	parameter of Gamma distribution	-
	β	parameter of Gamma distribution	m^{-1}
	Δt	time step	S
	μ	cloud fraction	-
	μ_{cr}	critical cloud fraction	-
	ε	turbulent dissipation rate	m^2s^{-3}
	$\Gamma(x,t)$	conservative function	-
	$ ho_a$	air density	kg m ⁻³
	$ ho_{\scriptscriptstyle W}$	liquid water density	kg m ⁻³
	σ	square of droplet radius	m ²
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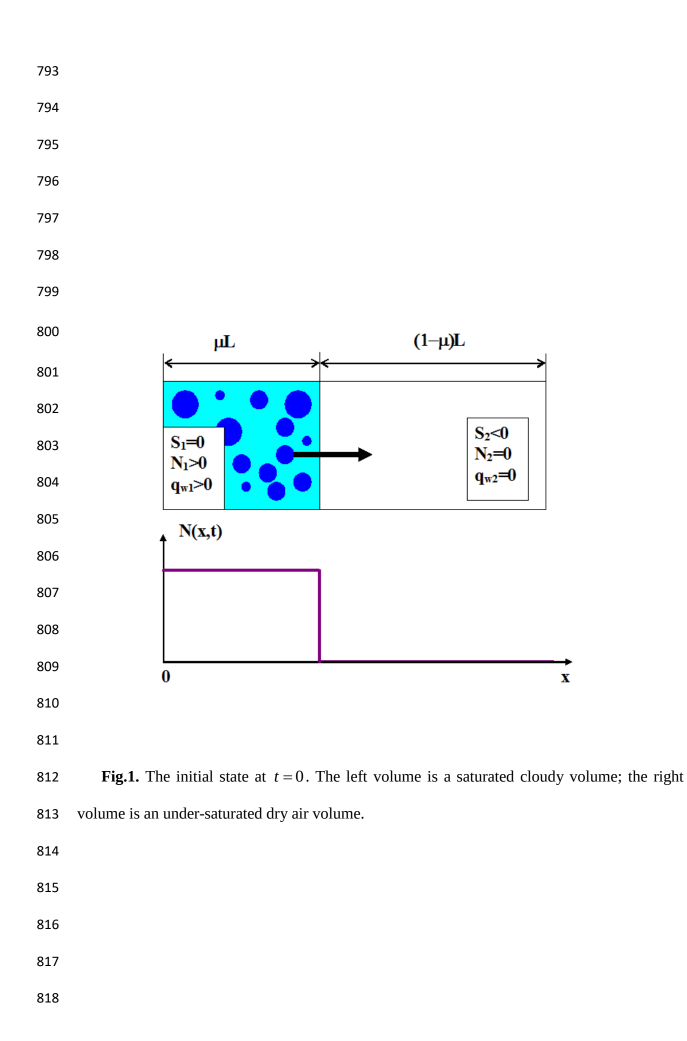
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Tab.1 Parameters of the initial Gamma distributions

DSD	N_0 , cm ³	α	β , μ m	Modal radius,	LWC, g/m ³
				μm	
Narrow	264.2	101.0	0.1	10.0	1.178
Wide	71.0	4.3	3.1	10.0	1.178
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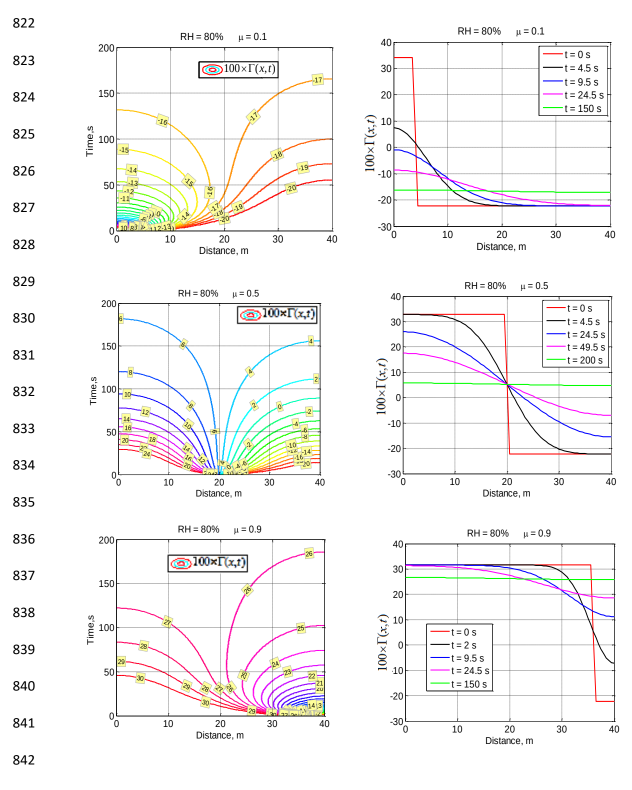


Fig. 2. Spatial-temporal variations of conservative function $100 \times \Gamma(x,t)$ for different cloud fractions μ and initial $RH_2 = 80 \%$.

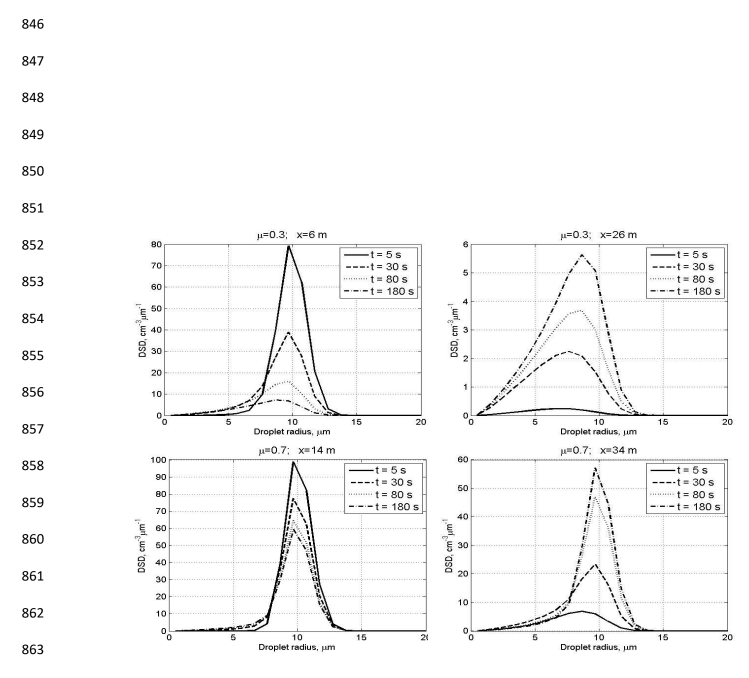
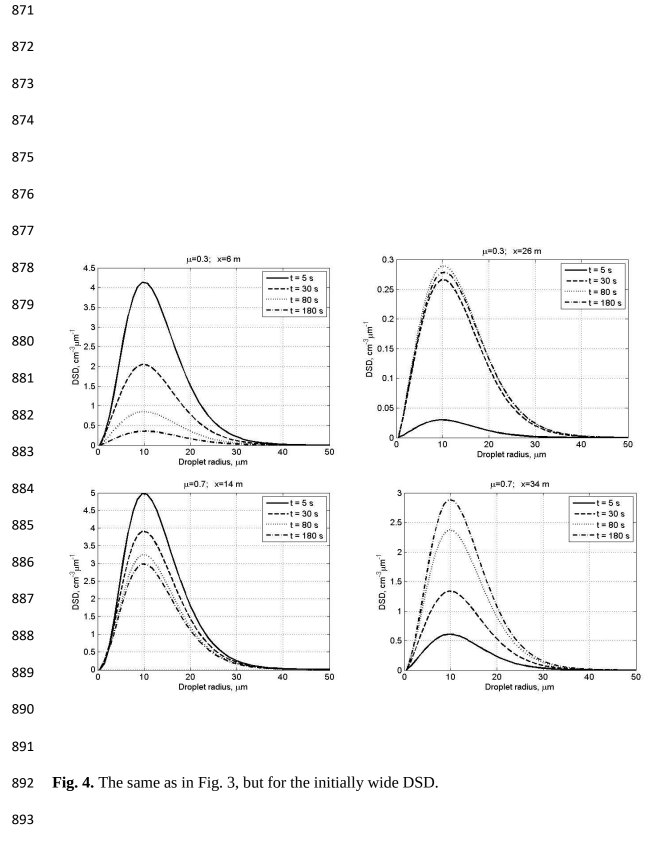


Fig. 3. Time evolution of DSD in the centers of the initially cloudy volume (left) and of the initially dry air volume (right) at initially narrow DSD. The initial mixing parameters are $RH_2 = 80 \%$, $T = 10^{\circ}C$, p = 828.8 mb and L = 40 m.



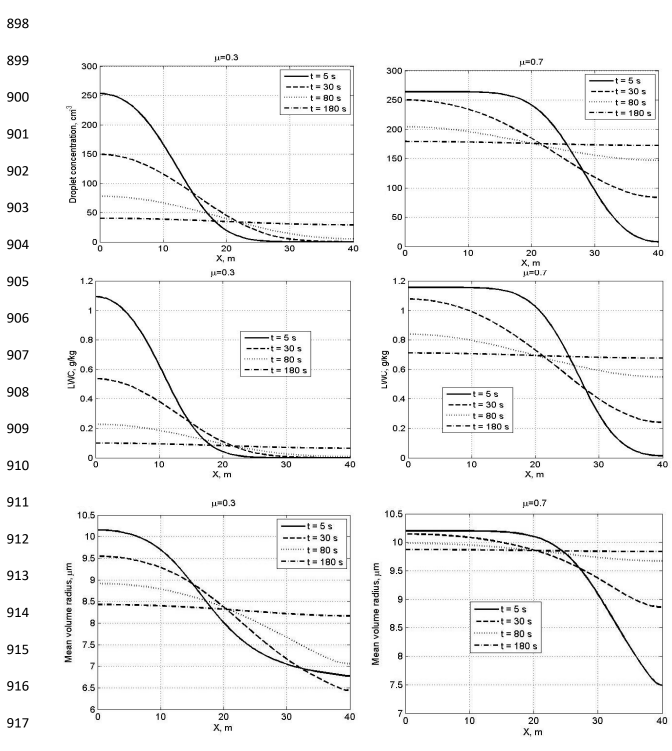


Fig. 5. Spatial dependences of droplet concentration, LWC and the mean volume radius within the mixing volume at different time instances at narrow initial DSD. The initial mixing parameters are $RH_2 = 80 \%$, $T = 10^{\circ}C$, p = 828.8 mb and L = 40 m.



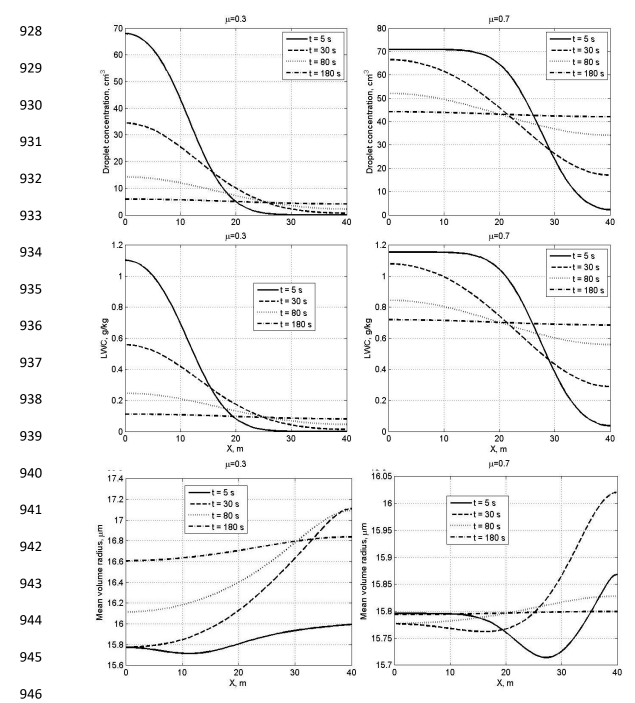


Fig. 6. The same as in Fig. 5, but for wide DSD

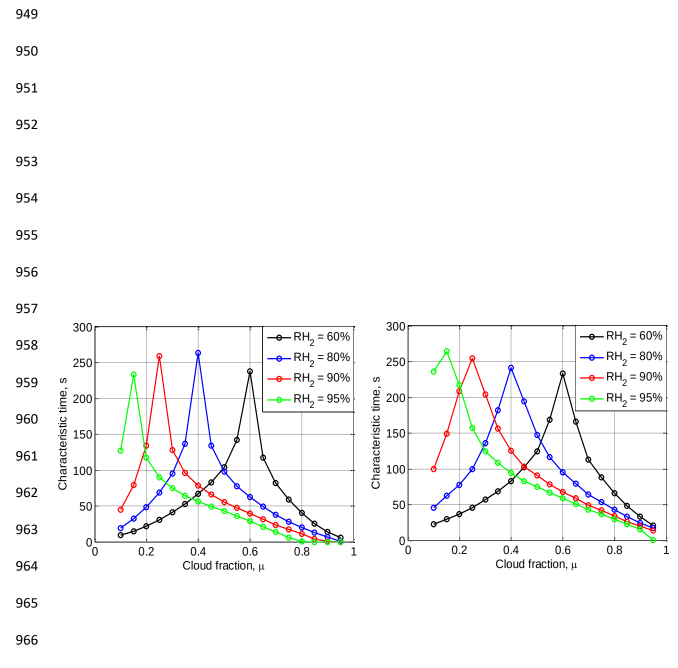


Fig. 7. Time required to reach the equilibrium state vs. the cloud fraction at different initial RH for the initially narrow DSD (left) and the initially wide DSD (right). Parameters of DSD are given in Tab. 1. The initial mixing parameters are $T = 10^{\circ}C$, p = 828.8 mb and L = 40 m.

- 971
- 972
- 973





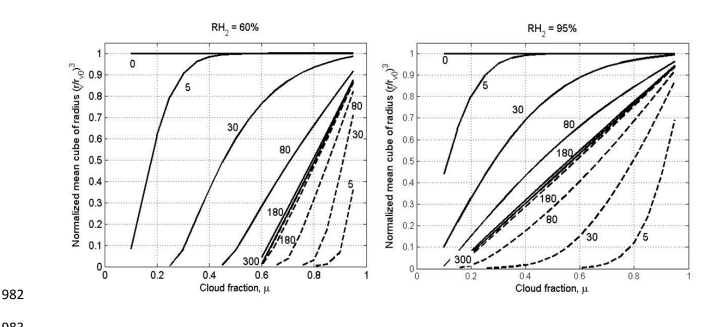




Fig. 8. Dependences of normalized cube of the mean volume radius on the cloud fraction at different time instances for x=0 (solid lines) corresponding to the initially cloud volume, and x = L (dash line) corresponding to the initially dry volume. The time instances in seconds are marked by numbers. The figure is plotted for the narrow initial DSD for two values of RH₂: 60% (left panel) and 95% (right panel). Parameters of DSD are given in Tab. 1. The initial mixing parameters are $T = 10^{\circ}C$, p = 828.8 mb and L = 40 m. Calculations performed within the range of $0.1 < \mu < 0.95$.



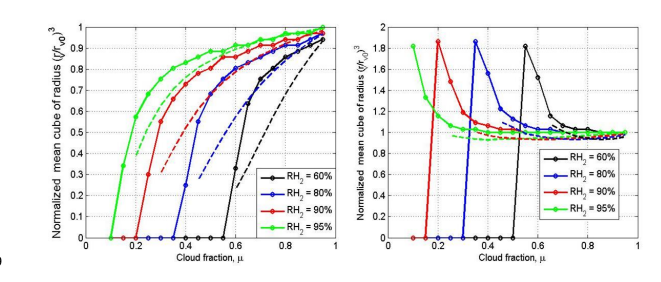


Fig. 9. Mixing diagrams. Normalized cube of the mean volume radius vs. the cloud fraction for initial narrow DSD (left) and initial wide DSD (right). The dependencies correspond to the equilibrium state. Parameters of initial DSD are presented in Tab. 1. Solid and dashed lines show the mixing diagrams for inhomogeneous and homogeneous mixing, respectively. The initial mixing parameters are $T = 10^{\circ}C$, p = 828.8 mb and L = 40 m.

