Increase of the aerosol hygroscopicity by aqueous mixing in a mesoscale convective system: a case study from the AMMA campaign

S. Crumeyrolle, L. Gomes, P. Tulet, A. Matsuki, A. Schwarzenboeck, and K. Crahan

1GAME/CNRM, METEO-FRANCE – CNRS, Toulouse, France
2Laboratoire de Meteorologie Physique, Clermont-Ferrand, France

Received: 6 March 2008 – Accepted: 23 April 2008 – Published: 29 May 2008
Correspondence to: S. Crumeyrolle (suzanne.crumeyrolle@cnrm.meteo.fr)
Published by Copernicus Publications on behalf of the European Geosciences Union.
Abstract

Aerosol properties were measured during an airborne campaign experiment that took place in July 2006 in West Africa within the framework of the African Monsoon Multidisciplinary Analyses (AMMA). The goal of the present study was to determine the main microphysical processes that affect the aerosols during the passage of a mesoscale convective system (MCS) over the region of Niamey in Niger. A main difference in the aerosol profiles measured before and after the passage of the MCS was found for a layer located between 1300 and 3000 m where the aerosol concentration has drastically decreased after the passage of the MCS. Concurrently, a significant increase of the cloud condensation nuclei fraction was also observed during the post-MCS period in the same layer. Moreover, the results of the elemental composition analyses of individual particles collected in this layer after the MCS passage have shown higher contributions of sulfate, nitrate and chloride to the total aerosol. A mesoscale atmospheric model with on-line dust parameterization and Lagrangian backtrajectories was used to interpret the impact of the MCS on the aerosol properties. The results of the simulation show that the MCS 1) generates dust particles at the surface in front of the system and washout particles behind, 2) modifies the aerosol mixing state through cloud processing, and 3) enhances CCN activity of particles through the coating of soluble material.

1 Introduction

The interactions between aerosols and clouds have a large influence on the role of aerosols in climate change. These interactions mainly depend on the particles characteristics (concentration, size, composition, hygroscopic properties, mixing state) and the type of clouds involved (continental or maritime, convective or stratiform, cold or warm). In addition, these interactions are complex because the aerosol characteristics can be modified by in-cloud processes. Indeed, the atmospheric aerosols affect
cloud formation through their role as cloud condensation nuclei (CCN). In return, the clouds affect the aerosol particles by changing their concentration in the atmosphere through various scavenging processes or by modifying their properties through various chemical reactions that take place in the drops (or in ice crystals).

There have been few field observations and measurements related to the impact of cloud formation on the evolution of aerosol properties. Levin et al. (1996) searching to identify the effects of dust and sulfate on cloud formation, found that dust particles were coated with sulfate due to in-cloud processing. A few results obtained from modeling studies mention the role of clouds in modifying aerosols. For example, using a numerical model, Yin et al. (2002) showed that insoluble mineral dust can become effective CCN after passing through a convective cloud. In order to study the processes involved in the interactions between convective clouds and mineral dust, Africa is probably a well designed region for obtaining field observations and measurements.

In Africa, precipitation mostly occurs in the form of mesoscale convective systems (MCS) during the monsoon season when air masses bring a variety of aerosol types from different sources. The air mass circulation is formed by south and north trade winds converging toward the Inter Tropical Discontinuity (ITD). The continental north easterly trade is called Harmattan and the south westerly trade is called monsoon flux. These two major fluxes create two distinct layers characterized by different aerosol properties: the monsoon layer (ML) and the Saharan Air Layer (SAL) (Carlson and Prospero, 1972; Prospero and Carlson, 1972). In the ML, the monsoon flux transports marine air masses over the continent where they are mixed with anthropogenic, biogenic and other sources of aerosols. The monsoon layer is affected by the land surface on diurnal time-scales, through convection and shallow cumulus clouds in the well mixed boundary layer. During their transport, mixed aerosols are modified and their properties are highly dependent on the mixing. North of the ITD and above the ML is the SAL, which can be characterized by high dust content, consistent with low visibility (Karyampudi et al., 1999). The SAL is decoupled from the surface below and is more closely linked to the desert regions (Parker et al., 2005). The presence of aerosol
in the SAL is connected to long range transport from Sahelian and Saharan regions. However, the formation of convective systems can entrain humid ML air into the SAL and dry SAL air into the ML. This mesoscale process erodes the thickness of the SAL and dries the monsoon layer (Parker et al., 2005). This process can also modify dust particles and enhance their CCN activity when they re-enter subsequent clouds.

The African Monsoon Multidisciplinary Analysis (AMMA) program was designed to improve our knowledge on the African monsoon and its variability. A detailed description on the project is available at www.amma-international.org/. One of the objectives of the project was to investigate the cloud-aerosol interactions and particularly the role of aerosols in the African monsoon system. Several intensive campaigns of aerosol and cloud measurements were carried out between 1 June and 25 August 2006 in the western African region by combining airborne and ground measurements.

The goal of the present study is to determine the main microphysical processes that affect the aerosols during the passage of a convective system, by comparing the vertical distribution and the physical and chemical properties of the aerosols before and after the convective system. For this purpose, airborne measurements were conducted in July 2006 over Banizoumbou in Niger during a Mesoscale Convective System mission of the AMMA experiment.

This paper describes the results of profile measurements of aerosol and CCN concentration, aerosol size spectra and chemical composition during this period. Then, a mesoscale model with on-line dust parameterization and Lagrangian backtrajectories was used to interpret the impact of the MCS on the aerosol properties. The synoptic conditions and the MCS description are presented in Sect. 2. The airborne sampling strategy is described in Sect. 3 as well as the data set of measured aerosol physical and chemical properties. A mesoscale simulation using an explicit representation of cloud and aerosols processes is then presented in Sect. 4.
2 Experimental strategy and instrumentation

The measurements were performed during the Special Observation Period #1a (SOP1a) of the AMMA experiment on the ATR-42 aircraft operated by the Service des Avions Français Instruments pour la Recherche en Environnement (SAFIRE). The aircraft was based at Niamey airport in Niger for the duration of the AMMA experiment and performed 11 research flights during this SOP of July 2006. Most of the flight patterns were centered on the surface measurement site of Banizoumbou, located 70 km east of Niamey. Sampling occurred on 1 and 2 July 2006, i.e. before and after the passage of a MCS over Niger. The environmental conditions being the same during both flights, any difference in the aerosol properties could be mainly attributed to the impact of the MCS.

2.1 Experimental strategy

Both flights were performed during early afternoon when the convective mixed layer is growing relatively slowly. The flight plans (Fig. 1) were similar, i.e. same geographic location and same period of the day (from 11:30 UTC to 15:30 UTC). During both flights, two vertical soundings were made, at the beginning and at the end of the flight, providing a rapid characterization of the inversion level and of the vertical profile of the thermodynamical and microphysical parameters. On 1 and 2 July, the ATR-42 flew above Banizoumbou along a cross flight-track with horizontal dimension of about 60 km (Fig. 1). The two axes of the cross flight-track were oriented North-South and West-East. Along these axes the ATR-42 flew at different altitudes, in the Boundary Layer (BL) and in the Saharan Air Layer (SAL). During the first flight (1 July), the ATR-42 made five legs on each axis, two of them in the BL (400 m and 850 m) and three in the SAL (1250 m, 2250 m, 2900 m). During the second flight (2 July), the ATR-42 made five legs on each axis, three of them in the BL (400 m, 650 m, 850 m) and two in the SAL (2550 m, 3200 m).
2.2 Instrumentation

Aerosol instrumentation was connected to the French community aerosol inlet (CAI) installed on the ATR-42. This isokinetic and isoaxial inlet has a 50% detection efficiency estimated at 2.5 µm. A condensation particle counter (CPC TSI model 3025) was used to measure total ambient aerosol concentration (CN) every 2 s for particles diameters ranging from 0.003 to 3 µm, with an efficiency of 100% for particles ranging from 0.01 to 2 µm and a relative uncertainty of 5%.

CCN measurements were made by using a static thermal-gradient diffusion chamber (model 100-B, University of Wyoming) which operated similarly to the chamber described by Snider and Brenguier (2000). CCN number concentrations were determined every minute at a supersaturation (SS) between 0.2 and 1% and CCN spectra were measured every five minutes. The instrument was calibrated just before the campaign. Its detection limit is approximately 50 cm$^{-3}$ and the overall measurement error is about ±15% at 1% SS and ±30% at 0.2% SS.

Aerosol size distributions were measured using two instruments. A scanning mobility particle sizer (SMPS) was used to measure the number distribution of aerosol particles with diameters from 0.02 to 0.3 µm. This instrument consisted of a differential mobility analyzer (DMA) as described by Villani et al. (2007) and a CPC (TSI model 3010) for particle detection after the DMA. An optical particle sizer (OPS GRIMM model 1.108) provide particle size distributions in the range 0.3 to 2 µm equivalent optical diameter. Both instruments were also calibrated before the campaign. Data collected were combined to provide a continuous size distribution between 0.02 and 2 µm every 2 min.

A two-stage low-volume impactor with 50% aerodynamic cutoff diameters of 0.2 and 1.6 µm was used to collect particles on electron microscope grids (Matsuki et al., 2003). Some grids were coated in advance with a thick layer of carbon in order to identify the particle shape and elemental composition (single particle analysis). In addition, some aerosol particle samples were collected on nitron-coated nitrocellulose films supported by Ni grids in order to identify nitrate as described by Mamane and Gottlieb (1992).
Sampling duration was between 2 and 5 min depending of the atmospheric load. The morphology and chemical constituents of individual particles were analyzed using a Scanning Electron Microscopy (SEM) with an energy dispersive X-ray spectrometer (EDS). The ATR-42 was also equipped for the measurements of wind, turbulent fluxes, and atmospheric state parameters.

3 Description of the situation on 1–2 July 2006

3.1 Synoptic conditions

Figure 2 represents the height of the geopotential at 850 hPa on 1 July 2006, 12:00 UTC, calculated from the ECMWF analysis. From this figure, two structures clearly appear: the north-west domain is dominated by high geopotential height (1580 m) while a low geopotential height (1470 m) is observed over Egypt and western Libya. The associated zonal pressure gradient is linked to strong northeasterly wind (>8 m s\(^{-1}\)), as seen over Niger and Tchad. Over the center of Niger, the northeasterly air masses of the Harmattan (between 3 and 5 m s\(^{-1}\)) and the southwesterly air masses of the monsoon flux (between 1 and 3 m s\(^{-1}\)) converge and depict the Intertropical Discontinuity (ITD).

Figure 3 is the ECMWF analysis on 1 July 2006, at 12:00 UTC, for the water vapor mixing ratio at 850 hPa. Larger values of the water mixing ratio (>12 g kg\(^{-1}\)) are associated with the monsoon flux while smaller values (<5 g kg\(^{-1}\)) characterize the Harmattan flux. This figure shows an intrusion of moist air corresponding to the advection by the monsoon flux around 14°N 5°E. This intrusion brings humidity and heat near the ITD close to where a convective system occurs at 15:00 UTC. The presence of the wind convergence, humidity and heat at low level allows the initiation and the propagation of a convective system.
3.2 MCS description

The initiation of the studied MCS occurs at the border between Niger and Nigeria (15.04° N and 5.91° E) around 15:00 UTC on 1 July 2006. This zone was highlighted by the ECMWF analysis to be favorable to convection. The MCS-tracking (Mathon et al., 2002) and Massachusetts Institute of technology (MIT) radar allowed us to watch the MCS formation and to follow its trajectory along West Africa (data available on the AMMA Operational Center, aoc.amma-international.org/). Figure 4 is a representation of the MCS-tracking and shows the evolution of the MCS at the time of its formation (Fig. 4a), its maximum activity (Fig. 4b), and one day after its formation (Fig. 4c). The MCS was over Banizoumbou (Niger) at midnight (2 July 2006, 00:00 UTC), and vanished two days later over Guinea.

3.3 Surface conditions

For the AMMA special observation periods, late June–early July can be characterized as a dry period (Janicot et al., 2008). The precipitation record at Banizoumbou (http://aoc.amma-international.org/dir.php?current=20060930&dir=prod_NIAMEY/observation/) indicates that the MCS passage on 1 July corresponds to the first intense precipitation (17 mm). Dry conditions and low vegetation coverage make the Sahelian-Saharan region prone to dust events especially when surface wind speed exceeds a certain threshold. This threshold wind speed mainly depends on surface roughness elements, grain size and soil moisture (Marticorena and Bergametti, 1995).

4 Results

The CN and CCN concentrations and the aerosol size distribution were first used to characterize the evolution of the hygroscopic properties of aerosols by comparing their microphysical properties before and after the MCS passage. For both flights, except during the time of the vertical sounding, the CCN and aerosol spectra were compared to estimate the efficiency of aerosol to serve as CCN. Thus, for the presentation of the following graphs, the red (and blue) color corresponds to data measured before (and after) the MCS passage. Moreover, it was observed that the MCS passage decreases the height of the BL as the inversion descended from 1500 m to 800 m after the passage of the MCS. Consequently, open symbols indicate data measured within the BL while solid symbols correspond to data measured above the boundary layer.

4.1 Aerosol concentration

Figure 5 shows concentrations for the total aerosol (CN) and the CCN (at a supersaturation of 0.6%, CCN$_{0.6}$) measured as a function of altitude during both flights. A strong vertical gradient of CN concentration is observed in the vertical profile. On 1 July, the total aerosol concentration is about 2000 cm$^{-3}$ within the BL and varies between 1500 and 500 cm$^{-3}$ within the SAL. On 2 July, the average concentration is about 2200 cm$^{-3}$ in the BL. Above the BL, the aerosol concentration decreases more rapidly than the previous day to about 500 cm$^{-3}$ at 3000 m. A main difference in the aerosol profiles is noted for a layer located between 1300 and 3000 m where the CN concentration has drastically decreased after the passage of the MCS, suggesting a cloud processing. In the BL, a large variability of the measurements is observed on both profiles, especially during the second flight, after the MCS passage.

The CCN$_{0.6}$ concentration profiles observed during both flights do not show such a clear difference after the MCS passage but a vertical gradient of CCN$_{0.6}$ concentration is also observed throughout the vertical profile. The CCN$_{0.6}$ number concentrations were relatively high in the BL with a mean value of 500 cm$^{-3}$ comparable with val-
ues typically measured in continental environments. Higher in altitude, the concentra-
tions resemble much more what is conventionally regarded as free troposphere values. Above 3200 m, the CCN$_{0.6}$ measurements were not available during the second flight because of a failure of the CCN chamber.

To quantify the changes of the CN and CCN$_{0.6}$ concentrations and characterize the relationship between CCN$_{0.6}$ and total aerosol population in the atmospheric column, the CCN$_{0.6}$/CN ratio was introduced as a tool to follow the intensity of the interaction between aerosols and clouds. When this ratio is 0, no activation of aerosols can occur to form cloud droplets while when this ratio is 1, all aerosol particles can activate to become droplets.

The variation with altitude of the CCN$_{0.6}$/CN ratio is shown on Fig. 6. In the BL, before the MCS passage, the value of the CCN$_{0.6}$/CN ratio ranged between 15 and 32%. In the SAL during the same day, this ratio was less than 20%. After the MCS passage, the CCN$_{0.6}$/CN ratio was less than 25% in the BL while, in the SAL, this ratio increased to 50%. Figure 6 shows a significant increase of the CCN$_{0.6}$ fraction within the SAL after the passage of the MCS, especially in the layer located between 1300 and 3000 m, while the ratio remained unchanged in the BL. Because the enhancement in CCN$_{0.6}$ activity is mainly a function of particle size and chemistry (Köhler, 1926), these properties will be discussed in more detail below.

### 4.2 Aerosol size distribution

Submicron aerosol size distribution were inverted using a software, developed by the LaMP (Laboratoire de Météorologie Physique, University of Clermont-Ferrand, France), taking into account charging probabilities, counting efficiencies of the CPC and transfer functions of the DMA. The number distributions measured during each flight by combining the SMPS and the OPS were averaged to determine a mean dis-
tribution for each layer (BL and SAL). From these mean number distributions, a mean volume distribution was calculated for each layer as well. The mean distributions were fitted by using multi-modal lognormal distributions that best represent the measured
distributions. A maximum of three modes were used to describe the number and volume distributions. The parameters of each mode of the lognormal fit (number or volume concentration, median diameter and geometric standard deviation) are shown in Tables 1 and 2, and the results are plotted on Fig. 7. The total aerosol concentrations estimated by integrating the size distributions from the SMPS and the OPS was lower than the CPC3025 measurements because of the higher detection limit (10 nm versus 3 nm).

Aitken ($D_p<0.1\ \mu m$), accumulation ($0.1<D_p<0.5\ \mu m$) and coarse ($D_p>0.5\ \mu m$) modes were present in both layers before and after the MCS passage. In addition, particles from a nucleation mode ($D_p<0.02\ \mu m$) were also observed in the SAL layer after the MCS passage (not shown). Airborne measurements have already found evidence that new particle formation may occur near cloud boundaries (Roberts et al., 2001).

In general, the shape of the size distributions was almost the same and the contribution of each mode remained unchanged after the passage of the MCS. In particular, for particles with diameter larger than 30 nm, no significant change of the aerosol size distribution appears in the SAL between both flights. This result suggests that the evolution observed in the SAL regarding the CCN fraction cannot be attributed to a particle growth in this layer due to the passage of the convective system. Therefore, since the particle size cannot explain the enhancement of the CCN ratio observed in the SAL, more knowledge on the chemical composition of the particles is necessary.

As shown by Fig. 7, the volume distribution correlates well to the OPS measurement, as the concentration of particles with diameter larger than $0.5\ \mu m$ increases significantly in the BL after the MCS passage. This observation suggests an increased concentration of dust that can be verified by an analysis of particle composition.

4.3 Single particle analysis

Aerosol samples were collected periodically throughout the flights for analysis of their composition using SEM-EDS. The distribution of the elemental composition of the particles as a function of size was determined by analyzing several tenths of particles...
from four samples. One was taken before the MCS passage in the SAL (at an altitude of 2790 m) and the three others after the MCS passage (one in the BL at 394 m and two in the SAL at 2493 and 3090 m). The analysis distinguished between fine (0.2 μm < D_p < 1.6 μm) and coarse fraction of particles (D_p > 1.6 μm) and the frequency of occurrence of the main elements was determined. It has to be noted that particles smaller than 0.2 μm were not analyzed for their chemical composition by EDS due to the low signal-to-noise ratio when using a beam energy low enough to avoid evaporation of the particles.

As expected, the most abundant component found in the BL after the MCS passage was mineral dust (47%). Most of the X-ray spectra showed an abundance of Al, Si and Fe characteristic of the presence of crustal components. None of these dust particles contained sulfur or chloride (which are common soluble elements measurable by the EDS analyzer) suggesting that the BL was rather enriched with fresh emitted dust. This dust abundance in the BL correlates with the increase of concentration of coarse particles observed in the BL on Fig. 7b. This result indicates that dust particles of the coarse mode emitted during the passage of the MCS are sedimenting in the BL after the passage of the MCS. However, dust is generally thought of as water insoluble. Therefore, its efficiency to serve as CCN is rather low. The processing cannot be fully described from the available observations but the presence of non-hygroscopic particles (like mineral dust) in the BL can explain that the CCN_{0.6} fraction remains the same in both flights. The other main components found on this sample were sulfate particles (26%), biomass burning particles (13%) and carbon particles (4%). The sea-salt particles were present in very low concentrations and the undetermined fraction of particles represented about 10%.

According to the single particle analysis of the fine fraction, a large difference of composition is observed between the samples that were taken on carbon-coated grids in the SAL during both flights. Before the MCS passage, most particles (48%) contain Al and Si with a high percentage occurrence of iron, suggesting a mineral composition. Among the other particles, the sulfate-containing particles represent only 12% of the
total number of particles observed. The samples taken after the MCS passage reveal a large change in the occurrence of components. Elements such as Al and Si were found with lower frequencies in the accumulation mode of particles. Mineral dust particles represented only 18% of the particles while the frequency of occurrence of sulfate particles increased markedly up to 43%. In addition, a significant number of dust particles contained sulfur (33%) or chloride (11%), suggesting that, contrary to what was observed in the BL, the surface properties of the mineral dust particles may have been altered by a coating of soluble elements.

Large differences were also observed between the samples that were taken on nitron-coated grids in the SAL, before and after the passage of the MCS. The particles of the coarse fraction collected on nitron-coated grids were analyzed to identify those containing nitrate (Mamane and Pueschel, 1980). Indeed, the nitron-coated substrate captures and, after octanol vapor exposition, crystallizes nitrate on the substrate surface. Nitrate, like sulfate or ammonium, is water soluble and can increase the efficiency of a particle to serve as CCN. However, several previous experimental analyses have indicated that sulfates are primarily found in the fine mode while nitrates are predominantly found in the coarse fraction (>1 µm) (Dentener et al., 1996; Murphy and Thompson, 1997; Gard et al., 1998; Zhuang et al., 1998). Figure 8a and b show photomicrographs of the nitron-coated substrates used to collect coarse particles on 1 July (at 2790 m) and on 2 July (at 3090 m), respectively. Figure 8b clearly shows the reactivity spot around large particles (visible at the center of the spot) surrounded by elongated fibers that are characteristic of the chemical reaction of nitrate with the nitron film. Figure 8a shows no such reaction of nitron substrate with the particles (Isawa and Ono, 1979; Mamane and Pueschel, 1980; Mamane and Gottlieb, 1992). Therefore, the single-particle analysis indicates the presence of nitrate on coarse particles (mainly mineral dust) collected during the post-MCS period, suggesting a coating of this ionic component. A modification of the surface properties of dust particles by coating of a highly water soluble component like nitrate (Seinfeld and Pandis, 1998) can enhance their hygroscopic properties (Levin et al., 1996). In addition, the high
frequency of occurrence of nitrate in the aerosols of the SAL suggests that most of the coarse particles are probably efficient CCN.

Levin et al. (1996) quoted different processes to explain the coating of soluble materials on core particles: coagulation of particles, gas to particles conversion, cloud processing. In Niger, the major source of nitrate are the marine air masses transported in the BL, from the southwest, by the monsoon flux. However, in this study, the particle coated with nitrate were collected at about 1500 m above the top of the BL. Thus, we believe that a physico-chemical mechanism was able, first, to lift up nitrate particles from the BL to the SAL, and secondly to mix particles and nitrates together.

Finally, all the observations described above have shown that strong aerosol modifications (concentration profile, CCN$_{0.6}$/CN, mixing state) have been caused by this mechanism. A main difference in the aerosol profiles is shown for a layer located between 1300 and 3000 m where the CN concentration has drastically decreased after the passage of the MCS. In this layer, a significant increase of the CCN$_{0.6}$ fraction has also been observed during the post-MCS period. Moreover, the results of the elemental composition analyses of individual particles collected in this layer after the MCS passage have shown a higher contribution of sulfate, nitrate and chloride to the total aerosol (in internal and/or external mixing with mineral dust), all of which being soluble and efficient CCN. We thus expect that MCS processing acted as a physico-chemical mechanism that is responsible for the enhancement of the CCN$_{0.6}$/CN ratio observed in the SAL during the post-MCS period. Thus, to complement the observations and interpret the results, a simulation exercise was carried out.

5 Numerical modeling

5.1 Model description

To interpret these results, the meso-scale, non hydrostatic atmospheric model MesoNH was used in this study. This model has been jointly developed by CNRM (Meteo...
France) and Laboratoire d’Aérologie (CNRS) (Lafore et al., 1998). MesoNH simulates small scale (LES type, horizontal resolution of a few meters) to synoptic scale (horizontal resolution of several tens of kilometers) and can be run in a two-way nested mode involving up to 8 nesting stages. Different sets of parameterization have been introduced for convection (Bechtold et al., 2001), cloud microphysics (Pinty and Jabouille, 1998; Cohard and Pinty, 2000), turbulence (Bougeault and Lacarrère, 1989), biosphere-atmosphere thermodynamic exchanges (ISBA) (Noilhan and Mahouf, 1996), urban-atmosphere interactions (Masson, 2000), lightning processes (Barthe et al., 2005), gaseous chemistry (Suhre et al., 1998; Tulet et al., 2003) and aerosol chemistry (Tulet et al., 2005).

Mineral dust emissions are parameterized by Grini et al. (2006). In this parameterization, the three dust aerosol populations proposed by Alfaro and Gomes (2001) are generated and transported by the ORILAM lognormal aerosol scheme (Tulet et al., 2005). Regarding emission processes, dust aerosols are mobilized using the Dust Entrainment and Deposition model (DEAD) (Zender et al., 2003) which calculates dust fluxes from wind friction speeds. The physical basis of the model is taken from Marticorena and Bergametti (1995) where dust fluxes are calculated as a function of saltation and sandblasting processes. Here, the emission of dust aerosols are calculated directly from ISBA surface parameters, and then sent to the atmosphere consistent with the fluxes of momentum, energy and humidity. Aerosol scavenging is explicitly determined according to a kinetic approach to calculate the aerosol mass transfer in cloud and rain droplets as defined by Seinfeld and Pandis (1997), Pruppacher and Klett (1978), and Tost et al. (2006). The in-cloud mass aerosol transfer into rain droplets by autoconversion and accretion processes have been introduced as described by Pinty and Jabouille (1998). The sedimentation of aerosol mass included in raindrops has been solved using a time splitting technique with an upstream differencing scheme of the vertical sedimentation raindrops flux. The release of aerosols into the air due to rain evaporation is assumed to be proportional to the water evaporated (Chin et al., 2000).
MesoNH uses the radiation code of ECMWF (Fouquart and Bonnel, 1980; Morcrette and Fouquart, 1986; ECMWF, 2004) which computes the radiative fluxes of shortwave and longwave radiations. Clouds and aerosols in the shortwave are taken into account using the Delta Eddington transformation (Joseph et al., 1976). Regarding the shortwave effect, a refractive index of the dust aerosols was assigned according to that measured over the ground site of Djougou (Benin) during the AMMA campaign (Mallet et al., 2008). To determine the origin of the simulated air masses, the model uses lagrangian backtrajectories as detailed by Gheusi and Stein (2002).

5.2 Simulation configuration

The simulation begins at 00:00 UTC on 29 June 2006, and ends at 00:00 UTC on 3 July 2006. Three two-way nested grid domains were performed. The large domain (36 km resolution) between 3.1°S and 31.7°N in latitude and 25.64°W and 35.64°E in longitude, gives a large scale synoptic view of west Africa. The first embedded domain (12 km resolution) is centered over Northwest Nigeria and covers a large part of the AMMA campaign area (latitudes 4.3°N and 17.6°N and longitudes 4.19°W and 16.24°E). The smallest embedded domain (3 km resolution) gives a fine scale view of the Niger between 11.80°N and 16.82°N in latitude and 0.10°E and 8.31°E in longitude. The vertical resolution is composed of 60 stretched vertical levels reaching the altitude of 34,000 m; 30 levels are located in the boundary layer between the surface and 2000 m.

Initialization and lateral boundary conditions of the large domain were taken from the ECMWF analysis. Vegetation types came from the ECOCLIMAP data base (Masson et al., 2003). The soil wetness index (SWI) was computed in offline mode using low level atmospheric data from the ECMWF forecast model merged with satellite precipitation

data from the EPSAT-SG product (REF) and downwelling shortwave and longwave 
atmospheric radiative fluxes from the LAND-SAF product (Trigo et al., 2008)\textsuperscript{3}. This 
forcing data was then used as an upper boundary condition to drive the ISBA land 
surface model (Noilhan and Mahouf, 1996) from 2002–2006 in order to obtain a soil 
moisture state which is more realistic than that obtained from the operational NWP 
model. Boone and deRosnay (2007) describe the forcing data and the methodology in 
more details.

5.3 Comparison between simulation and observations

Figure 9a gives the Meteosat infrared picture on 2 July 2006 at 00:00 UTC and Fig. 9b 
shows the brightness temperature (Chaboureau and Pinty, 2006) simulated by Meso-
NH on 1 July 2006 at 19:00 UTC. Both simulations and MSG images show the develop-
ment of a multiple convective system over the Niamey region. The two more intense 
systems are observed north and east of Banizoumbou. To follow their movement, the 
trajectory analysis has been superimposed to the satellite image. The northern MCS 
triggered east of the domain of Fig. 9b and propagated from north of Niamey to Burkina 
Faso (west of the domain). The second system observed east of Banizoumbou moved 
slowly and disappeared south of Niamey. In our simulation, the two modeled systems 
appear in the Niamey region 5 h sooner than those observed on the brightness temper-
ature map. The northern MCS evolves similarly to the observed one, but moves a bit 
more south, passing over Banizoumbou and Niamey whereas the observed trajectory 
of this system remains north of these locations. The MCS located east of Banizoum-
bou is simulated 0.5° too far east and its propagation is quite different to the observed 
one. Instead of disappearing, the system merges with the northern one when passing 
south of Niamey. It is reasonable to estimate that the cycle of life of these two main

\textsuperscript{3}Trigo, T. F., DaCamara, D. F., Viterbo, P., Roujean, J. L. Olesen, F., Camacho de Coca, F., 
systems is close enough to reality to use the simulation as a realistic tool to interpret the data. Nevertheless, the time delay of 5 h between simulation and observation will be taken into account in the following comparison.

The Atmospheric Radiation Measurement (ARM) ground site at Niamey airport (www.archive.arm.gov/) was an observation site to analyze the MCS passage and validate the simulation results. Figure 10 shows the evolution of temperature (a), pressure (b), wind speed (c) and accumulated rain water (d) observed and simulated by MesoNH. The 5 h of delay between the real and the simulated MCS’s passage are clearly retrieved on the surface field over Niamey by a decrease of the surface pressure and temperature, and an increase of surface wind and accumulated rainfall. From 20:00 UTC to 00:00 UTC on 1 July, the modeled and observed temperatures decrease (−14 °K and −9 °K, respectively) as the result of a MCS downdraft that generates a gust front over Niamey (respectively 15.3 m s⁻¹ and 14.8 m s⁻¹, Fig. 10c). On Fig. 10c, we clearly see the passage of both MCS, at 19:00 UTC and 23:00 UTC, when the averaged wind speed is about 15.3 m s⁻¹ and 14.2 m s⁻¹, respectively. The ARM data, in accordance with the MCS trajectories seen on Fig. 9, indicates only one MCS passage at 00:00 UTC over Niamey. Before and after the MCS passage, the observed and modeled wind speed are between 1 and 6 m s⁻¹. It is important to note that the temperature decrease and the surface wind speed generated by the gust front are similar in the observation and in the model. The modeled precipitation rate (3.4 mm) (Fig. 10d) confirms only one MCS passage over Niamey. The delay observed between measured and simulated precipitation rates indicates that only the northern MCS precipitated. The simulated precipitation rate is higher than the observed precipitation rate (2.2 mm) suggesting that Niamey is directly affected by the convective core of the system in the simulation and mostly by the stratiform part of the system in reality.

The modeled and the observed atmospheric pressures (Fig. 10b) are consistent during the 15 first hours of the 1 July 2006. Just before the MCS passage, the pressure value decreases to 984.3 hPa. From 1 July 18:00 UTC to 2 July 12:00 UTC, the modeled pressure is systematically underestimated. We can note that the MCS passage
perturbs the atmospheric pressure. 17 h after the MCS passage, the modeled pressure compares well again with the observed pressure.

The aerosol number concentration measured as a function of altitude with the SMPS and the OPS during the ATR flight on 2 July between 12:00 UTC and 16:00 UTC (i.e. 12 to 16 h after the MCS passage) were averaged and converted to mass concentration in order to give the mean profile of aerosol mass concentration for the zone considered (i.e. 0.5° around Banizoumbou; see Fig. 4). These data are compared to the vertical profile of dust mass concentration simulated over Banizoumbou on 2 July at 07:00, 08:00, 09:00 and 10:00 UTC, i.e. 12 to 16 h after the passage of the simulated MCS (Fig. 11a). A size cutoff of 2.5 μm similar to the 50% detection efficiency of the aerosol inlet was applied to the calculation in order to compare simulated aerosol profiles to airborne observations within a range of particles <2.5 μm. In the boundary layer, between the surface and 900 m of height, the observations highlight a well-mixed layer with a steady concentration value of 60 μg m⁻³. In the other hand, simulated dust mass concentrations range from 200 μg m⁻³ at 07:00 UTC (not shown) to 50 μg m⁻³ at 10:00 UTC within a boundary layer of 450 m in thickness. This corresponds to a mean temporal value of 100 μg m⁻³. This overestimation can be interpreted as the consequence of the difference between the BL height observed between 12:00 and 16:00 UTC and that simulated between 07:00 and 10:00 UTC. Indeed, in our simulation the aerosols present in the boundary layer are confined in the 450 first meters above the surface instead of being more diluted in a 900 m thick layer. Applying a dilution factor of 2 corresponding to the difference between both boundary layer thickness, the mean computed aerosol concentration (50 μg m⁻³) becomes comparable to the observation. Above the boundary layer, two distinctive layers are observed. The first one, between 900 and 2800 m, corresponds to a well-mixed layer with an aerosol mass concentration ranging between 27 and 35 μg m⁻³. The upper layer, separated from the intermediate layer by a maximum of 42 μg m⁻³ at 2800 m, is characterized by a linear decrease of the dust concentration with altitude. The simulation results provide a similar vertical structure with the presence of three layers (Fig. 11a). An intermediate
layer is also reproduced above the boundary layer with a quite constant concentration of aerosol of 25 μg m\(^{-3}\) at 700 m and 35 μg m\(^{-3}\) at 2000 m. Taking into account that this simulated layer is 400 m lower than the observed one, it is remarkable to note that the model is able to reproduce the same thickness of about 2000 m and the order of magnitude of the aerosol mass concentration. Above this layer, the simulation also reproduces the upper layer (corresponding to the SAL) with a decrease of the mass concentration with altitude. An examination of the aerosol vertical profiles simulated over various points around Banizoumbou (at 0.5° north, south, west and east of Banizoumbou) describing the limit of the area sampled by the aircraft is also shown on Fig. 11b. Small differences appear in the various simulated vertical structures, confirming that the whole region around Banizoumbou was characterized by the presence of three atmospheric layers 13 h after the MCS passage.

5.4 Air mass origin

The MesoNH tool allows us to determine the air mass origin using the lagrangian back-trajectories analysis of Gheusi and Stein (2002). The interest of this lagrangian approach is to transport position tracers that are able to precisely retrieve the trajectory of the simulated air mass (all implicit processes parameterized in a mesoscale model such as turbulence, convection, surface mixing, have been used to determine the evolution of tracers). Figure 12 shows the obtained backtrajectories ending at various altitudes (500 m, 1500 m, 2000 m, 3000 m and 5000 m) over Banizoumbou on 1 July 2006, 07:00 UTC, before the MCS passage. Two distinct groups are clearly visible. Group 1 is representative of air masses coming from Burkina-Faso that followed the monsoon flux and stayed in the BL. An ascent from the surface occurred near Banizoumbou. The monsoon flux mixed marine aerosols with continental pollution. Group 2 trajectories are characteristic of air masses originated from the Niger-Nigeria border (6.4° E) and that followed the Harmattan flux. Trajectory altitudes at 3000 m and 5000 m were stable during 8 h. Since the origin of the Harmattan flux is desert regions with few anthropogenic influence, these air masses mainly transported dust particles.
Prior to convective activity, these two air masses remained decoupled and established two well marked layers over Banizoumbou. Therefore, over Banizoumbou the upper layer was enriched by dust particles that are mostly hydrophobic, and in the lower layer (monsoon flux) the particles were assumed to be hydrophilic due to the mixing of marine and anthropogenic particles.

Figure 13 shows the air mass backtrajectories ending over Banizoumbou at the same altitudes as Fig. 12 on 2 July 2006, 07:00 UTC, 12 h after the MCS passage. The start points of these backtrajectories were on 1 July 2006, 18:00 UTC and the MCS dissipated at midnight over Burkina-Faso. The monsoon flux (group 1) and the Harmattan flux (group 2) are, respectively, restored in the lower part of the BL and in the SAL. But a new group of air masses (group 3) appears between 1000 m and 2800 m from a region located north of Banizoumbou where they have crossed the MCS a few hours earlier (see Fig. 9). During the first 3 h, one of the air masses falls 4500 m, indicating that the air mass follows the downdraft of the MCS. After the first 3 h, the altitude of the group 3 air masses is stabilized in a layer where the airborne measurements indicated a decrease of CN concentration (see Fig. 5) and an increase of the CCN/CN ratio (see Fig. 6).

5.5 MCS impact on aerosol dust distribution

Figure 14 is an horizontal view of the computed surface mass concentration of dust on 1 July at 19:00 UTC. This pattern was obtained by summing the three populations of particles emitted and transported by the model, and integrated over the whole domain during the passage of the MCS. The area situated north of Niamey and Banizoumbou is characterized by high surface dust concentrations reaching 10 000 µg m\(^{-3}\) and 5000 µg m\(^{-3}\), respectively. Chomette et al. (1999) used the infrared difference dust index and the reanalysis from the ECMWF to determine the threshold wind speed for soil erosion in Sahelian-Saharan regions. In Niger, around Niamey, they found that the wind speed must be higher than 6.5 m s\(^{-1}\) to generate emissions of dust particles. In Fig. 10c, the average surface wind speed over the whole domain is less than this...
threshold of 6.5 m s$^{-1}$ except around the MCS where a downdraft generates a gust front with surface wind speed exceeding 15 m s$^{-1}$. These surface winds are largely strong enough to move soil particles by saltation and generate high dust concentration in surface as simulated by the model in front of the MCS. On the contrary, behind the MCS, precipitation scavenges the particles and inhibits dust generation by increasing the soil moisture. Hence, the surface dust concentration decreases to about 500 $\mu$g m$^{-3}$ after the passage of the MCS.

The backtrajectories of Fig. 13 showed that the air mass sampled by the aircraft between 1000 m and 2800 m on 2 July, 12 h after the MCS passage, has crossed the MCS in a region situated north of Banizoumbou. In order to better understand the processes in which aerosols could be involved during the convection that took place north of Banizoumbou, a vertical cross section of the convective system is presented on Fig. 15. This cross section corresponds to the horizontal trace shown on Fig. 14. Figure 15a displays the vertical distribution of the cloud water mixing ratio (colored areas) and the mixing ratio of rain (isolines).

The simulation shows that the convective system was clearly mature with high concentrations of liquid and ice water (2 g kg$^{-1}$ at 5000 m and 4 g kg$^{-1}$ at 16 000 m, tropopause) and an important overshoot up to 18 000 m. The vertical wind speed reached 20 m s$^{-1}$ in the convective core. Under 5500 m, at altitudes sampled 12 h latter by the aircraft, important precipitations occurred with a maximum of 3.4 g kg$^{-1}$ at 3700 m. The simulation also shows that evaporation is important, with a maximum of 60% of precipitation reaching the surface. Such precipitation generates intense surface winds of 16 m s$^{-1}$ at the base of the convective core, the gust front.

In addition, Fig. 15b gives the vertical dust mass concentration between the surface and 4500 m along the same cross section. In the northern part of this cross section, the computed vertical profile is characterized by dust concentrations of about 500 $\mu$g m$^{-3}$ near the surface and 200 $\mu$g m$^{-3}$ between 600 and 3000 m. As shown before, MesoNH simulates high dust concentrations reaching 5000 $\mu$g m$^{-3}$ in front of the convective core, in a thin layer above the surface. An important fraction of this
surface dust amount (500–1000 µg m\(^{-3}\)) was lifted in the convective ascent. Where precipitation occurred, the dust mass concentration decreases to 150 µg m\(^{-3}\). This important horizontal gradient of aerosol concentration between scavenged and dry zones shows that an important mass of aerosols has been cloud-processed and evaporated as explained by Tulet et al. (2008). Indeed, it is reasonable to assume that the internal circulation of the MCS can mix long range transported dust from the SAL (northern part of the vertical cross section) with fresh dust generated by the MCS gust front and soluble material contained in the monsoon flux of the boundary layer (see surface back-trajectory of Sect. 5.4). This result is similar to the study of Levin et al. (1996) which determined three conditions that involve cloud processing: particles in an aqueous phase, presence of soluble element, and evaporation of droplets. Cloud processing is able to change the hygroscopicity of new particles by modifying their chemical composition (Meng and Seinfeld, 1994). Our results suggest that a cloud processing could occur in this MCS event, given that all conditions were present. First, various types of gas and particles provided by the monsoon flux are mixed in an aqueous environment with mineral dust produced by the MCS itself. The particles are then collected by cloud drops that have originally been nucleated by soluble materials (like nitrates or sulfates) brought by the monsoon flux. Upon evaporation (Fig. 15a), new aerosol particles having an increased ability to act as CCN are formed in the SAL from mineral and soluble compounds and left behind. Finally, the results of this simulation show that the African MCS’s create conditions which are highly favorable for cloud processing. Furthermore, this process can explain the observations made in the SAL after the MCS passage regarding the increase of the CCN/CN ratio and the chemical modification of aerosols by coating of nitrate and other soluble material on coarse particles.

6 Conclusions

This paper describes the increase of aerosol hygroscopicity by aqueous mixing in a mesoscale convective system observed during the AMMA experiment by using a com-
Combination of airborne observations and simulation exercises. Airborne measurements of aerosol characteristics were carried out 12 h before and 12 h after a mesoscale convective system that passed over Banizoumbou, in Niger, on 2 July 2006 at 00:00 UTC. The flight plans were the same in order to avoid differences in environmental conditions. Observations were then explored using a mesoscale model simulation in order to explain the changes observed regarding aerosol properties.

Before the convection event, air mass backtrajectories show that two distinctive layers have been identified over Banizoumbou. In the BL (at low level), air masses came from the monsoon flux. This layer is known to be associated with polluted gas and hygroscopic aerosols. Above the boundary layer, backtrajectories analyses show that the air masses came from the northeastern desert regions and contained mineral dust particles. Before the convection, both layers are disconnected and explain the difference in the CCN/CN ratio observed. Indeed, dust particles in the SAL are mostly hydrophobic while aerosols present in the BL are more hygroscopic.

After the convection event, a third intermediate layer located between about 1000 and 3000 m has been identified with air masses coming from a region north of Banizoumbou. In this new layer, the CN concentration has decreased and a significant increase by a factor of 4 of the CCN fraction is also observed during the post-MCS period in comparison with the same ratio measured in the SAL before the convection. Moreover, the results of the elemental composition analyses of individual particles collected in this layer after the MCS passage have shown a higher contribution of sulfate, nitrate and chloride to the total aerosol (in internal and/or external mixing with mineral dust), all of which being soluble and efficient CCN. On the other hand, in the BL, an increasing of the volume concentration of coarse particles (mode centered on 2.59 µm) was also noted. In addition, the single particle analysis indicates a dominance of freshly emitted dust particles known to be rather hygroscopic. Thus, the CCN/CN ratio is kept steady between the pre-MCS and post-MCS periods.

The simulation shows that this intermediate layer originates from a region where the convective process generated an aqueous mixing of gas, aerosols within the monsoon
flux, and mineral dust already present in the SAL or generated by the MCS gust front. In agreement with Levin et al. (1996) and Meng and Seinfeld (1994), the simulation results show that all conditions inside the MCS are highly favorable to a cloud processing. Indeed, it is reasonable to assume that the MCS creates a soluble element coating around dust particles that enhance their hygroscopic capacities.

Finally, this study shows that convective systems can mix aerosols from different origins and different layers in aqueous environment. Wet processes can modify externally mixed aerosols into an internal mixing of particles for which the hygroscopic characteristics increase. In addition, one peculiarity of the African MCS shown in this study, is its capability to generate its own dust particles in the gust front and modify their physical and chemical properties in the atmosphere. This study also shows that a realistic mesoscale modeling is a major tool to improve our knowledge on the complex aerosol-cloud processes such as those observed during the AMMA campaign.

Acknowledgements. This work has been supported by the African monsoon multidisciplinary analysis (AMMA) project. Based on a French initiative, AMMA was built by an international scientific group and is currently funded by a large number of agencies, especially from France, UK, USA and Africa. The authors wish to thanks the SAFIRE (Service des Avions Francais Instruments pour la Recherche en Environnement) for preparing and delivering the research aircraft (ATR-42 and Falcon-20). The authors are grateful to A. Boone (CNRM) for the ISBA scheme and C. Lac, I. Mallet for the Meso-NH assistance. Special Thanks to G. Roberts.
References


Boone, A. and deRosnay, P.: Towards the improved understanding of land-surface processes and coupling with the atmosphere over west africa, iLEAPS Newsletter, 3, 33–34, 2007. 10073


ECMWF: IFS documentation. Technical report, ECMWF, ecmwf.int/research/ifsdocs/, 2004. 10072
Roberts, G. C., Andreae, M. O., Zhou, J., and Artaxo, P.: Cloud condensation nuclei in the
Tulet, P., Crahan-Kaku, K., Crumeyrolle, S., and Gomes, L.: Mixing of dust aerosols into mesoscale convective system. An examination of the relative importance of downdraft generation and removal scavenging processes observed during the AMMA field campaign, International conference on cloud and precipitation, 2008. 10079

Table 1. Log-normal characteristics of the number spectra shown on Fig. 7a. C represents the concentration of the particle mode (cm\(^{-3}\)), \(\sigma\) is the standard deviation and \(D_p\) is the median diameter (nm).

<table>
<thead>
<tr>
<th>Number distribution</th>
<th>Before the MCS passage</th>
<th>After the MCS passage</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(C) (cm(^{-3}))</td>
<td>(\sigma)</td>
</tr>
<tr>
<td>Boundary layer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mode1</td>
<td>822</td>
<td>1.6</td>
</tr>
<tr>
<td>Mode2</td>
<td>201</td>
<td>1.5</td>
</tr>
<tr>
<td>Mode3</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>Saharan air layer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mode1</td>
<td>161</td>
<td>1.5</td>
</tr>
<tr>
<td>Mode2</td>
<td>360</td>
<td>1.6</td>
</tr>
<tr>
<td>Mode3</td>
<td>9</td>
<td>2</td>
</tr>
</tbody>
</table>
**Table 2.** Log-normal characteristics of the volume spectra shown on Fig. 7b. C represents the concentration of the particle mode ($\mu$m$^3$ cm$^{-3}$), $\sigma$ is the standard deviation and $D_p$ is the median diameter (nm).

<table>
<thead>
<tr>
<th>Volume distribution</th>
<th>Before the MCS passage</th>
<th>After the MCS passage</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C ($\mu$m$^3$ cm$^{-3}$)</td>
<td>$\sigma$</td>
</tr>
<tr>
<td><strong>Boundary layer</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mode1</td>
<td>0.4</td>
<td>1.6</td>
</tr>
<tr>
<td>Mode2</td>
<td>1.3</td>
<td>1.5</td>
</tr>
<tr>
<td>Mode3</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td><strong>Saharan air layer</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mode1</td>
<td>0.05</td>
<td>1.5</td>
</tr>
<tr>
<td>Mode2</td>
<td>1.6</td>
<td>1.6</td>
</tr>
<tr>
<td>Mode3</td>
<td>6.1</td>
<td>2</td>
</tr>
</tbody>
</table>
**Fig. 1.** Horizontal and Vertical ATR-42 trajectories (a and b) on 1 July 2006, at 12:00 UTC and (c and d) on 2 July 2006, at 12:00 UTC.
Fig. 2. ECMWF analysis of the height (m) of the geopotential at 850 hPa and wind direction at 850 hPa on 1 July 2006, at 12:00 UTC. The maximum vector corresponds to horizontal wind speed of 12.6 m s$^{-1}$. 
Fig. 3. ECMWF analysis of the water vapor mixing ratio (g kg\(^{-1}\)) and wind direction at 850 hPa on 1 July 2006, at 12:00 UTC. The maximum vector corresponds to horizontal wind speed of 12.6 m s\(^{-1}\).
Fig. 4. MCS tracking images, (a) 1 July at 15:00 UTC, (b) 2 July at 00:00 UTC, (c) 2 July at 12:00 UTC. Source: aoc.amma-international.org/observation/mcstracking.
Fig. 5. CN total concentration (line) and CCN$_{0.6}$ concentration (dotted line) as a function of altitude. Red color is for data collected on 1 July 2006, and blue one for those collected on 2 July 2006. The open symbols represent the data in the BL and solid ones represent the data in the SAL. Horizontal bars correspond to the variability on average concentrations ($\pm 1\sigma$).
Fig. 6. CCN$_{0.6}$/CN ratio as a function of altitude. Colors, symbols and horizontal bars are as in Fig. 5.
Fig. 7. Average number-size distribution (a) and volume-size distribution (b) observed by the SMPS and the OPS during 1 (red) and 2 (blue) July 2006, in the SAL (top) and in the BL (bottom).
Fig. 8. SEM photographs showing particles collected on the first impactor stage ($D_p > 1.6 \mu m$) on a nitron filter (a) sampling at 2790 m during 1 July 2006 flight and (b) sampling at 3090 m during 2 July 2006 flight.
Fig. 9. Comparison between (a) observed satellite image on 2 July 2006, at 00:00 UTC (source: aoc.amma-international.org/observation/mcstracking) and (b) modeled brightness temperature on 1 July 2006 at 19:00 UTC. On (a) (resp. b), the crosses correspond to the MCS initialisation, the black (yellow) lines correspond to the MCS trajectories and the black (yellow) triangles correspond to the MCS disappearance.
Fig. 10. Modeled (red line) and observed (black line) parameters (a) Temperature (°K) (b) Pressure (hPa) (c) Averaged wind speed (m s$^{-1}$) (d) Accumulated water (mm), from 1 July, 00:00 UTC to 2 July 18:00 UTC.
Fig. 11. (a) Comparison of modeled (lines) and observed (dots) vertical profiles of aerosol ($D_p < 2.5 \mu m$) mass concentration over Banizoumbou, on 2 July 2006, at 08:00 UTC (see text for details). (b) Comparison of modeled vertical profile of the aerosol mass concentration over Banizoumbou, 0.5° north of Banizoumbou, 0.5° south of Banizoumbou, 0.5° east of Banizoumbou and 0.5° west of Banizoumbou on 2 July 2006, at 08:00 UTC (12 h after the MCS passage).
Fig. 12. Backtrajectories ending over Banizoumbou on 1 July 2006, 07:00 UTC. Timestep: 1 h. (a) Horizontal projection. (b) West-east vertical projection.
Fig. 13. Backtrajectories ending over Banizoumbou on 2 July 2006, 07:00 UTC. Timestep: 1 h. (a) Horizontal projection. (b) West-east vertical projection.
Fig. 14. Surface dust concentration (µg m⁻³) on 1 July 2006, 19:00 UTC. Vectors correspond to the surface wind speed. The maximum vector represents 16.2 m s⁻¹.
Fig. 15. Meridional cross section of the convective system (see location in Fig. 14) on 1 July 2006, 19:00 UTC. (a) Colored area represents the mixing ratio for cloud and ice (g kg\(^{-1}\)) and the isoline corresponds to the mixing ratio for rain (g kg\(^{-1}\)). (b) Colored area represents the dust concentration (\(\mu g\) m\(^{-3}\)) and the isoline corresponds to the mixing ratio for rain (g kg\(^{-1}\)). Vertical scales are not the same.