Determining the hierarchical order by which the variables of sampling period, dust outbreaks occurrence, and sampling location, can shape the airborne bacterial communities in the Mediterranean basin

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Abstract. An NGS-based taxonomic analysis was carried out on airborne bacteria sampled at ground level in two periods (May and September) and two opposite localities on the North-South axis of the Sardinia Island. Located in a central position of the Mediterranean basin, Sardinia constitutes a suitable outpost to reveal possible immigration of bacterial taxa during transcontinental particle discharge between Africa and Europe. With the aim of verifying relative effects of dust outbreaks, sampling period and sampling site, on the airborne bacterial community composition, we compared air collected during dust-carrying meteorological events to that coming from wind regimes not associated to long-distance particle lifting. Results indicated that: (a) higher microbial diversity and richness (118 orders vs 65) and increased community evenness were observed in the campaign carried out in September in comparison to the one in May, irrespective of the place of collection and of the presence or absence of dust outbreaks. (b) During the period of standard wind regimes without transcontinental outbreaks a synchronous concerted turnover of bacterial communities across distant locations of the same island, accompanied as mentioned by a parallel rise in bacterial diversity and community evenness appears to have occurred. (c) changes in wind provenance could transiently change community composition in the locality placed on the coast facing the incoming wind, but not in the one located at the opposite side of the island; for this reason the community changes brought from dust outbreaks of African origin are observed only in the sampling station exposed to south; (d) the same winds, once proceeding over land appear to uplift bacteria belonging to a common core already present over the region, which dilute or replace those that were associated with the air coming from the sea or conveyed by the dust particulate, explaining the two prior points. (e) the hierarchy of the variables tested

in determining bacterial assemblages composition results: sampling period >> ongoing meteorological events > sampling location within the island.

40 1 Introduction

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With a total volume evaluated of 4.5x10¹⁸ m³, terrestrial lower atmosphere represents the most extended potential biome, followed by water, 1.3x10¹⁸ m³ (Gleick, 1993), and by soil with 6.2x10¹⁶ m³ (estimated on the basis of the deeper subsurface living bacteria currently described Szewzyk *et al.*, 1993). Concerning atmosphere, microbial cells and propagules, embody a particularly suitable conformation to take advantage of air utilization as an environment for survival and dispersal. Their movement can be favored by a natural mobile reservoir of physical solid carriers represented by the air-dispersed particulate matter. Such particles range between 0.2 and 100 um in size (Bernstein *et al.* 2004; Ryder et al., 2018) and average loads of 1-100 ug m⁻³ (Williams *et al.* 2002, Van Dingenen *et al.* 2004). It has been estimated that more than 5000 Tg of sea salt (Tegen *et al.* 1997) and 1000-2000 Tg of soil particles, passively uplifting and transporting live cells are released every year in the atmosphere giving rise to a widely heterogeneous material conveyed from different sources (Guang *et al.* 2009; Mc Tainsh 1989, Knippertz *et al.* 2009).

The tropical African and Asiatic belts, represent two amongst the major airlift dust sources (Prospero *et al.* 2002, Schepansky *et al.* 2007). Several studies underline that this phenomenon strongly contributes to a cosmopolitan microbial distribution (Favet *et al.* 2013, Griffin 2008, Yang *et. al.* 2008, Wainwright *et al.* 2003, Smith *et al.* 2010).

Moreover, the correlation between specific bacterial clades and particle size (Polymenakou *et al.* 2008) opened new hypotheses on differential dispersal of taxa in relation to the dust features. High amount of bacterial 'newcomers' have been pointed out in air samples collected in occasions of foreign dust outbreaks (Maki *et al.* 2014, Rosselli et al., 2015). Immigrant microorganisms classification (Sanchez de la Campa *et al.* 2013) and their effects on an autochthonous ecosystem have also been reported (Peter *et al.* 2014, Shine *et al.* 2000). Evidences of a correlation between aerosol-related biodiversity and seasons (Gandolfi *et al.* 2015) underlines the natural complexity related to this process, suggesting that effects may vary also depending on climatic periodicity. The impact of desert dust in transiently changing the airborne microbiota in cities has been described (Mazar *et al.*, 2016). In other works, the origin of dust was compared and storms from different origins exhibited distinct bacterial communities, with signature bacterial taxa (Gat *et al.*, 2017). Marked seasonal patterns in airborne microbiota have also been reported in long term studies (Cáliz *et al.*, 2018). Fungal taxa were also analyzed along with their and relationships with the chemical composition of the particulate (Lang-Yona *et al.*, 2020). The genes whose presence has been suggested to enable bacterial survival in the atmosphere, referred to as aeolian lifestyle, have been studied by metagenomics approaches and include UV-induced DNA damage repair, cell aerosolization,

Europe-Mediterranean air circulation routes offer an interesting case study when focusing on airborne bacteria. The system can be represented as a multidirectional network in which biological components and weather conditions are closely related (Lelived *et al.* 2002).

Extending for more than 30 degrees of latitude above the subtropical belt, Europe is crossed by middle-latitude and equatorial atmospheric systems. Mathematical models suggest that a considerable part of the air mass

aerotaxis, and thermal resistance (Aalismail et al., 2019).

movements has a Northern, Atlantic source in response to the pressure generated by the Azores high (Littmann, 2000). Southern winds from Africa, prone to carry desert sand, and potentially microbes, can be determined by specific climate conditions (Kostopoulou and Jones 2007, Benkhalifa et al, 2019). It has been estimated that, as a consequence, 80-120 Tg of dust per year are transported across the Mediterranean towards Europe (d'Almeida 1986; Dulac *et al.* 1996), reaching the higher troposphere layers (Alpert *et al.* 2004) and spilling over up to the far-Northern sides of the continent (Franzèn *et al.* 1991).

In order to track the biodiversity of these airways, the Italian island of Sardinia was chosen as ideal observatory point to collect airborne bacteria moving inside and outside Europe. Located in the middle of the Mediterranean Sea, this landmass is separated from Italy, France, Spain and Africa coastal baselines by distances of 120, 150, 230, and 100 nautical miles (NM) respectively (Fig. 1). Its geographical position facilitates the displacement of western high- and low-pressure air masses coming from Gibraltar and becoming the first and the last frontier for microbes entering or leaving Europe, respectively. In a prior study (Rosselli et al. 2015), we described a core microbiome in the bacteria cast upon the Sardinia island under different wind regimes through analyses of DNA from deposited particles. The analysis compared the trans-Mediterranean airflow with that of winds from Europe, and pinpointed a number of taxa which have records in clinical infections. In that investigation the sampling dates were all concentrated in a single period of six days (in February) and some variations of the airborne biota were observed in response to the opposite wind directions.



Fig. 1 Mediterranean area with Sardinia Island detail and sampling locations Sassari and Cagliari.

However, the most remarkable evidence was a prevailing constancy of the microbial composition in spite of the changing winds provenances. In the present study instead we analyzed a series of events featuring a starting dust outbreak, a 109 days-long period devoid of dust-carrying winds, and a second dust outbreak. The analyses were performed in two oppositely located stations: Cagliari, on the South-East side of Sardinia, facing the African side, and Sassari in the North-West, i.e. farthest from the dust-carrying winds. The sampled particulate was analyzed by NGS sequencing of the amplified 16S rRNA genes. The main goal of the project was to verify in which hierarchical order the different variables of (a) sampling period, (b) occurrence of dust-carrying outbreaks, and (c) sampling location, could act in determining airborne bacterial communities composition.

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2. Materials and Methods

2.1 Meteorological monitoring

Surveillance of the weather trends and conditions to anticipate dust outbreaks from Africa towards Sardinia and winds of interest was performed by routine checking of the MODIS satellite data and Meteosat imagery combined with the SKIRON forecasting model (Nickovic *et al.* 2001). Europe daily synoptic conditions were analyzed on the weather charts available from the www.eurometeo.com and www.metoffice.gov.uk websites. The origin and the trajectory of the dust carried by winds towards Italy were inferred by the NOAA HYSPLIT model (Hybrid Single Particle Lagrangian Integrated Trajectory Model) (Draxler *et al.* 2014; Rolph 2014).

Monitoring was aimed at predicting two distinct conditions: i) North-African high-pressure nuclei favoring Southern winds suitable to carry and deposit dust over Sardinia (dust-enriched events); and ii) North-European high-pressure nuclei, determining northern winds referred to as 'Controls' (dust-negative events).

In addition, PM10 concentration (particulate matter with a diameter of less than 10 µm) and meteorological data registered by the ARPAS (Regional Environmental Protection Agency of Sardinia) monitoring stations were taken into consideration in relation to the arrival of African air masses.

Information about wind direction and intensity (every 10 minutes), temperature and humidity (once per hour) were downloaded by the ISPRA website (http://www.mareografico.it/) and two sampling stations located in Cagliari (39.21°N, 9.11°E) and Sassari - Porto Torres (40.84°N, 8.40°E). Data covered a 7 months time-lapse, from March to September 2014, in order to obtain a nearly annual view to focus within the main weather instability period.

2.2 Sampling

The experimental design involved: 2 sampling sites at the opposite corners of the Sardinia island (Sassari vs. Cagliari), 2 sampling periods (May vs. September) 2 meteorological conditions (absence vs. presence of a dust outbreak). In each of these situations, two replicate samples were taken and processed independently throughout the DNA extraction step to be pooled before sequencing. Samples were collected on Teflon filters (Sartorius Stedim Biotech) by using a Skypost Tecora apparatus (compliant to the European legislation 96/62/gmeCE) processing 39 liters of air per minute. To constitute 'a sample' a continuous 24h-long air intake through the filters was performed.

In the case of the dust outbreaks the 24h sampling was further divided in two periods, by considering independently the first 12 hours and the second 12 hours. The number of resulting samples was therefore 12; namely the module of three conditions: (a) Control; (b) Dust h 1-12, (c) Dust h 12-24; multiplied by 2 sampling

periods and by the 2 sampling places, resulting in 3 x 2 x 2= 12 samples.

As technical note, the scope and meaning of 'controls' here was that of samples that could allow to individuate the shift between one condition and its adjacent one. In our cases, catching the sudden change of wind regime by sampling immediately before or after a dust outbreak. Therefore, the controls were thus defined as single 24h time points flanking the key dust events.

For each sample, date and atmospheric conditions are reported and fully described in the Results chapter and Supplementary Materials.

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2.3 DNA extraction and Sequencing

accession numbers ERX836645-56.

DNA was extracted using the E.Z.N.A.® Soil DNA Kit (Omega Bio-Tek Inc.) as described by the manufacturer. Quality and quantity of the extracted nucleic acid were measured using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific Inc.).

Amplification of the 16S-rRNA genes for sequencing was performed using the universal primers 27F-1492R (AGAGTTTGATYMTGGCTCAG and TACGGYTACCTTGTTACGACTT, respectively). PCR was carried out using Platinum® Taq High Fidelity DNA Polymerase (Life Technologies) in a PTC-200 Thermal Cycler (MJ Research Inc.) set as follows: 95°C for 5 min, (95°C for 0.5 min, 51°C for 0.5 min, 72°C for 2 min for 30 cycles), 72°C for 10 min and 4°C on hold. The amplification of the No Template Control (NTC) was negative. Next generation sequencing was carried out at the facilities of the Porto Conte Ricerche Srl (Alghero, Italy). Briefly, amplicons were quality-checked on an agarose gel and purified using the Agencourt® Ampure® XP PCR Purification Kit. One ng of DNA was processed using the Nextera XT DNA Sample Preparation Kit (Illumina Inc.) and sequenced using the HiScanSQ (Illumina Inc.) with 93bp x 2 paired-end reads. Sequences were submitted to the European Nucleotide Archive(ENA) inside the "Dust Metagenome" BioProject with the

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Reads were cleaned on the basis of quality and fragments of Nextera adapters removed by

Trimmomatic (Bolger et al. 2014) set at the value of 3 for leading and trailing trimming, and bases lower than 20 on a 4-base wide sliding window. Quality was confirmed by **FastQC** (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and reads were analyzed with Qiime1.9.0 (Caporaso et al. 2010). The OTU table was created using the pick_otus script with the Closed-reference OTU picking strategy with the Greengenes reference OTUs database clustered at 97% (ver. gg_13_8). The same script checked against chimeric sequences using the Broad Microbiome Utilities' 16S Gold reference database (version microbiomeutilr20110519). The OTU table was filtered based on the total observation count of an OTU at least of 3 and low abundance filtering of 0.005%. Finally, the OTU table was rarefied (subsampled) at 1109571 counts (equal to the sample with a lowest depth) for all the samples. Perl and the R-package Vegan were subsequently used for cladograms and distance-based clustering analyses, Ggplot, Plotrix and WindRose provided graphic support. Molecular data regarding bacterial species compositional differences across different treatments were analysed by multivariate analyses (Principal Coordinate Analysis, PCoA; Principal Component Analysis, PCA, Discriminant Analysis of Principal Components DAPC), and ecological indices calculation, using the Calypso online software tool (Zakrzewski et al., 2017). Prior to the analyses, the relative abundances of OTUs were equalized by applying the total sum of squares scaling (TSS) normalization followed by square root transformation.

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3. RESULTS

3.1 Meteorological events

To capture the air microbiota of Sardinia and to put in evidence taxa, which could be associated to specific events,

weather forecasts and other data on air circulation were regularly browsed to select suitable dates for the sampling.

Such events were intended as winds in northbound direction prone to carry dust from shores across the Mediterranean as opposed to calm air or slow flows from the opposite quadrant. This allowed to integrate boundary conditions and environmental variables to assess possible correlations between these and microbial community fluctuations (Fiamma, 2016).

Dust-carrying air masses moved over the Mediterranean on May 21st through May 22nd 2014 towards north-east, covering the entire Sardinia island. PM10 concentrations throughout the second half of May at both North and South collection sites displayed increases in correspondence of the dust event (Supplementary Figure S1). Incoming dust of African origin was equally evidenced by charts reporting wind fronts and pressure (Supplementary Figure S2), and images from satellite (Supplementary Figure S3). The itinerary of particles was reconstructed by plotting 3-day backward trajectories of the air mass using a NOAA HYSPLIT model (Figure 2) which tracked the North-African zone as the source of the convective motion responsible for the dust discharge on Italy observed on May 21st -22nd.

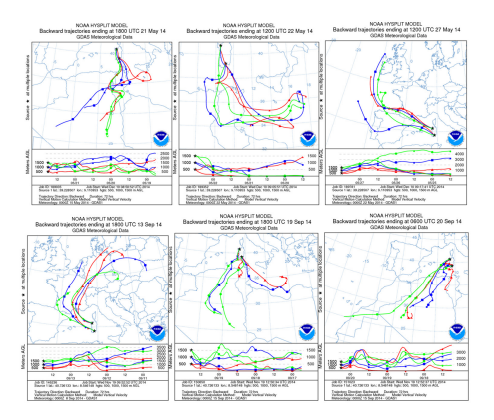


Fig. 2. Upper panel: 3-Day air mass backward trajectories calculated by the NOAA HYSPLIT model ending at 18:00 UTC May 21st, 12:00 UTC May 22nd and 12:00 UTC May 27th 2014 at both sampling sites. **Lower panel**: 3-Day air mass backward trajectories calculated as above, ending at 18:00 UTC September 13th, 18:00 UTC September 19th and 06:00 UTC September 20th 2014 at both sampling sites (credit to: ready.arl.noaa.gov/HYSPLIT.php).

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May 27th was selected as "clear day" featuring a weather not conducive anymore for air convection from Africa to Sardinia. Such conditions consisted in overall European low pressures as opposed to high pressures over Mauritania, Mali, Libya and Algeria (Supplementary Figures S2 and S3). Particle back-tracking supported the evidence of a slow flow of air masses only from north-western corners on May 27th (Fig.2, upper panel, rightmost image).

An outbreak of dust on Sardinia was recorded again in 2014 during the second fortnight of September. Low pressure from the north-western coast of Spain to Morocco was opposed to parallel high-pressure system that extended over North Africa (Libya, Algeria, and Tunisia) through Sicily. This circumstance caused the flow of dust-carrying air masses over the Mediterranean basin, reaching in particular Southern Italy and Sardinia. Air movement from the African continent made air temperature rise to values above the usual September means, with a peak on Sep. 20th in Sassari (northern sampling site) and on Sep. 21st in Cagliari (southern sampling site) (Supplementary Figure S4-a,b). In relation to this condition, from September 19th through the 21st a dust outbreak from Sahara flew over the Mediterranean and entirely covered Sardinia. The relative wind fronts and pressure

values are shown in Supplementary Fig. S5. Patterns of PM10 from daily records taken at both Sardinian sampling stations, also displayed a rise during the dust outbreak (Supplementary Fig. S4-c). Satellite imagery confirmed again the occurrence of incoming dust-loaded air masses from Northern Africa (Supplementary Fig. S6) consistent with their 3-day back-trajectories (Fig. 2 lower panel). Those confirmed that on September 19th - 20th air flows were from North African origin. About a week earlier instead, September 13th had featured low pressures on Italy while high pressures were recorded over the southern part of Morocco, Algeria and Mauritania. This picture was not permissive for any transport of air loads from Africa to Sardinia and the day was therefore considered as the "clear day" reference of the period. Air representative of the dust outbreak condition was thence sampled from Sep.19th through 20th, while the corresponding control air was collected on September 13th.

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3.2 Bacterial community composition

A synoptic view of the results at phylum/class level is shown in Fig. 3. Details on orders are shown in Table 1.

Complete data are available in the Supplementary Datasheet S1

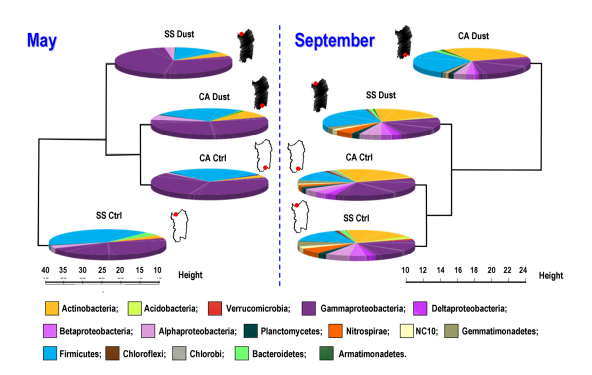


Fig. 3: Pie-charts of percent relative abundance and cluster analysis dendrograms (Euclidean distance method, complete linkage) constructed on the identified bacteria grouped at phyla/classes rank level, as stemming from the OTUs annotation of the DNA sequencing from the filtered atmosphere sampling. SS: Sassari station, Northern Sardinia; CA: Cagliari station, Southern Sardinia; 'Dust': collection during dust outbreaks under winds from

Africa, 'Ctrl': Control: collection upon under opposite wind front change, following (May) or preceding (September) the dust outbreak. The data available from the two 12h sub-periods (h 0-12, and h 12-24) collected during the dust events were merged together in the corresponding pie-charts to be compared at equal sampling duration with the 24h control samples. The Sardinia island contour map in white (Control) or black (Dust) and the red dot, pointing at the sampling site, add visual aids to each chart attributes. The squares providing a color legend to the taxa names follow a corresponding clockwise order of their possible presence on each pie-chart.

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In terms of conserved taxa the core of those observed more regularly included classes as Gammaproteobacteria, Bacilli (Firmicutes phylum) and orders as Actinomycetales (Actinobacteria phylum); their maxima were seen in the May samples where those reached percentages above 90%, while their minima appeared in the Sassari controls in September with values around 50%.

The Actinomycetales order was found in all samples; in particular it was featured in the south-facing station (Cagliari), and its numbers tended to double in relation to the dust events. The overall levels of relative abundance as well as richness within members of the Actinobacteria phylum increased from 5.66 % values, observed in May to 13 % in September. Particularly enriched were the Gaiellales and Solirubrobacterales order within the Thermooleophilia class. The orders within the Firmicutes phylum, that dominated the May samples, resulted Lactobacillales and Bacillales. Their relative abundances were higher in the Sassari (Notrhern Sardinia) control samples, than in those collected in the southern point of Cagliari, with values of 37 % vs. 12 % respectively. At the same time an unchanging level of 25% was recorded in both control and dust samples in the south-facing location. In the September samples the situation was different as in both controls those orders were below 15%, while during the dust outbreak it was 25 % in both Cagliari and Sassari stations.

The May sampling was also characterized by a large share of Gammaproteobacteria, a class reaching 75% of the dust-related spring samples in Sassari. In particular, Pseudomonadales and Enterobacteriales were constantly observed. Some taxa appearing as spring-signature cases were detected in the Alteromonadales with *Marinimicrobium*, *Marinobacter* and taxon

OM-60. For the fall period instead, Xantomonadales recurred with some genera in the Sinobacteraceae family, amounting from 1.2% to 2% respectively in the dust-free controls of Cagliari and Sassari. Within the Alphaproteobacteria class, Caulobacterales, with genera related to *Brevundimonas* were at 1% relative abundance level in the Cagliari samples collected during the dust episode of May. The Rhizobiales order was present in both seasons with a 3% peak in spring (Cagliari, dust-related), dropping to 1.5% in all fall analyses. In the same period Rhodospirillales showed a relative increase, particularly in the controls in Sassari where they reached 3.4%. The Burkholderiales (Class Betaproteobacteria) of the population were found at 1% in May within the dust-related sequences and at higher values, reaching 2.7% in Cagliari and 4% in Sassari, in the controls of September. Some groups appeared rather period -specific as the Mollicutes for May, while the Pirellulales order (in the Planctomycetes phylum) and the classes of Nitrospira and Gemmatimonadetes characterized the September sampling.

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Phylum	Class	Order	Mean	Mean
			percentage	percentage
			May	September
Proteobacteria	Gammaproteobacteria	Enterobacteriales	27.40	11.55
Proteobacteria	Gammaproteobacteria	Pseudomonadales	26.67	9.90
Firmicutes	Bacilli	Lactobacillales	18.67	15.96
Actinobacteria	Actinobacteria	Actinomycetales	5.66	13.36
Firmicutes	Bacilli	Bacillales	4.46	6.56
Proteobacteria	Gammaproteobacteria	Alteromonadales	2.77	0.76
Proteobacteria	Gammaproteobacteria	Xanthomonadales	1.51	2.63
Proteobacteria	Gammaproteobacteria	Aeromonadales	1.51	0.96
Proteobacteria	Alphaproteobacteria	Rhizobiales	1.35	1.59
Bacteroidetes	Sphingobacteria	Sphingobacteriales	1.00	1.04
Proteobacteria	Alphaproteobacteria	Rhodospirillales	0.06	2.02
Actinobacteria	Acidimicrobiia	Acidimicrobiales	0.05	2.01
Nitrospirae	Nitrospira	Nitrospirales	0.01	3.38
Actinobacteria	Thermoleophilia	Gaiellales	0.00	2.81
Actinobacteria	Thermoleophilia	Solirubrobacterales	0.00	2.37
Gemmatimonadetes	Gemm-1	Gemm-1	0.00	2.52

Table 1. Percent frequency of sequences belonging to the indicated orders in the averaged data of all samplings (Dust and control) of each sampling period (May or September). Data in which frequencies were higher than 1% in at least one of the two seasons are reported. These represent the 91.1% of the total sequences for the May sampling (on a total of 65 orders found) and 79.4% of the September sampling (on a total of 118 orders found).

The numerical effect of the different sampling period on bacterial communities is visible in Tab. 1, comparing in this case the mean relative abundances of the main orders in the two sampling months, grouped independently from site and meteorology events. Among the most evident phenomena confirmed by this single-variable grouping, the September campaign shows the enrichment in the Actinomycetales order and in a number of others that were below detection in the May sampling. In parallel, the diminution of the formerly dominant Enterobacteriales and Pseudomonadales, and the substantial stability of the Bacilli across the compared times were observed.

Besides the comparisons that included all OTUs to put in evidence community variations, in parallel we exploited an additional opportunity to detect possible dust-specific taxa. The rationale was to seek differential enrichment within the dust storm, by dissecting the process, during its progression, splitting its onset from its fully established stage. To this aim, we collected separately the filters of the first 12h of the event, and replaced them with new ones that collected air during the second lapse (hours 12 to 24). Thus the availability of two timeframes, both within the dust event, allowed to verify which OTUs would be incrementally enriched along with the progression of the stormy condition.

This allowed to better refine the bacterial deposition dynamics during the outbreaks. From the visual and physical points of view, an increase in the inflow of air particulate was observed for the 12-24 h period, confirming the differential level of deposition occurring in the maturity stage of the meteorological phenomenon.

This within-outbreaks set up was essentially aiming at individuating OTUs that would display high variation in relation to dust events in comparison to those who would not. The latter were considered to represent the common core of bacteria that were constantly present in samples, irrespective of the changing meteorological events. To apply this distinction, the criterion was to set a cutoff value with respect to the percent of variation occurring between the first 12 h of the collection time and the second half of it. The choice of this threshold was considered critical and, in order to ensure robust conclusions, we deemed necessary to require a considerable consistency of variation. Pointing at this objective, only the taxa which displayed a mean variation higher than ½ of the corresponding standard deviation were taken into account. The resulting level of variation in the two sampling stations is reported in Tab. 2 and the corresponding number of orders is displayed in Tab. 3. The Sassari (Northfacing) collection site was the one that in both seasons resulted to feature the highest number of significantly changing taxa. The identities of these are shown in Supplementary Fig.S7 (May event), and Supplementary Fig. S8 (September event). In the graphs, the first 12h lapse is plotted above the baseline and the second (12-24 h) is on the specular position below.

Sample	Avg. variation %	Min variation %	Max variation %
Sassari May - Dust	1.4	0.05	6.7
Sassari Sepember - Dust	1.3	0.4	5.4
Cagliari May - Dust	2.1	0.5	5.0
Cagliari September - Dust	4.7	1.1	11.4

Table 2: Extent of OTUs change across cell harvesting time during the same dust outbreak sampling. The percent variation (either increase or decrease) of a given OTUs abundance between the values found in the community obtained by the first 12h sampling and the ones resulting from the following 12h lapse was computed. The average, minimum and maximum percent variation between counts are shown. Only taxa displaying a difference in percentages higher than half of their standard deviation were selected for the present comparison.

Site and period	Total Orders	Selected Orders	% of total orders
Sassari May	56	16	28%
Sassari September	103	28	28%
Cagliari May	52	11	21%
Cagliari September	87	14	16%

Table 3. Community richness at order level of taxa occurring during dust events and of those displaying variations higher than half the standard deviation between the first 12h and the second 12h sampling period (selected orders). The percentage of orders selected upon this criterion over the total of the orders observed in samples collected during the dust events is indicated.

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As regards the ecological indexes characterizing the communities, species diversity and evenness values were calculated, and results are shown in Tab. 4. The difference that can be appreciated is mainly relative to the series of September samples, in which all had higher values for each of the indexes when compared to the May ones. Conversely, neither the presence of dust events nor the sampling location appeared to confer relevant differences in this respect.

The patterns of conservation and diversity involving the bacterial communities analyzed were subsequently inspected by multivariate approaches. Principal Component Analysis yielded an output (Fig. 4) that confirms how a separation of communities can be viewed only when considering the temporal factor (Fig.4. A), while the variables of dust vs. calm air, or the sampling location, led to plots with heavily overlapping patterns. The May vs. September divide occurs along the horizontal axis, i.e. the one explaining the highest fraction of variation (35%). The same phenomenon is reproduced with a higher support (54 %) in a parallel

Month	, Event	, Place		Simpson 1-D	Shannon H	Evenness
May	Dust	SS	h 1-12	0.771	2.062	0.151
May	Dust	SS	h 12-24	0.740	1.902	0.156
May	Dust	CA	h 1-12	0.833	2.175	0.183
May	Dust	CA	h 12-24	0.833	2.205	0.197
May	Ctrl	SS		0.794	2.064	0.164
May	Ctrl	CA		0.778	1.900	0.142
Sep.	Dust	SS	h 1-12	0.928	3.187	0.260
Sep.	Dust	SS	h 12-24	0.914	3.015	0.240
Sep.	Dust	CA	h 1-12	0.887	2.792	0.212
Sep.	Dust	CA	h 12-24	0.838	2.339	0.176
Sep.	Ctrl	SS		0.948	3.438	0.311
Sep.	Ctrl	CA		0.936	3.292	0.286
May: mean ±SD		0.79 ± 0.04	2.05 ± 0.13	0.17 ± 0.02		
September : mean ±SD		0.91 ± 0.04	3.01 ± 0.40	0.25 ± 0.05		

Table 4. Ecological diversity and evenness indices resulting from the sequence checklist analysis in the different samplings.

ordination approach, the principal Coordinate Analysis (Fig. S9A, Supplementary Material). In the same figure the main differences occurring in community structure between the two sampling periods are further explored by reporting the ecological indexes of Shannon species diversity (Fig. S9B) and community evenness (Fig. S9C) resulting from grouping the data and separating them only in relation to the sampling period variable, irrespective

of meteorology events and collection sites. The superiority of the September values in both indexes, and particularly for the taxa diversity, is supported by the significance of the p values of discrimination between samples thereby reported (p = 0.00584 for species diversity, and p = 0.026 for community evenness).



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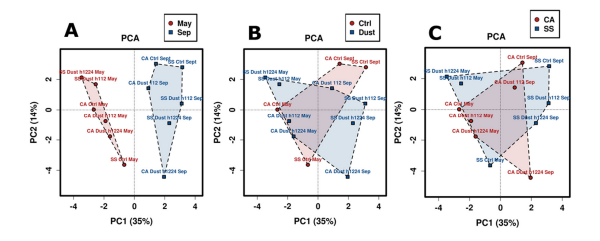


Fig. 4. Principal Component Analysis (PCA) ordination plot of the bacterial community compositional data. Polygons encompassing the positions of three different variables are drawn to visualize sampling month (A), ongoing meteorological event (B), or sampling location (C).

The higher strength of clustering of the sampling date groups with respect to the alternative ones (meteorological or geographical) was verified by running a Discriminant Analysis on the Principal Component ordination (DAPC) in which the data are first transformed by PCA, from which, clusters are subsequently identified using Discriminant Analysis, thus partitioning sample variance into the between-group and within-group components. Results are shown in Fig. 5. Besides confirming the sampling period as the strongest driver of community change, the analysis further shows that the dust vs. control clustering is acting more efficiently than the Sassari vs, Cagliari sampling site comparison. This allows to draw a hierarchical ranking of the three variables in shaping the bacterial airborne communities, in which, noting also the different scale of the horizontal axis (Discriminant function 1) adopted for the three graphs, the order results: Time >> Meteorology > Geography.

In order to determine which bacterial taxa were mostly accompanying/causing those changes in a statistically significant manner, and to rank their individual importance in this phenomenon, we run an analysis of the differentially featured taxa, testing both an ANOVA variance analysis and a non parametric Wilcoxon Rank test verification of the ranking. The two tools gave coherent scores and the results of the ANOVA output are shown in Supplementary Table S1. A total of 76 taxa were found featuring p values < 0.05, from which, upon applying a stringent Bonferroni-adjusted p value correction, six of those stood above the significance cutoff, and all within minimal false discovery rate values (FDR < 0.005). All of them were cases which were highly reduced in September in comparison to May. The taxa included as the most effective in explaining the differences (p value = 0.000019), the order Oceanospirillales, known as marine oil spill-associated bacteria (Cao et al, 2013), followed

by known animal parasites as the Coxiellaceae family (Lory, 2014), marine extremophiles as the Thiohalorhabdales (Tian et al., 2017), and three species of *Pseudomonas*, including the pathogenic *P. viridiflava* (Hu et al., 1998), the decontamination-associated *P. nitritireducens* (Wang et al., 2012) and *P. alcaligenes* which is reported also a human pathogen (Suzuki et al, 2013). The two corresponding analyses of differentially represented taxa by meteorology or by geography, i.e., grouping dust vs. calm air or Cagliari vs. Sassari sites did not yield any significantly supported cases under the Bonferroni-adjusted p values stringent condition (data not shown).

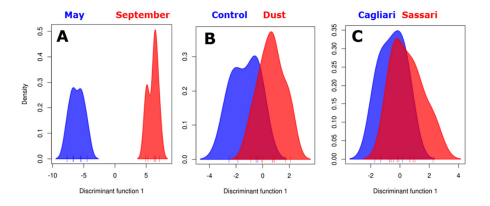


Fig. 5. Discriminant Analysis of Principal Components analysis. Group partitioning involved **A**: sampling month; **B**: ongoing meteorological event; **C**: sampling location.

In order to compare all communities with each other and extract further information on their degrees of divergence, the sequencing data were analyzed by individual comparisons across sites and dates. The results of each of the 66 pairwise combinations are shown in Tab. 5, displaying the Bray-Curtis similarity values between each couple of communities. Color-based conditional formatting applied to the values allows to appreciate how all the comparisons involving different seasons show the most divergent scores (red shades) in comparison to those within the same season, that show much more similarity, with few exceptions related to dust events and depending on the aspect faced by the collecting site with respect to the incoming wind direction.

Different season						
<u>Different place</u>			<u>Same place</u>			
SS D1-12 May	CA D1-12 Sep	0,476	SS D1-12 May	SS D1-12 Sep	0,348	
SS D1-12 May	CA D12-24 Sep	0,344	SS D1-12 May	SS D12-24 Sep	0,380	
SS D1-12 May	CA Ctrl Sep	0,449	SS D1-12 May	SS Ctrl Sept	0,355	
SS D12-24 May	CA D1-12 Sep	0,416	SS D12-24 May	SS D1-12 Sep	0,317	
SS D12-24 May	CA D12-24 Sep	0,321	SS D12-24 May	SS D12-24 Sep	0,372	
SS D12-24 May	CA Ctrl Sep	0,423	SS D12-24 May	SS Ctrl Sep	0,312	
CA D1-12 May	SS D1-12 Sep	0,471	SS Ctrl May	SS D1-12 Sep	0,450	
CA D1-12 May	SS D12-24 Sep	0,532	SS Ctrl May	SS D12-24 Sep	0,507	
CA D1-12 May	SS Ctrl Sep	0,388	SS Ctrl May	SS Ctrl Sept	0,420	
CA D12-24May	SS D1-12 Sep	0,481	CA D1-12 May	CA D 1-12 Sep	0,536	
CA D12-24May	SS D12-24 Sep	0,538	CA D1-12 May	CA D12-24 Sep	0,476	
CA D12-24May	SS Ctrl Sep	0,417	CA D1-12 May	CA Ctrl Sep	0,526	
SS Ctrl May	CA D 1-12 Sep	0,448	CA D12-24 May	CA D 1-12 Sep	0,530	
SS Ctrl May	CA D12-24 Sep	0,523	CA D12-24 May	CA D12-24 Sep	0,479	
SS Ctrl May	CA Ctrl Sep	0,469	CA D12-24 May	CA Ctrl Sep	0,525	
CA Ctrl May	SS D1-12 Sep	0,387	CA Ctrl May	CA D 1-12 Sep	0,450	
CA Ctrl May	SS D12-24 Sep	0,464	CA Ctrl May	CA D12-24 Sep	0,416	
CA Ctrl May	SS Ctrl Sep	0,355	CA Ctrl May	CA Ctrl Sep	0,453	
	Same season	(May) Cor	ntrol taken after du	st		
<u> </u>	<u>Different place</u>		Same place			
SS D1-12 May	CA D1-12 May	0,649	SS D1-12 May	SS D12-24 May	0,736	
SS D1-12 May	CA D12-24 May	0,627	SS D1-12 May	SS Ctrl May	0,545	
SS D1-12 May	CA Ctrl May	0,640	SS D12-24May	SS Ctrl May	0,522	
SS D12-24 May	CA D1-12 May	0,709	CA D1-12 May	CA D12-24May	0,802	
SS D12-24 May	CA D12-24 May	0,655	CA D1-12 May	CA Ctrl May	0,784	
SS D12-24 May	CA Ctrl May	0,713	CA D12-24 May	CA Ctrl May	0,791	
CA D1-12 May	SS Ctrl May	0,618				
CA D12-24 May	SS Ctrl May	0,679				
SS Ctrl May	CA Ctrl May	0,676				
Same season (September) Control taken before dust						
<u>Different place</u>		Same place				
SS D1-12 Sep	CA D 1-12 Sep	0,600	SS D1-12 Sep	SS D12-24 Sep	0,718	

SS D1-12 Sep	CA D12-24 Sep	0,604	SS D1-12 Sep	SS Ctrl Sep	0,705
SS D1-12 Sep	CA Ctrl Sept	0,656	SS D12-24 Sep	SS Ctrl Sep	0,616
SS D12-24 Sep	CA D 1-12 Sep	0,599	CA D 1-12 Sep	CA D12-24 Sep	0,554
SS D12-24 Sep	CA D12-24 Sep	0,655	CA D 1-12 Sep	CA Ctrl Sep	0,589
SS D12-24 Sep	CA Ctrl Sep	0,656	CA D12-24 Sep	CA Ctrl Sep	0,441
CA D 1-12 Sep	SS Ctrl Sep	0,523			
CA D12-24 Sep	SS Ctrl Sep	0,434			
SS Ctrl Sep	CA Ctrl Sep	0,705			

Table 5. Bray Curtis similarity values between the bacterial communities composition resulting from pairwise comparisons of all samples. Abbreviations: SS: Sassari; CA: Cagliari; D1-12: dust event, first 12 hour period; D12-24: dust event, second 12 hour period; Ctrl: control conditions (absence of dust events); Sep: September.

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4. DISCUSSION

Two technical aspects to be defined beforehand are the definitions of 'sample' and 'control' that apply to this type of studies. An atmospheric sampling is not to be regarded with the same conceptual metrics that would apply if one were to study water or soil environments, in whose cases, a sample could correspond to one liter or one gram taken respectively from those milieus. In air-filtering the operation is carried out continuously for days and one sample, in our tests, accumulated the content of 56160 liters of continuously changing atmosphere, which takes into account the variations that occur during all those hours, inclusive of the day/night shifts. One sample is

therefore not a 'point' but a built-in averaged replication protocol for the chosen window of events.

Second, as regards the definition of control in our study, it is not meant to represent a supposedly stable

community that could apply to an hypothetical average dust-free local scenario. As several studies cited in the introduction showed, even in the absence of dust outbreaks, the ambient state of the atmosphere is not stable either. Our goal therefore was not to compare a hypothetical *status quo* with an altered one. Since the airborne community composition does change daily even during periods that do not feature the dust carrying episodes, there is not a stable condition that could be considered as a durable control community. Even the evening and morning opposite breeze regimes that occur daily in coastal locations impart modifications in local airborne communities. The meaning of the 'control' here was instead meant to catch 1) the first possible timeframe after the stopping of a northbound dusty wind outbreak (it occurred in May) or 2) the latest possible timeframe of a situation before the onset of a dust-carrying change of wind regime (it occurred in September). Thence, in the latter event the control is not intended as a situation of calm that could represent a period of unknown length, but rather the time-zero sample of the dust event itself. While for the former case in May, the control was symmetrically defined as the earliest stage of quiet after the storm.

In the present study the filtered air particulate was analyzed in different seasons and under different wind regimes, using culture-independent DNA sequencing-based approaches targeting the species-diagnostic 16S-rRNA genes

from the air-carried bacterial community and an Illumina next generation sequencing platform. Sites were selected also because of their opposite positions facing Africa (Cagliari) or continental Europe (Sassari). The whole analysis was performed within a 7-month time lapse, March to September, chosen also as it offers higher probabilities of weather shifts favoring both northern- and southern-winds (Israelevich *et al.* 2012). This timeframe proved suitable to the scope as it was possible to exploit two episodes in which dust outbreaks carried by winds of African origin occurred and were preceded and followed by inversions of the air circulation offering control sampling periods with opposite features.

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Commenting on the taxonomical abundance shifts observed between May and September and trying to interpret the rise of some phyla and the drop of others (Fig.3; Tab. 1), a preliminary consideration needs to be recalled. The issue has to do with the distinction between actual population dynamics (ecologically-ruled) and mathematical effects of sampling from a 'ballot box' of objects, all of which compete for the constrained 100% format of results (probability-ruled). This caveat was put forward as early as statistics itself was born as a discipline (Pearson, 1897). There is in this respect a general problem in comparing communities at different time points (and even more with DNA-based methods with fixed total DNA amount processed), whose results are based on percent values (relative abundance of taxa). In that condition the multiplication of any determines obviously a reduction of the relative abundance of others, when those do not grow at equal or higher rate. Therefore, one given group could have been increasing, but its share in the sum could appear as if it had instead decreased, if a different group has increased faster. This consideration, applies inevitably for all metagenomics/metabarcoding surveys, and should be kept in mind for all kinds of interpretations about increases and decreases, which could be either real or apparent (when driven by a stronger change of a different group). As consequence, comparing different sampling points through time is linked to this inevitable constraint: the compositional nature of the datasets binds all relative frequencies to each other (Gloor et. Al., 2017). Therefore, since, as mentioned, the sum of them is bound to give always 100%, the decrease of a given species could be either apparent (driven by the increase of another), or real (due to its actual negative population dynamics). The problem is that the two causes can not be uncoupled by just comparing species frequencies at the two sampling periods. Moreover, as mentioned, even an actual increase of a given species could be masked by the parallel increase of another at a faster pace (or by its net immigration into the scene). For this reason, we consider with caution the possibility of looking at taxa fluctuations as indicative of their actual ecological outcomes. Having clarified that we will therefore limit to comment only the major phenomena that stand out from the comparison. The largest taxa trade-off that is apparent when comparing the two periods is the decline of Proteobacteria and the parallel rise of Actinobacteria. Trying to frame this within possibly seasonal parameters we can consider that the latter are typically relying on profuse spore formation from colonial growth, while the former are non-sporeforming bacteria either motile via flagella or gliding/swarming mechanisms. The basic life forms of the two groups predict therefore that proteobacteria would be more suited by wet seasons and vice versa for the soil-dwelling Actinobacteria. This phylum has been reported by other authors to reach its peaks in fall (Glöckner, et al., 2000). Being also a group of major litter decomposers their rise along with the end of the plants' vegetative season can be seen as compliant with their landscape and ecosystem cycles.

A further ecological aspect that can be commented is the higher diversity of the communities during the September sampling in comparison to May, independently from the dust events and from the sampling station location.

This phenomenon, besides the ecological values differences (Tab. 2, Tab. 4, Fig.S9, and Tab. S1) can be also appreciated visually, by comparing the left and the right pie charts in Fig.3, (featuring community composition at order-rank level, and the corresponding cluster analysis based on their relative percentages), and noticing the more complex color-coded pattern of the latter sampling, showing also a consistent similarity of most color sectors presence and proportions. It is not possible from these single-year data to deduce whether such increase could be part of a recurring seasonal phenomenon causing, cyclically, higher species richness after summer periods, or if what we observe could be part of a different pattern of stochastic variability.

Nevertheless, the overall partitions of systematic groups observed in a given sampling time, irrespective of dust outbreaks or sampling corner of Sardinia, share much more similarity within the samples of that period than with any of those collected in the other season. It appears that in general, air collected during dust discharge from a Saharan wind can account for less variation over its reference control sampling than the choice of sampling that site four months apart.

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In our prior work (Rosselli et al., 2015) we had studied community composition in the same Sardinian stations in a short period of winter (in late February) during and after a single dust-carrying event. In that study the main feature evidenced was the existence of a conserved core microbiome, encompassing 86-95 % of the taxa, to which the incoming dust would cause some detectable diversity variation but on a rather limited proportional scale. Such minor effect of the dust-lifting storms observed in winter is in fact confirmed in the present work in which the time of the year factor appears as the variable of major order in shaping community structure and richness. In that prior analysis of ours the existence of a common core microbiome of the investigated area was one of the suggested evidences. That concept was stemming from the analysis run in February, therefore towards the end of a winter period throughout which Europe experiences its minima in terms of temperature-driven air turbulence events and as consequence receives more limited influxes of air travelling from seas to land. In the present analysis, we observe that, in spite of the major changes brought about by the temporal factor, the two sampling stations at opposite corners of the 270 km-long island shared the closest level of community composition when they were compared at the same time (see CA Ctrl vs. SS Ctrl in Fig.3 Fig.4, Fig. S9, and Table 5). Moreover, this similarity was maintained in May even though the two control stations were compared after the dust outbreak. Foremost, those two distant sites achieved the maximum of community overlap in September, when controls were compared right before the next outbreak, after a 109 days-long period without such events. During that time the air microbiome of the whole area appears to have changed profoundly, but in a concerted fashion, leading to a high uniformity across the island. These data confirm the view of the prevalence of a core microbiome, as emerged in our 2015 report and add the evidence that such extended core community undergoes also a temporally related concerted turnover. Whether or not this could be also a seasonal (regularly recurrent) phenomenon, will have to

Literature reports have in this sense pointed out differences in airborne microbial composition between seasons; peaks of fungi causing invasive infections in humans were signaled in spring whereas higher proportions of allergenic fungi were observed in fall (Yamamoto et al. 2012).

be demonstrated by further research on this subject.

Consistent with the present data, a higher diversity of both fungal and bacterial airborne cells in late summer and early fall has been observed in United States-based surveys (Bowers et al. 2012, Bowers et al., 2013).

Hypotheses to explain the increase in circulating taxa widely observed in the fall sampling campaign can be formulated. In first instance one should consider whether there could have been a change in the prevailing winds

origin or direction across the period that encompasses the two sampling seasons. This can be evaluated upon inspecting publically available meteorology records showing the wind roses for the two sampled localities. These data, from March to November, for the Cagliari and Sassari weather stations, are shown in Supplementary figures Fig. S10 and Fig S13, respectively. In the Cagliari plots (southern Sardinia) it can be observed that between May and September there was basically no variation of the wind patterns, with the prevailing ones blowing towards North-West, with stable intensities. Likewise in the Sassari area (Fig. S13), although some fluctuations in the strength of the westbound winds can be seen, the dominant air motion throughout the period remained the one heading South. In essence these data allow to rule out that the change in community patterns could be due to major air-driven events of taxa immigration from other insular or continental sources.

In addition to the wind orientation and force, data from the two stations regarding temperature and humidity of the same winds can be analyzed (Supplementary Fig. S11, Fig. S12, Fig. S14, Fig. S15). Humidity values from May to September winds tend to be rather similar, whereas air temperatures increase in line with the summer progression. These data do not account by themselves for events of species enrichment either.

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Another aspect that can be verified is to compare the two periods in terms of PM10 particulate concentration; these are reported in Supplementary Fig. S1.C (May) and Fig. S4.C (September). Although there are obvious peaks of PM10 in correspondence with the dust outbreaks dates, the basal levels of PM10 concentrations before and after those, are rather similar in the spring and fall period. This rules out the possibility of a diversity rise as linked to a general increase of such small particles trafficking over the areas.

The observed data reveal that, while dust-associated winds can account for some specific limited ingression of taxa, a far more noticeable pattern appears consisting in a successional rise of taxa diversity. It is not yet possible to establish whether this occurrence could be linked to late summer in relation to the climatic conditions of the season. The second part of the summer, especially in the Mediterranean regions, is characterized by prolonged drought alternated to irregular thunderstorms. The income of a thunderstorm is accompanied by convective instability of the atmosphere and this phenomenon has been already pointed out as conducive to the emission and transport of fungal spores plumes (Burch and Levetin, 2002). A possible explanation for a richer pattern of airborne microbes after several weeks of prevailingly dry climate can be sought in the acknowledged fact that those seasonal conditions enhance the daytime height of the planetary boundary layer over Europe and continental US (Seidel et al. 2012), and that the ensuing low pressures foster the turbulence near ground and the overall convection, resulting in a frequent uplift of particles from land surfaces. In addition, it could also be postulated that the dryer and warmer summer conditions can eventually lead to partial cell dehydration in microbes lying at soil or vegetation surface, resulting in lighter cell weights more prone to be advantageously lifted by the local low layers air turbulence.

A further factor that can be hypothesized to have played a role in reducing the diversity of airborne community samples in May, comes from the analysis of the differentially featured taxa between the spring and the fall samplings (Tab. S1) where the strongest statistically significant differences were six taxa that resulted highly enriched in the former period and that, as cited above, included marine bacteria associated to oil spill-related oleovory phenotypes, extremophyles, and potential pathogens. These occurrences can be interpreted as possible clues for a transient event of water pollution around the sampled areas that could have impacted also on the overall airlifted microbial diversity.

In addition to the above, a series of considerations can be drawn upon inspecting the pairwise community difference analysis, whose similarity values are shown in Tab. 5. It also needs to be recalled that, in order to examine the effect of a dust-free period, in May the control (May 27th) was sampled after the dust event (May 21st -22^{nd}), while in September the control (Sep 13th) was taken before the new dust outbreak (Sep 19th -20^{th}). Therefore, the summer, within which communities could undergo dust-independent changes, is in fact framed between the two control points, chosen as representative of dust-free atmosphere following a wind direction reversal. Within those months there were no dust-carrying wind outbreaks from the African land. This enabled also to verify whether a relatively long period without dust intrusions could have allowed an overall homogenization of the bacterial airborne communities over the island Sardinia. Regarding Tab. 5, we can point out the main message that this color table is meant to convey, which is just the fact that the low similarity pairwise comparisons (conditionally formatted as red-yellow shaded cells) are almost all observed in the upper table. i.e. the between-seasons comparisons, while the green ones (higher similarities) are all distributed in the two sameperiod comparisons. The first consideration that stems from the global view of these data is once again that the most distant communities are those compared from different seasons (Tab.5, upper section). It is worth noticing in this respect that no particular difference appears when comparing communities between those collected from the same site (right panel in the upper section) or in the cross-comparison between the two different places. Moreover, in these samplings from different seasons, the effects of the dust events in comparison to calm air with dust-free wind regimes, is not apparent, being diluted in the major time-related divergence of the communities. When inspecting cases of the same season, the situation in May is representing a comparison picturing the recovery after the dust event, as the control follows the outbreak. One evident aspect in that is how the juxtapositions within the same place, feature the most similar cases (darkest shades of green) with the notable exceptions linked to the dust outbreak in Sassari, which is the North-facing station (notice the two yellow-shaded values). On the contrary, such divergence does not appear at all in the South-facing Cagliari site. One interpretation of this interesting difference is that in the May control, when the air flux reversed after the dust event, the wind blowing from the northern quadrant, was conveying in Sassari air masses that came straight from the sea; while on the opposite corner (Cagliari) instead, the same air had passed over the whole Sardinia. Thus, the Northern collection station received sea-sweeping air, bringing 'fresh' taxa, i.e. not belonging to the Sardinian land-related common bacterial core of the period, while the southern station of Cagliari received instead landsweeping air that had travelled all the way over the island latitudinal extension, and that therefore would have become mixed with the island-related core of biota. Thus, the sea-related entries would bring little contribution to the southern site communities after more than 100 miles of travelling and being diluted through the terra firma atmosphere. This would explain why, in May, shifting from dust outbreak to control in the Southern location, did not bring community divergence as it did in the Northern one. The imprint of the common Sardinian core on homogenizing communities when the dust preceded the control, is

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also testified by the left panel (same month of May but different places) resulting in all green shades of medium value, showing that in such situation there was little difference also between different places.

An independent and indirect confirm of this interpretation is given by the situation in September. In that case, the African dust-carrying event was set to be taken after the control; this originated a reversed situation in comparison to the one observed in May; this time the place where the dust-outbreak did not bring particular change was the Northern site, Sassari, as the northbound wind from African origin had supposedly already discharged its load while passing over the land of the island from which, at the same time it would have lifted a vast portion of landrelated common biota. Vice versa, in Cagliari, appreciable changes occurred in relation to the dust arrival, which support the view of air blown over the sea plus dust, as the elements causing changes due to the new kinds of bacteria that hit this side at the frontal south-facing port of entry of the island. The left panel of the section (same season, September, different place) further confirms this as: (a) the strongest drivers of community divergence (yellow to orange colors) are flagged by the two comparisons between the Sassari control and the Cagliari dust situations, and the second of those, in the 12-24 hours window of the dust event is progressively more divergent then that recorded during the first 12 h (0.434 vs. 0.523 similarity value). Moreover, the comparison between the two sites in the September control before dust, shows a rather high similarity (0.705), that is the highest among the September comparisons of different sites, which confirms that, before the dust outbreak, when both localities had experienced a long period devoid such phenomena, the two places had achieved a high degree of uniformity in spite of their distance. In that status, both communities were also profoundly different from their composition in May. A period of over 100 days without intrusions of dust-carrying northbound winds, appears to have accompanied an appreciable concerted change of the air-associated bacteria upon the Sardinian territory. Essentially it appears that when airborne dust has to cross longitudinally the entire large island, it reaches the Northern sampling site (Sassari) less charged with community-changing potential, and/or, that it must have lifted bacteria from of the Sardinian common core, thus causing little variation upon their discharge over a station on the same island. On the contrary, when landing on the south-facing outpost of Cagliari, coming straight form Africa and, until that moment, having travelled over the sea only with air-lifted transcontinental dust, those air masses delivered in the south outpost of Cagliari an appreciably novel community. The geographic position of the sampling sites in relation to the wind origin appears therefore to play a major role in the patterns outcome. This supports the view that, in case of dust outbreaks, Cagliari, in the south, is at the forefront of changes that are substantially attenuated before they reach Sassari; and vice versa, in case of reversed winds. A distance of >100 miles appears sufficient to absorb and buffer wind-borne taxa immigration in quantitative terms, from either side. In conclusion, the central goal of this study was to assess which variables (sampling period, dust outbreak vs.

homogenizing communities is stronger than the one conferred by occasional transcontinental discharges. These clues entail novel aspects for our better understanding of microbial transport and spread across territories, of the epidemiological patterns for clinically relevant taxa, and can foster the predictive modeling of overall environmental microbiology dynamics.

calm atmosphere, and north-facing vs. south-facing collection site) would be most effective in determining

airborne community divergence or homogenization. Data are supportive of time-related turnover phenomena, involving a pattern of diffuse contemporary colonization over large portions of land, whose effect in shaping and

Supplement.

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The supplement related to this article is available online at: https.....

Author contributions.

PC conceived the project; RM, AS, AC, and PD designed the experiments and supervised the project. RR, MF, MD, GP1 and GP2 carried out the analyses and interpreted the data. AS wrote the manuscript.

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Competing interests

The authors declare that they have no conflict of interest.

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