

Supplementary Materials for

Bioaerosols in the Amazon rain forest: Temporal variations and vertical profiles of Eukarya, Bacteria, and Archaea

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S1 Supporting Text

S1.1 Measurement location: The Amazon Tall Tower Observatory

Aerosol sampling and measurements were conducted at the Amazon Tall Tower Observatory

(ATTO, 2°08.602' S, 59°00.033' W, 130 m.s.l.), a research site in the Amazonian tropical

5 rainforest, 150 km northeast of the city of Manaus, Brazil (Andreae et al., 2015). Two 80 m

towers, a tall tower of 325 m height and several containers at ground level are equipped with

various instruments monitoring e.g. greenhouse gases and aerosols. A detailed site

description and instrumental setup is presented in (Andreae et al., 2015). The Amazon rain

forest is characterized by a pronounced seasonality. During the wet season, from February to

10 May, conditions at this location can be described as near-pristine. At that time, winds are

coming from northeast. Thus, air masses are traveling across large areas of undisturbed terra

firme forest before reaching the site (Pöhlker et al., 2019). The map in Figure S1 displays the

location of the ATTO site and 3-day backward trajectories. The trajectories prove, that

typical wet season conditions prevailed during the FISH-sampling period. Air masses mainly

15 traveled from northeastern direction across untouched rain forest towards the site. Moreover,

the modelled precipitation rates show several strong rain events above the forest leading to

particle scavenging. Consequently, the air masses which were sampled at ATTO in February

and March 2018, are assumed to be unaffected by any anthropogenic activities. Moreover,

dust plumes that originate in Africa typically during the wet season and influence aerosol

20 mixtures and properties in the Amazon basin, did not reach the site at that time (Moran-

Zuloaga et al., 2018; Swap et al., 1992). Accordingly, during the wet season FISH sampling,

conditions are temporarily referred to as clean, green ocean conditions, characterized by

natural biosphere atmosphere exchange only. As visualized in the time series (Figure S2),

precipitation, relative humidity, and temperature during sampling conform to the mean values

25 calculated for the past 36 years. A medium strong La Niña period apparently did not have

major impact on climatic conditions in the ATTO foot print region.

S1.2 Bioaerosol emission in the Amazon

Bioaerosols are defined as liquid or solid airborne particles of biological origin, which were released directly from their sources into the air (Després et al., 2012). The Amazon rain forest 5 is thought to be the most species-rich freshwater and terrestrial ecosystem in the world. Consequently, there are manifold potential bioaerosol sources including water and soil surfaces, animals, plants, microbial surface communities, and decaying biomaterial. (Artaxo et al., 1990; Löbs et al., 2020; Fröhlich-Nowoisky et al., 2016). Their emission can be effected by means of active processes (e.g. fungal spore release based on osmotic pressure 10 changes; Pringle et al., 2005; Trail et al., 2005; Elbert et al., 2007) or passive mechanisms (e.g. air currents or rain splash; Joung et al., 2017; Jones and Harrison, 2004). Bioaerosol emission rates in the Amazon are therefore dependent on factors such as relative humidity and precipitation, air temperature and wind speed (Fröhlich-Nowoisky et al., 2016, and 15 references therein). Previous studies investigated bioaerosols in tropical (and boreal) rainforests by use of various instrumentations and analysis techniques. The obtained number concentrations vary according to the different detections methods and sampling location (Table S3; Šantl-Temkiv et al., 2019, Artaxo 2020, under revision).

S1.3 Bioaerosols' role in the Amazon rain forest

From an ecological point of view, bioaerosols play an essential role in the reproduction and 20 biogeographic distribution within the Amazonian ecosystem. Moreover, they are supposed to influence its hydrological cycle by acting as ice nuclei (IN) or giant cloud condensation nuclei (GCCN; Pöschl et al., 2010; Tobo et al., 2013; Artaxo, 2020, under revision). Still, the effect aerosols in general and bioaerosols in particular have on regional and global atmospheric processes is still under discussion and one of the major uncertainties in 25 understanding the climate system (Pöschl et al., 2010; Ariya et al., 2009). Previous studies reported bioaerosol concentrations to range between $\sim 10^4$ and $\sim 10^6 \text{ m}^{-3}$ accounting for the

majority of coarse mode aerosols in the Amazon under pristine conditions (Table S3; Graham et al., 2003; Huffman et al., 2012; Whitehead et al., 2016; Moran-Zuloaga et al., 2018). Exhibiting strong diurnal cycles, fungal spores were reported to be the most frequent bioaerosol type (Elbert et al., 2007; Souza et al., 2019). Bacteria in contrast, are found to be released in lower concentrations (in the understory; Souza et al., 2019). As IN particle abundance was suggested as limiting factor for cloud ice formation in the Amazon, bioaerosol number concentrations and especially their vertical diffusion are highly important to advance modelling studies (Pöschl et al., 2010). Thus, quantification and identification of bioaerosol abundances and cycling in the Amazon is key to shed light on their role in the pre-industrial biosphere-atmosphere interaction.

S1.4 Airborne DNA Mass

Using number concentrations obtained by FISH, airborne DNA mass could be calculated.

The factor 609.7 g mol^{-1} included in the equation is the average mass of a base pair in bound form:

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$$m_{abp} = \frac{(m_{dAMP} + m_{dCMP} + m_{dGMP} + m_{dTMP})}{2} - (2 \cdot m_{H_2O})$$
$$= \frac{(331.22 \frac{\text{g}}{\text{mol}} + 307.2 \frac{\text{g}}{\text{mol}} + 347.22 \frac{\text{g}}{\text{mol}} + 306.2 \frac{\text{g}}{\text{mol}})}{2} - 2 \cdot 18 \frac{\text{g}}{\text{mol}} = 609.7 \frac{\text{g}}{\text{mol}}$$

20 m_{abp} = average mass of base pair in bound form

At 5 m height, $11.8 \pm 4.63 \text{ ng m}^{-3}$ ($11.6 \pm 4.53 \text{ ng m}^{-3}$ eukaryotic and $0.29 \pm 0.09 \text{ ng m}^{-3}$ bacterial), at 60 m, $4.49 \pm 1.14 \text{ ng m}^{-3}$ ($4.17 \pm 0.99 \text{ ng m}^{-3}$ eukaryotic, $0.26 \pm 0.10 \text{ ng m}^{-3}$ bacterial), and at 325 m, $1.20 \pm 0.44 \text{ ng m}^{-3}$ ($1.07 \pm 0.38 \text{ ng m}^{-3}$ eukaryotic, and

0.12 ± 0.05 ng m⁻³ bacterial) DNA was found (Table S2). As explained before, archaeal numbers suffer from insufficient statistics and so do the respective calculated DNA masses. As a result, the standard deviation exceeds the mean archaeal number concentration. As described in the article, these numbers are similar to DNA concentration found in other 5 forested ecosystems. Nevertheless, Helin et al. (2017) found also samples with up to 48 ng m⁻³, more than 4 times higher than measured in the Amazon during this project. The reason behind this is twofold: The boreal forest, which Helin et al. (2017) analyzed, mainly consists of pines and other vascular plants. As a result, bioaerosols are expected to include pollen with comparably big genome sizes. Furthermore, these pollen are emitted during a 10 fairly short time period, specifically in spring, causing a temporal peak in bioaerosol load and consequently, DNA concentration. When averaging DNA measurements in the boreal forest, Helin et al. (2017) found 8.60 ± 11.41 ng m⁻³, which is well comparable to the DNA concentrations we calculated for the Amazon forest.

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S2 Supplementary Figures and Tables

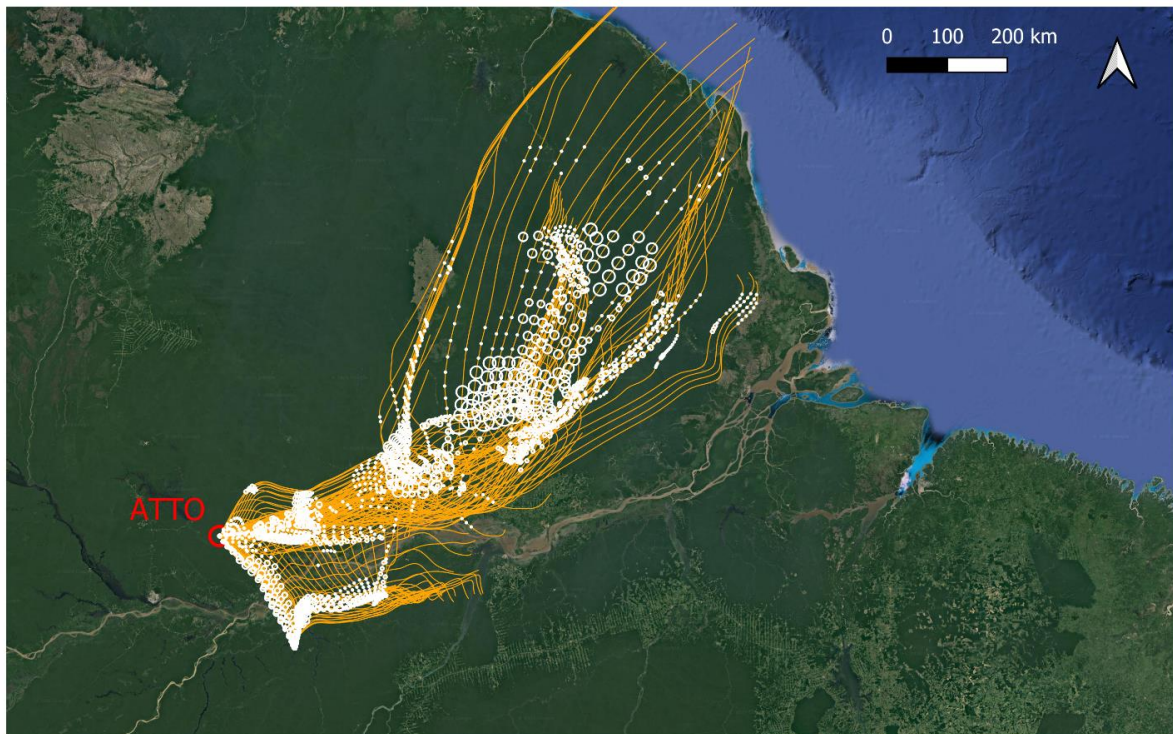


Figure S1. Map showing location of the ATTO site with ensemble of 3-day backward trajectories (yellow lines) using the Hybrid Single-Particle Lagrangian Integrated Trajectory model (HYSPLIT, NOAA-ARL) with meteorological input data from the Global Data Assimilation System (GDAS1, 1° resolution).

Trajectories were started at 200 m above ground every hour during the sampling period of this study. White circles represent precipitation obtained through GDAS (HYSPLIT model) for every hourly data point of the individual trajectories. The size of white circles shows the extent of precipitation en route (ranging from 0 to ~11 mm). The map shows that the trajectories during the sampling period mostly moved over the relatively untouched rain forest areas north of the Amazon River with some direct influence from the Amazon River valley itself. Precipitation along the trajectories shows that the transported air masses experienced relatively strong rain-related scavenging.

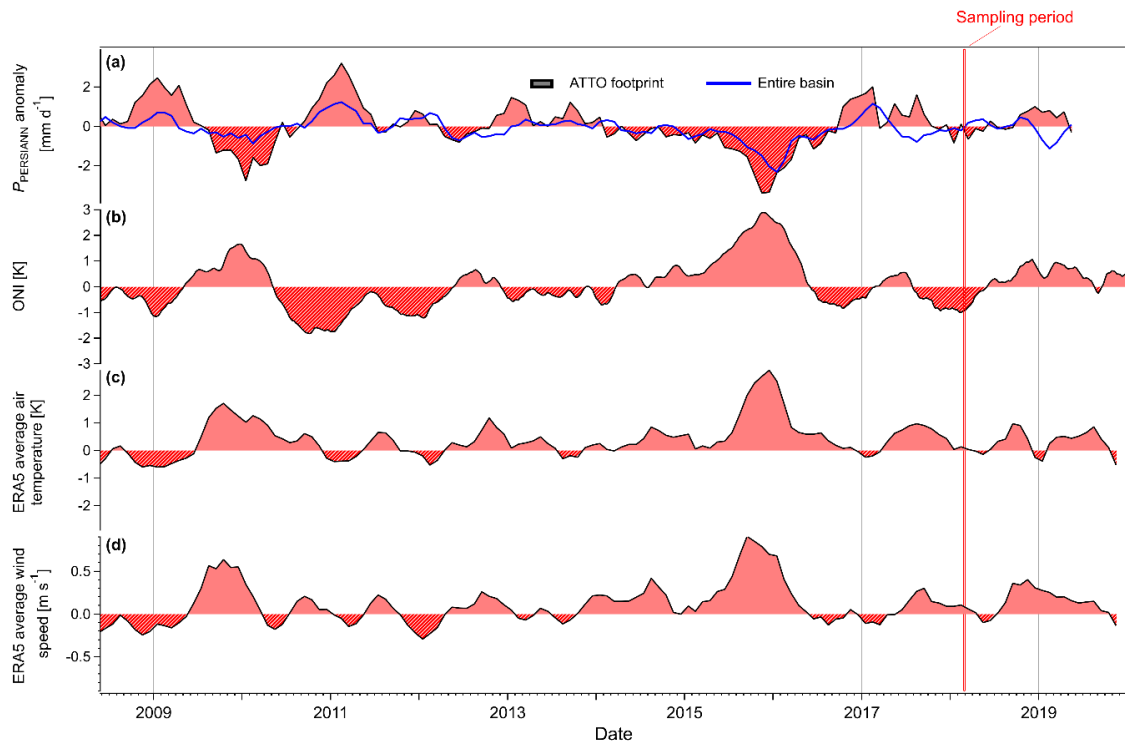


Figure S2. Time series of anomalies in precipitation, air temperature, and wind speed including the sampling period to characterize the overall atmospheric conditions. The analysis was conducted in close relation to the study by Pöhlker et al. (2019), where detailed information can be found. (a)

- 5 Anomaly in precipitation rate P (from the Precipitation Estimation from Remotely Sensed Information using the Artificial Neural Networks for Climate Data Record, PERSIANN-CDR, data product, reference time frame 1983-01-01 to 2019-12-31) for two regions: (i) the core region of the ATTO footprint (i.e., contour line with largest 0.5 % of air mass residence times) and (ii) for the entire Amazon watershed region (blue line). (b) Oceanic Niño index (ONI) for comparison with precipitation
10 variability and indicating El Niño vs. La Niña periods (i.e., El Niño influence is very strong for $ONI > 2.0$, strong for $2.0 > ONI > 1.5$, medium for $1.5 > ONI > 1.0$, and weak for $ONI > 1.0$. La Niña influence is strong for $-2.0 < ONI < -1.5$, medium for $-1.5 < ONI < -1.0$, and weak for $ONI > -1.0$). (c) Anomaly in daily averaged air temperature (2 m height) from ERA5 reanalysis product (reference time frame 1979-01-02 to 2019-12-31). (d) Anomaly in daily averaged wind speed (10 m height) from ERA5 reanalysis product (reference time frame 1979-01-02 to 2019-12-31).
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Table S1: Particle number concentrations obtained by FISH, relative fractions and raw counts (number of fluorescent signals identified under the microscope) for bioaerosols hybridized with a probe or stained with DAPI. Presented are numbers from samples collected at three heights at 5 or 6 consecutive days during approximately 23 h sampling time.

height	day	Archaea [·10 ⁴ m ⁻³]	A/DAPI [%]	particle counts	Bacteria [·10 ⁴ m ⁻³]	B/DAPI [%]	particle counts	Eukarya [·10 ⁴ m ⁻³]	E/DAPI [%]	particle counts	DAPI [·10 ⁴ m ⁻³]	particle counts	probe/ DAPI
5 m	1	0.35 ± 0.52	0.7	7 / 987	2.5 ± 1.9	4.6	50 / 999	49 ± 33	91	1041 / 1284	54 ± 14	1864	96
	2	0.31 ± 0.44	0.6	9 / 1272	7.5 ± 2.1	15	236 / 1142	25 ± 2.9	51	1151 / 1628	50 ± 29	4395	67
	3	0.19 ± 0.32	0.3	6 / 1644	14.3 ± 2.3	24	455 / 1660	36 ± 2.7	61	1341 / 1914	59 ± 15	1971	85
	4	0.22 ± 0.37	0.4	7 / 1510	5.6 ± 1.7	10	168 / 1539	41 ± 18	75	1039 / 1341	54 ± 19	5663	86
	5	0.15 ± 0.24	0.3	3 / 979	5.3 ± 2.7	11	196 / 2076	40 ± 19	82	1403 / 1687	48 ± 29	6522	94
	mean (1-5)	0.25 ± 0.38	0.5		7.0 ± 2.1	13		38 ± 15	72		53 ± 21		86
60 m	1	1.02 ± 1.08	4.0	39 / 867	2.4 ± 2.2	9.3	67 / 714	13 ± 2.9	49	688 / 1031	26 ± 15	1968	62
	2	0.38 ± 0.48	2.4	32 / 1032	2.3 ± 1.0	15	157 / 986	10 ± 2.1	62	653 / 920	16 ± 8.2	1788	79
	3	0.52 ± 0.76	2.1	25 / 1233	5.4 ± 1.9	22	321 / 1328	13 ± 2.6	53	578 / 987	24 ± 7.7	1926	77
	4	2.30 ± 2.09	11	112 / 1069	7.7 ± 2.2	38	330 / 902	11 ± 3.5	56	434 / 815	20 ± 8.0	2290	105
	5	3.14 ± 2.43	12	120 / 1031	6.3 ± 3.2	24	352 / 1430	18 ± 2.7	66	995 / 1384	27 ± 9.4	1598	101
	6	0.21 ± 0.43	0.5	4 / 751	14.9 ± 4.8	39	347 / 890	18 ± 5.7	48	497 / 1032	38 ± 14	958	87
	mean (1-6)	1.26 ± 1.21	5.3		6.5 ± 2.5	24		14 ± 3.3	56		25 ± 10		85
325 m	1	0.11 ± 0.26	1.5	6 / 361	0.4 ± 0.6	6.1	24 / 329	3.3 ± 0.8	47	126 / 267	7.0 ± 2.2	672	54
	2	0.02 ± 0.07	0.4	2 / 649	2.6 ± 1.2	54	243 / 459	1.4 ± 0.7	29	87 / 284	4.8 ± 2.0	638	83
	3	0.09 ± 0.24	0.9	5 / 630	4.1 ± 1.6	39	255 / 674	4.6 ± 1.6	44	198 / 427	10 ± 3.0	688	85
	4	0.09 ± 0.25	0.4	5 / 1018	6.1 ± 2.2	25	405 / 1317	5.1 ± 1.9	21	188 / 716	24 ± 12	2503	46
	5	0.21 ± 0.24	1.6	11 / 667	1.7 ± 1.1	13	96 / 785	3.3 ± 1.3	25	158 / 651	13 ± 3.7	1058	39
	mean (1-5)	0.10 ± 0.21	0.9		3.0 ± 1.3	27		3.5 ± 1.2	33		12 ± 4.6		61

Table S2: Diel airborne DNA mass concentration calculated for bioaerosol classes at different heights. Mean bioaerosol number concentrations obtained by FISH were multiplied with the calculated mean DNA mass per cell.

	Archaea mean ± sdev [ng m ⁻³]	Bacteria mean ± sdev [ng m ⁻³]	Eukarya mean ± sdev [ng m ⁻³]
sampling height			
5 m	0.01 ± 0.02	0.29 ± 0.09	11.6 ± 4.53
60 m	0.05 ± 0.05	0.26 ± 0.10	4.17 ± 0.99
325 m	0.00 ± 0.01	0.12 ± 0.05	1.07 ± 0.38

Table S3: Overview table of bioaerosol number and mass concentrations found in tropical and boreal forest systems.

Table S4: Chemicals and producing companies used for filter sample fixation, hybridization and microscopic visualization.

Chemical	product number	company
Acchromopeptidase	A3547-500KU	Sigma-Aldrich
Agarose LE	840001	Biozym
Blocking reagent	10447200	Roche
Citifluor AF1	17970-100	Citifluor
Diamidino-2-phenylindol-2HCl	18860.02	Serva
Ethanol	K928.5	Roth
Ethylendiaminetetraacetic acid	ED-100g	Sigma-Aldrich
Formaldehyde solution 37%	F8775-500mL	Sigma-Aldrich
Formamide	47671-1L-F	Roth
Lysozyme from chicken egg white	62970-5g-F	Sigma-Aldrich
Phosphate buffered saline 10x	79383-1L	Sigma-Aldrich
Sodium chloride	S7653-250g	Sigma-Aldrich
Sodium dodecyl sulfate	L3771-25g	Sigma-Aldrich
Tris HCl	15568-025	Invitrogen

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