## Dear editor,

We wish to thank you, the two anonymous referees, and Dr H. Moosmüller for useful comments on the manuscript. We have carefully revised the text to improve the clarity of the reading.

In particular we have made a small change to the paper title from

"Spectral- and size-resolved mass absorption efficiency of mineral dust aerosols in the shortwave: a simulation chamber study"

to

"Spectral- and size-resolved mass absorption efficiency of mineral dust aerosols in the shortwave spectrum: a simulation chamber study"

The detailed answers to the two anonymous referees are presented

# Anonymous Referee #1

The paper presents needed results on dust optical properties (mass absorption efficiency and absorption Angstrom exponent) for D<2.5um and D<10um particles, relating them to chemical composition. The samples analyzed are from 12 locations in northern Africa (5 samples), Namibia (1 sample), northern China (1 sample), the Middle East (2 samples), North America (1 sample), South America (1 sample) and Australia (1 sample). I have no substantial issues with the analysis or the paper. I recommend publication after addressing the minor points below:

1) In the Abstract and somewhere earlier in the paper (before the Results section) the location where the samples were collected from should be presented. Some information about sample collection is also needed – i.e.: a) Was only one sample collected at each location, or were multiple samples collected then combined? b) Were the samples collected from locations know to be preferential sources for atmospheric dust or was the collection location just random? The latter point is important, since it's know that atmospheric dust comes from preferential locations.

To address this point, the sentence in the result section has been rewritten as "The selection of these soils and sediments was made out of 137 individual top-soil samples collected in major arid and semi-arid regions worldwide and representing the mineralogical diversity of the soil composition at the global scale. As discussed in Di Biagio et al. (2017), this large sample set was reduced by a set of 19 samples their availability in sufficient quantities for injection in the chamber. Because some of the experiments did not produce enough dust to perform goodquality optical measurements, , in this paper we present a set of twelve samples distributed worldwide but mostly in Northern and Western Africa (Libya, Algeria, Mali, Bodélé) and the Middle East (Saudi Arabia and Kuwait). Individual samples from the Gobi desert in Eastern Asia, the Namib Desert, the Strzelecki desert in Australia, the Patagonian deserts in South America, and the Sonoran Desert in Arizona have also been investigated."

The first sentence of the abstract has been changed as "This paper presents new laboratory measurements of the mass absorption efficiency (MAE) between 375 and 850 nm for twelve individual samples of mineral dust from different source areas worldwide and in two size classes".

2) Abstract, pg. 2, lines 33-34: "The size-independence of AAE suggests that, for a given size distribution, the possible variation of dust composition with size would not affect significantly the spectral behavior of shortwave absorption." Either that or the composition simply DIDN'T vary with size for this set of samples – so I'd reword this a bit.

This correction has been accepted

3) pg. 2, line 39: Need to spell out AAOD this first time that you use it.

## This correction has been accepted

4) pg. 2, lines 40-41: "which is relevant to the development of remote sensing of lightabsorption aerosols from space." Not only from space! This approach has also been used extensively for AERONET (surface-based remote sensing) AAOD attribution.

The reviewer is right, the sentence has been rewritten as ", which is relevant to the development of remote sensing of light-absorption aerosols from space, and their assimilation in climate models."

5) pg. 3 line 56 contains a partial sentence ("In the solar spectrum (Boucher et al., 2013)").

This was reworded as "Albeit partially compensated by the radiative effect in the thermal infrared, the global mean radiative effect of mineral dust in the shortwave is negative both at the surface and the top of the atmosphere (TOA) and local warming of the atmosphere (Boucher et al., 2013)."

6) pg. 6, lines 162-163: "The total uncertainty, including the effects of photon counting and the deposit inhomogeneity, on the absorption coefficient measurement is estimated at 8%." Is there a basis/reference for this?

The references to the papers of Petzold et al. (2004) and Massabo' et al. (2013) have been added to the text.

7) pg. 7, line 170: It would be useful to give typical total masses and/or the fractional error/uncertainty in total aerosol gravimetric mass based on this error in filter mass.

This has been added in lines 173-175.

8) pg. 8, line 223 equation: When I read this my immediate question was: "How does this compare to the measure of total gravimetric mass?". You answer this question (appropriately) in the results section but I still think it would be useful to add a note here pointing to the fact the in your results discussion you found this agreed well with the total gravimetric mass.

The following sentence has been added "As it will be explained in the result section (paragraph 3.1), the values of MCdust estimated from Equation 4 were found in excellent agreement with the measured gravimetric mass on the filters"

9) pg. 10: Do you have any estimate / sense of how well the dust suspended by this "shaking" compares to the dust lofted by winds?

Prior to any scientific analysis, we have dedicated a lot of energy to investigate the realism of our dust generation system, both in terms of the composition and the size distribution of the dust aerosols. These results are reported in two papers:

Di Biagio, C., P. Formenti, S. A. Styler, E. Pangui, and J.-F. Doussin (2014), Laboratory chamber measurements of the longwave extinction spectra and complex refractive indices of African and Asian mineral dusts, Geophys. Res. Lett., 41, doi:10.1002/2014GL060213. – Figure 2 and discussion

But in particular in

 Di Biagio, C., Formenti, P., Balkanski, Y., Caponi, L., Cazaunau, M., Pangui, E., Journet, E., Nowak, S., Caquineau, S., Andreae, M. O., Kandler, K., Saeed, T., Piketh, S., Seibert, D., Williams, E., and Doussin, J.-F.: Global scale variability of the mineral dust long-wave refractive index: a new dataset of in situ measurements for climate modeling and remote sensing, Atmos. Chem. Phys., 17, 1901-1929, doi:10.5194/acp-17-1901-2017, 2017

where we have dedicated two paragraphs (5.1. Atmospheric representativity: mineralogical composition and 5.2 Atmospheric representativity: size distribution) to show how the composition and the size distribution of the generated dust are well representative of those of real dust in the atmosphere, which makes the laboratory experiments well suited for studying the dust optical properties.

In order to stress this point further, without repeating results already presented in these two publications, we have added the following sentence in paragraph 3 "Di Biagio et al. (2014; 2017) have demonstrated the realism of the generation system concerning the composition and the size distribution of the generated dust with respect to the properties of mineral dust in the atmosphere".

10) pg. 11, line 209: Mauritania is not listed as one of the sample site locations in Table 3. ?

For Mauritania, we only have chemical composition but not optical measurement results. The reference to this sample has been taken out of the paper.

11) pg. 11, lines 309-310 (an onward): The results here are said to agree well with that found for atmospheric aerosols in other studies, but the values in these studies is not given so this feels very hand-waving and unconvincing. Are you referring to the values given in Table 5? If so, please refer directly to them. If not, the comparison here needs to be more quantitative (discuss numbers from the literature vs. what is found here).

Additional text and values have been added in lines 321-392 to address this point

12) pg. 13, lines 353-354: If you have results for Niger why not show them?

We do not have optical results for Niger

13) pg. 14, lines 387-388: MAE doesn't vary linearly inversely with wavelength, it varies linearly inversely with the log of the wavelength (hence our ability to use the AAE relationship).

The reviewer is correct, this has been corrected as "and decrease in a linear way with the logarithm of the wavelength"

14) pg. 15, lines 429-434: A few things: a) "satisfactory" and "loose" are not quantitative terms, nor are they really appropriate for a scientific paper. What constitutes "satisfactory"? Best is to just give the correlations. b) The high correlation coefficients for PM2.5 are really driven by one high data point and so are probably not very robust.

These sentences have been reworded as "Regardless of the size fraction, the correlation between the MAE values and the percent mass of total elemental iron are higher at 375, 407 and 532 nm" and "At these wavelengths, linear correlations with the mass fraction of iron oxides are low in the  $PM_{10.6}$  mass fraction (R<sup>2</sup> up to 0.38-0.62), but higher in the  $PM_{2.5}$  fraction (R<sup>2</sup> up to 0.83-0.99)"

15) pg. 16, lines 455-456: How can fine-mode-only AAOD be GREATER than total aerosol AAOD?

Total and fine were inverted by mistake, this is now corrected

16) pg. 17, lines 487-488: As written this implies Solomon et al. varied SSA by 5%.

For, say, SSA of 0.9, that they varied SSA by 0.045. That is, the co-albedo (or absorption) was varied by 45% (0.1+/-0.045). This is the proper comparison to make to the variation in MAE that you calculate

The reviewer is right. The sentence has been corrected as "As a comparison, Solmon et al. (2008) showed that varying the single scattering albedo of mineral dust over western Africa by  $\pm$  5%, that is, varying the co-albedo (or absorption) by 45% (0.1 $\pm$  0.045) could drastically change the climate response in the region."

7) Overall: Some editing is needed for language throughout. Here I list some that stood out to me – all small stuff but editing would help readability:

pg. 3, line 67-68: "in the last ten years or so" (too casual for scientific writing)

removed

pg. 3, line 74: "A significant body of observations have been performed."

Replaced by "A significant number of observations have quantified"

pg. 8, lines 214-215: "The linear deconvolution, performed the Athena IFEFFIT free-ware analysis program (Ravel and Newville, 2005), provided with the proportionality factors alpha\_i representing the mass fraction of elemental iron to be assigned to the i-th standard mineral." (I found this sentence nearly impossible to follow)

Replaced by "The linear deconvolution has been performed with the Athena IFEFFIT freeware analysis program (Ravel and Newville, 2005). This provided with the proportionality factors i representing the mass fraction of elemental iron to be assigned to the i-th standard mineral."

pg. 10, line 253: "the chamber was evacuated by to" (delete "by")

Corrected

pg. 10, line 259: "dust particles produced was" -> "dust particles produced were"

The sentence was corrected as "The dust particles produced by the mechanical shaking, mimicking the saltation processing that soils experience when eroded by strong winds, were injected in the chamber by flushing the flask with N2 at 10 L min-1 for about 10-15 min, whilst continuing shaking the soil."

pg. 10, lines 279 "dust on filter for" -> "dust on the filter for"

Corrected as "dust on the filter membranes for subsequent chemical analysis"

pg. 10, line 280: "by placing the loaded filter holders": This reads as if you are placing LOADED FILTERS (vs holders with blank filters in them, which is what I assume you mean). Reword.

Corrected as "by placing the filter holders loaded with filter membranes"

pg. 11, line 206: "the origin of used dust samples". I think this should be "the origin of our dust samples", yes?

Corrected as "the origin of dust samples"

pg. 12, line 315: "Henceforth, and contrary to the soil samples"

pg. 12, lines 317-318: "could reflect more that of the parent sedimentary soil than not the other samples."

These two sentences were rewritten as "This powder is likely to be injected in the chamber with little or no size fractionation. Henceforth, the aerosol generated from it should have a closer composition to the original powder than the other samples.

pg. 12, line 335: "An the exception"

Corrected as "Exceptions are"

pg. 13, lines 353-354: "As a matter of fact, the number fraction of particles in the size classes above 0.5  $\mu$ m in diameter are different in the dust aerosol generated in the Alfaro et al. (2004) study with respect to ours."

Corrected as "As a matter of fact, the number fraction of particles in the size classes above 0.5  $\mu$ m in diameter is different in the dust aerosol generated in the Alfaro et al. (2004) study with respect to ours."

pg. 13, line 364: "On the contrary" -> "In contrast

Corrected

pg. 13, line 365: "These differences could yield either to difference in the"

Corrected as "These differences could either be due to difference in the chemical composition and/or in the total mass in the denominator of Equation 8."

pg. 15, lines 415-417: "using a power-law function fit as from Equation 2, provides with the values of"

Corrected as "using the power-law function fit (Equation 2)"

pg. 16, line 448: "The size-dependence, yielding significantly higher MAE values"

Corrected as "The size-dependence, yielding significantly higher MAE values in the fine fraction (PM2.5) than in the bulk (PM10.6) aerosol,"

pg. 17, line 470: "A closer look to observations" -> "A closer look at observations"

Corrected as "These differences could either be due to difference in the chemical composition and/or in the total mass in the denominator of Equation 8."

pg. 17, line 478: "our estimated MAE average at" -> "our average MAE values are" Corrected

pg. 17, line 490: "As Moosmuller et al." -> "As in Moosmuller et al."

# Corrected

pg. 18, line 503-504: "pointing out to the" -> "pointing out the

# Corrected

### Anonymous Referee #2

### GENERAL COMMENT

The manuscript presents important results from a carefully designed and conducted study on the light-absorbing properties of mineral dust from various origins. Dust samples collected at the different source regions have been re-suspended in an aerosol chamber and characterized with respect to microphysical, optical and chemical properties by state of the art methods. The study provides urgently needed knowledge on the multi-spectral light absorbing properties of mineral dust and for sure deserves publication in ACP. The manuscript is well structured, the methods are described in necessary detail and the referenced literature reflects the current state of knowledge. I recommend publication after the following minor revisions have been considered.

## SPECIFIC REMARKS

1. In the experimental protocol section, the potential impact of gravitational settling on the resuspended fraction of the dust samples is mentioned.

Given the instrumentation list, the size distributions of the airborne dust samples were monitored during the runs of the experiments. It appears obvious to control the change of the size distribution during the experiment time in the chamber. Since the mass concentration in the chamber decreased very rapidly after injection, whereas the chemical composition of the dust samples was determined from bulk samples, it would be important to know if the airborne fraction sampled for the determination of optical properties features the same chemical properties as the bulk samples. At least a discussion of this potential source of uncertainties should be presented, along with a plot showing the change of the size distributions during the experiment time. The current analysis starts from the assumption that the dust bulk properties represent also the properties of the sampled airborne fractions. However, is this really justified?

The reviewer is right when saying that the samples collected for the investigation of the chemical composition are time-integrated and henceforth might reproduce dust with varying size distributions. Examples of the time variability of the size distributions are provided by Di Biagio et al. (2017) - Figures 7 and 5S in the supplementary material. These figures show that, after the very strong initial depletion of particles larger than 10  $\mu$ m in diameter (when no sampling on filters was performed), the number concentration decreases at a rate, which is almost independent of size, suggesting that no significant distortion of the particle size distribution occurs after the most significant removal at the beginning of the experiment.

We also would like to stress that our generation system allows to generate a dust aerosol from a soil, and that this dust aerosol is injected in the chamber, not the soil. Henceforth, when talking about "bulk" we refer to the total aerosol fraction, sampled from the chamber without any size segregation other than that imposed by the cutoff the sampling lines. The fine fraction corresponds to the same dust aerosols, but sampled by an impactor with a 2.5 diameter cutoff.

2. In section 3.2, the variability of dust optical properties with particle size is discussed. The authors found no statistically significant size-dependence of the absorption Ångström exponent (AAE), whereas the absolute values of the mass absorption efficiencies (MAE) show large differeces between the PM2.5 and PM10 fractions with larger values for the fine mode fraction.

These findings imply that the relative chemical composition with respect to light-absorbing compounds does not change between the size fractions (similar AAE values), whereas the differences between the MAC values indicate that coarse mode particles contain more non-absorbing matter than fine mode particles (higher MAE values for smaller particles). This however, this is in contrast to the assumption that the chemical composition is uniformly distributed across the particle size distribution. Here, a detailed discussion is requested.

The reviewer's statement is in error as for a given mineral composition (given effective complex refractive index) the MAE depends strongly on size, decreasing with size at larger sizes. So having a smaller MAE for PM10 than for PM2.5, does not necessarily imply that PM10 contains less absorbing matter than PM2.5, but may just be due to the change in size distribution. This is shown by our calculations. An independent example of the size dependence is also given by Fig. 1 of Moosmuller et al. (2009).

Moosmüller, H., R. K. Chakrabarty, and W. P. Arnott (2009), Aerosol light absorption and its measurement: A review, Journal of Quantitative Spectroscopy and Radiative Transfer, 110(11), 844-878.

3. In section 4, it is discussed that the potential impact of light absorption by mineral dust may play an important role even after long range transport. Cited studies all refer to observations in China.

However, there is another detailed study on this effect available for the pollution plume of Dakar mixing with mineral dust which also includes the variation of the AAE during mixing

(Petzold et al., 2011). The authors may consider including this study.

This reference has been added to the manuscript.

MINOR COMMENTS

1. The list of references contains various references which are not cited in the manuscript. This should be checked, I found the following but there may be more: Anderson et al., 1998, Andrews et al., 2006, Arnott et al., 2005, Collaud Coen et al., 2010, Petzold et al., 2013.

# Corrected

2. Line 56: The sentence seems to be incomplete.

The sentence was corrected as "Albeit partially compensated by the radiative effect in the thermal infrared, the global mean radiative effect of mineral dust in the shortwave is negative both at the surface and the top of the atmosphere (TOA) and local warming of the atmosphere (Boucher et al., 2013)."

3. Line 97 - 98: The basic unit of mass concentrations is gm-3. Using this unit, then the unit of the combined property MAE is m2 g-1 as stated. In its current version this link is not clearly visible.

4. Line 102: The sentence seems to be incomplete.

The sentence was removed

5. Line 163: A reference for the uncertainty of the MWAA is required.

This has been added

6. Line 912: Please check for correct reference, there is no reference Petzold et al. (2008) in the list of references.

## Corrected

7. In Figure 4, regression lines may be shown as full line to improve their visibility.

Done

TYPOS

1. Line 108: It should read: "absorption Ångström exponent".

# Corrected

2. Line 138: Skip "with".

# Corrected

3. Line 165: It should read "deposited on a filter ...".

# Corrected

4. Line 245: It should read: "the uncertainty of values ...".

# Corrected

5. Line 253: Skip "by".

# Corrected

6. Line 338: It should read: "PM2.5 fraction".

# Corrected

7. Line 426 –427: I assume the Figures 4 are referenced here.

## Corrected

# REFERENCES

Petzold, A., Veira, A., Mund, S., Esselborn, M., Kiemle, C., Weinzierl, B., Hamburger, T., Ehret, G., Lieke, K., and Kandler, K.: Mixing of mineral dust with urban pollution aerosol over Dakar (Senegal): impact on dust physico-chemical and radiative properties, Tellus, 63B, 619-634, doi: 10.1111/j.1600-0889.2011.00547.x, 2011.

1	Spectral- and size-resolved mass absorption efficiency of mineral dust aerosols in
2	the shortwave <u>spectrum</u> : a simulation chamber study
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4	ard Pangui <sup>1</sup> , Servanne Chevaillier <sup>1</sup> , Gautier Landrot <sup>3</sup> , Meinrat O. Andreae <sup>4,11</sup> , Konrad Kandler <sup>5</sup> , Stuart
5	Piketh <sup>6</sup> , Thuraya Saeed <sup>7</sup> , Dave Seibert <sup>8</sup> , Earl <u>e</u> Williams <sup>9</sup> , Yves Balkanski <sup>10</sup> , Paolo Prati <sup>2</sup> , and Jean-
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### 24 Abstract

This paper presents new laboratory measurements of the mass absorption efficiency (MAE) between 25 375 and 850 nm for twelve individual samples of mineral dust from different source areas worldwide 26 and in of different origin in two size classes: PM<sub>10.6</sub> (mass fraction of particles of aerodynamic diameter 27 28 lower than 10.6  $\mu$ m) and PM<sub>2.5</sub> (mass fraction of particles of aerodynamic diameter lower than 2.5  $\mu$ m). EThe experiments have been were performed in the CESAM simulation chamber using generated min-29 30 eral dust generated from natural parent soils, and included optical and gravimetric analyses. The Rresults show that the MAE values are lower for the PM<sub>10.6</sub> mass fraction (range 37-135 10<sup>-3</sup> m<sup>2</sup> g<sup>-1</sup> 31 at 375 nm) than for the PM<sub>2.5</sub> (range 95-711 10<sup>-3</sup> m<sup>2</sup> g<sup>-1</sup> at 375 nm), and decrease with increasing wave-32 length as  $\lambda^{-AAE}$ , where the Angstrom Absorption Exponent (AAE) averages between 3.3-3.5, regardless 33 34 of size. The size\_independence of AAE suggests that, for a given size distribution, the possible variation of dust composition did not vary with size for this set of samples with size would not affect significantly 35 the spectral behavior of shortwave absorption. Because of its high atmospheric concentration, light -36 37 absorption by mineral dust can be competitive to with black and brown carbon even during atmospheric transport over heavy polluted regions, when dust concentrations are significantly lower than at emission. 38 39 The AAE values of mineral dust are higher than for black carbon (~1), but in the same range as light-40 absorbing organic (brown) carbon. As a result, depending on the environment, there can be some ambi-41 guity in apportioning the aerosol absorption optical depth (AAOD) based on spectral dependence, which 42 is relevant to the development of remote sensing of light-absorbing<del>ption</del> aerosols from space, and their assimilation in climate models. We suggest that the sample-to-sample variability in our dataset of MAE 43 44 values is related to regional differences of in the mineralogical composition of the parent soils. Particularly in the PM<sub>2.5</sub> fraction, we found a strong linear correlation between the dust light-absorption prop-45 erties and elemental iron rather than the iron oxide fraction, which could ease the application and the 46 47 validation of climate models that now start to include the representation of the dust composition, as well 48 as for remote sensing of dust absorption in the UV-VIS spectral region.

## 49 **1. Introduction**

50 Mineral dust aerosols emitted by wind erosion of arid and semi-arid soils account for about 40% of the 51 total emitted aerosol mass per year at the global scale (Knippertz and Stuut, 2014). The episodic but 52 frequent transport of intense mineral dust plumes is visible from spaceborne sensors<sub>a</sub> as their high con-53 centrations, combined with to their ability of to scattering and absorbing solar and thermal radiation, 54 give raise to the highest registered values of aerosol optical depth (AOD) on Earth (Chiapello, 2014). 55 The instantaneous radiative efficiency of dust particles, that is, their radiative effect per unit AOD, is of the order of tenths to hundreds of W  $m^{-2}$  AOD<sup>-1</sup> in the solar spectrum, and of the order of tenths 56 of W m<sup>-2</sup> AOD<sup>-1</sup> in the thermal infrared (e.g., Haywood et al., 2003; di Sarra et al., 2011; Slingo et al., 57 2006 and the compilation of Highwood and Ryder, 2014). In the solar spectrum, (Boucher et al., 2013). 58 59 Albeit partially compensated by the radiative effect in the thermal infrared, the global mean radiative 60 effect of mineral dust in the shortwave is negative both at the surface and the top of the atmosphere (TOA) and produces a local warming of the atmosphere (Boucher et al., 2013). Many-There are the 61 62 consequences numerous impacts of dust on the global and regional climate, that which ultimately feed 63 back on wind speed and vegetation and therefore on dust emission (Tegen and Lacis, 1996; Solmon et al., 2008; Pérez et al., 2006; Miller et al., 2014). Dust particles perturb the surface air temperature 64 65 through their radiative effect at TOA, can increase the atmospheric stability (e.g., Zhao et al. 2011) and 66 might affect precipitation at the global and regional scale (Solmon et al., 2008; Xian, 2008; Vinoj et al., 67 2014; Miller et al., 2014 and references therein).

All models show-indicate that the effect of mineral dust on climate has a great sensitivity to their 68 69 shortwave absorption properties of mineral dust (Miller et al., 2004; Lau et al., 2009; Loeb and Su, 2010; 70 Ming et al., 2010; Perlwitz and Miller, 2010). Absorption by mineral dust started receiving a great deal of interest-in the last ten years or so, when spaceborne and ground-based remote sensing studies (Dubo-71 vik et al., 2002; Colarco et al., 2002; Sinyuk et al., 2003) suggested that mineral dust was less absorbing 72 that than it had been suggested indicated by in situ observations (e.g., Patterson et al., 1977; Haywood et 73 al., 2001), particularly at wavelengths below 600 nm. Balkanski et al. (2007) showed that lowering the 74 75 dust absorption properties to an extent that reconciles them both with the remote-sensing observations and the state-of-knowledge of the mineralogical composition, allowed calculating the clear-sky dust 76 77 shortwave radiative effect of dust in agreement with satellite-based observations. A significant body number of observations has ve been performed in quantifyquantifieding the shortwave light-absorbing 78 79 properties of mineral dust, by direct measurements (Alfaro et al., 2004; Linke et al., 2006; Osborne et al., 2008; McConnell et al., 2008; Derimian et al., 2008; Yang et al., 2009; Müller et al., 2009; Petzold 80 et al., 2009; Formenti et al., 2011; Moosmüller et al., 2012; Wagner et al., 2012; Ryder al., 2013a; Utry 81 82 et al., 2015; Denjean et al., 2015c; 2016), and indirectly, by quantifying the amount and the speciation 83 of the light-absorbing compounds in mineral dust, principally iron oxides (Lafon et al., 2004; 2006;

84 Lazaro et al., 2008; Derimian et al., 2008; Zhang et al., 2008; Kandler et al., 2007; 2009; 2011; Formenti et al., 2014a; 2014b). 85

However, existing data are often limited to a single wavelength, which moreover are is not the same 86 87 identical for all experiments. Also, frequently they do not represent the possible regional variability of 88 the dust absorption, either because they are obtained from field measurements integrating the contribu-89 tions of different source regions, or conversely, by laboratory investigations targeting samples from a 90 limited number of locations. This might lead to biases in the data. Indeed, iron oxides in mineral dust, 91 mostly in the form of hematite (Fe<sub>2</sub>O<sub>3</sub>) and goethite (Fe<sub>(</sub>O)OH), have specific absorption bands in the UV-VIS spectrum (Bédidi and Cervelle, 1993), and have a variable content depending on the soil min-92

93 eralogy of the source regions (Journet et al., 2014).

94 Henceforth, iIn this study, eExperiments on twelve aerosol samples generated from natural parent top 95 soils from various source regions worldwide have beenwere conducted with a large atmospheric simu-96 lation chamber. we We present a new evaluation of the ultraviolet to near-infrared (375-850 nm) lightabsorbing properties of mineral dust by studying investigating the size-segregated mass absorption effi-97 ciency (MAE, units of m<sup>2</sup> g<sup>-1</sup>) and its spectral dependence, largely widely used in climate models to 98 calculate the direct radiative effect of aerosols. Experiments on twelve aerosol samples generated from 99 100 natural parent top soils from various source regions worldwide have been conducted with a large atmos-101 pheric simulation chamber.

#### 102 2. Instruments and methods

- 103 At a given wavelength,  $\lambda$ , the mass absorption efficiency (MAE, units of m<sup>2</sup> g<sup>-1</sup>) is defined as the ratio of the aerosol light-absorption coefficient  $b_{abs}(\lambda)$  (units of m<sup>-1</sup>), and its mass concentration (in  $\mu g m^{-3}$ )
- 105

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104

$$MAE(\lambda) = \frac{b_{abs}(\lambda)}{Mass \ Conc} \tag{1}$$

107

108 MAE values for mineral dust aerosol are expressed in. MAE values for mineral dust aerosol are expressed in  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup>. 109

The spectral dependence of the aerosol absorption coefficient  $b_{abs}(\lambda)$  is described by the power-law 110 111 relationship

113

$$b_{abs}(\lambda) \sim \lambda^{-AAE}$$
 (2)

114

where the AAE is the <u>Absorption</u> Ångström <u>Absorption</u> Exponent, representing the negative slope of  $b_{abs}(\lambda)$  in a log-log plot (Moosmüller et al., 2009)

117

$$AAE = -\frac{dln(b_{abs}(\lambda))}{dln(\lambda)}$$
(3)

119

118

### 120 **2.1. The CESAM simulation chamber**

The eExperiments in this work have been performed in the 4.2 m<sup>3</sup> stainless-steel CESAM (French acronym for Experimental Multiphasic Atmospheric Simulation Chamber) simulation chamber (Wang et al., 2011). The CESAM chamber has been extensively used in recent years to simulate, at sub and supersaturated conditions, the formation and properties of aerosols at concentration levels comparable to those encountered in the atmosphere (Denjean et al., 2015a; 2015b; Bréegonzio-Rozier et al., 2015; 2016; Di Biagio et al., 2014; 20176).

127 CESAM is a multi-instrumented platform, equipped with twelve circular flanges to support its analytical environment. Basic instrumentation comprises sensors to measure the temperature, pressure and relative 128 humidity within the chamber (two manometers MKS Baratrons (MKS, 622A and MKS, 626A) and a 129 130 HMP234 Vaisala® humidity and temperature sensor). The particle size distribution is routinely measured by a combination of (i) a scanning mobility particle sizer (SMPS, mobility diameter range 0.02– 131 0.88 µm), composed of a Differential Mobility Analyzer (DMA, TSI Inc. Model 3080) and a Conden-132 sation Particle Counter (CPC, TSI Inc. Model 3772); (ii) a SkyGrimm optical particle counter (Grimm 133 134 Inc., model 1.129, optical equivalent diameter range 0.25–32 µm); and (iii) a WELAS optical particle counter (PALAS, model 2000, optical equivalent diameter range 0.5–47 µm). Full details of operations 135 and data treatment of the particle counters are provided in Di Biagio et al. (20162017). 136

### 137 **2.2. Filter sampling**

Three filter samples per top soil sample were collected on different types of substrate based on the analysis to <u>be performed</u>. Sampling dedicated to the determination of the aerosol mass concentration by

gravimetric analysis and the measurement of the absorption coefficients by optical analysis was per-140 formed on 47-mm quartz membranes (Pall Tissuquartz<sup>™</sup>, 2500 QAT-UP). Two samples were collected 141 in parallel. The first quartz membrane sample ("total") was collected without a dedicated size cut-off 142 using an in-house built stainless steel sampler operated at 5 L min<sup>-1</sup>. However, as detailed in Di Biagio 143 et al. (20162017), the length of the sampling line from the intake point in the chamber to the filter en-144 145 trance was 50 cm, vielding resulting in with a 50% cut-off of the transmission efficiency at 10.6 µm in 146 particle aerodynamic diameter. This fraction is therefore indicated as PM<sub>10.6</sub> in the forthcoming follow-147 ing discussion. The second quartz membrane sample was collected using a 4-stage DEKATI impactor operated at the a flow rate of 10 L min<sup>-1</sup> to select the aerosol fraction of particles with aerodynamic 148 149 diameter smaller than 2.5 µm, indicated as PM<sub>2.5</sub>-here forth. Sampling for the analysis of the iron oxide content was performed on polycarbonate filters (47-mm Nuclepore, Whatman; pore size of 0.4 µm) 150 151 using the same sample holder than as used for the total quartz filters, and therefore referring corresponding to the PM<sub>10.6</sub> mass fraction. Samples were collected at a flow rate of 6 L min<sup>-1</sup>. All flow rates were 152 monitored by a thermal mass flow meter (TSI Inc., model 4140). These samples were also used to de-153 154 termine the elemental composition (including Fe) and the fraction of iron oxides in the total mass.

### 155 **2.3. The Multi-Wavelength Absorbance Analyzer (MWAA)**

The aerosol absorption coefficient,  $b_{abs}(\lambda)_a$  at 5 wavelengths ( $\lambda = 375, 407, 532, 635$ , and 850 nm) was measured by *in situ* analysis of the quartz filter samples using the Multi-Wavelength Absorbance Analyzer (MWAA), described in detail in Massabò et al. (2013; 2015).

159 The MWAA performs a non-destructive scan of the quartz filters on-at 64 different points, each  $\sim 1 \text{ mm}^2$ wide. It measures the light transmission through the filter as well as backscattering at two different angles 160 (125° and 165°). This is necessary to constrain the multiple scattering effects occurring within the par-161 162 ticle-filter system. The mMeasurements are used as input of to a radiative transfer model (Hänel, 1987; 1994) as implemented by Petzold and Schönlinner (2004) for the Multi-Angle Absorption Photometry 163 164 (MAAP) measurements. In this model, a two stream approximation is applied (Coakley and Chylek, 165 1975), in which the fractions of hemispherical backscattered radiation with respect to the total scattering for collimated and diffuse incident radiation are approximated on the basis of the Henyey-Greenstein 166 167 scattering phase function (Hänel, 1987). This approximation assumes a wavelength-independent asymmetry parameter (g) set to 0.75, appropriate for mineral dust (Formenti et al., 2011; Ryder et al., 2013b). 168

- 169 The total uncertainty, including the effects of photon counting and the deposit inhomogeneity, on the
- absorption coefficient measurement is estimated at 8% (Massabò et al., (2013)-

## 171 **2.4. Gravimetric analysis**

The aerosol mass deposited on the filters  $(\mu g)$  was obtained by weighing the quartz filter before and after 172 sampling, after a period of 48 hours of conditioning in a room with controlled atmospheric conditions 173 (temperature,  $T \sim 20 \pm 1$  °C; relative humidity, RH ~ 50 ± 5%). Weighting is performed with an analyt-174 175 ical balance (Sartorius model MC5, precision of 1 µg), and repeated three times to control the statistical variability of the measurement. Electrostatic effects are removed by exposing the filters, prior weighing, 176 177 to a de-ionizer. The error on-in the measured mass is estimated at 10  $\mu$ g, including the repetition variability. The aerosol mass concentration ( $\mu g m^{-3}$ ) is obtained by dividing the mass deposited on the filter 178 to the total volume of sampled air  $(m^3)$  obtained from the mass flowmeter measurements (+-5%). The 179

180 percent error on mass concentrations is estimated to 5%.

# 181 **2.5. Dust composition measurements**

## 182 **2.5.1. Elemental composition**

Elemental concentrations for the major constituents of mineral dust (Na, Mg, Al, Si, P, S, Cl, K, Ca, Fe, 183 Ti, Mn) were obtained by a Wavelength Dispersive X-ray fluorescence Fluorescence (WD-XRF) of the 184 185 Nuclepore filters using a PW-2404 spectrometer by Panalytical. Excitation X-rays are produced by a 186 Coolidge tube ( $I_{max} = 125 \text{ mA}$ ,  $V_{max} = 60 \text{ kV}$ ) with a Rh anode; the primary X-ray spectrum can be controlled by inserting filters (Al, at different thickness) between the anode and the sample. Each ele-187 ment was analyzed three times, with specific conditions (voltage, tube filter, collimator, analyzing crys-188 189 tal, and detector). Data collection was controlled by the SuperQ software provided with the instrument. The elemental mass thickness ( $\mu g \text{ cm}^{-2}$ ), that is, the analyzed elemental mass per unit surface, was ob-190 191 tained by comparing the elemental yields with a sensitivity curve measured in the same geometry on a 192 set of certified mono- or bi-elemental thin layer standards by Micromatter Inc. The certified uncertainty of the standard deposit ( $\pm$  5%) determines the lower limit of the uncertainty of the measured ele-193 194 mental concentrations, which ranges between 8% and 10% depending on the element considered-element. Thanks to the uniformity of the aerosol deposit on the filters, the atmospheric elemental concen-195 trations (µg m<sup>-3</sup>) were calculated by multiplying the analyzed elemental mass thickness by the ratio 196 197 between the collection and analyzed surfaces of each sample (41 and 22 mm, respectively), then di-198 vidinged by the total sampled volume (m<sup>3</sup>). Finally, concentrations of light-weight elements (atomic 199 number Z < 19) were corrected for the underestimation induced by the self-absorption of the emitted 200 soft X-rays inside aerosol particles according to Formenti et al. (20112010).

Additional XRF analysis of the quartz filters <u>was has been</u> performed both in the  $PM_{10.6}$  and the  $PM_{2.5}$ fractions, <del>so to verify the absence of biases between the experiments dedicated to the determination of</del> particle composition to <u>and</u> those where the optical properties where measured.

### 204 **2.6.2. Iron oxide content**

The content and the mineralogical speciation of <u>the</u> iron oxides, also defined as free-iron, <u>that isi.e.</u>, the fraction of iron <u>which-that</u> is not in the crystal lattice of silicates (Karickhoff and Bailey, 1973), was determined by XANES (X-ray absorption near-edge structure) in the Fe K-range ( $K_{\alpha}$ , 7112 eV) at the SAMBA (Spectroscopies Applied to Materials based on Absorption) beamline at the SOLEIL synchrotron facility in Saclay, France (Briois et al., 2011). The position and shape of the K pre-edge and edge peaks were analyzed as they depend on the oxidation state of iron and the atomic positions of the neighboring ions, mostly O<sup>+</sup> and OH<sup>-</sup>.

As in Formenti et al. (2014b), samples were mounted in an external setup mode. A Si(220) doublecrystal monochromator was used to produce a monochromatic X-ray beam, which was  $3000 \times 250 \,\mu\text{m}^2$ in size at <u>the</u> focal point. The energy range was scanned from 6850 eV to 7800 eV at a step resolution varying between 0.2 eV in proximity to the Fe-K absorption edge (at 7112 eV) to 2 eV in the extended range. Samples were analyzed in fluorescence mode without prior preparation. One scan acquisition lasted approximately 30 minutes, and was repeated three times to improve the signal-to-noise ratio.

218 The same analytical protocol was applied to five standards of Fe(III)-bearing minerals (Table 1), including iron oxides (hematite, goethite) and silicates (illite, montmorillonite, nontronite). The standard spec-219 tra were used to deconvolute the dust sample spectra to quantify the mineralogical status of iron. The 220 221 linear deconvolution was, performed with the Athena IFEFFIT freeware analysis program (Ravel and 222 Newville, 2005). This, provided with the proportionality factors,  $\alpha_{i-}$ , representing the mass fraction of 223 elemental iron to be assigned to the *i*-th standard mineral. In particular, the values of  $\alpha_{hem}$  and  $\alpha_{goe}$ 224 represent the mass fractions of elemental iron that can be attributed to hematite and goethite, and  $\alpha_{Fe ox}$ 225  $(\alpha_{hem} + \alpha_{goe})$ , the mass fraction of elemental iron that can be attributed to iron oxides.

226 **2.6.3. Calculation of the iron oxide content** 

The measured elemental concentrations obtained by X-ray Fluorescence (XRF) are expressed in the form of elemental oxides and summed to estimate the total mineral dust mass concentration  $MC_{dust}$  according to the equation from Lide (1992)

231 
$$[MC_{dust}] = 1.12 \times \begin{cases} 1.658[Mg] + 1.889[AI] + 2.139[Si] + 1.399[Ca] + 1.668[Ti] + 1.582[Mn] \\ + (0.5 \times 1.286 + 0.5 \times 1.429 + 0.47 \times 1.204)[Fe] \end{cases}$$
(4)

The relative uncertainty <u>on in  $MC_{dust}$ </u>, estimated from the analytical error <u>ion</u> the measured concentrations, does not exceed 6%. As it will be explained in the result section (paragraph 3.1), the values of  $MC_{dust}$  estimated from Equation 4 were found in excellent agreement with the measured gravimetric mass on the filters.

The fractional mass ratio (in percent) of elemental iron ( $MR_{Fe\%}$ ) with respect to the total dust mass concentration,  $MC_{dust-1}$  is then calculated as

239

240

$$MR_{Fe\%} = \frac{[Fe]}{[MC_{Dust}]} \times 100 \tag{5}$$

241

The mass concentration of iron oxides or free-iron ( $MC_{Fe ox}$ ), representing the fraction of elemental iron in the form of hematite and goethite (Fe<sub>2</sub>O<sub>3</sub> and FeOOH, respectively), is equal to

244

245

 $MC_{Fe ox} = MC_{hem} + MC_{goe} \tag{6}$ 

246

where  $MC_{hem}$  and  $MC_{goe}$  are the total masses of hematite and goethite. These can be calculated from the values  $\alpha_{hem}$  and  $\alpha_{goe}$  from XANES analysis, which represent the mass fractions of elemental iron attributed to hematite and goethite, as

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251

$$MC_{hem} = \frac{\alpha_{hem} \, x \, [Fe]}{0.70} \tag{7.a}$$

$$MC_{goe} = \frac{\alpha_{goe} \, x \, [Fe]}{0.63} \tag{7.b}$$

where the values of 0.70 and 0.63 represent the mass molar fractions of Fe in hematite and goethite, respectively. The relative errors of  $MC_{hem}$  and  $MC_{goe}$  are obtained from the uncertainties on-of the values of  $\alpha_{hem}$  and  $\alpha_{goe}$  from XANES analysis (less than 10%).

257 The mass ratio of iron oxides ( $MR_{Fe ox\%}$ ) with respect to the total dust mass can then be calculated as

258

$$MR_{Fe \ ox\%} = MC_{Fe \ ox} \times MR_{Fe \ \%} \tag{8}$$

- 260
- 261

### 262 **3. Experimental protocol**

At the beginning of each experiment, the chamber was evacuated by to  $10^{-4}$ - $10^{-5}$  hPa. Then, the reactor 263 was filled with a mixture of 80% N<sub>2</sub> and 20% O<sub>2</sub> at a pressure slightly exceeding the current atmospheric 264 265 pressure, in order to avoid contamination from ambient air. The eExperiments were conducted at ambient temperature and at a relative humidity <2%. As in Di Biagio et al. (2014; 20162017), dust aerosols 266 267 were generated by mechanical shaking of the parent soils, previously sieved to  $< 1000 \mu m$  and dried at 268 100 °C for about 1 h to remove any residual humidity. About 15 g of soil was placed in a Buchner flask and shaken for about 30 min at 100 Hz by means of a sieve shaker (Retsch AS200). The dust particles 269 produced wproduced by the mechanical shaking, mimicking the saltation processing that soils experi-270 271 ence when eroded by strong winds, as were then injected in the chamber by flushing the flask with N<sub>2</sub> at 10 L min<sup>-1</sup> for about 10-15 min, whilst continuing shaking the soil. Di Biagio et al. (2014; 2017) have 272 273 demonstrated the realism of the generation system concerning the composition and the size distribution 274 of the generated dust with respect to the properties of mineral dust in the atmosphere.

The Ddust was injected for about 10-15 minutes, and left\_remained\_suspended in the chamber for approximately 120 min thanks to the 4-wheel fan located in the bottom of the chamber body. Previous measurements at the top and bottom of the chamber showed that the fan ensures a homogeneous distribution of the dust starting approximately 10 minutes after the end of the injection (Di Biagio et al., 2014). To compensate for the air extracted from the chamber by sampling, a particle-free flow of N<sub>2</sub>/O<sub>2</sub>, regulated in real time as a function of the total volume of sampled air, was re-injected in the chamber. To avoid excessive dilution the flow was limited to 20 L min<sup>-1</sup>. Two experiments per soil type were conducted: a first experiment for sampling on the nuclepore polycarbonate filters (determination of the elemental composition and the iron oxide fraction) and *in situ* measurements of the infrared optical constants (Di Biagio et al., 20162017), and a second experiment <u>sampling</u> on total quartz filter and impactor for the study of dust MAE presented in this paper.

286 Figure 1 illustrates as typical example the time series of the aerosol mass concentration during the two experiments conducted for the Libyan sample. The comparison demonstrates the repeatability of the dust 287 288 concentrations, both in absolute values and in temporal dynamics. It also shows that the mass concen-289 trations decreased very rapidly by gravitational settling within the first 30 minutes of the experiment (see also the discussion in Di Biagio et al., (20162017)), after which concentrations only decrease by 290 291 dilution. The filter sampling was started after this transient phase, and then continued through the end of the experiments, in order to collect enough dust on the filter membranes for subsequent the chemical 292 analysis. Blank samples were collected before the start of the experiments by placing the loaded filter 293 294 holders loaded with filter membranes in line with the chamber and by flushing them for a few seconds with air coming from the chamber. 295

At the end of each experimental series with a given soil sample, the chamber was manually cleaned in order to remove carry-over caused by resuspension of particles deposited to the walls. Background concentrations of aerosols in the chamber vary between 0.5 and 2.0  $\mu$ g m<sup>-3</sup>, i.e., a factor of 500 to 1000 below the operating conditions.

### 300 **34. Results and discussion**

301 The geographical location of the soil collection sites is shown in Figure 2, whereas and the coordinates are summarized in Table 2. As discussed in Di Biagio et al. (2016), the selection of these soils and 302 303 sediments was governed by the need of representing the major arid and semi-arid regions worldwide, the need of taking into account the mineralogical diversity of the soil composition at the global scale, 304 and finally by their availability in sufficient quantities for injection in the chamber. When doing so, we 305 obtained a set of twelve samples distributed worldwide but mostly in Northern and Western Africa 306 (Libya, Algeria, Mali, Bodélé) and the Middle East (Saudi Arabia and Kuwait). Individual samples from 307 the Gobi desert in Eastern Asia, the Namib Desert, the Strzelecki desert in Australia, the Patagonian 308

309 deserts in South America, and the Sonoran Desert in Arizona have also been investigated. The selection 310 of these soils and sediments was made out of 137 individual top-soil samples collected in major arid and 311 semi-arid regions worldwide and representing the mineralogical diversity of the soil composition at the 312 global scale. As discussed in Di Biagio et al. (2017), this large sample set was reduced byto a set of 19 313 samples representing the mineralogical diversity of the soil composition at the global scale and based on their availability in sufficient quantities for injection in the chamber. Because some of the experiments 314 315 did not produce enough dust to perform good-quality optical measurements, -, in this paper we present a set of twelve samples distributed worldwide but mostly fromin Northern and Western Africa (Libya, 316 317 Algeria, Mali, Bodélé) and the Middle East (Saudi Arabia and Kuwait). Individual samples from the 318 Gobi desert in Eastern Asia, the Namib Desert, the Strzelecki desert in Australia, the Patagonian deserts

- 518 <u>Obbi desett in Eastern Asia, the Namio Desett, the Strzetecki desett in Australia, the Patagolitan desett</u>
- 319 in South America, and the Sonoran Desert in Arizona havewere also been investigated.

### 320 **34.1. Elemental composition and iron oxide content**

A total of 41 filters including 15 polycarbonate filters (12 samples and 3 blanks) and 25 quartz filters (12 for the total fraction, 10 for the fine fraction and 3 blanks) were collected for analysis.

The dust mass concentration found by gravimetric analysis varied between 50  $\mu$ g m<sup>-3</sup> and 5 mg m<sup>-3</sup>, in relatively good agreement with the dust mass concentrations.  $MC_{dust-3}$ , from (Equation 4), based on Xray Ray fluorescence FluorescenceXRF analysisaAnalysis: the slope of the linear regression between the calculated and the gravimetric values of  $MC_{dust}$  is 0.90 with R<sup>2</sup> = 0.86.

327 Di Biagio et al. (20162017) showed that clays are the most abundant mineral phases, together with quartz 328 and calcite, and that significant variability exists as function of the compositional heterogeneity of the 329 parent soils. Here we use the Fe/Ca and Si/Al elemental ratios obtained from X-ray FluorescenceXRF 330 analysis to discriminate the origin of used dust samples. These ratios have been extensively used in the 331 past to discriminate the origin of African dust samples collected in the field (Chiapello et al., 1997; 332 Formenti et al., 2011; Formenti et al., 2014a). The values obtained during our experiments are reported 333 in **Table 3.** There is a very good correspondence between the values obtained for the Mali, Libya, Algeria, Mauritania and (to a lesser extent) Morocco experiments to values found in environmental aerosol 334 335 samples by Chiapello et al. (1997) and Formenti et al. (2011; 2014a). These authors indicate that dust from local erosion of Sahelian soils, such as from Mali, have Si/Al ratios in the range of 2-2.5 and Fe/Ca 336 337 ratios in the range 3-20, depending on the time proximity to the erosion event. Dust from sources in the 338 Sahara, such as Libya and Algeria, show Si/Al ratios in the range of 2-3 and Fe/Ca ratios in the range

339 0.7-3, whereas dust from Morocco has Si/Al ratios around 3 and Fe/Ca ratios around 0.4. The only major difference is observed for the Bodélé experiment, for which the Fe/Ca ratio is enriched by a factor of 6 340 341 with respect to the values of 1 found during to the field observations (Formenti et al., 2011; Formenti et 342 al., 2014a). This could reflect the fact that the Bodélé aerosol in the chamber is generated from a sedi-343 ment sample and not from a soil. As a matter of fact, the Bodélé sediment sample is constituted consists of by a very fine powder which becomes very easily airborne. Henceforth, and contrary to the soil sam-344 345 ples, this This powder is likely to be injected in the chamber with little or no size fractionation. Henceforth, the aerosol generated from it . As a consequence, should have a closer the composition to the 346 original powder of the aerosol collected in the chamber could reflect more that of the parent sedimentary 347 soil than the not the other samples. On the other hand, Bristow et al. (2010) and Moskowitz et al. (2016) 348 349 showed that the iron content and speciation of the Bodélé sediments is very heterogeneous at the source 350 scale. For samples from areas other than non-northern African samples, the largest variability is observed 351 for the Fe/Ca values, ranging from 0.1 to 8, whereas the Si/Al ratio varied <u>only</u> between 2.5 and 4.8. In this case, values are available in the literature for comparison (e.g., Cornille et al., 1990; Reid et al., 352 1994; Eltayeb et al., 2001; Lafon et al., 2006; Shen et al., 2007; Radhi et al., 2010; 2011; Formenti et 353 al., 2011; 2014a; Scheuvens et al., 2013, and references within). Values in the PM<sub>2.5</sub> fraction are very 354 consistent with those obtained in the PM<sub>10.6</sub>: their linear correlation has a slope of 1.03 ( $\pm$  0.05) and a  $R^2$ 355 equal to 0.97, suggesting that the elemental composition is relatively size -independent. 356

The mass fraction of total Fe (*MC<sub>Fe</sub>*<sup>%</sup> from Equation 5), also reported in **Table 3**, ranged from 2.8 (Na-357 mibia) to 7.3% (Australia), values found for the Namibia and the Australia samples, respectively. This 358 These are in the range is in good agreement with of values reported in the literature, taking into account 359 360 that differences might be also due to the method (direct measurement/calculation) and/or the size fraction over which the total dust mass concentration is estimated (Chiapello et al., 1997; Reid et al., 1994; 2003; 361 362 Derimian et al., 2008; Formenti et al., 2001; 2011; 2014a; Scheuvens et al., 2013). The agreement of *MC<sub>Fe%</sub>* values obtained by the XRF analysis of polycarbonate filters (Equation 5) and those obtained 363 364 from the XRF analysis of the quartz filters, normalized to the measured gravimetric mass is well within 10% (that is, the percent error of each estimate). An the e-Exceptions are the samples of from Bodélé 365 and Algeria, for which the values obtained from the analysis of the quartz filters are significantly lower 366 367 than those obtained from the nuclepore filters (3.1% versus 4.1% for Bodélé and 4.3% versus 6.8% for 368 Algeria). We treat that as an additional source of error in the rest of the analysis, and add it to the total uncertainty. In the PM2.5 fraction, the content of iron is more variable, ranging from 4.4% (Morocco) to 369

33.6% (Mali), showing a size dependence. A word of caution on this conclusion as-is that the two estimates are not necessarily consistent in the way that the total dust mass is estimated (from Equation 4 for
the PM<sub>10.6</sub> fraction and by gravimetric weighing <u>form</u> the PM<sub>2.5</sub>).

373 Finally, between 11 and 47% of iron in the samples can be attributed to iron oxides, in variable propor-374 tions between hematite and goethite. The iron oxide fraction of total Fe in this study is on-at the lower end of the range (36-72%) estimated for field dust samples of Saharan/Sahelian origin (Formenti et al. 375 376 2014b). The highest value of Formenti et al. (2014b), obtained for a sample of locally-emitted dust col-377 lected at the Banizoumbou station in the African Sahel, is anyhow in excellent agreement with the value of 62% obtained for an experiment (not shown here) using a soil collected in the same area. Likewise, 378 379 the proportions between hematite and goethite (not shown) are reproduced, showing that goethite is more 380 abundant than hematite. The mass fraction of iron oxides ( $MR_{Fe \ ox^{9/6}}$ ), estimated from Equation 8 and 381 shown in Table 3, ranges between 0.7% (Kuwait) to and 3.6% (Australia), which is in the range of 382 available field estimates (Formenti et al., 2014a; Moskowitz et al., 2016). For China, our value of MRFe <sub>ox%</sub> is lower by almost a factor of 3 in comparison with compared to that obtained on the same dust of the 383 384 same origin sample by Alfaro et al. (2004) (0.9% against 2.8%), whereas on a sample from Niger (how-385 ever not considered in this study) our estimates and that by Alfaro et al. (2004) agree perfectly agree (5.8%). A possible underestimate of the iron oxide fraction for samples other than those from the Sahara-386 387 Sahel area could be due to the fact that - opposite to the experience of Formenti et al. (2014b) - the linear deconvolutions of the XANES spectra were not always satisfactory (see Figure S1 in the supplementary). 388 389 This resulted in a significant residual between the observed and fitted XANES spectra. IndeedIn fact, the mineralogical reference for hematite is obtained from a soil from Niger (Table 1) and might not be 390 391 fully suitable for representing aerosols of different origins. Additional differences could arise from dif-392 ferences in the size distributions of the generated aerosol. As a matter of fact, the number fraction of 393 particles in the size classes above 0.5 µm in diameter is are different in the dust aerosol generated in the 394 Alfaro et al. (2004) study with respect compared to ours. In the study by Alfaro et al. (2004), the number 395 fraction of particles is lowest in the 0.5-0.7 size class and highest between 1 and 5 µm. On the contraryIn 396 contrast, in our study the number fraction is lowest in the 1-2 µm size range and highest between 0.5 397 and 0.7 µm. These differences could vield either be due to differences in the chemical composition 398 and/or to a difference in the total mass in the denominator of Equation 8.

# **399 34.2. Spectral and size\_-variability of the mass absorption efficiency**

400 The spectral mass absorption efficienciesy (MAE) at 375, 407, 532, 635, and 850 nm for the PM<sub>10.6</sub> and 401 the PM<sub>2.5</sub> dust fractions are summarized in Table 4 and displayed in Figure 3. Regardless of particle size, the MAE values decrease with increasing wavelength (almost one order of magnitude between 375 402 403 and 850 nm), and display a larger variability at shorter wavelengths. The MAE values for the PM<sub>10.6</sub> range from 37 (± 3)  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup> to 135 (± 11)  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup> at 375 nm, and from 1.3 (± 0.1)  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup> to 15 404 (± 1) 10<sup>-3</sup> m<sup>2</sup> g<sup>-1</sup> at 850 nm. Maxima are found for the Australia and Algeria samples, whereas the minima 405 406 are for Bodéelé and Namibia, respectively at 375 and 850 nm. In the PM2.5 fraction, the MAE values range from 95 (± 8)  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup> to 711 (± 70)  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup> at 375 nm, and from 3.2 (± 0.3)  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup> to 36 407  $(\pm 3)$  10<sup>-3</sup> m<sup>2</sup> g<sup>-1</sup> at 850 nm. Maxima at both 375 and 850 nm are found for the Morocco sample, whereas 408 the minima are for Algeria and Namibia, respectively. The MAE values for mineral dust resulting from 409 410 this work are in-relatively in good agreement with the estimates available in the literature (Alfaro et al., 2004; Linke et al., 2006; Yang et al., 2009; Denjean et al., 2016), reported in Table 5. For the China 411 Ulah Buhn sample, Alfaro et al. (2004) reported 69.1 10<sup>-3</sup> and 9.8 10<sup>-3</sup> m<sup>2</sup> g<sup>-1</sup> at 325 and 660 nm, respec-412 tively. The former is lower than the value of 99 10<sup>-3</sup> m<sup>2</sup> g<sup>-1</sup> that we obtain by extrapolating our measure-413 414 ment at 375 nm. Likewise, our values for the Morocco sample are higher than reported by Linke et al. (2006) at 266 and 660 nm. Conversely, the agreement with the estimates of Yang et al. (2009) for mineral 415 416 dust locally re-suspended in Xianghe, near Beijing (China) is very good at all wavelengths between 375 and 880 nm. As expected, the MAE values for mineral dust resulting from this work are almost one 417 418 order of magnitude smaller than for other absorbing aerosols. For black carbon, MAE values are in the range of 6.5–7.5 m<sup>2</sup> g<sup>-1</sup> at 850 nm (Bond and Bergstrom, 2006; Massabò et al., 2016), and vary-decrease 419 in a linear way with the logarithm of the inversely with wavelength. For brown carbon, the reported 420 MAE range between 2.3–7.0 m<sup>2</sup> g<sup>-1</sup> at 350 nm (Chen and Bond, 2010; Kirchstetter et al., 2004; Massabò 421 et al., 2016), 0.05–1.2 m<sup>2</sup> g<sup>-1</sup> at 440 nm (Wang et al., 2016) and 0.08–0.72 m<sup>2</sup> g<sup>-1</sup> at 550 nm (Chen and 422 Bond, 2010). 423

The analysis of **Table 4** indicates that, at every wavelength, the MAE values in the PM<sub>2.5</sub> fraction are equal or higher than those for PM<sub>10.6</sub>. The PM<sub>2.5</sub>/PM<sub>10.6</sub> MAE ratios reach values of 6 for the Mali sample, but are mostly in the range 1.5-3 for the <u>remaining other</u> aerosols. <u>The v</u>Values decrease with wavelength up to 635 nm, whereas at 850 nm they have values comparable to those at 375 nm. The observed size\_-dependence of the MAE values is consistent with the expected behavior of light absorption of particles in the Mie and geometric optical regimes that <u>concern are relevant for</u> the two size fractions. Light\_-absorption of particles of size<u>s</u> smaller or equivalent to <u>the</u> wavelength is proportional to their

431 bulk volume, whereas for larger particles absorption occurs on their surface only (Bohren and Huffmann, 1983). On the other hand, the size-resolved measurements of Lafon et al. (2006) show that the proportion 432 433 (by volume) of iron oxides might be higher in the coarse than in the fine fraction, which would counteract 434 the size-dependence behavior of MAE. To validate the observations, we calculated the spectrally-re-435 solved MAE values in the two size fractions using the Mie code for homogeneous spherical particles 436 (Bohren and Huffmann, 1983) and the number size distribution estimated by (Di Biagio et al., 437  $\frac{2016}{2017}$  and averaged over the duration of filter sampling. We estimated the dust complex refractive index as a volume-weighted average of a non-absorbing dust fraction having the refractive index of 438 439 kaolinite, the dominant mineral in our samples (see Di Biagio et al., 20162017), from Egan et Hilgeman 440 (1979) and an absorbing fraction estimated from the mass fraction of iron oxides and having the refractive index of hematite (Bedidi and Cervelle, 1993). The rResults of this calculation indicate that the 441 442 observed size-dependent behavior is well reproduced at all wavelengths, even in the basic hypothesis 443 that the mineralogical composition does not change with size. The only exception is 850 nm, where at 444 times, PM<sub>2.5</sub>/PM<sub>10.6</sub> MAE ratio is much higher than expected theoretically. We attribute that to the rela-445 tively high uncertainty affecting the absorbance measurements at this wavelength, where the signal-to-446 noise ratio is low. Indeed, the two sets of values (MAE in the PM2.5 fraction and MAE in the PM10.6 447 fraction) are not statistically different according to a two-pair t-test (0.01 and 0.05 level of confidence), 448 confirming that any attempt of differentiation of the size -dependence at this wavelength would require 449 a stronger optical signal.

450 The analysis of the spectral dependence, using a the power-law function fit as (from Equation 2), pro-451 vides with the values of the Angstrom Absorption Exponent (AAE), also reported in **Table 4**. Contrary 452 to the MAE values, there is no statistically significant size\_dependence of the AAE values, ranging from 453 2.5 ( $\pm$  0.2) to 4.1 ( $\pm$  0.3), with an average of 3.3 ( $\pm$  0.7), for the PM<sub>10.6</sub> size fraction and between 2.6 ( $\pm$ 454 0.2) and 5.1 ( $\pm$  0.4), with an average of 3.5 ( $\pm$  0.8), for the PM<sub>2.5</sub> fraction. Our values are in the range of 455 those published in the open-literature (Fialho et al., 2005; Linke et al., 2006; Müller et al., 2009; Petzold 456 et al., 2009; Yang et al., 2009; Weinzierl et al., 2011; Moosmüller et al., 2012; Denjean et al., 2016), 457 shown in **Table 5**. AAE values close to 1.0 are found for urban aerosols where fossil fuels combustion 458 is dominant, while AAE values for brown carbon (BrC) from incomplete combustion are in the range 459 3.5-4.2 (Yang et al., 2009; Chen et al., 2015; Massabò et al., 2016).

Finally, **Figure 4** shows correlations between <u>the MAE</u> values in the PM<sub>10.6</sub> fraction (Figure <u>34</u>.a) and in the PM<sub>2.5</sub> fraction (Figure <u>34</u>.b) and the estimated percent mass fraction of iron and iron oxides 462 ( $MC_{Fe\%}$  and  $MC_{Fe ox\%}$ ), respectively. Regardless of the size fraction, t<sup>+</sup> he correlation between the MAE values and the percent mass of total elemental iron are satisfactory. Higher higher correlations are ob-463 tained at 375, 407 and 532 nm<sub>.</sub>, and in the PM<sub>2.5</sub> fraction, where a linear correlation with  $R^2$  up to 0.94 464 are obtained. Best correlations are obtained when forcing the intercept to zero, indicating that elemental 465 iron fully accounts for the measured absorption. At these wavelengths, linear correlations with the mass 466 fraction of iron oxides are loose low in the PM<sub>10.6</sub> mass fraction ( $R^2$  up to 0.38-0.62), but again satisfac-467 toryhigher in the PM<sub>2.5</sub> fraction ( $R^2$  up to 0.83-0.99), where, whoever however, one should keep in mind 468 that they have been established only indirectly by considering the ratio of iron oxides to elemental iron 469 470 independent of size. At 660 and 850 nm, little or no robust correlations are is obtained, often based on 471 very few data points and with very low MAE values. It is noteworthy that, in both size fractions, the 472 linear correlation yields a non-zero intercept is obtained, indicating a contribution from minerals other 473 minerals butthan iron oxides account to for the measured absorption.

### 474 **45**. Conclusive remarks

In this paper, we reported new laboratory measurements of the shortwave mass absorption efficiency (MAE) of mineral dust of different origins and as a function of size and wavelength in the 375-850 nm range. <u>Results Our results have beenwere</u> obtained in the CESAM simulation chamber using <u>generated</u> mineral dust <u>generated</u> from natural parent soils, <u>in combination with and</u> optical and gravimetric analysis on extracted samples.

Our results can be summarized as follows: at 375 nm, the MAE values are lower for the *PM*<sub>10.6</sub> mass fraction (range 37-135  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup>) than for the PM<sub>2.5</sub> fraction (range 95-711  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup>), and vary opposite to wavelength as  $\lambda^{-AAE}$ , where AAE (Angstrom Absorption Exponent) averages between 3.3-3.5 regardless of size fraction. These results deserve some conclusive concluding comments:

484 The size\_dependence, vielding\_characterized by significantly higher MAE values in the fine fraction (PM2.5) than for the in the bulk (PM10.6-) aerosol, indicates that light\_absorption by min-485 eral dust can be important even during atmospheric transport over heavily polluted regions, when 486 487 where dust concentrations are significantly lower than at emission. This can be shown by comparing the aerosol absorption optical depth (AAOD) at 440 nm for China, a well-known mixing 488 489 region of mineral dust and pollution (e.g., Yang et al., 2009; Laskin et al., 2014; Wang et al., 2013), as well as offshore western Africa where large urban centers are downwind of dust 490 491 transport areas (Petzold et al., 2011). Laskin et al. (2014) reports that the average AAOD in China 492the area is of the order of  $0.1_{5}$  for carbonaceous absorbing aerosols (sum of black and brown493carbon; Andreae and Gelencsér, 2006). This is lower or comparable to the AAOD of 0.17 and4940.11 at 407 nm (fine and total and fine mass fractions, respectively) that we obtain derive by a495simple calculation (AAOD = MAE x MCdust x H), where from MAE are the values estimated in496this study, (, MCdust-, the dust mass concentrations typically observed in the areaurban area of497Beijing during dust storms (Sun et al., 20012005), and H, a scale height factor of 1 km).

- 498 The spectral variability of the dust MAE values, represented by the AAE parameter, is equal in 499 the PM<sub>2.5</sub> and PM<sub>10.6</sub> mass fractions. This suggests that, for a given size distribution, the possible 500 variation of dust composition with size does not affect in a significant way the spectral behavior 501 of the absorption properties. Our average value for AAE is  $3.3 \pm 0.7$ , higher than for black carbon, 502 but in the same range than as light-absorbing organic (brown) carbon. As a result, depending on the environment, there can be some ambiguity in apportioning the AAOD based on spectral de-503 pendence. Bahadur et al. (2012) and Chung et al. (2012) couple the AAE and the spectral de-504 505 pendence of the total AOD (and/or its scattering fraction only) to overcome this problem. Still, 506 Bahadur et al. (2012) show that there is an overlap in the scatterplots of the spectral dependence of the scattering and absorption fractions of the AOD based on an analysis of ground-based re-507 mote sensing data for mineral dust, urban, and non-urban fossil fuel over California. A closer 508 look should be taken at\_to-observations in mixing areas where biomass burning aerosols may 509 have different chemical composition and/or mineral dust has heavy loadings should be given in 510 511 order to generalize the clear separation observed in the spectral dependences of mineral dust and 512 biomass burning (Bahadur et al., 2012). This aspect is relevant to the development of remote 513 sensing retrievals of light-absorption by aerosols from space, and their assimilation in climate 514 models (Torres et al., 2007; Buchard et al., 2015; Hammer et al., 2016).
- There is an important sample-to-sample variability in our dataset of MAE values for mineral dust aerosols. At 532 nm, our <u>estimated\_average\_MAE values\_average toare\_at</u>  $34 \pm 14 \text{ m}^2 \text{ g}^{-1}$  and 78  $\pm 70 \text{ m}^2 \text{ g}^{-1}$  in the PM<sub>10.6</sub> and PM<sub>2.5</sub> mass fractions, respectively. Figure 3, showing the correlation with the estimated mass fraction of elemental iron and iron oxides, suggests that this variability could be related to the regional differences of the mineralogical composition of the parent soils. These observations lead to <u>different considerations\_further conclusions</u>. To start with, our study reinforces the need for regionally-resolved representation of the light\_-absorption properties of

mineral dust in order to improve the representation of its effect on climate. As a matter of fact, 522 the natural variability of the absorption properties that we obtain from our study is in the range 523 524 50-100%, even when we limit ourselves to smaller spatial scales, for example those of from north 525 Africa (samples from Libya, Algeria, Mali and Bodélé). This is far above the  $\pm$  5% sensitivity factor used by Solmon et al. (2008) to vary the single scattering albedo (as a proxy of absorption) 526 of mineral dust over western Africa, and to show how this could drastically change the climate 527 response in the region. As a comparison, Solmon et al. (2008) showed that varying the single 528 scattering albedo of mineral dust over western Africa by  $\pm$  5%, that is, varying the co-albedo (or 529 absorption) by 45% (0.1±0.045) could drastically change the climate response in the region. 530

The question is then "how to represent this regional variability?" As-Like Moosmüller et al. 531 (2012) and Engelbrecht et al. (2016), we found that elemental iron is a very good proxy for the 532 533 MAE, especially in the  $PM_{2.5}$  fraction, where iron-bearing absorbing minerals (hematite, goe-534 thite, illite, smectite clays) would beare more concentrated. In the coarse fraction, Ca-rich minerals, quartz, and feldspars could also play a role, and that could result in the observed lowered 535 correlation (although adding a term proportional to elemental Ca does not ameliorate-improve 536 the correlation-result in the present study). The correlation of the spectral MAE values with the 537 iron oxide fraction is satisfactory but rather noisy, also owing to some uncertainty in the quanti-538 539 fication of iron oxides from X-Ray Absorption absorption measurements. In this case, the intercept is significantly different from zero, again indicating that a small but elear distinct fraction 540 541 of absorption is due to minerals other than iron oxides. There are contrasting results on this topic: Alfaro et al. (2004) found an excellent correlation between MAE and the iron oxide content, 542 whereas Klaver et al. (2011) found that the single scattering albedo (representing the capacity of 543 an aerosol population to absorb light with respectin relation to extinction) was almost independ-544 545 ent on the mass fraction of iron oxides. Moosmüller et al. (2012) disagreed, pointing out to the uncertainty in the correction procedure of the measurement of absorption by Klaver et al. (2011). 546 As a matter of fact, Klaver et al. (2011) and Alfaro et al. (2004) used the same correction proce-547 dure. It is more likely that the lack of correlation found in Klaver et al. (2011) is due to the fact 548 that other minerals other than iron oxides contribute to absorption, in particular at their working 549 550 wavelength (567 nm), where the absorption efficiency of iron oxides starts to weaken. Clearly, the linear correlation between elemental iron in mineral dust and its light-absorption properties 551

552 could ease the application and validation of climate models that are now starting to include in-553 cluding the representation of the mineralogy (Perlwitz et al., 2015a; 2015b; Scanza et al., 2015). 554 Also, they this would facilitate detecting source regions based on remote sensing of dust absorp-555 tion in the UV-VIS spectral region (e.g., Hsu et al., 2004). However, such a quantitative relation-556 ship cannot be uniquely determined from these studies, including the present one, which use 557 different ways of estimating elemental iron, iron oxides, and the total dust mass. A more robust 558 estimate should be obtained from by estimating the imaginary parts of the complex refractive indices associated to-with these measurements of absorption, and their dependence on the min-559 560 eralogical composition.

### 561 Author contributions

L. Caponi, P. Formenti, D. Massabò, P. Prati, C. Di Biagio, and J. F. Doussin designed the chamber 562 563 experiments and discussed the results. L. Caponi and C. Di Biagio realized conducted the experiments 564 with contributions by M. Cazaunau, E. Pangui, P. Formenti, and J.F. Doussin. L. Caponi, D. Massabò 565 and P. Formenti performed the full data analysis with contributions by C. Di Biagio, P. Prati and J.F. Doussin. L. Caponi, P. Formenti and S. Chevaillier performed the XRF measurements. P. Formenti and 566 567 G. Landrot performed the XAS measurements. D. Massabò performed the MWAA and the gravimetric 568 measurements.-M. O. Andreae, K. Kandler, T. Saeed, S. Piketh, D. Seibert, and E. Williams collected the soil samples used for experiments. L. Caponi, P. Formenti, D. Massabò and P. Prati wrote the man-569 570 uscript with comments from all co-authors.

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### 932 Table captions

**Table 1.** Characteristics of the standards used for the quantification of the iron oxides in the XAS anal-ysis.

935 **Table 2.** <u>Summary Geographical of information on the soil samples used in this work.</u>

**Table 3.** Chemical characterisation of the dust aerosols in  $PM_{10.6}$  and  $PM_{2.5}$  (in parentheses) size fractions. Columns 3 and 4 <u>report-give</u> the Si/Al and Fe/Ca elemental ratios obtained from X-Ray Fluorescence analysis. The uncertainty o<u>f</u>n each individual value is estimated to be 10%. Column 5 <u>reports</u> <u>shows</u> *MR<sub>Fe%</sub>*, the fractional mass of elemental iron with respect to the total dust mass concentration (uncertainty 10%). Column 5 reports *MR<sub>Fe%</sub>*, the mass fraction of iron oxides with respect to the total dust mass concentration (uncertainty 15%). For PM<sub>2.5</sub> the determination of the Si/Al ratio is impossible due to the composition of the filter <u>medium-membranes (quartz)</u>.

Table 4. Mass absorption efficiency (MAE,  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup>) and Ångström Absorption Exponent (AAE) in the PM<sub>10.6</sub> and PM<sub>2.5</sub> size fractions. Absolute errors are in brackets.

Table 5. Mass absorption efficiency (MAE,  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup>) and Ångström Absorption Exponent (AAE) of from the literature data discussed in the paper

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## 948 Figure captions

Figure 1. Time series of aerosol mass concentration in the chamber for the two companion experiments
(Libyan dust-sample). Experiment 1 (top panel) was dedicated to the determination of the chemical
composition (including iron oxides) by sampling on polycarbonate filters. Experiment 2 (bottom panel)
was dedicated to the determination of the absorption optical properties by sampling on quartz filters.

**Figure 2.** Locations (red stars) of the soil and sediment samples used to generate dust aerosols.

Figure 3. Spectral dependence of the MAE values for the samples investigated in this study in the PM<sub>10.6</sub>
(left) and in the PM<sub>2.5</sub> (right) mass fractions.

**Figure 4.** Illustration of the links between the MAE values and the dust chemical composition found in this study. Left column, from top to bottom: linear regression between <u>the MAE values between in the</u> range from 375 to and 850 nm and the fraction of elemental iron with respect to relative to the total dust mass ( $MR_{Fe\%}$ ) in the PM<sub>10.6</sub> fraction; Middle column: same as left column but respect to for the mass

- 960 fraction of iron oxides <u>relative</u> to the total dust mass ( $MR_{Fe ox\%}$ ) in the PM<sub>10.6</sub> size fraction; Right column:
- 961 same as left column but in the  $PM_{2.5}$  size fraction.

**Table 1.** Characteristics of the standards used for the quantification of the iron oxides in the XAS anal-

964	ysis.
	-

	Standard	StoechiometricStoichiometric Formula	Origin
	Illite of Puy	$(Si_{3.55}Al_{0.45})(Al_{1.27}Fe_{0.36}Mg_{0.44})O_{10}(OH)_2(Ca_{0.01}Na_{0.01}K_{0.53}X(I)_{0.12})$	Puy, France
	Goethite	FeO OH	Minnesota
	Hematite	Fe <sub>2</sub> O <sub>3</sub>	Niger
	Montmorillonite	$(Na,Ca)_{0,3}(Al,Mg)_2Si_4O10(OH)_2 \cdot n(H_2O)$	Wyoming
	Nontronite	Na0.3Fe2(Si,Al)4O10(OH)2·nH2O	Pennsylvania
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Geographical area	Sample	Desert area	Geographical coordinates
	Morocco	East of Ksar Sahli	31.97°N, 3.28°W
Sahara	Libya	Sebha	27.01°N, 14.50°E
	Algeria	Ti-n-Tekraouit	23.95°N, 5.47°E
Sahal	Mali	Dar el Beida	17.62°N, 4.29°W
Sallel	Bodélé	Bodélé depression	17.23°N, 19.03°E
Middle Feet	Saudi Arabia	Nefud	27.49°N, 41.98°E
Wilddie East	Kuwait	Kuwaiti	29.42°N, 47.69°E
Southern Africa	Namibia	Namib	21.24°S, 14.99°E
Eastern Asia	China	Gobi	39.43°N, 105.67°E
North America	Arizona	Sonoran	33.15 °N, 112.08°W
South America	Patagonia	Patagonia	50.26°S, 71.50°W
Australia	Australia	Strzelecki	31.33°S, 140.33°E

**Table 2.** Geographical information on the soil samples used in this work.

971 **Table 3.** Chemical characterisation of the dust aerosols in  $PM_{10.6}$  and  $PM_{2.5}$  (in parentheses) size frac-972 tions. Columns 3 and 4 give the Si/Al and Fe/Ca elemental ratios obtained from X-Ray Fluorescence 973 analysis. The uncertainty of each individual value is estimated to be 10%. Column 5 shows MRFe%, the 974 fractional mass of elemental iron with respect to the total dust mass concentration (uncertainty 10%). 975 Column 5 reports MRFe%, the mass fraction of iron oxides with respect to the total dust mass concentra-976 tion (uncertainty 15%). For PM2.5 the determination of the Si/Al ratio is impossible due to the composi-977 tion of the filter-membranes (quartz) 978 Chemical characterisation of the dust aerosols in PM<sub>10.6</sub> and PM<sub>2.5</sub> (in parentheses) size fractions. Col-979 umns 3 and 4 report the Si/Al and Fe/Ca elemental ratios obtained from X-Ray Fluorescence analysis. 980 The uncertainty on each individual value is estimated to be 10%. Column 5 reports MR<sub>Fe%</sub>, the fractional 981 mass of elemental iron with respect to the total dust mass concentration (uncertainty 10%). Column 5 982 reports MR<sub>Fe ox%</sub>, the mass fraction of iron oxides with respect to the total dust mass concentration (un-983 certainty 15%). For PM2.5 the determination of the Si/Al ratio is impossible due to the composition of

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the filter medium.

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Geographical area	Sample	Si/Al	Fe/Ca	MC <sub>Fe%</sub>	MC <sub>Fe-ox%</sub>
	Morocco	3.12 ()	0.24 (0.28)	3.6 (4.4)	1.4 (1.8)
Sahara	Libya	2.11 ()	1.19 (1.12)	5.2 (5.6)	3.1 (3.4)
	Algeria	2.51 ()	3.14 (4.19)	6.6 (5.4)	2.7 (2.2)
Sahal	Mali	3.03 ()	2.99 (3.67)	6.6 (33.6)	3.7 (18.7)
Sallel	Bodélé	5.65 ()	12.35 ()	4.1 ()	0.7 ()
Middle East	Saudi Arabia	2.95 ()	0.29 (0.27)	3.8 (5.1)	2.6 (3.5)
	Kuwait	3.15 ()	0.89 (1.0)	5.0 (13.6)	1.5 (4.2)
Southern Africa	Namibia	3.41 ()	0.11 (0.10)	2.4 (6.9)	1.1 (3.1)
Eastern Asia	China	2.68 ()	0.77 (0.71)	5.8 (13.6)	0.9 (2.5)
North America	Arizona	3.30 ()	0.95 ()	5.3 ()	1.5 ()
South America	Patagonia	4.80 ()	4.68 (4.64)	5.1 ()	1.5 ()
Australia	Australia	2.65 ()	5.46 (4.86)	7.2 (11.8)	3.6 (5.9)

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		PM <sub>10.6</sub>					
Geographical area	Sample	375 nm	407 nm	532 nm	635 nm	850 nm	AAE
	Morocco	()	()	()	()	()	()
Sahara	Libya	89 (11)	75 (9)	30 (5)	()	()	3.2 (0.3)
	Algeria	99 (10)	80 (10)	46 (7)	16 (3)	15 (3)	2.5 (0.3)
Sahal	Mali	()	103 (18)	46 (12)	()	()	()
Saner	Bodélé	37 (4)	25 (3)	13 (2)	6(1)	3 (1)	3.3 (0.3)
Middle Feat	Saudi Arabia	90 (9)	79 (8)	28 (3)	6 (1)	4 (1)	4.1 (0.4)
Middle East	Kuwait	()	()	()	()	()	2.8 (0.3)
Southern Africa	Namibia	52 (7)	49 (7)	13 (3)	5 (2)	1 (2)	4.7 (0.5)
Eastern Asia	China	65 (8)	58 (7)	32 (4)	8 (2)	7 (2)	3 (0.3)
North America	Arizona	130 (15)	99 (12)	47 (7)	21 (4)	13 (4)	3.1 (0.3)
South America	Patagonia	102 (11)	80 (9)	29 (4)	17 (2)	10 (2)	2.9 (0.3)
Australia	Australia	135 (15)	121 (13)	55 (7)	26 (4)	14 (3)	2.9 (0.3)

**Table 4.** Mass absorption efficiency (MAE,  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup>) and Ångström Absorption Exponent (AAE) in the PM<sub>10.6</sub> and PM<sub>2.5</sub> size fractions. Absolute errors are in brackets.

		PM <sub>2.5</sub>					
Geographical area	Sample	375 nm	407 nm	532 nm	635 nm	850 nm	AAE
	Morocco	107 (13)	88 (11)	34 (6)	14 (3)	15 (4)	2.6 (0.3)
Sahara	Libya	132(17)	103 (14)	33 (7)	()	()	4.1 (0.4)
	Algeria	95(8)	71 (11)	37 (7)	12 (5)	12 (5)	2.8 (0.3)
Sahel	Mali	711 (141)	621 (124)	227 (78)	()	()	3.4 (0.3)
	Bodelé	()	()	()	()	()	()
	Saudi Arabia	153 (18)	127 (15)	42 (7)	8 (4)	6 (4)	4.5 (0.5)
Middle East	Kuwait	270 (100)	324 (96)	()	54 (52)	()	3.4 (0.3)
Southern Africa	Namibia	147 (36)	131 (32)	31 (21)	6 (16)	3 (15)	5.1 (0.5)
Eastern Asia	China	201 (30)	176 (26)	89 (17)	14 (10)	23 (10)	3.2 (0.3)
North America	Arizona	()	()	()	()	()	()
South America	Patagonia	()	()	()	()	()	2.9 (0.3)
Australia	Australia	335 (39)	288 (33)	130 (19)	57 (11)	36 (9)	2.9 (0.3)

	Geo- graph ical area	Sample	266 nm	325 nm	428 nm	532 nm	660 nm	880 nm	106 4 nm	AAE
		Morocco*								2.25-
		Morocco, PM <sub>2.5</sub> <sup>£</sup>								2.0-6.5
		Morocco, submicron <sup>#</sup>	1100			60			30	4.2
	Sa-	Egypt, submicron <sup>#</sup>	810			20				5.3
	hara	Tunisia <sup>\$</sup>		83			11			
		Saharan, transported <sup>µ</sup>								$2.9\pm0.2$
		Saharan, transported $(PM_{10})^{\%}$			37	27%%	15%%%			2.9
		Saharan, transported $(PM_1)^{\%}$			60	40%%	30%%%			2.0
	Sahel	Niger <sup>\$</sup>		124			19			
	East-	China <sup>\$</sup>		69			10			
	ern Asia	China <sup>&amp;</sup>		87 <sup>&amp;</sup> &	50 <sup>&amp;&amp;&amp;</sup>	27 <sup>&amp;&amp;&amp;</sup> &	13	1		3.8
	Ara-									
	bian									
	renin-									
	N/NF									
	Af-	Various locations <sup>@</sup>								2.5-3.9
	rica,									
	Cen-									
	tral									
	Asia									
996	* Müller et	al. (2008)								
997	<sup>t</sup> Petzold et	al. ( <del>2008</del> <u>2009</u> )								
998	<sup>#</sup> Linke et a	(2006)								
1000	$\mu$ Fialho et	al. (2004) al. (2005)								
1000	<sup>%</sup> Denjean	et al. (2005); <sup>%%</sup> at 528 nm, <sup>%%</sup>	% at 652 nr	n						
1002	<sup>&amp;</sup> Yang et a	ıl. (2009); <sup>&amp;&amp;</sup> at 375 nm, <sup>&amp;&amp;&amp;</sup> at	470 nm, <sup>&amp;</sup>	<sup>&amp;&amp;&amp;</sup> at 5	90 nm					
1003	<sup>@</sup> Mossmül	ler et al. (2012)								
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**Table 5.** Mass absorption efficiency (MAE,  $10^{-3}$  m<sup>2</sup> g<sup>-1</sup>) and Ångström Absorption Exponent (AAE)995from the of-literature data discussed in the paper

**Figure 1.** <u>Time series of aerosol mass concentration in the chamber for two companion experiments</u> (<u>Libyan dust</u>).<u>Time series of aerosol mass concentration in the chamber for the two companion experi-</u> <u>ments (Libya sample</u>). Experiment 1 (top panel) was dedicated to the determination of the chemical composition (including iron oxides) by sampling on polycarbonate filters. Experiment 2 (bottom panel) was dedicated to the determination of the absorption optical properties by sampling on quartz filters.

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**Figure 2.** Locations (red stars) of the soil and sediment samples used to generate dust aerosols.

Figure 3. Spectral dependence of the MAE values for the samples investigated in this study in the PM<sub>10.6</sub>
(left) and in the PM<sub>2.5</sub> (right) mass fractions.



1021	Figure 4. <u>Illustration of the links between the MAE values and the dust chemical composition found in</u>
1022	this study. Left column, from top to bottom: linear regression between the MAE values in the range from
1023	375 to 850 nm and the fraction of elemental iron relative to the total dust mass (MR <sub>Fe%</sub> ) in the PM <sub>10.6</sub>
1024	fraction; Middle column: same as left column but for the mass fraction of iron oxides relative to the total
1025	dust mass (MR <sub>Fe ox%</sub> ) in the PM <sub>10.6</sub> size fraction; Right column: same as left column but in the PM <sub>2.5</sub> size
1026	fraction.
1027	Illustration of the links between the MAE values and the dust chemical composition found in this study.
1028	Left column, from top to bottom: MAE values between 375 and 850 nm versus the fraction of elemental
1029	iron with respect to the total dust mass (MR <sub>Fe%</sub> ) in the PM <sub>10.6</sub> fraction; Middle column: same as left
1030	column but versus the mass fraction of iron oxides to the total dust mass (MRFe ox%) in the PM10.6-size
1031	fraction; Right column: same as left column but in the PM2.5 size fraction. The linear regression lines
1032	between MAE and MR <sub>Fe%</sub> and MAE and MR <sub>Fe Ox%</sub> are reported in each plot.
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