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Size-Resolved Dry Deposition Velocities of Dust Particles: In Situ Measurements and Parameterizations Testing

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Abstract

Bulk and size-resolved particle concentrations were measured at 2.1 and 6.5 m above the soil surface during an intense dust deposition event that occurred in June 2006 in Niger. Bulk concentration measurements were performed using two tapered element oscillating microbalance instruments, and the size-resolved particle concentrations (from 0.3 to 20 μm) using two optical particle counters. The deposition fluxes derived from the bulk concentrations and those derived from the size-resolved ones are in very good agreement. The largest deposition fluxes are recorded when the dust concentrations are maximal. The temporal evolution of the dust deposition flux follows that of the dust concentration. This is not the case of the dry deposition velocities that are most of the time controlled by the wind friction velocity. The results also show that large particles are strongly sensible to rebound when the wind friction velocity is the highest. Size-resolved dry deposition velocities are compared with the rare measurements of dust deposition velocities over bare sandy soils and are confronted to existing parameterizations of the dry deposition velocity. The parameterization of Zhang and Shao (2014, https://doi.org/10.5194/acp-14-12429-2014) appears to be the only one able to reproduce satisfyingly the measured dry deposition velocities on sandy soils in the 1- to 10-μm particle size range. Indeed, unlike others, this scheme considers the desert sandy surfaces as rough surfaces, allowing the interception of dust particles by the sand grains and/or by the small roughness elements present on the surface.

1. Introduction

Wet and dry deposition are the only processes able to remove air-suspended dust particles from the atmosphere. However, dry deposition is the predominant deposition process for dust due to its mass-size distribution dominated by large particles and to the scarcity of precipitation in the source regions as well as along its main transport routes. By comparing simulations performed with seven dust models, the AEROCOM (AEROsol interComparison) project suggested that dry deposition contributes 65–80% of the total dust deposition (Textor et al., 2007). A precise quantification of this sink is thus necessary to constrain the mass budget in dust transport models (e.g., Bergametti & Foret, 2014).

Most deposition data reported in the literature correspond to direct deposition measurements performed with collectors that separate, or not, wet from dry deposition. These data when acquired over a long period of time (e.g., Bergametti et al., 1989; Heimburger et al., 2013; Loï-Jé-Pilot & Martin, 1996; Marticorena et al., 2017; Prospero et al., 2010; Uematsu et al., 1985; Vincent et al., 2016) provide information on the seasonality and on the orders of magnitude of dust deposition fluxes but rarely on the size of the deposited dust. They allow comparisons with the deposition simulated by dust transport models over periods ranging from 1 week to months. However, although a large fraction of the emitted dust mass is dry deposited along the first 1,000 km of transport (e.g., Schütz, 1979), only a limited number of deposition measurements were performed close to the dust source regions (for more details, see Marticorena et al., 2017, and references therein). In addition, very few of these measurements provided relevant information on the dust dry deposition only mainly because of the technical difficulties one must overcome to perform these direct measurements. Indeed, inertial impaction and interception are, with gravitational settling, the key processes controlling the deposition of large particles, and the efficiency of these processes depends strongly on the characteristics of the collecting surface. Unfortunately, the dry collection vessels of the measurement networks are far from being able to mimic correctly the microscale roughness features of...
the most frequent (bare soil, grassland, forest, or water) real deposition surfaces (e.g., Goossens, 2010; Sow et al., 2006).

In dust transport models, dry deposition is typically calculated from a combination of dry deposition velocities and particle size distributions but, as mentioned by Huneeus et al. (2011), “such estimates are prone to large uncertainties which are typically quoted as plus/minus a factor of three (Duce et al., 1991) but which could well be larger.” Most of the dry deposition parameterizations currently used in transport models (e.g., Davidson & Friedlander, 1978; Giorgi, 1986; Petroff & Zhang, 2010; Slinn, 1982; Wesely et al., 1985; Zhang et al., 2001; Zhang & He, 2014; Zhang & Shao, 2014) are based on deposition data obtained in wind tunnel experiments. Indeed, until recently, field experiments allowing the comparison of dry deposition velocity parameterizations with actual in situ data were rare. Ideally, the most efficient method to derive dust dry deposition velocities from in situ measurements would be the eddy correlation technique (e.g., Businger, 1986). It requires measurements of the dust concentration at a frequency large enough (>1 Hz) to separate the turbulent fluctuations of the concentration from the mean value. Unfortunately, most of the instruments currently able to measure particle concentrations at such a frequency are efficient in a size range limited to particles smaller than 1–2 μm in diameter and the deposition velocities derived from this method generally concern only the submicron particle sizes. Moreover, most of the dry deposition velocities derived from the eddy correlation technique were obtained over more or less vegetated surfaces (e.g., Fowler et al., 2009; Petroff et al., 2008). The first attempts to apply such methods to the dry deposition of dust in semiarid regions were those of Porch and Gillette (1977) in Texas and those of Lamaud et al. (1994) in Niger, but again both remained limited to particles in the ~0.1- to 1-μm particle size range.

The alternative to eddy correlation methods consists in using the flux gradient method. The main difficulty with this method consists in measuring the particle concentrations at different heights with a sufficient accuracy because the concentration gradient may be very low (Businger, 1986). This explains why the gradient method was rather used for measuring dust emission fluxes, this type of situations producing generally stronger concentration gradients (e.g., Gillette, 1977; Nickling & Gillies, 1989; Sow et al., 2009) than deposition events.

In this paper, we use data obtained in Niger in June 2006 during a deposition event to determine the dry deposition flux of dust particles. This event was huge and consecutive to dust emission occurring only 12 hr before, providing the opportunity to measure large concentration gradients in different size classes and discuss the role of the different factors in controlling the dust deposition flux and the dry deposition velocity. We then compare the dry deposition velocities derived from our measurements with the ones simulated using two dry deposition parameterizations based on different approaches (Zhang et al., 2001; Zhang & Shao, 2014).

2. Background on Dry Deposition Concepts and Modeling

Different physical processes interact for controlling the dry deposition of dust particles. Among these processes, gravitational settling, turbulent diffusion, and surface collection are generally the most important, and their effects are usually accounted for by the means of equivalent resistances (gravitational resistance, aerodynamic resistance, and surface collection resistance, respectively) that oppose to the deposition.

Gravitational settling is a process operating as long as a particle remains air-suspended and depends mainly on the size and density of particles. A particle falling freely in still air is subject to gravity and aerodynamic drag forces: When these forces are in equilibrium, the particle reaches the so-called particle terminal velocity or gravitational settling velocity that is given by Stokes formula:

\[ V_s = \frac{d_p^2 (\rho_p - \rho_a) g C_u}{18 \mu_a} \]  

where \( d_p \) (in m) is the particle diameter; \( \rho_p \) and \( \rho_a \) (in kg/m\(^3\)) are the particle and air densities, respectively; \( g \) is the gravity acceleration (in m/s\(^2\)); \( \mu_a \) (in kg·m\(^{-1}\)·s\(^{-1}\)) is the air dynamic viscosity; and \( C_u \) is the Cunningham correction factor that accounts for the slipping effect affecting the finest particles, especially the submicron ones.
\[ Cu = 1 + \left( \frac{2 \lambda_a}{d_p} \right) \left( 1.257 + 0.4 \exp \left( -0.55 \frac{d_p}{\lambda_a} \right) \right) \]  

(2)

where \( \lambda_a \) (in m) is the air mean free path:

\[ \lambda_a = 2 \mu_a \left( P \sqrt{\frac{\gamma}{\pi RT}} \right) \]  

(3)

where \( R \) (in J·K\(^{-1}\)·kg\(^{-1}\)) is the gas constant, \( T \) (in K) is the temperature, and \( P \) (in Pa) is the pressure.

When a particle approaches a surface, an aerodynamic resistance and a surface resistance add their effects to the gravitational settling for controlling its dry deposition. The current dry deposition parameterizations mostly differ in the way these resistances, particularly the surface one, are parameterized.

The aerodynamic resistance, \( r_a \), depends on the characteristics of the flow in the atmospheric surface layer. Theoretically, in neutral conditions (i.e., no vertical gradient of potential air temperature), for a steady state and horizontally homogeneous flow and for a given surface, a simple similarity hypothesis allows the description of the vertical distribution of the wind velocity. The vertical gradient of the wind velocity \( du/dz \) is assumed to depend only on the height, \( z \), above the surface as described by a log law (e.g., Panofsky & Dutton, 1984), for \( z \gg z_0 \):

\[ u(z) = \frac{u_*}{k} \ln \left( \frac{z}{z_0} \right) \]  

(4)

where \( u(z) \) is the averaged horizontal component of the wind velocity (in m/s) measured at height \( z \) (in m), \( u_* \) is the wind friction velocity (in m/s), \( z_0 \) is the aerodynamic roughness length (in m), and \( k \) is the von Karman constant. The precise value of this constant is still in debate but is generally taken as 0.4 (e.g., Foken, 2006).

A quite general expression of \( r_a \) has been proposed by Zhang and Shao (2014) for smooth surface and neutral stability conditions:

\[ r_a(z) = \frac{C S c}{k u_*} \ln \left( \frac{z}{z_0} \right) \]  

(5)

where \( S c \) is the Schmidt number (i.e., the ratio of the turbulent viscosity to the particle eddy diffusivity) and \( C \) is an empirical constant. Most dry deposition parameterizations impose a numerical value to the term \( C S c \) that ranges from about 0.6 to 1 (e.g., Slinn & Slinn, 1980; Zhang et al., 2001). For their part, Zhang and Shao (2014) compute \( S c \) by taking the trajectory-crossing effect (e.g., Yudine, 1959) into consideration and thus only assume a numerical value to \( C (0.45) \). These authors also use an expression slightly modified in the case of rough surfaces (the constant \( C \) is suppressed, and a characteristic height of the roughness elements is used instead of \( z_0 \)).

The surface resistance, \( r_s \), depends strongly on the properties of the surface that control its capability to trap definitively the particles. The three main processes influencing the surface resistance are Brownian diffusion, interception, and inertial impaction. The first one concerns mainly the submicron particles, whereas the other two are most important for the deposition of particles in the 1- to 10-μm size range. The surface resistance is the sum of the inverse of these three components weighted by a rebound function to account for the non-perfect sticking of the particles when they hit the surface.

In dry deposition parameterizations, the efficiency of the collection attributable to Brownian motion is expressed as a more (e.g., Petroff et al., 2008) or less (Zhang et al., 2001) sophisticated expression of Schmidt’s number. The particle inertial impaction, that is, the collection of particles on the roughness elements present on a given surface due to the inertia of particles depends on the particle Stokes number, \( S t \), and on the obstacle geometrical characteristics:

\[ S t = \frac{\tau_p u_*}{A} \]  

(6)

where \( A \) is the characteristic dimension of the roughness elements and \( \tau_p \) is the particle relaxation time:
\[ \tau_p = \frac{V_s}{g} \]  

(7)

The collection of particles by interception occurs when the particle passes an obstacle at a distance shorter than its physical dimensions. In dry deposition parameterizations, various expressions are used to describe interception, but all express their efficiency as being directly proportional to the size of the particle and inversely proportional to that of the roughness elements (e.g., Giorgi, 1986; Slinn, 1982; Zhang et al., 2001; Zhang & Shao, 2014).

In almost all dry deposition parameterizations, the terms constituting the surface resistance are weighted by a term accounting for the immediate bounce-off of particles upon deposition and their reentrainment by the airflow. Only particles having a sufficient kinetic energy are liable to rebound when they hit the surface, generally those larger than 5 \( \mu m \) in diameter. The estimation of the rebound of atmospheric particles over natural surfaces is not a well-resolved question. The most common parameterization used to express the fraction of particles sticking to the surface, \( R \), is that derived by Slinn (1982) from an adjustment of the data obtained by Chamberlain (1966, 1967) who studied the deposition of Lycopodium spores and other small particles on natural and sticky grass:

\[ R = \exp\left(-b \sqrt{St}\right) \]  

(8)

Various values of \( b \), generally ranging between 0.2 and 2 are found in the literature (e.g., Giorgi, 1986; Slinn, 1982; Zhang et al., 2001; Zhang & Shao, 2014).

3. Materials and Methods

3.1. Data Acquisition

The experiment was part of the Special Observing Period 1 of the African Monsoon Multidisciplinary Analysis (AMMA) program and took place close to Banizoumbou (13.54°N, 2.66°E; altitude: 237 m), a small village located about 60 km east of Niamey (Niger). The site is a Sahelian remote place mainly devoted to agriculture, that is, a mosaic of cultivated fields and fallows. The agricultural field on which the experiment was set up was one of the largest in the area. The sandy soil was almost bare: At the time of the experiment (mid-June), most of the plant residues from the previous year have been decomposed or grazed. Indeed, as observed by Abdourhamane Touré et al. (2011), the crop residue estimated at the end of the dry season was very low on this field, less than 100 kg/ha, covering about 1% of the surface.

The measurement station consisted of two instrumented platforms: a container buried underground and a tower located 3.8 m north of the container. The two platforms were installed near the center of the field in order to ensure a maximal fetch in the dominant wind directions. The fetch varied between 190 m in the northeast direction and 575 m in the south direction.

3.1.1. Aerosol Measurements

3.1.1.1. Aerosol Sampling

The design of the station was thought to perform dust flux measurements. Thus, the aerosol sampling flux was performed using two identical inlet systems called IPC (isokinetic particle collector) positioned at 2.1 and 6.5 m above the soil surface for the container and the tower, respectively. The inlets were designed for the AMMA field campaigns to avoid any size distribution bias due to particle collection.

They consist of a sampling head (56-mm diameter) in which the airflow is kept constant at 50 m\(^3\)/h by a rotating pump controlled by a volume flow controller inserted in the airflow. In order to optimize the collection of large particles, the sampling head is equipped with a wind vane maintaining the opening always faced to the wind. The sampling head is followed by a 110-mm divergent where particles are decelerated and transported into a vertical cylindrical chamber (150-mm diameter, 2.7 m high) in which seven sampling lines are inserted. These seven sampling lines can be connected to different online instruments or offline collectors. The diameter of each sampling line has been designed to match the nominal flow rate required by the connected instrument. Calculations indicate that particles of aerodynamic diameter of about 40 \( \mu m \) are transported with 50% efficiency at the entrance of the sampling lines. These two sampling heads had
been carefully intercompared at the beginning of the AMMA campaigns. For details, see Rajot et al. (2008) and Sow et al. (2009).

### 3.1.1.2. Particle Concentration Measurements

Two tapered element oscillating microbalances (TEOM®, model 1400a; Patashnick & Rupprecht, 1991) were used to measure simultaneously on each platform the total suspended particle mass concentration with a flow rate of 3 L/min. The sample stream was preheated to 50 °C before entering the mass transducer so that the measurement was always performed at very low and constant humidity. The selected averaging time for data acquisition was 2 min.

### 3.1.1.3. Particle Size Measurements

The number size distribution (cm$^{-3}$) was also measured simultaneously on each platform by using two optical particle counters (OPCs; model 1.108, GRIMM® Aerosol Technik GmbH & Co., Ainring, Germany). These OPCs allowed the measurement of the number concentration in 15 optically equivalent diameter size classes over a large particle size range (0.3–0.4, 0.4–0.5, 0.5–0.65, 0.65–0.8, 0.8–1.0, 1.0–1.6, 1.6–2.0, 2.0–3.0, 3.0–4.0, 4.0–5.0, 5.0–7.5, 7.5–10, 10–15, 15–20, and >20 μm). Aerosols were sampled inside the OPCs at a controlled volume flow rate of 1.2 L/min, and two optical sensors provided near-real-time particle number concentration measurements at a maximum logging rate of 0.17 Hz (or 6 s). Averaging time was 1 min. We derived from the OPC measurements the volume of particles by size classes by assuming that particles are spherical. Because of very low and noisy counts, we do not use the last channel (>20 μm in diameter).

These measurements of the number size distribution were performed with maximum care. Prior to the experiment itself, we cross-calibrated the two OPCs channel by channel by using them to measure the size-resolved concentration of ambient particles at the same height and for a duration of 20 days. In all channels except the coarsest one, an excellent correlation was found for data averaged over 2-min time steps ($R = 0.93$ for the coarsest channel and $R > 0.98$ for all the others). The slopes of the regression ranged between 0.87 and 1.13, depending on the channel. We also intercompared the two OPCs once they were located at their final heights by selecting periods of the day (mainly afternoons) between 1 and 15 June 2006 without dust emission but with enough convection to assume a well-mixed surface layer. The results also presented similar correlation coefficients and slopes of the regressions close to those determined prior to the experiment except for two channels (3–4 and 4–5 μm) for which the slopes previously found were lower (0.896 and 0.947 instead of 1.037 and 1.128). Thus, the slopes determined in gradient positions were subsequently used to make the concentrations measured at the two levels consistent.

At 2.1 m, a nephelometer (TSI®, model 3563) was used to measure with a 2-min temporal resolution and at three wavelengths (450, 550, and 700 nm) the aerosol scattering coefficients ($\sigma_{\text{neph}, \lambda}$ in m$^{-1}$) integrated between 7° and 170°. At a given wavelength, these scattering coefficients are extensive quantities that increase with the aerosol concentration and also depend on its particle size distribution and refractive index. Another important aerosol characteristic derived from nephelometer measurements is the spectral dependence of the scattering coefficients. This dependence is an intensive parameter in the sense that it depends on the aerosol-relative size distribution and refractive index but contrary to $\sigma_{\text{neph}, \lambda}$ not on the aerosol concentration. In the visible spectrum, $\sigma_{\text{neph}}$ can be described as a power law of $\lambda$:

$$\sigma_{\text{neph}, \lambda} = \alpha A_{\text{neph}}^{-A_{\text{neph}}}$$

(9)

where $\alpha$ is a constant and $A_{\text{neph}}$ is the nondimensional Angström exponent. We compute $A_{\text{neph}}$ from the nephelometer measurements at 450 and 700 nm:

$$A_{\text{neph}} = - \log \left( \frac{\sigma_{\text{neph}, 450}}{\sigma_{\text{neph}, 700}} \right) / \log(450/700)$$

(10)

Values of $A_{\text{neph}}$ were used to provide a complementary information on the particle size since the Mie theory shows that $A_{\text{neph}}$ increases when the size of the aerosol particles decreases (for details, see Alfaro et al., 2003; Anderson & Ogren, 1998).
3.1.1.4. Consistency of the Aerosol Measurements

We tested the consistency of the 2-min measurements performed with the TEOM® and OPC instruments. Figure 1 shows a very good agreement between the volume deduced from the OPC measurements (i.e., the sum of the volumes computed by channel) and the particulate matter concentration as measured by the TEOM® instruments ($R > 0.98$ for $n = 602$). The slope of the regression, $a = 2.266 \text{g/cm}^3$, corresponds to the density of the dust particles, a density close to that of the principal minerals (quartz, clays, calcite, feldspars, etc.) constituting mineral dust. The densities derived specifically from the 2.1 and 6.5 m couples of OPC and TEOM® data are very close (2.264 and 2.259 g/cm$^3$, respectively, i.e., $\pm 0.22\%$). These values of the dust density are also consistent with those obtained by Sow et al. (2009; 2.38 g/cm$^3$ $\pm 0.24$) on the same site, using the same method, but during dust emission events. Note that these densities are “equivalent” mass densities. Indeed, they are derived from (i) volumes based on the assumption that dust particles are spherical while various studies suggest that the aspect ratio for African dust particles is close to $1.5–1.7$ (e.g., Chou et al., 2008; Kandler et al., 2007; Klaver et al., 2011) and (ii) on diameters that are optically equivalent particle diameters. In the following, the equivalent mass density of 2.27 g/cm$^3$ is used to convert number concentrations into mass concentrations.

3.1.2. Wind Erosion Measurements

A saltiphone (Eijkelkamp, Giesbeek, The Netherlands) located 5 m from the isokinetic particle collectors was used for identifying the periods during which saltation occurred. This instrument works according to the acoustic measuring principle (Spaan & van den Abeeleb, 1991). Saltating grains are counted at 9 cm above the soil surface, and the digital output signal is cumulated over a 10-s time step and is registered by a data logger.

3.1.3. Meteorological Measurements and Data Processing

3.1.3.1. Instrumentation

The wind and temperature profiles were measured from a meteorological mast equipped with a wind vane (W200P Vector Instrument®), five anemometers (A100R Vector Instrument®), and four temperature probes (ASPTC, Aspirated Shield With Fine Wire Thermocouple type Chromel Constantan; Campbell®) positioned above ground level at 0.35, 0.8, 1.5, 2.32, 4.7 m and 0.59, 1.28, 2.11, and 4.48 m, respectively. The data were scrutinized every 10 s and averaged over a 1-min period.

3.1.3.2. Data Processing

As mentioned in section 2, in neutral conditions, the vertical gradient of the wind velocity can be described using equation (4). In nonneutral conditions, correction functions are added to this equation to account for diabatic states:

$$u_z = \frac{u^2}{C_3 k} \ln \frac{z}{z_0} - \psi_m \left( \frac{z}{L} \right) + \psi_H \left( \frac{z}{L} \right)$$

(11)

where $\psi_m$ is the momentum stability function and $L$ is the Monin-Obukhov length (Monin & Obukhov, 1954), reflecting the thermal stability of the surface boundary layer:

$$L = \frac{u^2}{k \theta} \frac{T_s}{T_0}$$

(12)

where $T_0$ is the surface temperature (in K) and $T_s$ is the scaling temperature (in K) that characterizes the production of eddy by buoyancy.

A formulation similar to equation (11) can be written for describing the vertical profile of temperature:

$$\Delta T = T_z - T_{z_2} = \frac{T_r}{k} \left[ \ln \frac{z_2}{z_1} - \psi_H \left( \frac{z_2}{L} \right) + \psi_H \left( \frac{z_1}{L} \right) \right]$$

(13)

where $\psi_H$ is the heat stability function.

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Figure 1. Comparison between the total suspended particulate mass measured by tapered element oscillating microbalance (TEOM®; g/m$^3$) and the total particulate volume derived from the number size distribution measured by optical particle counter (OPC; cm$^3$/m$^3$) assuming spherical particles. Data are averaged over 2 min and cover the period 0716–1800 UTC on June 18. Blue circles: measurements at 6.5 m; gray circles: measurements at 2.1 m.
Different forms of the stability functions have been derived and used in the literature. Following the reviews of Högström (1988) and Foken (2006), we use the formulations obtained by Paulson (1970), Dyer and Hicks (1970), and Hicks (1976):

For unstable conditions,

\[ \psi_m = 2 \ln \left( \frac{1 + x}{2} \right) + \ln \left( \frac{1 + x^2}{2} \right) - 2 \arctan(x) + \frac{\pi}{2} \]  
\[ \psi_H = 2 \ln \left( \frac{1 + y}{2} \right) \]  
\[ x = \left( 1 - 15 \frac{z}{L} \right)^{1/4} \text{ and } y = \left( 1 - 15 \frac{z}{L} \right)^{1/2} \]  

and for stable conditions (Webb, 1970),

\[ \psi_m = \psi_H = -\frac{z}{L} \]  

Practically, \( u^*, z_0, T^*, L \) were computed using the iterative routine optimized by Frangi and Richard (2000) for fitting the above equations to the wind speed and temperature profile measurements. These computations are performed over durations of 20 min (with a time slipping of 5 min) because it has been shown that such a duration is required for integrating the major time scales of turbulence occurring in the surface boundary layer (e.g., Dupont et al., 2018; Wieringa, 1993). During the deposition event (18 June 2006, 0730–1730 UTC, i.e., local time minus 1 hr) the surface wind direction was south-southwest \((205^\circ \pm 30^\circ)\). This allows the fetch to be equal to 88 times the height of the higher particle sampler \((i.e., 575 \text{ m}/6.5 \text{ m})\). The aerodynamic roughness length of the surface is very low and allows satisfying easily the limiting value \( \frac{z}{z_0} \geq 100 \) above which the profiles are valid \((\text{the height of lower anemometer being } 0.35 \text{ m})\).

The fitting procedure is applied only when the wind speed is larger than 1 m/s. Furthermore, following Marticorena et al. (2006), the quality of the inversion procedure is considered as satisfying only when the difference between the computed and measured profiles is less than 5% for wind speed and less than 0.05 K for temperature. The results of this inversion allow the computation of additional parameters useful for describing the stability conditions, that is,

\[ R_i = \frac{z_m}{L} \text{ for } R_i < 0 \text{ and } \frac{z_n}{L} = \frac{R_{izn}}{1 - 5R_{izn}} \text{ for } 0 \leq R_{izn} < 0.2 \]  
\[ \text{with } z_m = (z_{\min} - z_{\max})^{0.5} \]  

where \( C_p \) is the air isobaric heat capacity \((J \cdot kg^{-1} \cdot K^{-1})\).

### 3.1.3.3 Air Mass Trajectories

Forward air mass trajectories were computed using the National Oceanic and Atmospheric Administration (NOAA) HYbrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Rolph et al., 2017; Stein et al., 2015). We selected the matrix option that allows running a grid of trajectories evenly spaced around a point source. We also used the HYSPLIT dispersion model to trace the dust plume. Both models used the 1° × 1° latitude-longitude grid, National Centers for Environmental Prediction’s Global Data Assimilation System wind field reanalysis.
4. The 17 June 2006 Dust Emission Event and Its Subsequent Deposition

4.1. Meteorological Situation

In the Sahel, most of the wind erosion events occur from May to mid-July when the protection of the soil surface by vegetation and crop residues is the lowest and episodes of high wind velocities the most frequent (e.g., Abdourhamane Touré et al., 2011; Bergametti et al., 2017; Bielders et al., 2004; Rajot, 2001). At that time, in West Africa, most of the humidity comes from the South Tropical Atlantic Ocean and is transported northward by the African monsoon. This input of water vapor allows the formation of convective systems of various sizes. When sufficiently developed and well organized, these convective systems are called mesoscale convective systems. Most of them have sizes ranging from tens to hundreds of kilometers (Laing & Fristch, 1993; Marsham et al., 2008; Roberts & Knippertz, 2012), generate gust fronts induced by convective cold pools, and are generally associated with precipitation.

During the experiment, such a convective mesoscale event affected the Banizoumbou region on 17 June 2006 at 2145 UTC. A sudden change in wind direction (from southwest to north-northeast) was observed when the mesoscale convective system arrived. At the same time, the wind speed rapidly increased to 20 m/s (Figure 2) and the saltiphone immediately started recording a huge saltation event. The saltation stopped completely about 30 min after the precipitation had started although the wind speed remained quite high (>10 m/s) until 2245 UTC. Then, the wind speed decreased progressively and the wind direction shifted gradually back to its original south-southwest direction. Significant precipitation (about 7 mm) occurred mainly between 17 June 2150 UTC and 18 June 0140 UTC.

On 18 June, the surface wind speed was rather low (<5 m/s) in the early morning but increased progressively to reach about 7–8 m/s between 0800 and 1030 UTC. This increase of the wind speed in the morning is a well-established pattern in this region and is associated with an increase of turbulence by surface heating, allowing downward mixing of momentum from the nocturnal low-level jet (e.g., Lothon et al., 2008). However, no saltation was observed at that time, probably because the soil was moist (e.g., Bergametti et al., 2016). Then, when the wind direction turned again to north-northwest (≈1300 UTC), the wind speed decreased progressively to reach 2 m/s in the late afternoon. On 18 June, the sky remained cloudy as can be seen on the Moderate Resolution Imaging Spectroradiometer Terra and Aqua images taken respectively at 0950 and 1255 UTC (not shown).

As mentioned above, the wind direction reversed during the night and this allowed the air mass loaded with dust particles to pass again over the Banizoumbou site during daytime on 18 June. The forward trajectories starting from the Banizoumbou region on 17 June at 2300 UTC confirm that the air mass loaded with dust during the erosion event came back to the site on 18 June (Figure 3). The dispersion model clearly illustrates first the southeastward transport of the dust plume and then its turning back to the northwest in the early morning (see Movie S1 in the supporting information).

4.2. Particle Concentrations and Gradient

To protect the aerosol instruments (TEOM® and OPC), the sampling was stopped just after the beginning of the rain. When the rain stopped, the
aerosol inlets and the aerosol instruments were cleaned and turned on again at 0630 UTC (TEOM®) and 0715 UTC (OPC) on 18 June, that is, just in time to record the arrival of the dust plume on the site. At about 0700 UTC, the TEOM® instruments registered an abrupt change in the particulate matter concentration that increased from about 120 μg/m³ to more than 880 μg/m³ in less than 1 hr (Figure 4). Then, the particle mass concentration progressively decreased until 1230 UTC to stabilize at around 80–100 μg/m³.

The negative difference of concentration between the TEOM® instrument measuring at 2.1 m and that measuring at 6.5 m clearly shows that this dust episode corresponded to a huge deposition event. The whole day long, the particulate matter concentration remained significantly larger at 6.5 m than at 2.1 m. The difference was as high as 60 to 100 μg/m³ at the beginning of the event and remained higher than 15 μg/m³ during most of the day (Figure 5).

4.3. Particle Size Distribution

The changes in particle concentrations are concomitant with the changes in the particle size distribution. The fact that the Angström coefficient measured by the nephelometer suddenly became very low and even negative (≈−0.4) around 0700 UTC clearly indicated the arrival of a dust air mass characterized by the presence of coarse particles (Figure 6). Afterward, this coefficient increased slowly until midday to reach about 0.15, a value slightly positive but still indicative of the presence of large particles although finer than those carried by the air masses in the early morning.

The OPC measurements allow describing more quantitatively this evolution of the particle size distribution (Figure 7). Particles having a diameter between 3 and 10 μm dominate (>90%) the particle mass size distribution from 0730 to 1130 UTC. After that and until 1730 UTC, in agreement with the evolution of the Angström coefficient, the abundances of the largest particle size classes decrease slowly and the mass size distribution is dominated by the particles in the 1.5- to 5-μm particle size range. The contribution of the submicron particles to the mass size distribution is very low in the early morning (<3%) but progressively increases without exceeding 13%. The presence of a significant amount of large particles indicates that the dust cloud reaching the site in the early morning is not very aged. When the day progresses, the dusty air masses arriving on the site are more aged and a significant part of their coarsest particles has been probably lost on the way.

5. Results

5.1. Meteorological Conditions During Dust Deposition

5.1.1. Stability Conditions

The wind friction velocity, $u_*$, and the temperature scale, $T_*$, can be used to trace the origin (dynamical and/or thermal) of the turbulence controlling the atmospheric stability. Their time evolution on 18 June and that of the sensible heat flux, $H_s$, are reported in Figure 8.

The wind friction velocity slowly increases from 0600 to 0800 UTC, stabilizes around 0.3–0.4 m/s until midday and then decreases to reach 0.2 m/s at the end of the afternoon. As mentioned before, the higher values of the wind friction velocity in the morning likely result from the strengthening of the surface wind speed in the early morning due to the downward mixing of momentum from the nocturnal low-level jet. However, on 18 June, this enhancement of the surface wind speed is moderate (less than 8 m/s; see Figure 2). Indeed, at that time, the sensible heat flux is close to zero, and this does not favor vertical mixing.

In fact, the sensible heat flux exhibits a nonclassical diurnal pattern: It remains very low (maximum 60 W/m²) during all along the 18 June 2006
with a long stagnation period (0600–1100 UTC) corresponding roughly to the period during which the dust concentration was the largest. Figure 9 shows that the diurnal cycle of the temperature is different on 18 June than it was on the previous or following days. In particular, a sudden and abrupt decrease in temperature (~13 °C in less than 1 hr) occurred on 17 June 2200 UTC, that is, when the mesoscale convective event reached the Banizoumbou experimental site and rain started. Moreover, as mentioned before, the sky remained cloudy on June 18 and the atmosphere very dusty, thus strongly limiting the incoming solar flux at the surface. Thus, the surface temperature was not able to reach as high values as those observed during the morning and early afternoon on the previous or following days.

Figure 10, directly inspired from Affre et al. (2000), summarizes these observations and provides an overall view of the stability conditions encountered during 18 June. Five-minute periods are classified according to the scales of speed ($u_*$) and temperature ($T_*$). In this figure, a network of hyperbolae represents the isovalues of sensible heat flux, whereas the isovalues of the stability parameter $1/L$ correspond to a network of parabola. The isolines corresponding to low absolute values of $1/L$ are located close to $T_* = 0$ corresponding to neutrality. Stability or instability increase with absolute values of $1/L$ and larger values are observed close to the x-axis. Taking into account that our experimental field had a very smooth sandy surface, values of $1/L$ ranging from $<0.02$ to $<0.05$ and $H_s$ was maximum (~50–60 W/m²), but $u_*$ remained sufficiently high (~25–30 cm/s) to produce the dynamical turbulence necessary to counterbalance the low production of thermal turbulence.

Most of the time, dry deposition occurred during neutral or almost neutral stability conditions. Thus, the number of rejected 5-min periods (i.e., periods not satisfying the conditions described in 3.1.3) is null and the stability corrections we had to apply were very limited for the whole day, giving good confidence in the retrieved dynamical parameters.

5.1.2. Roughness Length

The aerodynamic roughness length, $z_0$, is an important parameter that characterizes the capability of the surface to trap the wind momentum. Even if it can only be precisely determined from wind velocity measurements, $z_0$ is linked to the density and height of the roughness elements present on the surface (e.g., Lettau, 1969; Marticorena et al., 1997, 2006).

Because $z_0$ is much more sensitive to doubts and stability corrections, we restricted its determination to the period during which the stability conditions can be considered as neutral or quasi-neutral ($<0.02 < 1/L < 0.01$). These conditions were observed between 0647 UTC and 1217 UTC, the period during which $z_0 = 0.00076 ± 0.00044 m (n = 75)$. This value of the aerodynamic roughness length is typical of the roughness lengths measured in situ or in wind tunnel for bare or almost bare sandy surfaces (e.g., Greeley et al., 1991, 1997; Lancaster, 2004; Marticorena et al., 2006; Xian et al., 2002; Zhang et al., 2014).

5.2. Dust Dry Deposition

5.2.1. Gradient Method for Dust Particle Fluxes

Inside the boundary layer, one can define an atmospheric layer close to the surface where the fluxes vary by less than 10% of their magnitude with height (e.g., Rossby & Montgomery, 1935; Stull, 1988). This surface layer is thus assumed to be a constant flux layer under the assumptions of steady state and horizontal homogeneity. In this layer, the vertical flux of particles, $F_c$, can be separated into two contributions: one linked to the action of gravity on particles, the other resulting from the action of the turbulent and molecular diffusion. This latter flux is connected to the average concentration gradient via the turbulent diffusivity coefficient. The constant vertical flux is then as follows:
\[
F = -K_p \frac{\partial C}{\partial z} - V_s C
\]  
(21)

where \(K_p\) is the eddy diffusivity or turbulent transfer coefficient (in \(m^2/s\)) of the particles, \(\frac{\partial C}{\partial z}\) is the particles concentration gradient (in \(kg \cdot m^{-3} \cdot m^{-1}\)) in the vertical direction \(z\), and \(V_s\) is the gravitational settling velocity as expressed by equation (1). By convention, a downward flux (i.e., deposition) is negative.

As shown by equation (1), the gravitational settling is independent of the concentration gradient, and thus, the measurement of the vertical concentration gradient only yields the diffusive deposition flux, \(F_{\text{diff}}\):

\[
F_{\text{diff}} = -K_p \frac{\partial C}{\partial z}
\]  
(22)

For applying equation (22), it is necessary to assume a similarity between the eddy diffusivity of particles and those of momentum or heat. The relationship between these quantities is well described far from the roughness of the surface, which is to say at an altitude greater than about 100 times the aerodynamic roughness length, \(z_0\) (e.g., Businger, 1986). However, this similarity depends on the particle size, large particles having a greater inertia and being more affected by the gravity force. Thus, they are more prone to follow trajectories different from those of the air parcels. After analyzing various models accounting for this trajectory-crossing effect, Shao (2008) concluded that the eddy diffusivities of particles smaller than 20 \(\mu m\) in diameter and of scalars can reasonably be considered identical when turbulence is sufficiently strong.

Equation (22) can be integrated between two measurement heights \(z_1\) and \(z_2\) (\(z_2 > z_1\)):

\[
F_{\text{diff}} = -\frac{u_* \Delta z}{\ln \left( \frac{z_2}{z_1} \right)} \frac{k}{\Psi_{m,z} + \Psi_{m,\xi}} \frac{\Delta C}{\Delta z}
\]  
(23)

where \(\Delta C\) (in \(kg/m^3\)) is the concentration difference between the heights \(z_2\) and \(z_1\) and \(\Psi_{m,z}\) is the stability correction for momentum transfer at each height.

Shao (2008) pointed out that if the dust flux, \(F\), is constant with height, the diffusive dust flux, \(F_{\text{diff}}\), is not because of the gravitational settling term. Thus, the diffusive dry deposition velocity, \(V_{\text{diff}}\) at a height \(z\), is

\[
V_{\text{diff}}(z) = \frac{-F_{\text{diff}}(z)}{C(z)}
\]  
(24)

\(V_{\text{diff}}(z)\) being downward positive.

The large difference (Figure 5) observed between the particle concentrations measured at 2.1 and 6.5 m allows using equations (23) and (24) to compute the diffusive dust deposition fluxes and the diffusive dry deposition velocities, respectively. Because the evolution of the vertical profiles with height are generally close to logarithmic, the geometric height is commonly used as the height characterizing the profile, that is, the height where the tangent to the profile is equal to the gradient (Businger, 1986). In our case, the geometric height, \(z\), is equal to 3.69 m.

Practically, the diffusive deposition fluxes derived from the bulk concentration measurements are directly computed from the difference in particle concentrations measured by the two TEOM® instruments. In perfect timing with the computation of both \(u_*\) and the stability functions, this computation is performed over...
20-min periods and with a slipping time step of 5 min. The diffusive vertical fluxes derived from the OPC measurements are computed for each size class and then summed to obtain the total diffusive deposition fluxes. The computations are performed for the same 20-min periods as those used for the computation of the diffusive deposition fluxes from the bulk concentration measurements.

5.2.2. Dust Deposition Fluxes

The diffusive dust deposition flux \( F_{\text{diff}} \) deduced from the TEOM* and OPC measurements is reported in Figure 11 for the period 0732–1747 UTC. The time evolution of the deposition flux follows that of the dust concentration: The maximum of the deposition is observed around 0800 UTC, that is, when the dust concentration reaches its maximum and the progressive decrease of the \( F_{\text{diff}} \) is very similar to the decrease of the dust concentration all along the day. The TEOM* and OPC measurements describe a similar deposition pattern, but the maximum deposition flux derived from the TEOM* measurements is slightly lower than the one derived from the OPC measurements (10.8 and 14.6 \( \mu \text{g} \cdot \text{m}^{-2} \cdot \text{s}^{-1} \), respectively).

The contribution of the different size classes to \( F_{\text{diff}} \) for different time periods of 18 June are computed from OPC size-resolved data (Figure 12). When the deposition is the highest (0737–0827 UTC), 70% of \( F_{\text{diff}} \) is controlled by the particles in the medium size range (3–7.5 \( \mu \text{m} \)) when these particles account for only about 45% of the mass size distribution at that time (Figure 6). Surprisingly, the contribution of larger particles (7.5–20 \( \mu \text{m} \)) to deposition is, at that time, less (~28%) than their relative contribution to the mass size distribution (~41%) and their supposedly higher deposition velocity might suggest. At midday (i.e., 1042–1142 and 1247–1347 UTC), the largest particles largely dominate the deposition with a contribution to the total deposition exceeding 70% when the medium size range particles contribute only ~15% to the total flux. This is also surprising because for this period the contribution of the largest particles to the mass size distribution is about 2 times smaller than it was during the morning (0737–0827 UTC) period. At the end of the afternoon (1637–1737 UTC), the contribution of the different size classes to the deposition is more homogeneous, with a rebalancing of the contributions of the large, medium, and small particles in phase with the evolution of the mass size distribution observed at that time (Figure 6).

This comparison of the temporal patterns of the diffusive dry deposition fluxes and mass size distributions clearly confirms that particle size is not the only driver of the dry deposition flux. Other factors such as wind friction velocity also play a significant role in deposition processes as shown by many authors from in situ or wind tunnel measurements (e.g., Lamaud et al., 1994; Zhang & Shao, 2014) and during model intercomparisons (e.g., Kahn & Perlinger, 2017).

5.2.3. Dry Deposition Velocities

Figure 13 reports the diffusive dry deposition velocities derived from the TEOM* and OPC measurements. These diffusive dry deposition velocities were computed by dividing \( F_{\text{diff}} \) by the geometric mean of the particle concentrations between 2.1 and 6.5 \( \mu \text{m} \) for each 20-min averaging period.

The time evolutions of the dry deposition velocities and of the dry deposition fluxes are very different: The maximum in \( V_{\text{diff}} \) (~4.5 cm/s) is not observed when the dry deposition fluxes are the highest but around midday, consistently with the dominant contribution of the large particles to the deposition fluxes at that time. At the maximum of deposition (~0800 UTC), \( V_{\text{diff}} \) is only 1.9 cm/s.

The temporal evolution of \( V_{\text{diff}} \) during 18 June is very different for the 3- to 5-\( \mu \text{m} \) and 7.5- to 15-\( \mu \text{m} \) particle size classes (Figure 14). The diffusive dry deposition velocity for 3- to 5-\( \mu \text{m} \) particles roughly follows the general trend of the friction velocity with a maximum around 0800 UTC and then a more or less continuous decrease, but the evolution of \( V_{\text{diff}} \) for larger particles only matches that of the wind friction velocity after approximately 1045 UTC. Indeed, between 0727 and 1030 UTC the diffusive dry...
deposition velocity of the 7.5- to 15-μm particles is very low (−2.1 cm/s) and even lower than that of particles in the 3- to 5-μm particle size range at the same time (−3.4 cm/s). The maximum values of \( V_{\text{diff}} \) for the 7- to 15-μm particles (−10–12 cm/s) are observed between 1042 and 1157 UTC, that is, not when the wind friction velocity is maximum.

### 5.2.4. Rebound of Large Particles

For a given size class, the main driver of the dry deposition velocity is the wind friction velocity: An increase in \( u_* \) leads to an increase of the surface processes (inertial impaction, interception, and Brownian diffusion) governing the diffusive dry deposition of particles and thus to an increase of the dry deposition velocity. Figure 15 shows that the diffusive dry deposition velocities of fine- and medium-size particles, that is, those <7.5 μm in diameter, are well correlated with \( u_* \). For a given size class higher dry deposition velocities are observed for higher wind friction velocities. But Figure 15 also shows that particles >7.5 μm in diameter have not the same behavior: If a relative good covariation of \( V_{\text{diff}} \) and \( u_* \) is observed until the wind friction velocity reaches about 0.3 m/s, the diffusive dry deposition velocity of these large particles drops sharply for higher \( u_* \) values.

The only process for which the wind friction velocity has a size-selective negative effect on \( V_d \) is rebound, that is, the immediate bounce-off of particles upon deposition and their reentrainment by the airflow. Indeed, various authors (e.g., Paw U, 1983; Paw U & Braaten, 1992; Wu et al., 1992a) have shown that a minimum velocity exists, called the critical rebound velocity, for an impinging particle of a given size for which rebound occurs: Only particles having a sufficient kinetic energy are liable to rebound when they hit the surface. Since the kinetic energy is proportional to the mass of the particle and to the square of its velocity, the particles having a large diameter and moving inside a relatively high velocity airflow are the most affected by this process. When the critical rebound velocity is exceeded, the fraction of particles able to rebound increases rapidly as the particle velocity increases.

Thus, it is highly probable that differences in rebound efficiency explain the observed patterns of the diffusive dry deposition velocities of the different size classes during early morning on 18 June 2006. Indeed, fine- and medium-size particles are not or only little affected by rebound because of their low mass, whereas large particles are, and particularly in the early morning when the friction velocity is the highest. Then, around 1000 UTC, \( u_* \) decreases and becomes probably lower than the critical rebound velocity, the rebound becomes minimal and the dry deposition velocity of the large particles comes back to a temporal pattern similar to that of the fine- and medium-size particles, that is, a covariation with \( u_* \) (Figure 14).

As mentioned in section 2, the estimation of the fraction of rebound of atmospheric dust particles over natural surfaces is not a well-resolved question. Despite Slinn (1982) mentioning that the formulation he proposed to account for rebound (equation (8)) is not satisfying, most of the dry deposition schemes developed during the last 30 years (Giorgi, 1986; Zhang et al., 2001; Zhang & Shao, 2014, among others) used equation (8)
with only changes in the value attributed to the parameter $b$ (generally taken as 1 or 2).

A rough estimate of the rebound can be attempted using the TEOM® data that are less scattered than those of the individual channels of the OPC are. Figure 16a shows that the trend between $V_{\text{diff}}$ derived from TEOM® data and $u^*$ is very similar to those observed for the 7.5- to 10-μm and 10- to 15-μm size classes of OPC (Figure 15). More interestingly, it shows that for large particles, $V_{\text{diff}}$ is nicely correlated with $u^*$ until the wind friction velocity reaches about 33.5 cm/s as can also be seen in Figure 14. Once this value is exceeded, the dry deposition velocity drops very rapidly.

To estimate the fraction of rebound, we introduce the concept of “ideal deposition velocity,” $V_{\text{diff, ide}}$, defined as the dry deposition velocity of the particles if no rebound occurred. This ideal deposition velocity is obtained by extrapolating above 33.5 cm/s the relationship linking $V_{\text{diff}}$ to $u^*$ for $u^* < 33.5$ cm/s. Then, a rough estimate of the fraction of rebound, $F_R$, can be derived from the relative difference between $V_{\text{diff, ide}}$ and $V_{\text{diff}}$:

$$F_R = \frac{V_{\text{diff, ide}} - V_{\text{diff}}}{V_{\text{diff, ide}}}$$

The results (Figure 16b) suggest that rebound could affect up to 80% of deposited (large) particles in the early morning and that this proportion decreased progressively to less than 20% around midday. This also explains why

1. only particles having a diameter lower than about 7.5 μm are well correlated with $u^*$ all along the day,
2. the diffusive dry deposition velocity of particles 7.5–15 μm is lower in the early morning than that of 3–5 μm particles (Figure 14), and
3. the deposition flux is mainly controlled by particles in the 3- to 7.5-μm particle size range between 0737 and 0827 UTC (Figure 12) even though this is the period of the day during which large particles are most abundant.

Our estimates of the fraction of rebound deduced from the measurements are in quite good agreement with the experimental values obtained in wind tunnels by Wu et al. (1992b) for the rebound of 8.6- and 10-μm uranine particles or by Paw U (1983) for glass microbeads, Lycopodium spores, and ragweed pollen. They also confirm that rebound is zero when $u^*$ is below the critical rebound velocity in disagreement with equation (8).

### 5.3. Comparison With Dry Deposition Schemes

As mentioned above, the dry deposition of particles involved different resistances that oppose to deposition. In most of the parameterizations, dry deposition is generally represented as a combination of these resistances in which gravitational settling operates in parallel to the other resistances that operate in series like in an electrical circuit (Hicks et al., 1987; Sehmel, 1980; Slinn, 1982; Wesely & Hicks, 2000; Zhang et al., 2001). However, while this analogy of dry deposition to electrical resistances has many practical advantages, it does not satisfyingly represent the way by which the various deposition processes interact. Indeed, as mentioned in section 5.2.1, contrary to the other resistances that are concentration gradient dependent, gravitational settling is not and thus cannot be assimilated to a voltage. This has been clearly stated by Verkatram and Pleim (1999) who showed that the representation based on electrical analogy is not mass conservative even if it introduces only little difference in the estimations of the dry deposition velocities.

There is a general agreement on how the gravitational and aerodynamic resistances can be represented in dry deposition schemes. In fact, most of the dry deposition parameterizations differ on the parameterization of
the collection resistance on surfaces that are not totally smooth (e.g., Petroff & Zhang, 2010; Slinn, 1982; Slinn & Slinn, 1980; Zhang et al., 2001; Zhang & Shao, 2014).

Recently, Zhang and Shao (2014) proposed a new deposition scheme in which (i) the dry deposition velocity is directly derived from the particle concentration equation and (ii) the surface collection process is accounted for by using the drag partition theory (e.g., Arya, 1975; Raupach, 1992; Shao & Yang, 2005, 2008). One of the specifics of this parameterization that directly concerns deposition of dust particles is that desert bare surfaces (like sandy or silty soils) are not considered as totally smooth surfaces. This allows the interception of dust particles by micro-roughness elements, while interception is set to zero in most other dry deposition schemes that consider desert surfaces as smooth surfaces (e.g., Zhang et al., 2001). Moreover, dry deposition velocity measurements of dust particles were performed in wind tunnel over sandy, silty loam, and Gobi surfaces by the same team (Zhang et al., 2014) in order to serve as a basis for their parameterization.

Thus, we compared our in situ measurements to (i) the wind tunnel data obtained by Zhang et al. (2014) and to (ii) the dry deposition velocities predicted by the parameterizations of Zhang et al. (2001) and Zhang and Shao (2014). Indeed, as mentioned before, these dry deposition schemes are significantly different in the way by which both the deposition processes and their interaction are represented. We do not recall here the details of these parameterizations that are well described in the original papers.

The comparison is made on $V_d$, that is, $V_{diff} + V_s$ where $V_s$ is calculated according to equation (1) and added to the measured $V_{diff}$.

For applying these schemes, the values of some parameters need to be estimated. In the scheme proposed by Zhang et al. (2001), parameters or constants are ascribed to different land use categories. We selected for

![Figure 14.](image)

**Figure 14.** Diffusive dry deposition velocity of particles for the 3- to 5-$\mu$m (red circles) and 7.5- to 15-$\mu$m (gray circles) particle size ranges and wind friction velocity (blue line) during 18 June 2006.

![Figure 15.](image)

**Figure 15.** $V_{diff}$ versus $u^*$ for different size classes on 18 June 2006 (only $V_{diff}$ for which the difference between $C_{6.5} - C_{2.1}$ is >5% are reported; Table S1 in the supporting information provides the number of data used for each channel).
Zhang et al. (2014) are also reported.

Tunnel over sand (blue circles) and sandy loam (green circles) surfaces by period 1007

circles correspond to the 0737

Comparison between measured dry deposition velocities (red

Figure 17.

(a) Evolution of \( V_{\text{diff}} \) derived from TEOM® measurements as a function of \( u_* \) for \( u_* < 33.5 \text{ cm/s} \) (blue circles) and \( u_* > 33.5 \text{ cm/s} \) (red circles). (b) Fraction of rebound estimated as the relative difference between \( V_{\text{diff}} \) deduced from measurements and \( V_{\text{diff}, \text{c}} \) computed using the linear regression of Figure 16a.

Figure 16. (a) Evolution of \( V_{\text{diff}} \) derived from TEOM® measurements as a function of \( u_* \) for \( u_* < 33.5 \text{ cm/s} \) (blue circles) and \( u_* > 33.5 \text{ cm/s} \) (red circles). (b) Fraction of rebound estimated as the relative difference between \( V_{\text{diff}} \) deduced from measurements and \( V_{\text{diff}, \text{c}} \) computed using the linear regression of Figure 16a.

The dry deposition velocities measured for particles greater than 10 \( \mu \text{m} \) in diameter, red circles correspond to the 0737–1002 UTC period and the red squares to the period 1007–1757 UTC. Only \( V_{\text{diff}} \) for which the difference between \( C_{0.5} \) and \( C_{0.1} \) is >5% are accounted for. Dry deposition velocities measured in a wind tunnel over sand (blue circles) and sandy loam (green circles) surfaces by Zhang et al. (2014) are also reported.

these parameters or constants the values recommended for the “desert” (land use category 8) category, except for \( z_0 \) for which we used the value we measured in the field (0.00076 m, see section 5.1). The Zhang and Shao (2014) scheme requires defining the height \( (h_c) \) and the frontal area index \( (\lambda) \) of the roughness elements. As for the application of the Zhang et al. (2001) scheme, we used the \( z_0 \) we measured in the field and then selected a \( \lambda C_0 / h_c \) ratio and \( \lambda \) as close as possible to those reported by Zhang and Shao (2014) for their sand surface (1.35–1.5 and 0.125, respectively). Thus, we selected \( h_c = 6 \times 10^{-4} \text{ m} \). Assuming as Zhang and Shao said that the obstacles are spherical and that \( h_c \) is equal to half their diameter, \( d_v \) leads to \( \lambda = 0.125 \). Thus, with these assumptions, the obstacles on the field in Banizoumbou should have an averaged diameter of about \( 1.2 \times 10^{-3} \text{ m} \), a reasonable estimate if we take into account the large dominance of coarse sand in the Banizoumbou soil (Lafo et al., 2014) and the low abundance of remaining crop residues.

For each scheme, the computations have been made assuming neutral stability conditions. Because the conditions were neutral or near neutral on 18 June 2006 (Figure 10), this choice has no impact on the comparison. The computations were performed for three different wind friction velocities (0.2, 0.3, and 0.4 \text{ cm/s}) covering roughly the range of wind friction velocities observed during the deposition event. The computations were made for \( z = 3.69 \text{ m} \). For purpose of comparison, the wind tunnel data obtained by Zhang et al. (2014) were corrected for height using the method described in their paper.

Figure 17 shows that the measured dry deposition velocities are in good agreement with the dry deposition velocities measured in a wind tunnel by Zhang et al. (2014) over sand and sandy loam for wind friction velocities ranging between 0.3 and 0.49 \text{ cm/s}. Our data also fit well with the Zhang and Shao (2014) scheme, especially for particles in the 0.6- to 10-\( \mu \text{m} \) particle size range, while the Zhang et al. (2001) scheme underestimates by almost 1 order of magnitude the dry deposition velocities for the same particle size range.

The better agreement observed with Zhang and Shao (2014) is due to the fact that this scheme treats bare sandy surfaces as slightly rough surfaces. This explains the increase of the deposition velocity in the 1- to 10-\( \mu \text{m} \) size range as compared to Zhang et al. (2001) who consider the desert surfaces as smooth surfaces and thus set the interception of particles to zero.

This strongly suggests that roughness elements present over the bare soil, even how small, favor significantly the interception of dust particles. Our in situ measurements confirm, in complement to the wind tunnel measurements of Zhang et al. (2014), that this approach accounts probably better for the processes occurring over bare soils than the Zhang et al. (2001) scheme does.

The dry deposition velocities measured for particles greater than 10 \( \mu \text{m} \) are slightly higher than those both schemes predict. Because these particles were subjected to important rebound in the early morning when \( u_* \) was the highest, we distinguished between data obtained for high and low \( u_* \) (red circles and red squares, respectively; Figure 17). We observed that the dry deposition velocities of these particles are closer to the predictions of the dry deposition scheme for high \( u_* \), that is, when the rebound is efficient. On the opposite, for lower \( u_* \) the measured dry deposition velocities are significantly higher than those predicted by dry deposition
schemes. In fact, equation (8) does not account for the existence of a critical velocity for rebound and thus generates rebound whatever the value of $u_*$. For example, in the Zhang and Shao (2014) scheme, rebound accounts for 30–40% for particles in the 10- to 20-$\mu$m particle size range even at low wind friction velocities ($\sim 0.2$/m s). This increases the surface resistance and decreases the dry deposition velocity that tends toward the gravitational settling velocity. Our results clearly suggest that the parameterization of rebound in dry deposition schemes is not satisfying and should be reexamined.

6. Conclusions

A huge deposition event that occurred in Southwestern Niger in June 2006 was sampled over an almost bare surface. The large concentration gradients measured during this event allowed to use with a good confidence the flux gradient method to determine the diffusive deposition fluxes and associated dry deposition velocities. For that, we first compared the data obtained from instruments providing both the bulk and size-resolved deposition fluxes and found a good agreement between the two types of measurements. The stability conditions were near or neutral almost all along the 10 hours the deposition event lasted. The results show that the dust concentrations drive the diffusive dry deposition fluxes: deposition is maximum ($\sim 12$ $\mu g$-$m^{-2}$-$s^{-1}$) when the dust concentration is the highest ($\sim 900$ $\mu g$/m$^3$). This is not the case for the dry deposition velocities. Indeed, for a given particle size class, the dry deposition velocities are mainly controlled by the dynamical conditions as scaled by the wind friction velocity. Indeed, dust particles smaller than 7.5 $\mu$m in diameter follow quite well the temporal evolution of the wind friction velocity with higher diffusive dry deposition velocities observed when $u_*$ is the highest. For larger particles, the behavior is somewhat different: These particles follow the evolution of the wind friction velocity only when $u_*$ was below a threshold ($\sim 33$ cm/s in our case). Above this value, the diffusive dry deposition velocity of large particles dropped sharply when $u_*$ increased: We show that the rebound is responsible for such an abrupt decrease of the dry deposition velocity of particles larger than $\sim 10$ $\mu$m in diameter. For these dust particles, rebound can be so important that their diffusive dry deposition velocity can become lower than that of medium-size dust particles. Moreover, our results suggest that the way by which rebound is accounted for in dry deposition parameterizations is not satisfying, especially because no threshold for rebound is introduced in the existing parameterization and that the dependence of rebound to $u_*$ is too smooth in this parameterization.

The dry deposition velocities we derive from our in situ measurements by summing the diffusive and gravitational deposition velocity are in good agreement with the wind tunnel measurements performed by Zhang et al. (2014). They confirm that dry deposition velocities of dust over bare soils is significantly greater in the 1- to 10-$\mu$m size range than previously expected from “classical” dry deposition schemes. Thus, the comparison we performed between our in situ dry deposition velocities and two dry deposition schemes based on different modeling shows that only the Zhang and Shao (2014) scheme satisfactorily matches our in situ determinations of the dry deposition velocities for the 1- to 10-$\mu$m particle size range. This good agreement is mainly due to a different parameterization of the surface collection resistance in this parameterization. In particular, the interception process accounts for the contribution of the micro-roughness elements present in the surface elements to the dust trapping. Our in situ results confirm the wind tunnel measurements performed by Zhang et al. (2014) that surfaces exhibiting low roughness like desert surfaces cannot be treated as smooth surfaces in regard to the dust dry deposition.

This enhancement of the dry deposition velocity compared to other existing deposition schemes is important because it represents at least half an order of magnitude for the 1- to 10-$\mu$m size range that is by far the dominant size range of the dust mass size distribution.

References


