

The 7–13 March 2006 major Saharan outbreak: Multiproxy characterization of mineral dust deposited on the West African margin

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[1] Mineral dust deposits were collected at Mbour, Senegal, throughout the spring of 2006 and especially during the well-documented March 7–13 large Saharan dust outbreak. During this 7-day period, significant changes in mass flux, grain-size, clay mineralogy and Sr and Nd isotopic compositions were recorded, indicating distinct provenances for the dust transported and deposited during and outside the event. All these terrigenous proxies, as well as freshwater diatom taxa, also showed significant temporal variations during the outbreak, implying contributions from at least two different provenance regions. Tri-dimensional back-trajectories and satellite imaging enabled us to link those distinct signatures to regions increasingly to the southeast within a large area covering Mauritania, Mali and southern Algeria, identified by the Total Ozone Mapping Spectrometer (TOMS) as the main source of the prominent winter/spring plume over the tropical Atlantic. The multiproxy characterization of the March 7–13 dust fall therefore enables us to typify the terrigenous signature of two different regions supplying dust off West Africa, and provide valuable clues for the interpretation of Northeastern Tropical Atlantic Ocean dust sedimentary records in terms of changes in provenance regions and transport systems. Additionally, because dust deposition data are scarce, flux and grain size data obtained in this study, among other parameters such as clay assemblages, provide important constraints for atmospheric transport models and dust deposition budget estimates in this area.

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1. Introduction

[2] It is well established that wind-blown mineral dust plays an active role in the Earth climate system [e.g., *Maier et al.*, 2010]. During its transport, dust modifies scattering and absorption properties of the atmosphere as well as cloud characteristics [*Sassen et al.*, 2003], ultimately affecting the

radiative balance of the Earth and the hydrological cycle [*Arimoto*, 2001]. Dust may also affect climate indirectly via the boosting of the biological productivity in surface waters in iron-depleted areas [*Duce et al.*, 1991; *Martin et al.*, 1991] and/or of the export of organic matter down the water column, both of which having some impact on the uptake of atmospheric CO₂ by marine ecosystems [*Bopp et al.*, 2003]. Yet, despite the potential impact of mineral dust on global climate, dust remains one of the less well-quantified elements of the climate system [*Forster et al.*, 2007].

[3] Dust deposition in particular is poorly constrained. Even in the vicinity of major arid and semi-arid areas, such as the Sahara-Sahel region – the world's largest dust source [*Prospero et al.*, 2002] – there are virtually no data available in the literature. Dust deposition quantification is essential for assessing the role of dust on the ocean realm, however, and for validation of dust transport models [*Tegen*, 2003]. The characterization of present-day dust deposits signatures may also be useful for the interpretation of the marine sedimentary records, especially in the North Eastern Atlantic Tropical Ocean, where most of the terrigenous fraction is of aeolian origin.

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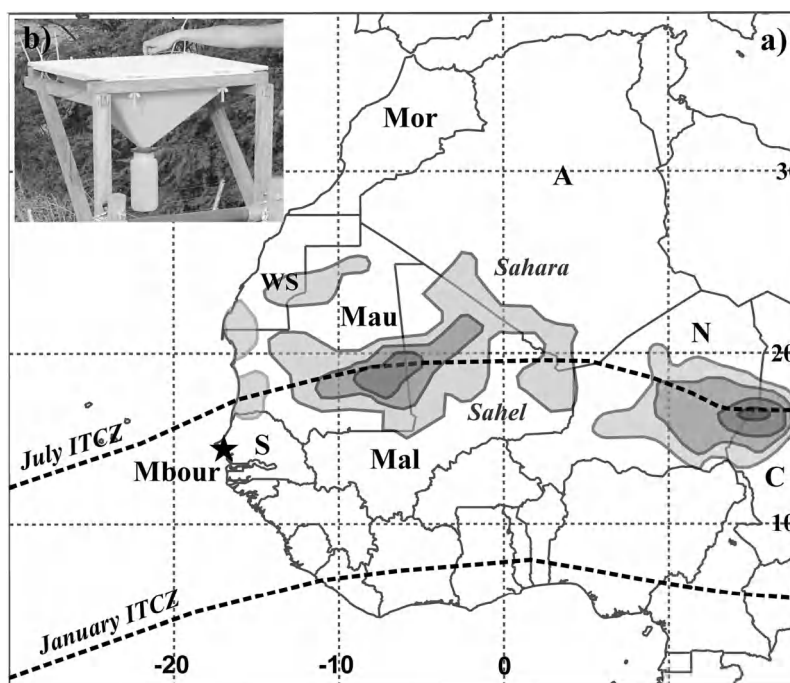


Figure 1. (a) Location of the Mbour site (star). The gray areas represented the >15 annual mean Aerosol Index derived from TOMS after Goudie and Middleton [2001]. Dotted lines represent the winter (January) and summer (July) ITCZ position. Countries are distinguished by letters: A (Algeria), C (Chad), Mal (Mali), Mau (Mauritania), Mor (Morocco), N (Niger), S (Senegal), WS (Western Sahara). (b) Picture of our Capyr-type trap.

[4] The African Multidisciplinary Monsoon Analysis (AMMA) field campaign in spring 2006 was partly devoted to improving our current knowledge of the dust particle atmospheric cycle in this region [Redelsperger *et al.*, 2006]. During March 2006, dust activity was very pronounced over a large part of West Africa, and especially throughout the March 7–13 period, when the formation of an intense dust haze was observed over the region [Slingo *et al.*, 2006; Tulet *et al.*, 2008; Thomas and Gautier, 2009]. Here we present a multiproxy characterization of the terrigenous signature of the dust that fell over Mbour, Senegal, during the spring 2006 AMMA field campaign, with focus on the March 7–13 event, which is regarded as a typical example of a winter monsoon season mineral dust event [Derimian *et al.*, 2008; Marticorena *et al.*, 2010].

2. Methods

[5] The sampling site was located at the southern end of the city of Mbour (~80 km south of Dakar; Figure 1a), on the premises of the Institut de Recherche pour le Développement (IRD) ecological center. It is a vegetated area and car traffic on the site is minimal, reducing any influence from locally produced-dust. Dust was collected using a Capyr-type reversed pyramid-shaped PVC collector [Goossens and Rajot, 2008] installed at about 8 m above ground on the rooftop terrace of the IRD research building facing the Atlantic Ocean. This collector has the advantage of being made entirely of plastic (Figure 1b), minimizing potential contamination of the dust samples with chemical elements of interest. The original Capyr-type collector,

however, has relatively lower efficiency (up to a mean factor of ~ 3) than other sampling devices due to its poorer aerodynamic shape [Goossens and Rajot, 2008]. For this reason, the 2500 cm² sampling area was covered in this study with a 1-cm thick honeycomb-like grid in order to reduce wind turbulence within the collector. This adjustment should have limited post-depositional remobilization, thus improving the collector's efficiency. Dust deposited in the device was recovered weekly from February 23 until May 29, 2006 by thorough rinsing with ultra-pure water. Temporal sampling resolution was increased around the March 7–13 outbreak (Table 1).

[6] Subsequent treatments were carried out in clean laboratory facilities. In order to remove the calcium carbonate component without altering clays, samples were leached using an acetic acid (HOAc) – sodium acetate buffer at pH ~ 4.5 [Biscaye, 1965]. Although there are studies that demonstrate that distal dust can be relatively coarse [e.g., Glaccum and Prospero, 1980], modern dust collected over the Atlantic Ocean at the latitude of our dust collector has been observed to be $<30 \mu\text{m}$ [Stuut *et al.*, 2005]. Therefore, samples were wet-sieved at $30 \mu\text{m}$ in order to prevent contamination by potential large ‘local’ mineral particles and vegetation detritus. Mass fluxes were calculated for the <30 and $<73 \mu\text{m}$ fractions, but all the following analyses were performed on the $<30 \mu\text{m}$ fraction.

[7] Grain-size measurements were carried out using a Malvern Mastersizer 2000 laser diffractometer as described by Sperazza *et al.* [2004] and the mean modal size was calculated from the volume size distribution of the dust deposits. Semiquantitative clay mineralogical composition

Table 1. Results of the Multiproxy Characterization of the Spring 2006 Dust Deposits Collected at Mbour^a

Sampling Start	Sampling End	Mass Flux 0–30 μm ($\text{mg}\cdot\text{m}^{-2}\cdot\text{d}^{-1}$)	Mass Flux 0–73 μm ($\text{mg}\cdot\text{m}^{-2}\cdot\text{d}^{-1}$)	Grain Size Mode ^b (μm)	Smectite ^c (%)	Kaolinite ^c (%)	Illite ^c (%)	Palygorskite ^c (%)	Chlorite ^c (%)	I/K	$^{87}\text{Sr}/^{86}\text{Sr}$ $\pm 2\sigma$ ($\times 10^{-6}$)	$^{143}\text{Nd}/^{144}\text{Nd}$ $\pm 2\sigma$ ($\times 10^{-6}$)	$\epsilon_{\text{Nd}}(0)$ $\pm 2\sigma$	<i>Hantschia</i> <i>Amphioxys</i> ^d ($10^6 \text{ v}\cdot\text{g}^{-1}$)	<i>Aulacoseira</i> <i>Golzeana</i> ^d ($10^6 \text{ v}\cdot\text{g}^{-1}$)
2/23/2006 10:22	3/3/2006 9:54	85	92	16.6	34	27	19	5	15	0.70	0.714741 \pm 5	0.511952 \pm 5	–13.4 \pm 0.1	0.36	0.00
3/3/2006 10:10	3/7/2006 16:45	103	116	16.8	25	27	26	5	16	0.96	0.715007 \pm 5	0.511960 \pm 5	–13.2 \pm 0.1	0.58	0.00
3/7/2006 16:55	3/9/2006 9:20	261	279	14.3	32	27	24	8	8	0.89	0.720475 \pm 6	0.511942 \pm 2	–13.6 \pm 0.1	1.96	0.39
3/9/2006 9:30	3/10/2006 10:47	209	244	n.d.	50	20	17	6	7	0.84	0.719425 \pm 6	0.511949 \pm 5	–13.4 \pm 0.1	n.d.	n.d.
3/10/2006 11:01	3/13/2006 9:45	285	298	14.3	63	22	8	3	5	0.35	0.717348 \pm 7	0.511973 \pm 4	–13.0 \pm 0.1	0.00	2.73
3/13/2006 10:00	3/20/2006 9:53	106	151	15.8	37	34	11	5	13	0.32	0.713595 \pm 9	0.511967 \pm 4	–13.1 \pm 0.1	1.20	0.72
3/20/2006 10:07	3/27/2006 10:35	123	157	18.6	48	25	9	6	12	0.36	0.715038 \pm 5	0.511900 \pm 5	–14.4 \pm 0.1	1.09	0.44
3/27/2006 10:44	4/10/2006 10:09	61	77	16.8	43	28	12	5	12	0.44	0.713867 \pm 5	n.d.	n.d.	0.37	1.30
4/10/2006 10:20	4/17/2006 11:04	137	179	16.8	30	33	17	5	14	0.51	0.712445 \pm 6	n.d.	n.d.	0.20	1.64
4/17/2006 11:14	4/24/2006 10:20	65	86	17.5	36	34	8	8	15	0.23	0.711954 \pm 6	n.d.	n.d.	0.00	1.22
4/24/2006 10:33	5/1/2006 12:21	12	47	15.7	35	29	8	10	18	0.28	n.d.	n.d.	n.d.	0.00	0.84
5/1/2006 12:05	5/8/2006 11:45	113	148	16.7	40	28	9	9	14	0.32	0.712448 \pm 6	n.d.	n.d.	0.00	0.42
5/8/2006 12:05	5/15/2006 10:38	75	132	16.0	33	32	10	9	17	0.31	0.711005 \pm 6	n.d.	n.d.	0.00	0.00
5/15/2006 10:53	5/22/2006 10:36	44	98	16.8	41	27	13	9	11	0.48	0.713233 \pm 6	n.d.	n.d.	0.00	0.00
5/22/2006 10:50	5/29/2006 11:00	76	103	12.7	29	45	9	7	11	0.20	0.714322 \pm 6	n.d.	n.d.	0.00	0.00

^aAbbreviation: n.d., no data.^bAnalytical error less than 0.4 μm .^cAnalytical error less than 5%.^dUnit is 10^6 valves per gram of sediment.

of the dust particles was carried out by X-ray diffraction (XRD). Non-oriented aggregates of the $<30\ \mu\text{m}$ carbonate-free fractions were deposited on silicon disks and measured by XRD on a Bruker D4 Endeavor set (with a Lynxeye fast detector; Cu X-ray tube; 30 kV voltage; 30 mA intensity). Two XRD runs were performed routinely for each sample: (1) air-dried sample, and (2) ethylene-glycol vapor saturation for 12 h. Each clay mineral was characterized as described by *Bout-Roumazeilles et al.* [1999] and semiquantitative estimation of clay mineral abundances (%) was performed using the Macintosh MacDiff[®] 4.2.5 software (R. Petschick, <http://www.geologie.uni-frankfurt.de/Staff/Homepages/Petschick/RainerE.html>).

[8] For strontium (Sr) and neodymium (Nd) isotope analyses, aliquots of about 20 mg were dissolved in PFA Savillex[®] beakers using a mixture of hydrofluoric acid (HF) and nitric acid (HNO₃). Sr and the rare earth elements (REE) were separated using standard cation exchange column chromatography procedures. Nd was purified from the REE fraction by cation exchange using α -hydroxyisobutyric acid (HIBA) as eluent. Sr and Nd isotopic compositions were measured by Thermal Ionization Mass Spectrometry (TIMS) on a ThermoFisher TRITON at the Max Planck Institute for Chemistry, Mainz, Germany. $^{87}\text{Sr}/^{86}\text{Sr}$ ratios were normalized to $^{86}\text{Sr}/^{88}\text{Sr} = 0.1194$ and replicate measurements of the NIST SRM 987 standard yield an average $^{87}\text{Sr}/^{86}\text{Sr} = 0.710262$ ($2\text{SD} = 8 \times 10^{-6}$, $n = 19$). $^{143}\text{Nd}/^{144}\text{Nd}$ ratios were normalized to $^{146}\text{Nd}/^{144}\text{Nd} = 0.7219$ and measurements of La Jolla Nd standard yielded $^{143}\text{Nd}/^{144}\text{Nd} = 0.511844$ ($2\text{SD} = 5 \times 10^{-6}$, $n = 8$). The $^{143}\text{Nd}/^{144}\text{Nd}$ ratios are reported as ε_{Nd} , the relative deviation from the Bulk Silicate Earth $^{143}\text{Nd}/^{144}\text{Nd}$ ratio of 0.512638 in parts per 10000. Analytical blanks were always less than 0.1% of Sr and Nd analyzed and can be considered negligible.

[9] Diatom species were identified and counted using a Olympus BX51 phase contrast microscope at a magnification of $\times 1000$ at University Bordeaux-1, France, following procedures described by *Rathburn et al.* [1997].

3. Results and Discussion

3.1. The 7–13 March 2006 Major Dust Event: The Context Over Mbour

[10] Westward Saharan dust transport across West Africa and the Northeastern Tropical Atlantic Ocean occurs nearly all year-round but is most prominent through the spring and the summer seasons [*Swap et al.*, 1996]. During the summer, when the ITCZ sits at its northernmost position and the West African monsoon provides rainfall to the Sahel region, dust transport occurs at high altitudes (5–6 km) in the so-called Saharan Air Layer [*Prospero and Carlson*, 1981], that is above the trade wind inversion along the African coast where it goes nearly unnoticed at ground level [*Bory et al.*, 2002]. Saharan outbreaks affecting the West African margin and the Northeastern Tropical Atlantic Ocean mainly occur during the winter and spring seasons (November to May), when dust is transported within the Harmattan (i.e., dry, winter monsoon continental trade winds) below 1.5–3 km altitude [*Chiapello et al.*, 1995]. The position of the low-level dust plume over the Atlantic Ocean moves northward throughout the winter monsoon season following the migration of the ITCZ [*Swap et al.*, 1996]. At the latitude of

Senegal, most dust events are recorded at the beginning of the year, often sweeping large parts of West Africa before reaching the Atlantic Ocean and generally peaking around February–March [e.g., *Caquineau et al.*, 2002; *Knippertz and Fink*, 2006].

[11] This was the case in 2006: significant dust activity began on March 5 initiated by a cold front in the lee of the Atlas Mountains, and developed further between March 7 and March 13, as the dust haze progressed across West Africa toward the Atlantic Ocean [*Slingo et al.*, 2006; *Tulet et al.*, 2008; *Thomas and Gautier*, 2009]. During this period, the aerosol optical thickness (AOT) at 440 nm measured by the Mbour Sun photometer jumped from its background level around 0.5 to values above 2.5 (Figure 2a). The AOT peaked first on March 9, and then again on March 11, implying a two-phased event in the atmospheric column. This was associated with changes in the Ångström exponent $\alpha_{440/870}$ (used in this case as an indicator of the particle-sizes), which dropped to its lowest level, indicating the presence of a “pure” mineral dust event (i.e., free of black carbon from biomass burning; see *Derimian et al.* [2008] for details).

[12] On the ground, at the Mbour station, the 7–13 March event led to the highest daily mean atmospheric particulate concentration ($<10\ \mu\text{m}$) recorded during the 2006 dry season (Figure 2b), but also the most important of the following three years of monitoring (2006–2008) by Tapered Element Oscillating Microbalance (TEOM) (see *Marticorena et al.* [2010] for details). As expected, this huge dust outbreak resulted in an important deposition event at our site. The terrigenous deposition flux recorded for the March 7–13 period averages $250\ \text{mg}\cdot\text{m}^{-2}\cdot\text{d}^{-1}$, which represents a 2- to threefold increase compared to the mean spring background level (Figure 2b and Table 1). Comparison of the $<30\ \mu\text{m}$ fraction and the total (0–73 μm) deposition fluxes confirms that most of the dust deposited in this area is $<30\ \mu\text{m}$ (Figure 2b). To our knowledge, there are no available deposition data in the literature with which we could compare our results. The deposition fluxes we obtained, however, are of the same order of magnitude as downward lithogenic fluxes in the water column estimated from sediment traps moored in the nearby Northeastern Tropical Atlantic [*Ratmeyer et al.*, 1999; *Bory and Newton*, 2000; *Bory et al.*, 2002]. Fluxes measured at Mbour, however, are higher, consistent with the fact that the mooring locations are farther afield from the dust sources compared to our site. As indicated by the AOT, two distinct phases were also recorded on the ground, as seen in both atmospheric concentration and deposition flux data.

3.2. Terrigenous Signature: March 7–13 Dust Event Versus Spring Background

[13] All dust deposits display a well-sorted unimodal grain-size distribution (Figure 3), which is a well-known characteristic of wind-blown sediments [*Krumbein and Pettijohn*, 1938]. The 2006 dry season mean modal size ranges from 12.7 to 18.6 μm (Figure 2c and Table 1). These values are comparable to those obtained for dust collected off NW Africa (9 to 18 μm) [*Stuut et al.*, 2005]. During the March 7–13 dust event, it is noteworthy that the mode decreased from $\sim 16\ \mu\text{m}$ to $\sim 14\ \mu\text{m}$, coinciding with a sharp decrease in the local wind intensity (Figure 2c). This

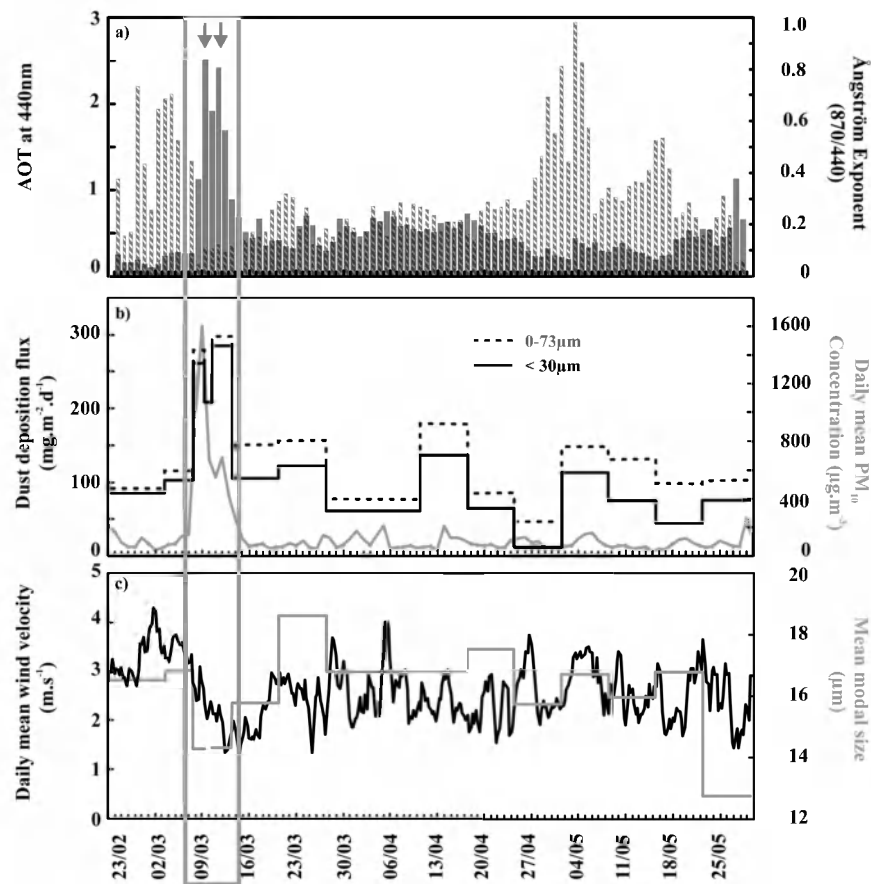


Figure 2. (a) Aerosol optical thickness at 440 nm (solid bars) and Ångström exponent ($\alpha_{440/870}$) (dash bars) from Feb. Twenty-three to May 25 2006 (AERONET database at <http://aeronet.gsfc.nasa.gov/>). (b) Dust deposition flux: $<73 \mu\text{m}$ (dash line) and $<30 \mu\text{m}$ (solid line) carbonate-free fractions and daily mean PM_{10} concentration ($\mu\text{g}\cdot\text{m}^{-3}$) [Marticorena *et al.*, 2010]. The gray frame indicates the March 7–13 dust event. (c) Daily mean surface wind velocity ($\text{m}\cdot\text{s}^{-1}$) [Marticorena *et al.*, 2010] and mean modal size of the dust deposits.

reduction in air velocity during the event excludes significant local contribution at that time, and must have instead favored deposition of fine aeolian particles transported within the large-scale plume from Saharan-Sahelian sources.

[14] Clay mineralogy and Sr and Nd isotopic compositions have been shown to vary with dust provenance, and are therefore useful tracers for source discrimination [Grousset and Biscaye, 2005; Bout-Roumzeilles *et al.*, 2007; Meyer *et al.*, 2011]. Outside the March 7–13 dust event, the clay mineralogy of the dust deposited at Mbour consisted mainly of smectite and kaolinite, with additional contribution of illite, chlorite and palygorskite (Table 1). During the March 7–13 outbreak, there was a significant increase in smectite, with proportions rising above 60% (Figure 4a). There was also an important change in the ratio between illite and kaolinite (I/K), which reached 0.9 compared to a background range of 0.2–0.5 for the rest of the sampling period (Table 1). These changes in the mineralogical composition of the dust were accompanied by a marked shift in Sr isotopic composition. The $^{87}\text{Sr}/^{86}\text{Sr}$ ratios suddenly became more radiogenic during the March 7–13 event, with measured values up to 0.720 (Figure 4b). By contrast, the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios background remained within

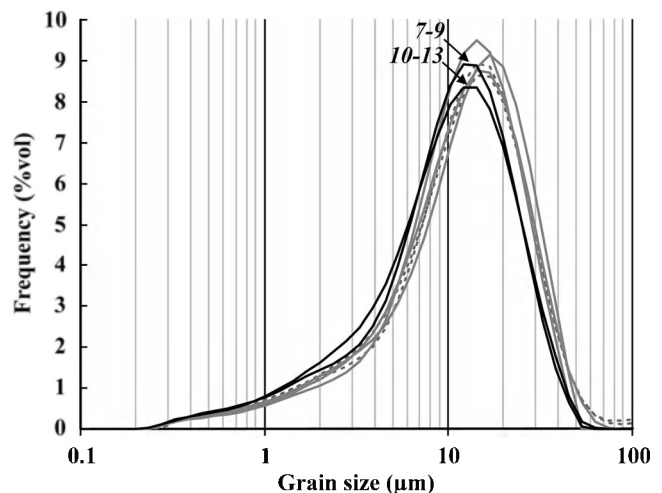


Figure 3. Comparison of grain-size distributions of mineral dust deposited in the sediment trap before the event (dashed gray lines), during the event (dark lines) and after the event (gray lines).

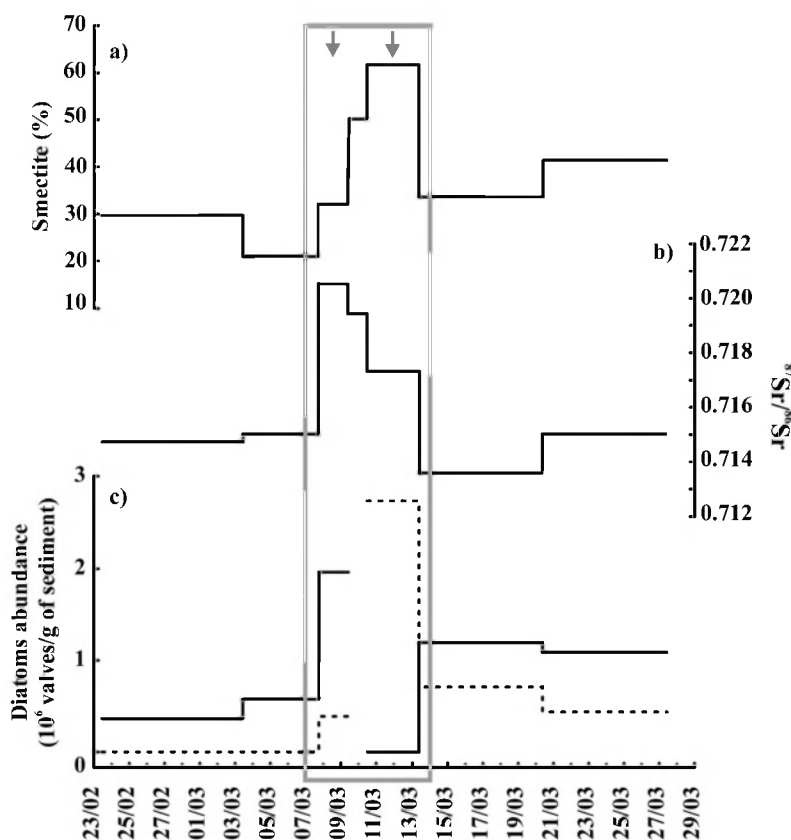


Figure 4. (a) Smectite content (%) and (b) $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of the dust deposits throughout the March 7–13 event 2006. (c) Fresh-water diatoms *Hantzshia amphioxys* (solid line) and *Aulacoseira granulata/gotzeana* (dash line) relative abundances. The gray frame highlights the March 7–13 dust event.

the 0.711–0.715 range, supposedly representing the signatures of local and/or regional contributions. It is noteworthy that the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio in terrigenous sediments is generally sensitive to grain size, with higher $^{87}\text{Sr}/^{86}\text{Sr}$ generally found in smaller size classes [Grousset *et al.*, 1998]. However, the shift in mean modal size of the dust is too subtle to account for the large difference in $^{87}\text{Sr}/^{86}\text{Sr}$ ratio we observe. Thus, the major dust outbreak of March 7–13 can be clearly fingerprinted in $^{87}\text{Sr}/^{86}\text{Sr}$ in addition to changes in clay-minerals assemblages. Taken together, these data imply a shift in dust provenance regions, along with an increased dust flux, during the March event relative to that of the spring of 2006 background period.

3.3. Change in Dust Provenance Within the Event

[15] The increased sampling resolution around the March 7–13 outbreak enabled us to document the temporal variability of the terrigenous signal throughout the event. The distinct two phases identified in the atmospheric column by optical data were captured at the ground site at Mbour as well, as readily shown by the dust deposition flux record (Figure 2b). Noticeably, the two stages were accompanied by changes in the nature/characteristics of the dust itself. For instance, the relative abundance of smectite increased from ~34 to 63% during the 7-day event whereas the I/K ratio decreased from 0.9 to 0.4 (Figure 5a and Table 1). The $^{87}\text{Sr}/^{86}\text{Sr}$ ratios exhibit a decrease from ~0.720 during the

early phase to ~0.717 toward the end, while ε_{Nd} displayed a moderate change from –13.6 to –13.0. When plotted against each others, the ε_{Nd} and the $^{87}\text{Sr}/^{86}\text{Sr}$ of the three samples collected during the event define a hyperbola (Figure 5b), suggestive of a two-component mixing.

[16] In an attempt to identify the main atmospheric transport patterns and regions of dust activity that led to the large deposition episode at Mbour, tri-dimensional back trajectories were calculated using the HYSPLIT model of NOAA. These back trajectories (Figure 6) show that air masses reaching Mbour during the March 7–13 event swept through Northern Algeria (Atlas region), where the dust storm was initiated. These air masses then progressed across southern Algeria, northern Mali and southeastern Mauritania, that is right through the second most important region of dust emission identified from the Total Ozone Mapping Spectrometer (TOMS) satellite data over the Sahara/Sahel region [Goudie and Middleton, 2001]. This is consistent with Goudie and Middleton's suggestion that the dust transported by the Harmattan from the Bodélé depression in Chad—the most important source area in the world—may not be the source of the prominent winter plume over the Northeastern tropical Atlantic. It is quite apparent, though, that during the event, air mass pathways shifted progressively to the southeast, reaching lower latitudes and longitudes further to the east; indeed, the trajectory ending on March 13 even passes over Northwest Niger (Figure 6). This

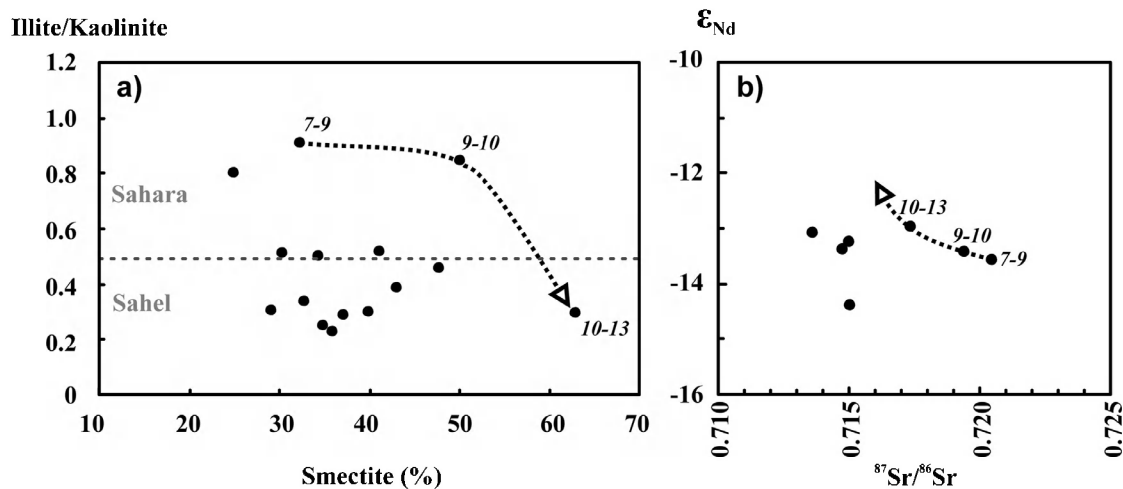


Figure 5. (a) Illite/kaolinite ratio versus smectite content (%). The dotted line indicates the approximate limit between Saharan and Sahelian I/K values as described by *Scheuven et al.* [2009]. (b) ϵ_{Nd} versus $^{87}Sr/^{86}Sr$ (error bars less than or equal to symbol size) during the study period. For Figures 5a and 5b, samples from the March 7–13 event are labeled with the corresponding sampling dates. Arrows underline the chronological order of the samples.

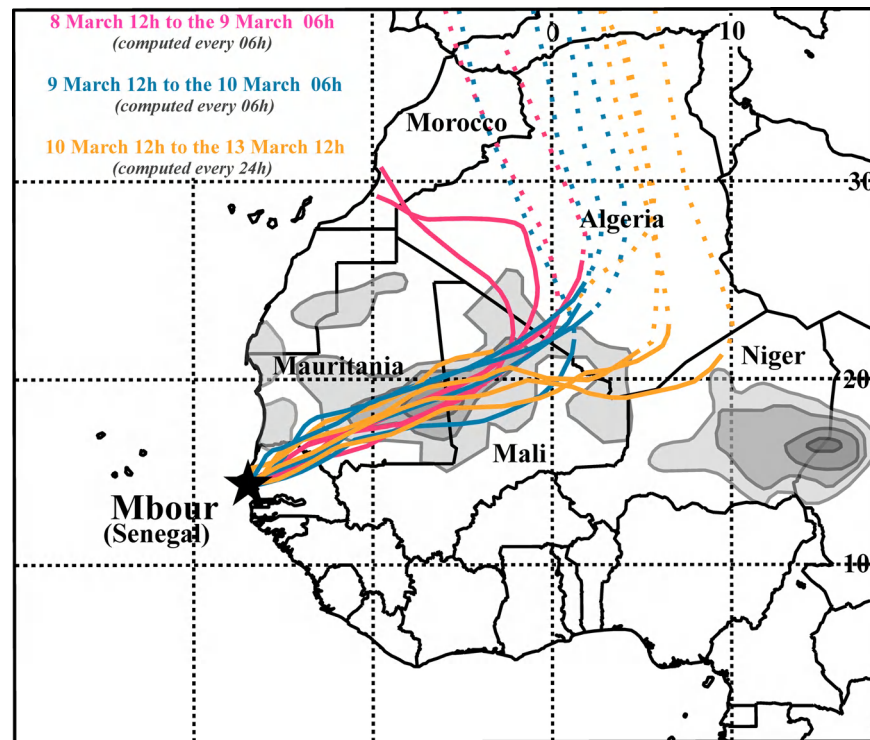


Figure 6. Tri-dimensional air masses back trajectories computed to end at approximately 400 m AGL above the Mbour station during the March 7–13 sampling period. Trajectories' duration varies from 3 days at the beginning of the time series to 6 days toward the end (durations were adapted so that trajectories reached the continent's hedges). Air mass latitude higher than 1500 m (dashed lines) are distinguished from lower transport (solid lines). Calculation was made using the HYSPLIT model (R. R. Draxler and G. D. Rolph, HYSPLIT (HYbrid Single-Particle Lagrangian Integrated Trajectory) Model, 2011, access via NOAA ARL READY Website <http://ready.arl.noaa.gov/HYSPLIT.php>; G. D. Rolph, Real-time Environmental Applications and Display system (READY) Website, <http://ready.arl.noaa.gov>). Background map shows the annual mean Aerosol Index (AI) values >15 from TOMS data (1980–93, 1997–2000) (adapted from *Goudie and Middleton* [2001]).

observation is in agreement with the change in dust transport direction already identified on MSG/SEVIRI satellite data [Tulet *et al.*, 2008; Marticorena *et al.*, 2010]. In fact, these images show that the initial dust plume, which formed in northern Algeria, first moved southwestward producing further dust emissions along its path before reaching Mbour on March 8. Then, between March 9 and 10, satellite images indicate a change in the direction of the plume, with dust being transported increasingly westward, in a narrower atmospheric corridor sweeping through the Sahelian band.

[17] Clay mineralogy signatures recorded during the March 7–13 event were compared with those of West African soils, loess and aerosols available from the literature. Although chemical and mineralogical fractionations may accompany grain-size sorting during deflation and transport processes, and may thus complicate direct comparison between aerosols and soils/sediments characteristics [Schütz and Seibert, 1987; Sabre *et al.*, 1997], such a comparison supports a southeastern shift in provenance areas supplying dust to Mbour during the event. Specifically, the Western Sahara region is depleted in smectite compared to Sahelian sources [Avila *et al.*, 1997]. Furthermore, the I/K ratio of mineral dust has been shown to decrease overall with decreasing latitude, as well as eastward across the Saharan-Sahelian stretch [Caquineau *et al.*, 2002]. This is consistent with the fact that higher chemical weathering generally results in higher abundances of kaolinite, which mainly forms through hydrolysis processes [Chamley, 1989]. Thus, more kaolinite is expected to be found in the Sahelian band [Caquineau *et al.*, 1998], which was subjected to a higher precipitation regime (summer West African monsoon) during the Quaternary. Southern Saharan and Sahelian regions are thus typified by low I/K ratios (typically <0.5) as well as by particularly low amounts of palygorskite [Scheuven *et al.*, 2009; Formenti *et al.*, 2010], which both characterize the end of the March 2006 event.

[18] Sr and Nd isotope ratios of dust deposited at Mbour during the event fall within the range of values found in the “Sub-Saharan/Sahelian” zone (Chad, Niger, central Mali and Mauritania) defined by Scheuven *et al.* [2009]. This author suggested that this area can be further subdivided into a “western” area with higher $^{87}\text{Sr}/^{86}\text{Sr}$ ratios (>0.720) and lower ε_{Nd} (around –15), and an “eastern” area with lower $^{87}\text{Sr}/^{86}\text{Sr}$ ratios (<0.720) and higher ε_{Nd} values (around –12). This geographical variation in $^{87}\text{Sr}/^{86}\text{Sr}$ and ε_{Nd} must largely reflect the fact that the “northwestern” area contains old Precambrian outcrops compared to the “southeastern” area, which is characterized by younger, Paleozoic terrains [Konate *et al.*, 2003]. According to this subdivision, the Sr and Nd isotope ratios measured in our samples (Figure 5b and Table 1) also support a west-to-east shift within the Sub-Saharan/Sahelian regions that supplied Mbour during the March 2006 event.

[19] Microscopic examination of the fresh-water diatoms in the samples indicates significant changes in the proportions of two taxa, with greater abundances of *Hantzshia amphioxys* during the early phase and greater occurrences of *Aulacoseira granulata/gotzeana* during the later phase of the dust event (Figure 4c and Table 1). These two freshwater diatom taxa are usually encountered in deflation basins, wadi valleys or hydrographic paleolakes throughout the Sahara and Sahel [Gasse *et al.*, 1989]. The very low

abundances of these diatoms in the dust samples argue for a remote origin. *Hantzshia amphioxys*, which occurs in various freshwater biotopes, is able to survive subaerial conditions and is common in Northern Mauritania [Gasse *et al.*, 1989]. The other observed taxon, *Aulacoseira granulata*, is common in lacustrine environments from East Africa [Gasse, 1986]. Changes in diatom dominance thus also corroborate a southeastern migration of the dust deflation zones throughout the dust event.

[20] The changing dust composition that we observe therefore seems to reflect differing mixtures. Dust deposited at Mbour during first phase of the event (March 8–10) likely reflects provenances from the northern and western part of this large Mauritania-Mali-Algeria area. On the other hand, the dust deposited at the end of the event (March 11–13) reflects additional inputs from more southeastern, Sahelian/sub-Saharan areas. Back trajectories presented here and earlier investigations [Slingo *et al.*, 2006; Thomas and Gautier, 2009], emphasize the fact that major Saharan dust events, such as the March 7–13 outbreak, involve a large geographical domain within which it is unrealistic to accurately pin-point the numerous contributing sources down to a local scale. Nevertheless, our data do show that it is still possible to identify different regional dust signatures associated with changing regional provenance during a major event.

3.4. Implications

[21] Major aeolian events, such as the March 7–13 one, are thought to account for most of the total annual dust depositional flux to the ocean [Guerzoni *et al.*, 1997]. For this reason, data obtained in this study provide valuable constraints for developing realistic atmospheric transport models, as well as estimates of dust deposits composition (grain size, mineralogy) and budgets (estimated deposition flux; see Methods for our Cypyr-type trap efficiency) for the Western African region. For instance, the comparison of the clay mineralogy of the dust deposited at Mbour during the event shows mismatches with mineralogy modeled from optical measurements for the same event (mainly in terms of clay minerals proportions [Thomas and Gautier, 2009]), underlying the need for field measurements in order to improve model calibrations.

[22] Owing to their significant influx to the ocean, major Saharan-Sahelian dust events have an overriding influence on the terrigenous marine sedimentation in the Northeastern Tropical Atlantic [Rea, 1994]. This conclusion is supported by the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios of the dust deposited during the 7–13 March 2006 outbreak. The corresponding ratios (0.717–0.720; Figure 5b) overlaps with the 0.718–0.722 range published for surface sediments, on the similar <30–40 μm size fraction, from the Cape Verde – Canaries Islands latitudinal band in the Northeastern Tropical Atlantic Ocean [Grousset *et al.*, 1998; Cole *et al.*, 2009]. Conversely, the $^{87}\text{Sr}/^{86}\text{Sr}$ ratios found outside the main event (~0.711–0.715), have no counterparts in surface sediments. Therefore, the multiproxy characterization of present-day dust deposited on the Atlantic margin during major events, combined with reliable estimates of dust emission and transport patterns involved, provides essential background information for the interpretation of marine sedimentary records in the Tropical Atlantic. Knowledge of particles provenance is indeed crucial information if we are to

interpret reliably the terrigenous signal in marine sediments in terms of changes in paleo-environmental conditions in sources areas and/or wind transport patterns.

4. Conclusion

[23] Dust deposited at Mbour, Senegal, was sampled during the spring 2006. The 7–13 March 2006 outbreak resulted in a large dust fall, whose mineralogical and Sr isotopic composition contrast with that of the background signal. Dust compositional changes were also observed between the beginning and the end of the outbreak, with marked differences in clay mineralogy, Sr and Nd compositions, and freshwater diatom taxa. Back trajectories and satellite images enabled us to link those distinct signatures to two different provenance regions. These regions lie within the large area identified by Goudie and Middleton [2001] as the main source of the prominent winter/spring plume over the tropical Atlantic—encompassing Mauritania, Mali and southern Algeria and some adjacent areas. Clearly, this survey will need to be repeated to capture other major events and should ideally be extended to other locations on the West African shoreline. Our mineralogical and isotopic fingerprinting of dust provenance and transport systems in West Africa, however, constitutes a first step toward a present-day “calibration” of the dust proxies measured in marine sediment records of the North-Eastern Tropical Atlantic. Since measurements of dust deposition are scarce, the flux and grain size data obtained in this study, among other parameters, provide much needed constraints for atmospheric transport models and dust deposition estimates in this area.

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