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Key Points:

- Three types of iron-containing particles were observed including freshwater diatoms that comprised 38% of particles between 10 and 18 μm
- Aspect ratios increased with particle size suggesting asphericity may aid in transporting supercoarse particles
- Low density and high aspect ratios of diatoms can increase residence times in surface waters and the likelihood of iron dissolution

Supporting Information:

- Supporting Information S1

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Atmospheric Transport of North African Dust-Bearing Supermicron Freshwater Diatoms to South America: Implications for Iron Transport to the Equatorial North Atlantic Ocean

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Abstract The equatorial North Atlantic Ocean (NAO) is a nutrient-limited ecosystem that relies on the deposition of long-range transported iron (Fe)-containing aerosols to stimulate primary productivity. Using microscopy, we characterized supermicron and supercoarse mode African aerosols transported to the western NAO in boreal winter/spring. We detected three particle types including African dust, primary biological aerosol particles, and freshwater diatoms (FDs). FDs contained 4% Fe by weight due to surficial dust inclusions that may be susceptible to chemical processing and dissolution. FDs were typically larger than dust particles and comprised 38% of particles between 10 and 18 μm in diameter. The low density, high surface-area-to-volume ratio, and large aspect ratios of FD particles suggest a mechanism by which they can be carried great distances aloft. These same properties likely increase the residence time of FDs in surface waters thereby increasing the time for Fe dissolution and their potential impact on marine biogeochemical cycles.

Plain Language Summary Atmospheric aerosols from Africa are transported by the trade winds to the western equatorial North Atlantic Ocean every winter and spring and can contain nutrients, such as iron (Fe). In this study, we measured the size and composition of supermicron (diameter (d) > 1 μm) aerosols collected at a site on the northeast coast of South America. Using electron microscopy, we found three distinct Fe-containing particle types: mineral dust, freshwater diatoms from African paleolakes, and pollen grains; all three particle types extended into the supercoarse mode (d > 10 μm). Particle asphericity increased with increasing particle size and could explain in part the long-range transport of supercoarse particles. Electron mapping of freshwater diatoms also revealed surficial Fe-rich inclusions. Once deposited in the ocean, the asphericity and light density of freshwater diatom particles likely increases their residence time and therefore, the time for Fe dissolution in the surface ocean compared to dust.

1. Introduction

There is great interest in the atmospheric transport of mineral dust because of the role that associated nutrients such as iron (Fe) play in biogeochemical cycles (Mahowald et al., 2005; Mills et al., 2004; Moore et al., 2013; Okin et al., 2011; Rizzolo et al., 2017). The transport distance of dust is determined in part by the particle size, density, and shape with larger, denser, and more spherical particles predicted to settle more quickly out of the atmosphere (Huang et al., 2020; Mallios et al., 2020). Consequently, particles larger than 5 μm , which includes supercoarse mode particles (diameter > 10 μm), are often ignored in climate studies (Kok et al., 2017; Myriokefalitakis et al., 2018). However, field studies have shown that large particles can be transported far from their source (Betzer et al., 1988; Denjean et al., 2016b; Glaccon & Prospero, 1980; Prospero et al., 1970; Ryder et al., 2019; Weinzierl et al., 2009, 2017). Recently, Adebisi and Kok (2020) show the need to better characterize the properties and size distribution of supermicron particles because of their

importance to various climate properties. There is relatively little research that has quantified the role of large particles deposited to remote oceans, but recent work suggests that large particles are important (Jickells & Moore, 2015; van der Does et al., 2018; Wang et al., 2015).

African dust is transported to the western equatorial North Atlantic Ocean (NAO) during the boreal winter and spring (Barkley et al., 2019; Prospero et al., 1981, 2020). The equatorial NAO is a nitrogen (N)-limited system, and the atmospheric deposition of Fe stimulates N-fixers that relieve N-limitations, increasing primary productivity and leading to the sequestration of atmospheric carbon dioxide (Mahowald, 2011; Mills et al., 2004; Moore et al., 2013; Okin et al., 2011; van der Does et al., 2016). Dust is thought to be the primary source of Fe delivered to this ecosystem with much of this dust delivered as supermicron particles (Moran-Zuloaga et al., 2018), but other aerosol particle types have also been shown to be important for Fe deposition (Buck et al., 2010; Hamilton et al., 2020; Ito et al., 2019; Scholes & Andreae, 2000; Trapp et al., 2010). Dust has a residence time in the surface ocean on the order of a few weeks with larger particles having shorter residence times than smaller particles, suggesting larger particles may have a limited impact on marine productivity unless they contain forms of Fe that are readily soluble (Gaston, 2020; Jickells & Moore, 2015).

Dust transport during winter and spring occurs primarily in the Saharan Air Layer (SAL), an elevated hot, drier layer of air located at an altitude of approximately 1.5–3 km (Ansmann et al., 2009; Carlson & Prospero, 1972; Chiapello et al., 1995; Prospero et al., 2020; Tsamalis et al., 2013). During transport, dust can mix with clouds, sea-spray aerosol, and sulfur (S)-containing emissions in the marine boundary layer (MBL) (Fraund et al., 2017; Wu et al., 2019). Interactions between dust and trace acidic gases can induce chemical reactions that enhance Fe solubility (Ingall et al., 2018; Paris & Desboeufs, 2013; Paris et al., 2010; Spokes & Jickells, 1995). The size and location of Fe-bearing compounds on aerosols is important with smaller particles and surficial forms of Fe undergoing the most efficient dissolution and chemical reactions (Baker & Jickells, 2006; Gaston, 2020; Journet et al., 2008; McDaniel et al., 2019; Shi et al., 2009). Recent work has shown the presence of surficial Fe nanoparticles attached to larger dust particles that may be an important source of soluble Fe (Lafon et al., 2006; Moskowicz et al., 2016).

In this study, we present measurements of aerosol samples collected at Cayenne, French Guiana, a site located on the North Atlantic coast that is impacted by the long-range transport of African dust (Barkley et al., 2019; Prospero et al., 1981, 2020). Using scanning electron microscopy (SEM), we measured particle size and aspect ratios (ARs; length to width ratios), which is a measure of particle sphericity. Our results show the presence of other types of Fe-containing supermicron particles in addition to mineral dust. Surprisingly, many of these particles were determined to be supercoarse mode in size. Using energy dispersive X-ray spectroscopy (EDX) and elemental mapping, we determined the size and spatial location of Fe inclusions. From these results, we discuss the mechanisms of the transport of supermicron and supercoarse mode Fe-containing particles from Africa and their implications on marine biogeochemical cycles.

2. Materials and Methods

2.1. Sample Collection

Our sampling site is located north of Cayenne, French Guiana (4.92°N, 52.31°W) on a small peninsula at 67 masl. Samples were collected on 20 cm by 25 cm Whatman-41 cellulose filters with an average airflow rate of 0.75 m³ min⁻¹. Filters are exposed directly to the atmosphere under a Plexiglas “hat”; the cutoff diameter is estimated to be 80–100 μm or greater. We analyzed 13 samples collected between December 15, 2015 and March 31, 2016. Our analysis focuses on supermicron particles, which preferentially deposit near the surface of the filters. These supermicron particles are more easily identified by SEM than submicron particles, which tend to penetrate deeper within the filter prior to being captured. Future work will utilize electron microscopy grids to capture finer particles.

2.2. Air Mass Histories

Air mass back trajectories (AMBTs) were computed using the Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Rolph et al., 2017; Stein et al., 2015). AMBTs show that air masses originate from North Africa during the boreal winter and spring (Barkley et al., 2019). We also include three AMBT frequency plots shown in the supporting information (SI) Figures S1a–S1c initialized at 500, 1,000, and 2,000 masl, respectively. Each AMBT was run for 240 h at regular intervals of 12 h from December 15, 2015 to March 31, 2016. We emphasize that HYSPLIT cannot be used to determine an exact dust source location because we do not know at what point along the trajectory the dust was injected into the atmosphere and carried to altitudes significant for long-range transport.

2.3. Scanning Electron Microscopy (SEM) and Energy Dispersive X-ray Spectroscopy (EDX)

SEM-EDX analysis yields simultaneous information about the size, morphology, and chemical composition of aerosols. A 1 cm by 1 cm piece of cellulose filter was gently torn with gloved hands and placed on a carbon tab on an aluminum SEM stub for SEM-EDX analysis. The samples were coated with palladium (Pd) in a Cressington-108 Sputter Coater and imaged with a Phillips XL-30/ESEM-FEG at 20 kV and 1.6 nA (Figure 1). An EDX spectrum of a blank filter can be found in Figure S2. The EDX spectra shown in Figure 1 and EDX elemental maps (Figure 2) were obtained with a FEI Helios 650 Nanolab Dualbeam at 20 kV and 0.8–1.6 nA. EDX analysis was performed on a different microscope that was fitted with a light element detector. The corresponding larger field-of-view images corresponding to the spectra in Figure 1 are shown in the SI (Figure S3). Elemental maps provide spatial information regarding individual elements. These maps were analyzed with GENESIS software EDX version 5.10. Each particle analyzed by EDX was magnified to fill the width of the scanning display window (Falkovich et al., 2001).

2.4. Particle Sizing and Aspect Ratios (AR_{perp})

Particles were measured using ImageJ (<https://imagej.nih.gov/ij/>). The longest dimension was taken as the diameter, d_{max} . Of the 2,206 particles analyzed, 2,153 had a d_{max} greater or equal to 1.0 μm . d_{max} was then converted to projected area diameter, d_{pa} . Of the total particles, 2,102 particles had a d_{pa} greater or equal to 1.0 μm . Aspect ratios (ARs), which provide a measure of particle sphericity, were computed by dividing d_{max} by the width at the largest point that is perpendicular to d_{max} , which hereafter is called AR_{perp} (Huang et al., 2020). More details regarding these calculations can be found in the SI.

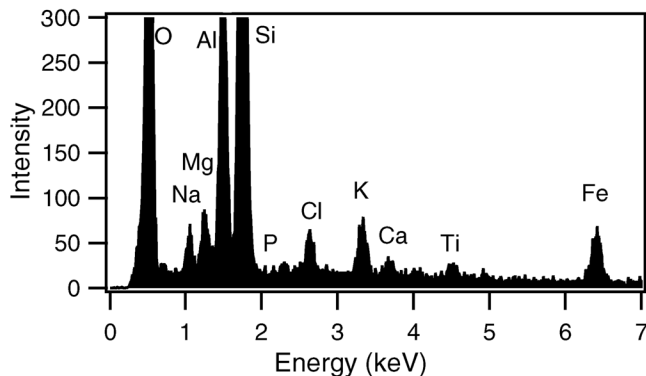
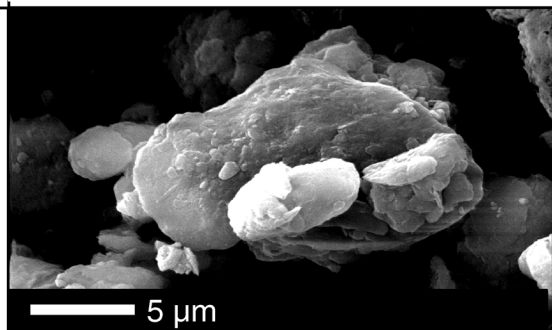
3. Results and Discussion

3.1. Transported Fe-Containing Supermicron Particle Types

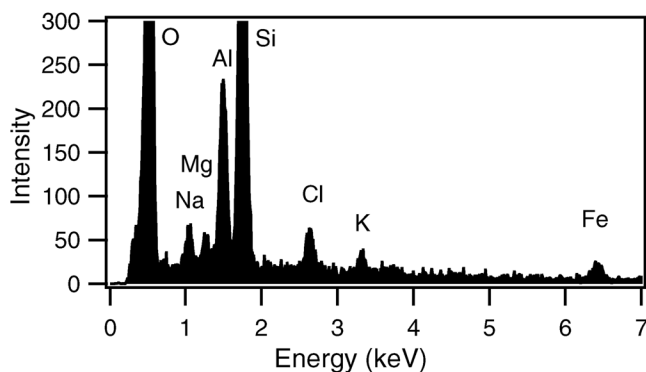
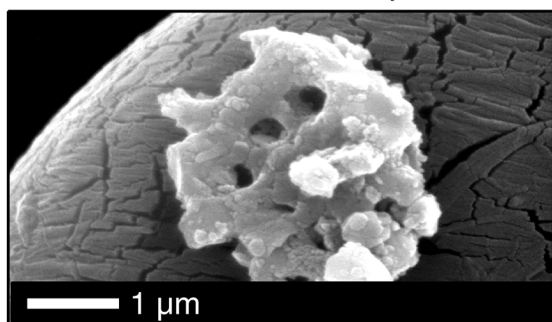
Three distinct particle types were observed: African dust (Figure 1a), freshwater diatoms (FDs) internally mixed with dust (Figure 1b and 1c), and primary biological aerosol particles (PBAPs) (Figure 1d). To the right of each representative SEM image is an EDX spectrum depicting the elemental composition of each particle type. Particles were categorized by morphology, size, and chemical composition and were similar to supermicron particles previously observed when African dust is transported to South America (Artaxo et al., 1988; Moran-Zuloaga et al., 2018; Pöschl et al., 2010; Rizzolo et al., 2017; Worobiec et al., 2007; Wu et al., 2019).

Dust comprised 95.5% of all particles analyzed by number (Figure 1a). These particles contained a large fraction of aluminum (Al) and silicon (Si), likely from aluminosilicate minerals, that were incorporated within every dust particle except quartz particles, which were composed entirely of Si and oxygen (O) and few in number (<1%). Additional elements detected include magnesium (Mg), which was found in 16% of particles. Calcium (Ca) was found in 29% of particles. Chlorine (Cl) was found in 11% of all particles and sodium (Na) was found in 20%. Cl and Na in dust particles is likely the result of sea-spray coagulation with dust during transport in the MBL. However, small amounts of halite are found in North African soils, which could contribute some Na and Cl to our samples (Scheuven et al., 2013). Previous work on particles

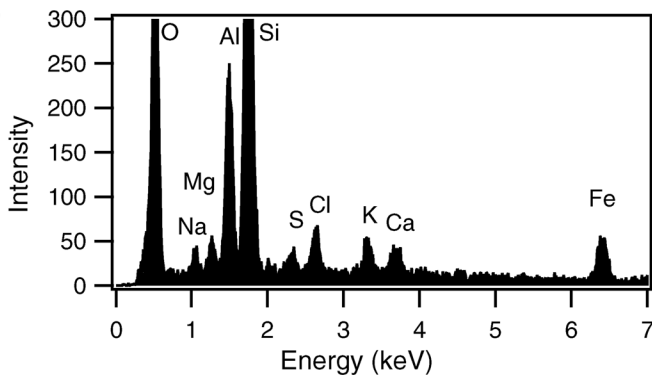
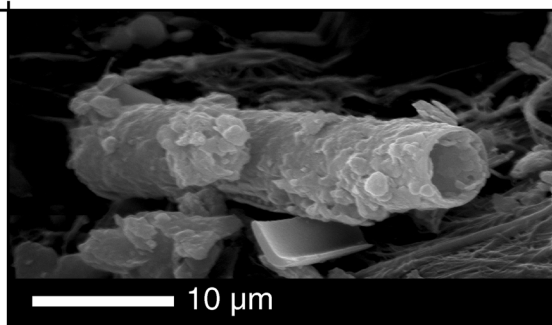
(a) Mineral dust



(b) FDs with dust: example 1



(c) FDs with dust: example 2



(d) PBAPs

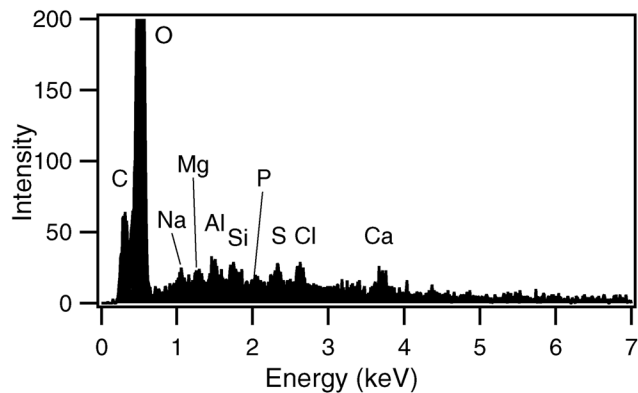
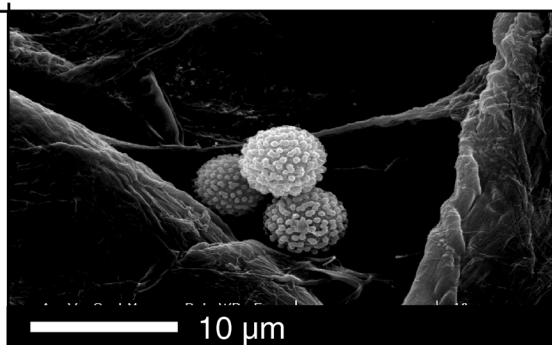


Figure 1. Representative scanning electron microscopy (SEM) images showing particle types (left) and energy dispersive X-ray spectroscopy (EDX) spectra (right) of each particle type: (a) mineral dust; (b) and (c) freshwater diatoms (FDs) internally mixed with dust; (d) primary biological aerosol particles (PBAPs). A cellulose filter fiber is shown behind the particle in (b). The field-of-view images corresponding to the particle spectra are shown in Figure S3.

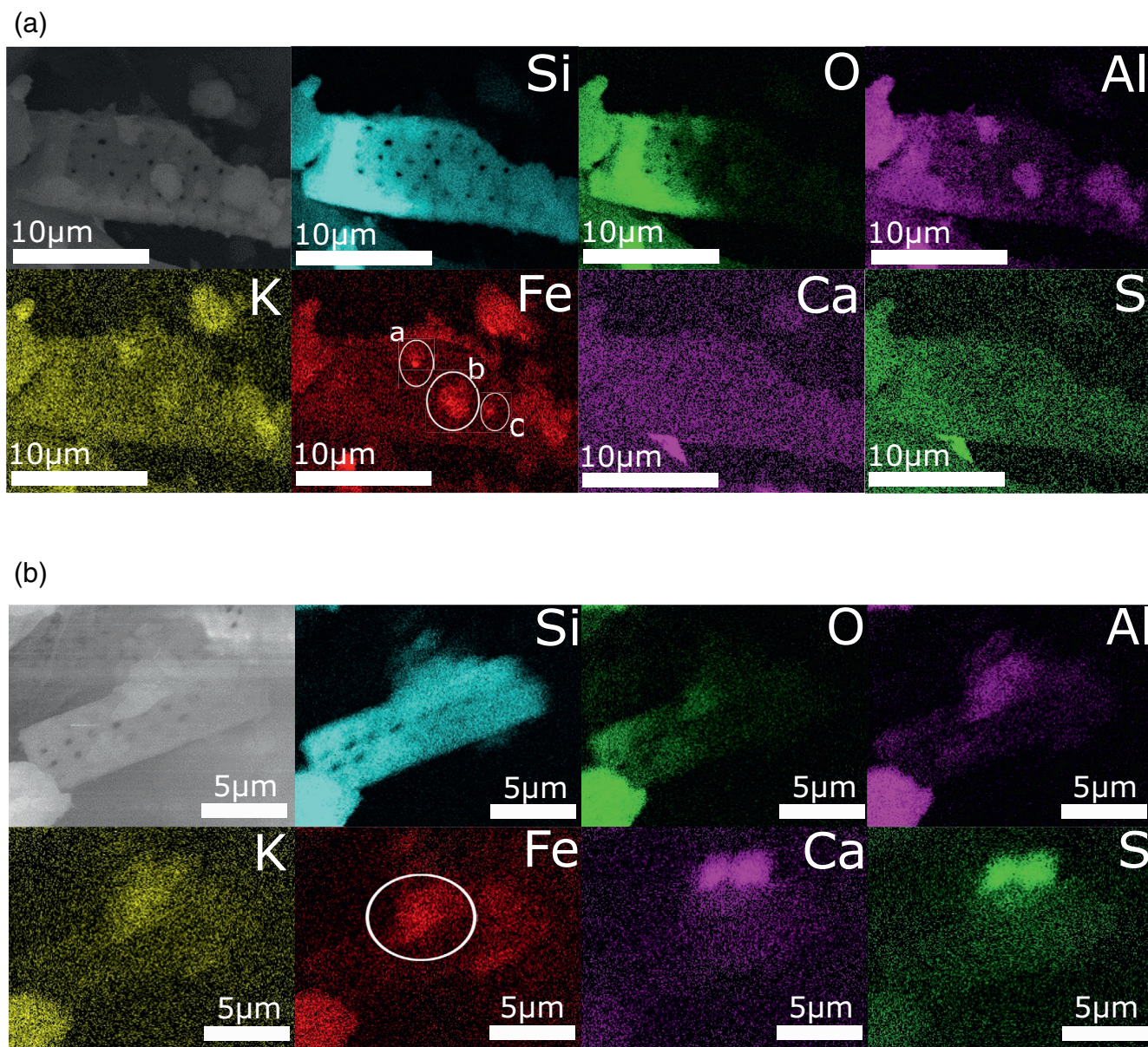


Figure 2. Panels (a and b) show two examples of scanning electron microscopy (SEM) images and corresponding energy dispersive X-ray spectroscopy (EDX) elemental maps of silicon (Si), oxygen (O), aluminum (Al), potassium (K), iron (Fe), calcium (Ca), and sulfur (S) on representative “FDs with dust.” Increased brightness indicates more of the element. The white circles emphasize the Fe-containing dust inclusions.

collected within the Amazon Basin has also shown similar internal mixtures of dust and sea salt (Adachi et al., 2020; Artaxo & Hansson, 1995; Formenti et al., 2003; Fraund et al., 2017; Worobiec et al., 2007).

In this study, we also include a new class of particles transported to South America: FDs derived from African paleolakes internally mixed with dust (hereafter “FDs with dust”), which were identified primarily by morphology. FDs with dust comprised 3.5% of all particles and were identified by their regular pattern of holes characteristic of their frustules or by their tubular shape (Figure 1b and 1c). Because these particles were internally mixed with mineral dust, their composition as detected by EDX was similar to that of mineral dust. We excluded a marine source for these diatoms for the following reasons: (i) greater than 95% of the FD particles were internally mixed with dust including dust inclusions on the interior of the diatom frustule that would not be expected if dust were simply deposited on the diatoms during sample collection (see Figure S4 and S5 for examples), (ii) FDs were not present during nondust transport days, and (iii) FD

species in our Cayenne samples match those collected in sediment traps in the Atlantic Ocean near the coast of Africa (Gasse et al., 1989; Romero et al., 2003), Barbados (Reid, 2003), and the Bodélé Depression (Bristow et al., 2009) (see Figure S6 for SEM images of diatomite from the Bodélé Depression). FD species in our samples include *Aulacoseira granulata*, which was the most common FD found in our samples, *Stephanodiscus rotula*, *Hantzschia amphioxys*, and the genus *Cylotella*.

The detected FDs in our samples are thought to originate from African paleolakes (Bakker et al., 2019; Ben-Ami et al., 2010; Prospero et al., 2002; Washington & Todd, 2005). Soil from paleolakes is comprised of diatomite, a mixture of low density FDs (0.8 g cm^{-3}) and higher density authigenic minerals (2.7 g cm^{-3}) (Bristow et al., 2010; Conrad & Lappartient, 1991). FDs have been collected in Caribbean aerosols (Reid, 2003) and in open-ocean sediment cores in the NAO, which have been used to assess changes in the aridity of the Sahara (Gasse et al., 1989; Pokras & Mix, 1987; Skonieczny et al., 2019). However, to our knowledge, our work is the first to show the transport of diatomite from African paleolakes to South America. Previous studies have suggested that multiple paleolakes within the Sahara including the Bodélé Depression and other paleolakes in the western Sahara are active dust sources that can transport mineral aerosols to South America (Bakker et al., 2019; Prospero et al., 2002). These paleolakes could explain the detection of FDs in our samples. However, a recent study by Yu et al. (2020) suggests that dust from the Bodélé Depression is not transported to South America as frequently as previously thought, and that FDs internally mixed with dust may originate from other African sources (Bozlaker et al., 2018; Jewell et al., 2020; Kumar et al., 2014; Pourmand et al., 2014). We are unable to definitively pinpoint sources in this work without further geochemical analysis.

The remaining particles (1%) were classified as long-range transported PBAPs (Figure 1d). While these particles were not found in great abundance, previous research has shown that PBAPs are important to the global aerosol budgets (Artaxo et al., 1998; Mahowald et al., 2008; Pöschl et al., 2010). They were either spherical or ellipsoidal, which distinguished them from dust particles. They were composed primarily of carbon (C) and O along with trace amounts of phosphorus (P), potassium (K), sulfur (S), Ca, Na, and Cl—a composition similar to previously measured PBAPs in the Amazon (China et al., 2018; Graham et al., 2003; Wu et al., 2019).

3.2. Elemental Mapping of Fe-Containing Particles and Their Mixing State

Fe was associated with all particle types except PBAPs (Figure 1). Of the particles characterized as dust, 85% of particles contained Fe, a result similar to a previous study (Falkovich et al., 2001). Dust particles contained an average of 12% Fe by weight while FDs with dust contained an average of 4% Fe by weight with Fe abundance scaling with the size and amount of dust inclusions. Figure S7 further highlights this point.

To characterize Fe-rich dust inclusions on FD particles, we performed elemental mapping, which shows the spatial distribution of individual elements based on the characteristic X-rays emitted by different elements. Figure 2 shows two examples of SEM images of FDs with dust in the top left panel and associated elemental maps in the other panels. Additional elemental maps are shown in Figure S8. As expected, the Si and O maps outline the silicate frustule structure of the FD. The map of Al in Figure 2a shows that the FD itself does not contain any Al; rather, dust inclusions associated with the FDs contain Al. Fe is also associated with the dust inclusions attached to the surface of the FDs similar to previous work that has shown surficial Fe oxides on larger, transported dust particles (Lafon et al., 2006; Moskowitz et al., 2016). Fe-containing dust inclusions on FDs were measured, and the longest axis is reported as their diameter. In Figure 2a, there are three Fe-rich dust inclusions labeled a–c; the Fe inclusion in (a) is $<0.1 \mu\text{m}$ in diameter; $2.6 \mu\text{m}$ in (b); and $1.5 \mu\text{m}$ in (c). Figure 2b shows a larger Fe inclusion with a diameter of $4.1 \mu\text{m}$. These Fe-rich inclusions are smaller than the FDs that they are internally mixed with and are mainly found on the particle surface, which increases their susceptibility to chemical reactions, photoreduction, ligand complexation, and dissolution that can enhance Fe solubility (Baker & Jickells, 2006; Cwierntny et al., 2008; McDaniel et al., 2019; Paris & Desboeufs, 2013; Shi et al., 2011; Zhu et al., 1997).

In addition to Si, O, Al, and Fe, we also performed elemental mapping for S, Ca, and K. As shown in Figure 2, we observed only small amounts of S on dust suggesting limited reactions of supermicron dust with

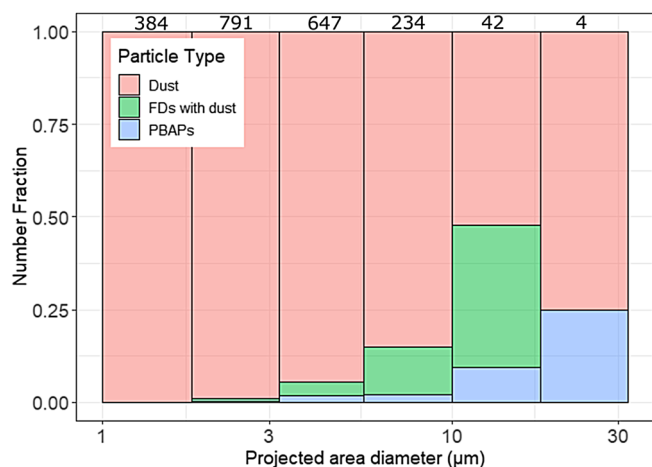


Figure 3. Relative fraction of each particle type: Dust (pink), Freshwater Diatoms (FDs) with dust (green), and Primary Biological Aerosol Particles (PBAPs; blue) as a function of projected area diameter (d_{pa}). The numbers in grey at the top show the number of particles in each size bin. Size bins are equally spaced on a log normal distribution with 4 bins per decade: 1.0–1.8, 1.8–3.2, 3.2–5.6, 5.6–10, 10–18, and 18–32 μm .

S-containing compounds typically found in the MBL, such as sulfuric acid or methanesulfonic acid. EDX analysis revealed that 16% of particles contained S, which is a lower fraction than observed in the central Amazon Basin (Wu et al., 2019). However, our observations are consistent with summertime dust measurements in marine and island environments (Denjean et al., 2016a; Kandler et al., 2018). In the central Amazon, there are more sources and emissions of S-containing species (e.g., SO_2 from pollution and biogenic sources) (Andreae et al., 1990; Wu et al., 2019). Therefore, we suggest that dust particles containing S-compounds observed in the central Amazon mainly underwent chemical reactions during transit from the coast to the interior of the Amazon, a distance on the order of 1,000 km. Elemental mapping results show that Ca and S are located in the same position on the dust particles (Figure 2a and 2b), indicating that S is associated with Ca, most likely in the form of gypsum ($\text{CaSO}_4 \cdot 2\text{H}_2\text{O}$). Gypsum could be a product of the chemical reaction of sulfur dioxide (SO_2) on the surface of calcite minerals (Andreae et al., 1986; Glaccum & Prospero, 1980; Ma et al., 2013) and reactions between calcite and sulfate from sea spray that occur in the atmosphere or subsequently after capture on the filter (Glaccum & Prospero, 1980). Other studies of long-range transported African dust using coupled SEM-EDX analysis also show gypsum from continental soil minerals (Coz et al., 2009; Falkovich et al., 2001), which could also explain our observations.

Elemental mapping also shows the presence of K on FDs and dust inclusions (Figure 2). From EDX spectra, K was found in about 62% of all mineral dust particles and was likely associated with K-feldspar minerals (Rizzolo et al., 2017; Scheuven et al., 2013). It is possible that some of the K is due to the co-transport of biomass burning (Ansmann et al., 2009). Coagulation of biomass burning and dust has been observed over Africa and has been suggested to enhance the bioavailability of Fe associated with dust (Paris & Desboeufs, 2013; Paris et al., 2010). We did not observe any visual evidence of the coagulation of dust with biomass burning, a result similar to Kandler et al. (2011); however, we suggest additional measurements to further probe the presence of internally mixed biomass burning and dust in transported particles.

3.3. Particle Size Distributions and Asphericity

The relative number fraction of each particle type as a function of size is shown in Figure 3. Raw counts as a function of d_{pa} are shown in Figure S9. Mineral dust dominates each size bin in the supermicron fraction and accounts for an average of 98% of particles, by number, in the three smallest size bins (1.0–5.6 μm), which is within the range of previous studies of long-range transported African dust measured in the Caribbean (Denjean et al., 2016b; Maring et al., 2003; Reid, 2003) and the Amazon (Artaxo et al., 1998; Moran-Zuloaga et al., 2018). FDs with dust and PBAPs comprise an increasing fraction by number at larger sizes, including supercoarse mode particles ($d_{pa} > 10 \mu\text{m}$). FDs with dust accounted for 38% of all supercoarse mode particles analyzed from 10 to 18 μm and the sizes of FDs with dust were significantly larger than dust (p value < 0.005). The d_{pa} of PBAPs ranged from 2.4 to 19 μm , which is similar to previous measurements of PBAPs in the Amazon (Graham et al., 2003; Worobiec et al., 2007). Overall, FDs and PBAPs comprised nearly 50% of supercoarse mode particles by number between 10 and 18 μm in diameter.

We also measured the aspect ratio (AR_{perp}) of particles transported to Cayenne (Figure 4 and Table S1) using a subset of particles. An increasing value of AR_{perp} indicates increased particle asphericity while a value of one indicates a spherical particle. Figure 4 shows that for all particle types, average AR_{perp} increases with increasing particle size, which is in agreement with other field studies of transported African dust (Kandler et al., 2011; Ryder et al., 2018; Saxby et al., 2018). The average AR_{perp} of all dust particles is 1.6, which is similar to previously reported values of AR_{perp} for boreal summertime long-range transported African dust (Huang et al., 2020). Interestingly, the average AR_{perp} of all FDs with dust was 2.5, significantly larger than

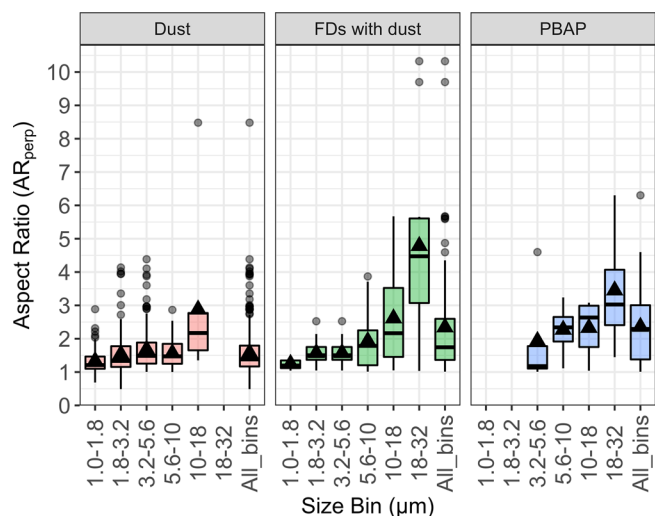


Figure 4. Aspect ratios (AR_{perp}) for dust (pink), FDs with dust (green), and PBAPs (blue). The center horizontal line of the box and whisker plot shows the median and the colored area shows the interquartile range. Whiskers show minimum and maximum values with grey dots representing outliers. The black triangles represent the mean of the data in each size bin. “All_bins” represents the AR across all size bins for each particle type. Gaps in the data represent either no particles or only one particle in the size bin. Note that the x axis is plotted as d_{max} .

pure dust particles (p value < 0.005), which is likely due to their tubular shapes (Figure 1c). Additionally, the median AR_{perp} of just the supercoarse mode FDs with dust was 3.7 with median values up to AR_{perp} of almost five clearly showing the increase in asphericity with particle size. PBAPs had an average AR_{perp} of 2.2. The elevated AR_{perp} and asphericity observed for all particle types that increase with particle diameter may explain the surprising long-range transport of supercoarse mode particles from Africa to the coast of French Guiana.

4. Conclusions and Atmospheric Implications

Our research highlights the diversity of Fe-containing particles transported to South America, including supercoarse mode particles, and serves as an important stepping-stone toward understanding the potential significance of these particles for marine biogeochemical cycles. Because most biogeochemistry models assume that supercoarse mode particles deposit close to their sources (Mahowald et al., 2014; Myriokefalitakis et al., 2018), our work sought to elucidate potential mechanisms that could explain large particle transport to our site. First, we found that particle asphericity (AR_{perp}) increases as a function of particle diameter, suggesting that particle shape reduces the settling velocity of large particles, particularly for unfragmented FDs whose tubular structure is still intact. The low density of FDs also enhances their atmospheric lifetime. Assuming FDs with dust represent a 1:1 mixture of FDs and dust, FDs with dust would have a 35% slower settling velocity when compared to similarly sized dust particles (Huang et al., 2020).

The same physical properties that decrease the atmospheric settling velocity for FDs with dust likely also decrease their sinking velocity in the water column. We estimate that FDs with dust have a 79% slower sinking velocity in surface water compared to dust based on particle shape, aspect ratio, and density (McDonnell & Buesseler, 2010). Details of these calculations are in the SI. A slower sinking velocity in the water column would enhance the residence time of FDs in surface waters compared to dust and increase the timescale for dissolution of the surficial Fe-rich inclusions that were observed on these particles. Additionally, a slower sinking velocity could increase the ability of FDs with dust to be carried by surface currents, increasing the spatial impact of FDs with dust.

In this study, we suggest that the transport of FDs with dust may facilitate delivery of Fe to remote marine ecosystems because of their low settling velocities in air and seawater. Even though FDs with dust accounted for 3.5% of the total particles analyzed by number fraction, they are consistently as large or larger than dust particles and comprise 38% of supercoarse mode particles by number between 10 and 18 μm , contain Fe-rich dust inclusions that extend Fe-containing dust into the supercoarse mode, and contain an average of 4% Fe by weight suggesting their potential impact on marine biogeochemical cycles. FDs found in open-ocean sediment cores in the NAO have also been linked to an increase in African aridity (Gasse et al., 1989; Pokras & Mix, 1987). Based on our results in present-day samples of FDs internally mixed with dust, FDs with dust could also have been a source of Fe to the equatorial NAO during previous climate regimes. We suggest additional studies that better quantify the contribution of FDs to the transported aerosol burden including both the wet and dry deposition rates of these particles to determine whether diatoms are indeed preferentially transported across the equatorial NAO during the dust transport season.

Data Availability Statement

Datasets in this study are available at the University of Miami’s Data Repository (under DOI <https://doi.org/10.17604/jmtk-yb12>).

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