Bioaerosols and atmospheric ice nuclei in a Mediterranean dryland: community changes related to rainfall

Kai Tang1, Beatriz Sánchez-Parra1,2, Petya Yordanova1, Jörn Wehking1,a, Anna T. Backes1, Daniel A. Pickersgill1, Stefanie Maier2, Jean Sciare3, Ulrich Pöschl1, Bettina Weber1,2, and Janine Fröhlich-Nowoisky1

1 Multiphase Chemistry Department, Max Planck Institute for Chemistry, P.O. Box 3060, 55020 Mainz, Germany
2 Institute of Biology, University of Graz, Holteigasse 6, 8010 Graz, Austria
3 Climate and Atmosphere Research Center, The Cyprus Institute, 2121 Nicosia, Cyprus

These authors contributed equally to this work.

Correspondence: Janine Fröhlich-Nowoisky (j.frohlich@mpic.de)

Received: 21 July 2021 – Discussion started: 9 August 2021
Revised: 25 October 2021 – Accepted: 20 November 2021 – Published: 5 January 2022

Abstract. Certain biological particles are highly efficient ice nuclei (IN), but the actual contribution of bioparticles to the pool of atmospheric IN and their relation to precipitation are not well characterized. We investigated the composition of bioaerosols, ice nucleation activity, and the effect of rainfall by metagenomic sequencing and freezing experiments of aerosol samples collected during the INUIT 2016 campaign in a rural dryland on the eastern Mediterranean island of Cyprus. Taxonomic analysis showed community changes related to rainfall. For the rain-affected samples, we found higher read proportions of fungi, particularly of Agaricomycetes, which are a class of fungi that actively discharge their spores into the atmosphere in response to humidity changes. In contrast, the read proportions of bacteria were reduced, indicating an effective removal of bacteria by precipitation. Freezing experiments showed that the IN population in the investigated samples was influenced by both rainfall and dust events. For example, filtration and heat treatment of the samples collected during and immediately after rainfall yielded enhanced fractions of heat-sensitive IN in the size ranges larger than 5 µm and smaller than 0.1 µm, which were likely of biological origin (entire bioparticles and soluble macromolecular bio-IN). In contrast, samples collected in periods with dust events were dominated by heat-resistant IN active at lower temperatures, most likely mineral dust. The DNA analysis revealed low numbers of reads related to microorganisms that are known to be IN-active. This may reflect unknown sources of atmospheric bio-IN as well as the presence of cell-free IN macromolecules that do not contain DNA, in particular for sizes < 0.1 µm. The observed effects of rainfall on the composition of atmospheric bioaerosols and IN may influence the hydrological cycle (bioprecipitation cycle) as well as the health effects of air particulate matter (pathogens, allergens).

1 Introduction

Primary biological aerosol particles, or short bioaerosols, are a subset of atmospheric aerosol particles directly emitted from the biosphere into the atmosphere. They comprise living and dead microorganisms (e.g., bacteria), dispersal units (e.g., fungal spores, plant pollen), fragments (e.g., plant debris), and viruses (Després et al., 2012; Whon et al., 2012; Fröhlich-Nowoisky et al., 2016; Smets et al., 2016; Yahya et al., 2019; Pöhlker et al., 2021). Bioaerosols can serve as nuclei for cloud droplets and ice crystals influencing the formation of precipitation, and some of them are important allergens and pathogens (Brown and Hovmøller, 2002; D’Amato et al., 2007; Andreae and Rosenfeld, 2008; Steiner et al., 2015; Müller-Germann et al., 2017; Fröhlich-Nowoisky et al., 2016; Lang-Yona et al., 2018; Huang et al., 2021). The abundance and composition of bioaerosols are determined by local emission sources, meteorological conditions, and long-range transport (Jones and Harrison, 2004; Franzetti et al., 2011; Bowers et al., 2012, 2013;
Rainfall and the associated increase in humidity can lead to increased emission of biological aerosol particles (Heo et al., 2014; Wright et al., 2014; Rathnayake et al., 2017). For example, raindrops hitting leaf or soil surfaces can disperse bacteria and other bioparticles by splashing (Hirst, 1953; Madden, 1997; Gilet and Bourouiba, 2014; Perryman et al., 2014; Kim et al., 2019; Gregory et al., 1959; Joung et al., 2017). Furthermore, the increase in relative humidity can trigger the emission of fungal spores by active discharge mechanisms (Zobeli, 1964; Lacey, 1996; Elbert et al., 2007), and pollen grains can rupture under moist conditions, releasing sub-pollen particles (Taylor et al., 2002, 2004; Taylor and Jonsson, 2004; Miguel et al., 2006; Steiner et al., 2015).

The release of bioaerosols, which can serve as cloud condensation nuclei (CCN) or ice nuclei (IN), can in turn influence the evolution of clouds and precipitation, thus closing a feedback cycle known as bioprecipitation (Sands et al., 1982; Möhler et al., 2007; Morris et al., 2014; Steiner et al., 2015; Fröhlich-Nowoisky et al., 2016). Especially over vegetated regions, in marine environments, or under remote conditions, bioparticles might represent a significant fraction of CCN and IN (Andreae and Rosenfeld, 2008; Pöschl et al., 2010; Pöhler et al., 2012; Burrows et al., 2013; Wilson et al., 2015). As IN of biological origin (bio-IN) can trigger freezing at higher temperatures than mineral IN, they are expected to play a role at cloud temperatures $> -15^\circ C$ (Hooge and Möhler, 2012; Fröhlich-Nowoisky et al., 2016; Coluzza et al., 2017; DeMott and Prenni, 2010).

Increased concentrations of atmospheric IN during and after rainfall have been linked to bioaerosols (e.g., Huffman et al., 2013; Prenni et al., 2013; Schumacher et al., 2013; Tobo et al., 2013; Bigg et al., 2015; Hara et al., 2016b), but the sources and composition of bioparticles, their contribution to the pool of atmospheric IN, and their relation to rainfall are still not well characterized. In this study, we investigated the effects of a short period with rain events on bioaerosol and atmospheric IN composition in a rural dryland. Aerosol samples of boundary layer air were collected on the Mediterranean island of Cyprus in April 2016. The taxonomic composition of bioaerosols was determined by shotgun metagenomic sequencing and analysis. Ice nuclei were measured using the high-throughput Twin-plate Ice Nucleation Assay (TINA). Additional filtration and heat treatment experiments were performed to narrow down the IN size ranges and to investigate the heat sensitivity of the IN.

2 Materials and methods

2.1 Aerosol sampling

Aerosol sampling was performed during the INUIT-BACCHUS-ACTRIS campaign in April 2016 at the Cyprus Atmospheric Observatory, which is a rural background station that operates under the co-operative program for monitoring and evaluating the long-range transmission of air pollutants in Europe (EMEP) and the European research infrastructure for the observation of Aerosols, Clouds and Trace Gases Research Infrastructure (ACTRIS) networks while, at the same time, it is a designated regional Global Atmosphere Watch (GAW) station. The station (35.038692° N, 33.057850° E) is located close to the villages of Agia Marina (∼ 630 inhabitants) and Xyliatos (∼ 150 inhabitants) and has an elevation of 532 m above sea level. The site lies at the northeastern foothills of the Troodos Mountains. The nearest main urban agglomeration is at least 35 km away. The vegetation at the measurement site is dominated by evergreen shrubs and small trees, blending at higher elevations of the Troodos Mountains into oak and pine forests (Fall, 2012).

Total suspended particle samples were collected on glass-fiber filters (Type MN 85/90 BF, 150 mm diameter, retention capacity of 0.5 µm; Macherey-Nagel GmbH & Co. KG, Germany) with a DHA-80 high-volume aerosol sampler (DIGITEL Elektronik AG, Hegnau, Switzerland; volumetric flow rate 1000 L min$^{-1}$, sampling time 24 h) positioned at the rooftop of the Cyprus Atmospheric Observatory station (∼ 5 m above the ground). Blank samples were taken at regular intervals (∼ 3 d intervals) to detect potential contamination by the aerosol sampler or by filter mounting. For this, filters were mounted in the sampler as for regular sampling, but the sampler was turned on either not at all (mounting blank) or only for 5 s (start-up blank).

Prior to sampling, all glass-fiber filters were pre-baked at 330 °C for 8 to 10 h in aluminum bags to remove any biological material. The filter holders were disinfected with Bacillol® AF (Bode Chemie GmbH, Hamburg, Germany) before use. After sampling, the loaded filters were removed from the filter holders, placed into baked (330 °C, 8 to 10 h) aluminum bags, stored at −20 °C, and stored at −80 °C after international transport on dry ice. The aerosol filter samples were cut into 16 equal pieces using a self-made cutting tool, made of sharp guided blades stuck to a round aluminum frame (140 mm diameter and approximately 1 cm thickness). Prior to cutting, the cutting tool was sterilized by dipping it into ethanol (96 %) and flaming. The filter sample aliquots were transferred into sterile 50 mL tubes (Greiner Bio-One, Kremsmünster, Austria) and stored at −80 °C until analysis. To avoid contamination, filter handling and processing were carried out under sterile conditions on a clean bench.

Based on the monitored environmental conditions as detailed below, nine aerosol samples were selected for metagenomic sequencing and ice nucleation measurements. A list of the investigated aerosol samples with sampling details is given in Table A1. Note, that the measurements refer to atmospheric particles that were not embedded in or scavenged by raindrops.

Gandolfi et al., 2013; Cao et al., 2014; Archer et al., 2020).
2.2 Meteorological conditions

The meteorological conditions during the sampling campaign in April 2016 were measured at the Cyprus Atmospheric Observatory by the Department of Labour Inspection of Cyprus. The data, comprised of hourly mean values for temperature, relative humidity, rainfall, wind speed, and regulatory mass concentrations for PM$_{10}$ and PM$_{2.5}$, were imported into R version 3.6.3; (R-Development-Core-Team, 2011) and the coordinated universal time (UTC) timestamps were converted to the local eastern European time (EET; UTC+2).

Figure A1 shows the meteorological conditions and aerosol mass concentrations during collection of the aerosol samples. Temperature and relative humidity (RH) exhibited some variation throughout the campaign. A strong increase in RH is noticeable on 9 April 2016, followed by a period of high RH (up to 80%) and a temperature drop until 14 April 2016 corresponding to samples CY19–CY26. Except for this 5 d period, the daily temperature peaks were in the range of 25 to 30 °C. A period of higher RH was again registered starting on 25 April 2016 (CY48).

April 2016 was mainly dry, with only three rain events. One rain event was registered on 12 April during the last 3 h of collection of sample CY20, and two small rain events occurred during collection of CY23 on 13 April. Increased PM concentrations, corresponding to dust events (Schrod et al., 2016), were measured from 9 to 12 April (CY18 and CY19) and from 15 to 18 April (CY31 and CY32).

2.3 DNA extraction

Two filter sample aliquots from opposite sides of each air filter sample were extracted with the ChargeSwitch™ Forensic DNA Purification Kit (Invitrogen, Thermo Fisher Scientific Inc., Waltham, MA, USA) according to the manufacturer’s instructions with the following modifications: incubation at 55 °C for 60 min and elution of DNA with 100 µL elution buffer. Moreover, the DynaMag™-2 magnet instead of MagnaRack™ was used. DNA extracts of the two filter pieces of each sample were pooled and purified with the PowerClean Pro DNA Clean-Up Kit (MO BIO Laboratories, Inc., QIA-GEN, Hilden, Germany) according to the manufacturer’s instructions. To detect potential contamination, four mounting blanks and four start-up blanks were included in the DNA extraction. Additionally, four baked filters were included as filter blanks, and four extractions without any filter served as extraction blanks. DNA concentrations were determined with the DeNovix dsDNA Ultra High Sensitivity Assay (DeNovix Inc., DE, USA) and the Qubit® 3.0 Fluorometer (Thermo Fisher Scientific, USA). DNA was not detected in the mounting, start-up, filter, and extraction blanks, indicating that no contamination occurred during sample handling and extraction.

2.4 Shotgun metagenome sequencing

From each DNA extract, 1 ng of DNA was sequenced on Illumina HiSeq 3000 (output 2 × 150 bp paired-end sequencing reads) and Illumina HiSeq 2500 sequencers (output 2 × 150 bp paired-end sequencing reads) at the Max Planck Genome Centre in Cologne, Germany. Illumina TruSeq adapters were used for the sequencing library. The data output per run was 17.5 GB reads. In total 4.14 × 10^8 reads were obtained from all samples after quality filtering, with an average of 4.6 × 10^7 reads per sample.

2.5 Taxonomic classification

To identify ribosomal small subunit (SSU, 16S and 18S) and large subunit (LSU, 23S and 28S) reads and to assign taxonomy and operational taxonomic units (OTUs), the MGnify pipeline 4.1 with MAPseq (1.2.2) and the SILVA database (SSU and LSU version 132) remapped to an eight-level taxonomy was used (Mitchell et al., 2018; Matias Rodrigues et al., 2017; Quast et al., 2013). In total, 8.41 × 10^5 SSU and 1.09 × 10^6 LSU reads were obtained with averages of 9.34 × 10^4 SSU and 1.21 × 10^5 LSU reads per sample. Reads identified as chloroplast and mitochondrial ribosomal gene fragments were excluded from downstream analysis. Preliminary taxonomic analysis displayed unusually high numbers of SSU reads and inconsistencies between SSU and LSU results for CY26. Analysis of read abundance revealed that, in contrast to the other samples, the data of CY26 contained a high number of reads starting with sequences of degenerated random 16S primers, which are usually used in fingerprinting techniques (Sakallah et al., 1995). Thus, all 16S rRNA reads from sample CY26 were excluded.

All downstream analyses were performed in R version 3.6.3 (R-Development-Core-Team, 2011). The base R functions were used for data processing and plotting. The SSU and LSU files were imported into R using the biomformat package (McMurdie and Paulson, 2019). Additionally, the InterPro matches provided by the MGnify pipeline (Mitchell et al., 2018) were specifically analyzed for viral, allergenic, and bacterial ice-nucleating proteins.

2.6 Freezing experiments and data analysis

For ice nucleation measurements, aqueous extracts of two pieces of each air filter sample were analyzed using the Twin-plate Ice Nucleation Assay (TINA) (Kunert et al., 2018). The extracts were prepared with pure water, which was obtained from a Barnstead™ GenPure™ xCAD Plus water purification system (Thermo Fisher Scientific, Braunschweig, Germany), autoclaved at 121 °C for 20 min, and filtered through a sterile 0.1 µm pore diameter polyethersulfone (PES) vacuum filter unit (VWR International, Radnor, PA, USA). For extract preparation, 20 mL of the pure water was added to each filter piece in sterile 50 mL tubes (SPL Life Sciences
The tubes were vortexed at 2700 rpm for 15 min and another 15 min upside down (Vortex-Genie 2, Scientific Industries, Inc., USA). Afterwards, the aqueous extracts were transferred into new sterile 50 mL tubes. For IN characterization, 1 mL aliquots of the aqueous extracts were treated as follows: (i) filtration through a 5 µm pore diameter filter (Acrodisc, PES, Pall GmbH, Dreieich, Germany), (ii) filtration through a 0.1 µm pore diameter filter (Acrodisc), (iii) incubation at 98 °C for 1 h, (iv) filtration through a 5 µm pore diameter filter and incubation at 98 °C for 1 h, and (v) filtration through a 0.1 µm pore diameter filter and incubation at 98 °C for 1 h.

Of each untreated and treated aqueous extract, 96 aliquots of 3 µL were pipetted by a liquid handling station (epMotion 5073, Eppendorf, Hamburg, Germany) into 384 well plates (Eppendorf). The plates were placed in the TINA instrument, and, after 1 min of equilibration at 0 °C, they were cooled to −28 °C with a continuous cooling rate of 1 °C min⁻¹. Pure-water samples (autoclaved and 0.1 µm filtered) served as negative controls in the individual experiments. Additionally, aqueous extracts of two filter pieces of one filter blank, one mounting blank, and one start-up blank served as negative controls for the aerosol sampling process (Sect. 2.1). As shown in Fig. B1, the blank samples showed freezing events starting around −20 °C with most droplets freezing at around −25 °C. Thus, the observed temperature range for the aerosol samples was 0 to −20 °C.

The freezing temperature data of the individual measurements were binned into 0.2 °C intervals, which corresponds to the temperature uncertainty in the measurement (Kunert et al., 2018). For each aerosol sample, the data of the 96 droplets of the two filter pieces were combined to make 192 droplets for further calculations. All calculations and plotting were performed with R version 3.6.3 (R-Development-Core-Team, 2011). The fraction of frozen droplets (f ice), the cumulative number of IN (N v) per liter of air, and corresponding errors were calculated as described in Vali (1971) and Kunert et al. (2018).

3 Results and discussion

3.1 Overall community composition and community changes

The composition of the airborne community was determined by taxonomic assignment of ribosomal RNA gene SSU (small subunit) and LSU (large subunit) reads. In total, we detected 3000 operational taxonomic units (OTUs) for SSU and 4167 OTUs for LSU reads. Note that the SSU OTUs do not include data from sample CY26 (see Sect. 2.5). The obtained OTUs, however, can be seen as lower estimates of species richness, as MGnify assigns OTUs based on a reference database to different taxonomic levels. This leads to numerous OTUs at higher taxonomic levels, which comprise reads from multiple species that cannot be identified to lower taxonomic levels. Thus, for downstream analysis and intersample comparison, read proportions were used.

Figure 1 shows the read proportions and taxonomic assignments on the domain and kingdom level for the individual samples. The taxonomic composition, in particular the read proportion of fungi and bacteria, changed with sample CY23, indicating a change in the airborne community. Sample CY23 was collected after a 3 h rain event, which corresponds to the collection end of CY20 (Fig. A1). Moreover, two additional rain events occurred during sampling of CY23. This sample shows the highest proportion of fungal reads (> 60 %) and lowest read proportion of Bacteria (< 20 %) and Viridiplantae (< 10 %). Samples collected up to 4 d after the rain events (CY26 and CY31) also show higher proportions of fungal reads (up to 50 %) and lower read proportions for Bacteria and Viridiplantae than the other samples. Metazoa and Archaea read proportions were < 7 % and < 1 %, respectively, in all samples (Fig. 1).

The high proportion of fungal reads in CY23 may result from increased concentrations of fungal spores following rainfall, as many fungal spores are dispersed by rain splash, rain tap and puff, or other spore discharge mechanisms that require water (e.g., Hirst, 1953; Hirst and Stedman, 1963; Gregory et al., 1959; Lacey, 1996; Jones and Harrison, 2004; Crandall and Gilbert, 2017; Kausarud et al., 2005; Kaygusuz and Faruk Çolak, 2017). Bacteria are often associated
with larger particles such as soil dust and plant fragments or form aggregates with each other (Shaffer and Lighthart, 1997; Polymenakou et al., 2008; Reche et al., 2018). As larger particles are more efficiently removed by precipitation than smaller particles, because the latter tend to follow the airstream around the raindrops (Pranesha and Kamra, 1997; Seinfeld and Pandis, 2016), an efficientwashout is assumed for bacteria and larger particles. This facilitates the return of airborne bacteria and other biological particles to the Earth’s surface as well as deposition in new habitats (Peter et al., 2014; Reche et al., 2018), which possibly explains the reduced read proportions for Bacteria, Viridiplantae, and unidentified Eukaryota in samples CY23 and CY26.

Figure 1 also shows, for the individual samples, that the proportions of SSU and LSU reads assigned to Archaea, Bacteria, Fungi, and Metazoa are comparable, indicating that both genes can be used as representative markers for these groups. In contrast, Viridiplantae and unidentified Eukaryota show differences in the SSU and LSU read proportions in samples CY14–CY20 and more equal read proportions in all other samples. This indicates a change in the airborne community due to the rain events and a lack of plant LSU sequences in the SILVA reference database.

Unlike pro- and eukaryotes, for which ribosomal RNA genes can be used for identification, viruses do not contain common genetic markers (Edwards and Rohwer, 2005). Analysis of the InterPro matches, however, revealed the presence of various double- and single-stranded DNA and RNA viruses in the aerosol samples. Of the virus-related InterPro matches ~44% were assigned to viruses that infect bacteria and archaea (e.g., Siphoviridae, Podoviridae, Myoviridae, and Microviridae), which cover ~80% of the reads that were assigned to virus-related matches. Viruses infecting animals and humans (e.g., Adenoviridae, Baculoviridae, Flaviviridae, Herpesviridae, Poxviridae) also exhibited ~44% of the matches corresponding to ~4% of reads. Plant viruses such as Geminiviridae, Caulimoviridae, and Potyviridae were less abundant with ~7% of the matches and <1% of reads. The remaining ~4% of the virus-related InterPro matches (~15% reads) either could not be assigned to hosts or were from fungi- and amoeba-infecting viruses. The detection of RNA viruses (e.g., Flaviviridae, Reoviridae, Retroviridae) can be explained by the integration of these viruses into the genomes of their host cells and their persistence as proviruses (Edwards and Rohwer, 2005).

### 3.1.1 Composition and community changes in Fungi, Viridiplantae, and Metazoa

The composition and community changes in Fungi, Viridiplantae, and Metazoa were further analyzed on the class level. Figure 2 shows that most fungal reads could be attributed to different classes and phyla of Ascomycota and Basidiomycota. Large fractions of Ascomycota reads (up to 40%) could not be attributed to any Ascomycota class. All samples show higher read proportions for Ascomycota than for Basidiomycota. Sample CY23, however, shows, with around 38% for SSUs and 32% for LSUs, the highest Basidiomycota read proportions of all samples. Of all Basidiomycota reads of sample CY23, ~79% (~30% of fungal reads) were assigned to the class Agaricomycetes. The Agaricomycetes read proportion is also enhanced in sample CY26 (~73% of Basidiomycota, ~8.5% of Fungi). The Agaricomycetes are wood-degrading fungi, and many of them form fruiting bodies, which release spores in response to humidity changes (Zoberi, 1964; Elbert et al., 2007; Hassett et al., 2015; Löbs et al., 2020). Analysis of the InterPro matches revealed higher read counts for the allergen Alt a 1 from Alternaria alternata (Dothideomycetes) for samples collected during and after the rainfall period (Table C1). Alternaria is a typical airborne fungus, for which the onset of rain can lead to a period of increased spore concentrations through a tap-and-puff mechanism (Hirst and Stedman, 1963; Lacey, 1996).

For the Viridiplantae (green plants), most reads (up to 92%) could not be classified on the class level (Fig. 3), indicating a lack of reference sequences in the SILVA database. All samples show large proportions of reads assigned to the Streptophyta, which include land plants and green algae. Between ~8% and ~28% of the sample reads were assigned to the class Liliopsida of the Tracheophyta (vascular plants), which comprises lilies, grasses, palms, orchids, and other monocotyledon plants. Samples CY23, CY26, CY31, and CY32 show higher differences in SSU and LSU proportions for unidentified Viridiplantae than the other samples. Although the limited taxonomic classification does not allow specific discussion, this change in proportions suggests a community change in plant-derived aerosol particles, which can be pollen, moss and fern spores, achene fibers, bark, seeds, algae, or fragments like leaf tissues. Pollen, however, is suggested to be the most abundant plant bioaerosol in the atmosphere (Rittenour et al., 2012). The effects of precipitation on pollen concentrations depend primarily on rainfall intensity. While decreased pollen concentrations are associated with more intensive rain events of at least 5 mm h−1 (Ribeiro et al., 2003; Rathnayake et al., 2017; Kluska et al., 2020), increased pollen concentrations have been measured immediately before and during rainfall (Puls and Von Wahl, 1991; Kluska et al., 2020). The pollen emission at the onset of rain can be caused by raindrops hitting the stamina of the flowers which are ready for pollen emission (Puls and Von Wahl, 1991).

As various airborne pollen species are important allergens, the InterPro matches were analyzed for allergenic protein-encoding reads. As detailed in Table C1, 12 possible allergen types were found, all from plants that are common in Cyprus. The most abundant allergens were the Amb allergen type, Lol p 1, Bet v 1 type, and Par j 1 and Par j 2. The Amb allergen type was found in all samples. Amb type allergens are major allergens in the pollen of ragweed (Am-
**Fungi**

Figure 2. Taxonomic composition of Fungi. Taxonomic classification and proportion of SSU and LSU reads of Fungi at the class level.

**Viridiplantae**

Figure 3. Taxonomic composition of Viridiplantae. Taxonomic classification and proportion of SSU and LSU reads of Viridiplantae at the class level.

*brosia artemisiifolia*, Asteraceae) and Cupressaceae trees (e.g., Mediterranean cypress, mountain cedar), which are abundant in Cyprus (Wolf et al., 2017; Gucel et al., 2013). The allergen Lol p 1 was found in all samples, except CY19. Lol p 1 is the major allergen of the perennial ryegrass (*Lolium perenne*, Poaceae), which is common in Cyprus and flowers in springtime (Gucel et al., 2013; Herridge et al., 2021), which was the period of sampling. The Bet v 1 type allergens can be found in members of various plant families such as Betulaceae, Fabaceae, and Rosaceae, which are all common in Cyprus. Samples CY26 and CY48 had 63 and 20 higher read counts, respectively, for Bet v 1 type allergens than all other samples (0–9 reads). The allergens Par j 1 and Par j 2 from *Parietaria judaica* exhibited the highest read counts (154) in CY31. *Parietaria judaica*, a weed of the Urticaceae family and found throughout Cyprus, has a flowering time in spring and autumn and is one of the most important allergens in the Mediterranean area (Colombo et al., 2003; Gucel et al., 2013; Ozturk et al., 2013; Ciprandi et al., 2018). The detected plant allergens allow an additional insight into the origin of the airborne plant particles in the taxonomic analysis of ribosomal reads but show no clear trends with regard to rain.

Metazoa show a diverse picture on the class level with high proportions of up to 60% of unidentified reads as well as differences in the proportions of SSU and LSU reads (Fig. 4). The identified metazoan classes comprised microbial metazoa (e.g., nematodes; Bik, 2019) and larger organisms (e.g., insects, spiders, snails). The analysis of the InterPro matches revealed reads encoding for allergens mainly from wasps (Ves 5 and Ves 5 type) and bees (Api m 3) but also from mites (e.g., Der f 7), nematodes (ABA-1), and mammals (e.g., Fel d 1) as listed in Table C1. All samples show a high proportion of reads assigned to Insecta and Arachnida (includes spiders and mites) and matches to their allergens (Fig. 4, Table C1). Two samples (CY14 and CY19) additionally show larger proportions of nematode reads and matches for nematode allergens. Nematodes are abundant in soil and...
aquatic environments or can be parasites in plants or animals (Seesao et al., 2017). They can be readily dispersed by the wind as living organisms or in resting states, which they go into to survive unfavorable conditions such as dehydration (Nkem et al., 2006; Vanschoenwinkel et al., 2008; Carroll and Viglierchio, 1981; Ptatscheck et al., 2018). The higher read proportions for Gastropoda (snails and slugs) in samples CY26, CY31, and CY48 might be explained by the presence of tissue or skin fragments in the air, due to their dispersal or transport by birds during the nesting phase (Pearce et al., 2010; Shikov and Vinogradov, 2013).

### 3.1.2 Composition and community changes in Bacteria and Archaea

Figure 5 shows the taxonomic composition of Bacteria on the class level, which is a typical composition of bacterial communities in the air (e.g., Tong and Lighthart, 2000; Bowers et al., 2011; Jeon et al., 2011; Bowers et al., 2012; Robertson et al., 2013; Smith et al., 2013; Yoosop et al., 2013; Jang et al., 2018; Mescioglu et al., 2019). Large proportions of reads were assigned to the Actinobacteria and Firmicutes. At a closer look, the Firmicutes were the dominating prokaryotic phylum, except in samples CY23 and CY26. This result agrees well with Mazar et al. (2016) and Mescioglu et al. (2019), who also found high relative abundances of Firmicutes in aerosol samples from the Mediterranean area. Taxa within the Firmicutes might be linked to anthropogenic influences, as they are commonly found in soil and in the human gut microbiome (Kho and Lal, 2018; Almeida et al., 2019). An animal waste treatment company close to the sampling station might be an important local emission source for Firmicutes.

Interestingly, for samples CY23 and CY26, the proportion of unidentified Bacteria reads is 40% to 50% more than twice as high than for all other samples (<20%). This might indicate that, for example, soil bacteria were emitted and aerosolized by the impact of raindrops on soil (Joung et al., 2017). The emitted bacterial taxa, however, were not represented in the reference database and remained unidentified. Overall, the results are consistent with the findings of Zhen et al. (2017), who reported temporary shifts in the airborne bacterial community depending on rain intensity but did not further specify the community structure.

Figure 6 shows that most archaea reads were assigned to the Euryarchaeota, and a small proportion was assigned to the Thaumarchaeota. Similarly to bacteria, high proportions of up to 50% of unidentified reads for samples CY23 and
CY26 might indicate some emission of archaea by the impact of raindrops on soil or water surfaces. Within the Euryarchaeota, Halobacteria reads dominated. Halobacteria belong to the extreme halophilic Archaea that can be found in hypersaline environments (> 10% salt up to saturation) such as salt lakes or saline sediments (Grant and Ross, 1986; Fendrihan et al., 2007; Oren, 2014). The various salt lakes (e.g., Larnaca Salt Lake) in Cyprus might be a source of the detected airborne Halobacteria. Moreover, the detection of methanogens indicates possible anthropogenic influences, as methanogens are found not only in terrestrial and aquatic environments but also in the intestines of larger organisms (Liu and Whitman, 2008; Söllinger and Urich, 2019). Thus, they might stem from the animal waste treatment company in Cyprus, which is close to the station; from wastewater possibly at sewage treatment plants; or from marine pollution by hotels or boats. Moreover, airborne Methanobacteria have been found during fertilization periods (Fröhlich-Nowoisky et al., 2014; Wehking et al., 2018).

### 3.2 Ice nucleation activity

Figure 7 provides an overview of the freezing abilities for the individual samples. Figure 7a shows the fraction of frozen droplets; Fig. 7b shows the median temperatures $T_{50}$ (i.e., the temperature at which 50% of the droplets were frozen); Fig. 7c shows the ice nuclei (IN) concentrations calculated per liter volume of sampled air (from the data shown in Fig. 7a). The individual samples initiated freezing between $-6.3$ and $-11.3$ °C. The frozen fraction curves show differences in shapes and slopes as well as different $T_{50}$ values. The curve of sample CY23, which was collected after and during the rain events, approximates a straight line with $T_{50} = -16.7$ °C. Contrastingly, the samples CY26–CY32, which were collected after the rain events, all show a similar convex curve shape and higher median freezing temperatures ($T_{50}$ is $-13.9$ to $-15.5$ °C). All other samples, including sample CY48, which was collected 13 d after the last rain event, show concave curves and lower $T_{50}$ values ($-17.3$ to $-18.3$ °C).

Figure 7c shows that the individual samples differ in their IN concentrations at given temperatures. For example, at $-15$ °C, values between $10^{-1}$ and $10^{-2}$ IN per liter of sampled air were found for the investigated samples. Overall, the obtained IN concentrations span about 2 orders of magnitude for temperatures $>-15$ °C and about 1 order of magnitude for temperatures $<-15$ °C. These results are in good agreement with Gong et al. (2019), who reported IN concentrations on the same order of magnitude for samples collected in Cyprus in April 2017.

The results indicate heterogeneity of the IN compositions in the samples. Higher initial freezing temperatures and $T_{50}$ values can be an indication of the presence of bio-IN, as many bio-IN are known to nucleate at temperatures

---

**Figure 5.** Taxonomic composition of Bacteria. Taxonomic classification and proportion of SSU and LSU reads of Bacteria at the class level. SSU reads of CY26 were excluded because of contamination by 16S rRNA gene amplicon sequences.
Figure 6. Taxonomic composition of Archaea. Taxonomic classification and proportion of SSU and LSU reads of Archaea at the class level.

Figure 7. Ice nucleation activity. Fraction of frozen droplets ($f_{\text{fro}}$) (a) and cumulative number of IN ($N_{\text{v}}$) per liter of air (b) vs. temperature ($T$) for all aerosol samples. The colors represent individual samples; the error bars were calculated using the counting error and the Gaussian error propagation.

To further characterize the IN of the different aerosol samples, filtration and heat treatment experiments were performed with the aqueous extracts. Figure 8 shows that filtration through a 5 µm and a 0.1 µm filter reduced the ice nucleation activity in all samples, indicating that all samples contained IN larger than 5 µm, IN between 0.1 and 5 µm, and IN smaller than 0.1 µm. For samples CY14, CY23, CY26, CY31, and CY48, however, the results show only a small fraction of IN between 0.1 and 5 µm while many IN were smaller than 0.1 µm. Of all samples, CY18 and CY19 show the lowest percentage of frozen droplets after 0.1 µm filtration and, thus, the lowest numbers of IN smaller than 0.1 µm in the observed temperature range.

Heat sensitivity can indicate biological IN, as heat can reduce ice nucleation activity of many bio-IN (Pouleur et al., 1992; Henderson-Begg et al., 2009; Morris et al., 2013; Fröhlich-Nowoisky et al., 2015; Pummer et al., 2015; Hara et al., 2016a; Kunert et al., 2019). Certain bio-IN, however, such as Lysinibacillus, a heat-resistant isolate of Xanthomonas, and various plant pollen, as well as inorganic particles are heat resistant to temperatures $>90^\circ$C (Pummer et al., 2012; Hill et al., 2014; Failor et al., 2017). Heat treatment at 98$^\circ$C for 1 h reduced the ice nucleation activity of the individual samples to a different extent (Fig. 8). For example, the ice nucleation activity of samples CY20–CY32 was significantly reduced in the different size fractions after
heat treatment (Fig. 8d–h), indicating the presence of heat-sensitive IN and thus presumably bio-IN of different sizes. Many fungal spores and bacterial aggregates are larger than 5 µm, whereas single bacterial cells or cell fragments can be smaller than 5 µm, and ice-nucleating macromolecules are typically in the submicrometer range (Fröhlich-Nowoisky et al., 2016; Pummer et al., 2012, 2015; Šantl-Temkiv et al., 2015; Felgitsch et al., 2018). The samples CY23 and CY26, which were collected during and after the rain events, appear to have the largest fraction of heat-sensitive IN larger than 5 µm. In contrast, heat only slightly affected samples CY18 and CY19 (Fig. 8b–c), indicating a significant fraction of heat-resistant IN in these samples, possibly due to particles from the dust events which were registered during the collection of these samples (Fig. A1). For samples CY14 and CY48, the ice nucleation activity of 5 and 0.1 µm filtrates was reduced by heat, while the unfiltered extract was only slightly affected (Fig. 8a, i). The results suggest that, in these samples, many heat-resistant IN were larger than 5 µm, whereas many heat-sensitive IN were smaller than 0.1 µm or, for CY14, also between 0.1 and 5 µm.

3.3 Possible sources of biological ice nuclei

The results of the freezing experiments indicate the presence of bio-IN in different size ranges. These bio-IN may be present as living or dead cells, fungal hyphae and spores, pollen, cell fragments, or detached macromolecules or associated with plant particles or soil organic matter (Schnell and Vali, 1976; Pummer et al., 2012; Fröhlich-Nowoisky et al., 2015; Šantl-Temkiv et al., 2015; O’Sullivan et al., 2016; Hill et al., 2016; Conen and Yakutin, 2018). Note that not all of these IN contain DNA, for example ice-nucleating macromolecules or cell membrane fragments with attached IN proteins, and they are thus not all covered by DNA analysis. The largest fraction of bio-IN > 5 µm was detected in CY23 and CY26. For these samples, the sequencing results show a higher proportion of reads assigned to fungi, in particular Agaricomycetes (Figs. 1, 2), suggesting a possible contribution of these fungi to the increase in IN. Systematic surveys suggest that fungal ice nucleation activity, in particular at temperatures > −15 ºC, is restricted to a limited number of fungal species (Fröhlich-Nowoisky et al., 2015; Haga et al., 2013, 2014; Iannone et al., 2011; Jayaweera and Flanagan, 1982; Kieft and Ahmadjian, 1989; Kunert et al., 2019; Morris et al., 2013; Pouleur et al., 1992; Pummer et al., 2013, 2015). The class Agaricomycetes contains approximately 21 000 fungal species (Hibbett et al., 2014), but only a handful has been tested for ice nucleation activity so far (Pummer et al., 2013; Haga et al., 2014). Further surveys of ice nucleation properties of Agaricomycetes and other fungi are necessary.

To identify possible bio-IN sources, the SSU and LSU data were analyzed for known IN fungi and bacteria that are active at temperatures > −15 ºC. In total, 4 fungal genera and 13 bacterial genera that contain IN-active species were detected, but all exhibited read proportions < 1 % and species identification was partly not possible. Note that not all species and strains of the identified genera are IN-active, and, especially in bacteria, not all cells express the IN gene (Lindow et al., 1982; Kunert et al., 2019). Moreover, some recently isolated IN-active fungi and bacteria such as Cryptococcus sp. and Brevibacterium sp. are not identified to the species level (Beall et al., 2021). Thus, the following microorganisms represent possible sources of IN but provide no proof of to what extent they were IN-active in the air.

Among the fungi were Cryptococcus sp., Fusarium sp., the soil fungus Mortierella alpina, and five species of the plant-pathogen rust fungi Puccinia (P. arachidis, P. hordei, P. horiana, P. striiformis, P. triticina). Spores of Puccinia striiformis (stripe rust or yellow rust) and P. triticinae have initial freezing temperatures of −7 and −9 ºC, respectively (Morris et al., 2013). Although several other rust species are known as ice nucleators, for P. arachidis (peanut rust), P. hordei (barley leaf rust), and P. horiana (chrysanthemum white rust), ice nucleation activity has not yet been reported. In contrast to Puccinia spp., for which the spores are IN-active, the IN of M. alpina and Fusarium spp. are cell-free proteinaceous macromolecules and can trigger freezing at temperatures as high as −5 ºC or −1 ºC, respectively (Pouleur et al., 1992; Richard et al., 1996; Fröhlich-Nowoisky et al., 2015; Kunert et al., 2019). The macromolecules might be released into the surrounding environment and contribute to atmospheric IN.

Among the potential IN-active bacteria were the well-known genera and species Erwinia, Pantoea agglomerans, Pseudomonas, and Xanthomonas as well as the recently identified Brevibacterium, Idiomarina, Lysinibacillus, Planococcus maritimus, and Psychrobacter (Maki et al., 1974; Govindarajan and Lindow, 1988; Hill et al., 2014; Failor et al., 2017; Beall et al., 2021). The genus Pseudomonas, which includes a range of IN-active species, was detected in all samples. Most of the reads (~94 %) assigned to the genus Pseudomonas, however, could not be identified to the species level. The remaining 58 reads were distributed over 12 species (1 to 14 reads per species) but did not include P. syringae, a common bacterial ice nucleator. Analysis of the InterPro matches for IN-active proteins, however, revealed one InterPro match (IPR000258, bacterial ice nucleation, octamer repeat) for in total 24 reads from six of the nine samples (1 to 12 reads per sample), indicating the presence of a bacterial IN-active species. Recently, ice nucleation activity was found in Halococcus morrhuae and Haloferax sulfurifontis, two species of Haloarchaea, which initiated the freezing of water droplets at temperatures as high as −18 ºC (Creamean et al., 2021). In our data set, however, only one read from CY48 was assigned to the genus Halococcus and Haloferax was not detected.
Figure 8. Effects of filtration and heat treatment on ice nucleation activity. Fraction of frozen droplets ($f_{\text{ice}}$) vs. temperature ($T$) for all aerosol samples (a–i). The color represents individual treatments; the error bars were calculated using the counting error and the Gaussian error propagation.

4 Conclusions

Our results show that rainfall in a rural dryland can lead to short-term changes in bioaerosol and IN composition. Particle washout and fungal spore discharge in response to humidity changes appear to play a major role in the observed composition changes, in particular for fungi and bacteria. The results of the freezing experiments suggest a rain-related enhancement of bio-IN $> 5$ and $< 0.1 \mu m$. Microorganisms that are known to be IN-active, however, could have only rarely been detected. This may reflect unknown sources of atmospheric bio-IN and the presence of cell fragments or cell-free IN macromolecules that do not contain DNA, as is known to be the case for many IN-active fungi. Similar observations of increased bioaerosol, fungal spore, and IN concentrations in relation to rainfall have been reported for different ecosystems, indicating a strong coupling of rainfall and particle emission (Huffman et al., 2013; Prenni et al., 2013; Schumacher et al., 2013; Tobo et al., 2013; Wright et al., 2014; Bigg et al., 2015; Heo et al., 2014; Gosselin et al., 2016; Rathnayake et al., 2017). The release of bioaerosols during and after rain may play an important role in the spread and reproduction of microorganisms and may contribute to the atmospheric transmission of pathogenic and allergenic particles. In view of the relevance of bioparticles as pathogens and allergens as well as nuclei for clouds and precipitation (bioprecipitation cycle) (Bigg et al., 2015; Morris et al., 2014, 2017) and potential future developments in the Anthropocene (Crutzen and Stoermer, 2000; Pöschl and Shiraiwa, 2015; Rodriguez-Caballero et al., 2018), further investigations are required to understand the sources, properties, and short-term dynamics of bioaerosols and bio-IN in various ecosystems. The bioaerosol communities need to be compared with the communities of cloud water and precipitation samples, as differences in the community compositions can provide insights into potential selection processes of specific bioaerosols and thus may provide hints to potential but previously unknown IN-active microorganisms (Els et al., 2019). Additionally, further investigations are necessary for the identification of the IN macromolecules and for the detection and quantification of these IN in soil, precipitation, and atmospheric samples.
Appendix A: Aerosol samples and meteorological conditions

Table A1. Overview of air filter samples. Sample names, sampling periods, and sampled air volumes at standard conditions (0°C, 1013 hPa).

<table>
<thead>
<tr>
<th>Sample</th>
<th>Start of sampling</th>
<th>End of sampling</th>
<th>Sampling time</th>
<th>Sampled air volume</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Date</td>
<td>Time [LT]</td>
<td>Date</td>
<td>Time [LT]</td>
</tr>
<tr>
<td>CY14</td>
<td>7 April 2016</td>
<td>09:41</td>
<td>8 April 2016</td>
<td>09:41</td>
</tr>
<tr>
<td>CY18</td>
<td>9 April 2016</td>
<td>09:52</td>
<td>10 April 2016</td>
<td>09:52</td>
</tr>
<tr>
<td>CY19</td>
<td>10 April 2016</td>
<td>09:54</td>
<td>11 April 2016</td>
<td>09:52</td>
</tr>
<tr>
<td>CY20</td>
<td>11 April 2016</td>
<td>09:54</td>
<td>12 April 2016</td>
<td>09:52</td>
</tr>
<tr>
<td>CY23</td>
<td>12 April 2016</td>
<td>10:02</td>
<td>13 April 2016</td>
<td>09:56</td>
</tr>
<tr>
<td>CY26</td>
<td>13 April 2016</td>
<td>10:01</td>
<td>14 April 2016</td>
<td>10:01</td>
</tr>
<tr>
<td>CY31</td>
<td>16 April 2016</td>
<td>10:07</td>
<td>17 April 2016</td>
<td>10:07</td>
</tr>
<tr>
<td>CY32</td>
<td>17 April 2016</td>
<td>10:07</td>
<td>18 April 2016</td>
<td>10:07</td>
</tr>
<tr>
<td>CY48</td>
<td>25 April 2016</td>
<td>10:01</td>
<td>26 April 2016</td>
<td>10:01</td>
</tr>
</tbody>
</table>

Figure A1. Meteorological conditions and aerosol mass concentrations: 1 h averages of temperature (T), relative humidity (RH), rainfall (Rain), wind speed (WS), and aerosol mass concentrations of PM\(_{10}\) and PM\(_{2.5}\) during the sampling campaign.
Appendix B: Freezing experiments with pure water and sampling blanks

Figure B1. Negative controls of the freezing experiments. Fraction of frozen droplets ($f_{\text{ice}}$) vs. temperature ($T$) for all water samples (a) and different types of sampling blanks that served as negative controls (b). For each type of sampling blank, two filter pieces were measured. The error bars were calculated using the counting error and the Gaussian error propagation.
Appendix C: InterPro matches and read counts for allergenic proteins

Table C1. Protein-encoding reads for allergens. Overview of InterPro matches and read counts for allergenic protein-encoding reads. IPR indicates the InterPro accession number.

<table>
<thead>
<tr>
<th>IPR</th>
<th>Allergens</th>
<th>Species/family</th>
<th>CY14</th>
<th>CY18</th>
<th>CY19</th>
<th>CY20</th>
<th>CY26</th>
<th>CY31</th>
<th>CY32</th>
<th>CY48</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fungi</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IPR032382</td>
<td>Alt a 1</td>
<td>Alternaria alternata</td>
<td>0</td>
<td>8</td>
<td>0</td>
<td>15</td>
<td>23</td>
<td>23</td>
<td>16</td>
<td>7</td>
</tr>
<tr>
<td>Viridiplantae</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IPR005611</td>
<td>Amb V</td>
<td>Ambrosia sp.</td>
<td>4</td>
<td>9</td>
<td>0</td>
<td>3</td>
<td>4</td>
<td>3</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>IPR018082</td>
<td>Amb</td>
<td>Asteraceae, Cupressaceae</td>
<td>30</td>
<td>56</td>
<td>31</td>
<td>31</td>
<td>22</td>
<td>86</td>
<td>31</td>
<td>11</td>
</tr>
<tr>
<td>IPR024949</td>
<td>Bet v 1 type</td>
<td>Betulaceae, Apiaceae, Rosaceae, Asparagaceae, Fabaceae, Solanaceae</td>
<td>0</td>
<td>9</td>
<td>4</td>
<td>9</td>
<td>8</td>
<td>63</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>IPR005795</td>
<td>Lol p 1</td>
<td>Lolium perenne</td>
<td>5</td>
<td>17</td>
<td>0</td>
<td>24</td>
<td>2</td>
<td>24</td>
<td>12</td>
<td>10</td>
</tr>
<tr>
<td>IPR005453</td>
<td>Lol p 2</td>
<td>Lolium perenne</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>IPR006040</td>
<td>Ole e 1</td>
<td>Olea europaea</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>IPR035506</td>
<td>Pollen allergen/Os</td>
<td>Oryza sativa</td>
<td>1</td>
<td>4</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>IPR000528</td>
<td>Par j 1, Par j 2</td>
<td>Parietaria judaica</td>
<td>11</td>
<td>2</td>
<td>5</td>
<td>0</td>
<td>14</td>
<td>20</td>
<td>154</td>
<td>0</td>
</tr>
<tr>
<td>IPR002411</td>
<td>Amyl inhib</td>
<td>Poaceae</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>IPR002914</td>
<td>Lol p 5, Pha a 5, Poa p 9, Phl p 5b, Phl p 6</td>
<td>Poaceae</td>
<td>0</td>
<td>3</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>4</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>IPR006106</td>
<td>Soft/Tryp amyl inhib CS</td>
<td>Poaceae</td>
<td>0</td>
<td>14</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>IPR006105</td>
<td>Tryp amyl inhib CS</td>
<td>Poaceae</td>
<td>0</td>
<td>15</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Metazoa</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IPR032487</td>
<td>ABA-1, DVA-1</td>
<td>Ascarididae, Dictyocaulidae</td>
<td>4</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>IPR002116</td>
<td>Api m 3</td>
<td>Apis mellifera</td>
<td>0</td>
<td>0</td>
<td>19</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>21</td>
</tr>
<tr>
<td>IPR002450</td>
<td>Bos d 2, Can f 2</td>
<td>Bovidae, Canidae</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>IPR020234</td>
<td>Der f 7</td>
<td>Dermatophagoides furiniae</td>
<td>10</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>12</td>
</tr>
<tr>
<td>IPR020306</td>
<td>Der p 5, Blo t 5, Blo t 21</td>
<td>Dermatophagoides pteronyssinus, Blomia tropicalis</td>
<td>0</td>
<td>17</td>
<td>0</td>
<td>3</td>
<td>0</td>
<td>6</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>IPR006178</td>
<td>Fel d 1 (chain 1)</td>
<td>Felis silvestris</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>IPR015332</td>
<td>Fel d 1 (chain 2)</td>
<td>Felis silvestris</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>IPR010629</td>
<td>NSP</td>
<td>Pieris rapae</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>6</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>IPR018244</td>
<td>Ves 5 (CS)</td>
<td>Vespidae</td>
<td>34</td>
<td>149</td>
<td>138</td>
<td>72</td>
<td>221</td>
<td>188</td>
<td>387</td>
<td>80</td>
</tr>
<tr>
<td>IPR001283</td>
<td>Ves 5</td>
<td>Vespidae</td>
<td>48</td>
<td>143</td>
<td>59</td>
<td>48</td>
<td>159</td>
<td>175</td>
<td>377</td>
<td>60</td>
</tr>
<tr>
<td>IPR002413</td>
<td>Ves 5 type</td>
<td>Vespidae, Formicidae</td>
<td>5</td>
<td>21</td>
<td>9</td>
<td>2</td>
<td>27</td>
<td>48</td>
<td>91</td>
<td>2</td>
</tr>
</tbody>
</table>
Data availability. The metagenomic data have been deposited at https://www.ebi.ac.uk/metagenomics/ (Yordanova, 2021) under MGYA00166510 (CY14), MGYA00199054 (CY18), MGYA00166484 (CY19), MGYA00166504 (CY20), MGYA00166527 (CY23), MGYA00166526 (CY26), MGYA00166493 (CY31), MGYA00199055 (CY32), and MGYA00166518 (CY48). The IN data are available at Edmond – the Open Access Data Repository of the Max Planck Society, under https://doi.org/10.17617/3.6r (Tang, 2021).

Author contributions. JFN, UP, and BW designed the study. PY performed the sample collection and DNA extraction. KT performed the freezing experiments. BSP, DP, JW, JFN, KT, and PY analyzed the taxonomic data. BSP and JFN analyzed the InterPro matches. KT and ATB analyzed the IN data. All authors discussed the results. JFN wrote the manuscript with contributions from all co-authors.

Competing interests. The contact author has declared that neither they nor their co-authors have any competing interests.

Disclaimer. Publisher’s note: Copernicus Publications remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Acknowledgements. We thank the entire INUIT-BACCHUS-ACTRIS campaign team for their co-operation and support; the Cyprus Department of Labour of Inspection for the provision of meteorological and PM data; Nina-Maria Kropf, Naama Lang-Yona, and Steven Lelieveld for technical assistance; Nadine Bothen, J. Alex Huffman, and Mira L. Pöhlker for helpful discussion.

Financial support. This research has been supported by the Max Planck Society (MPG), the Horizon 2020 ACTRIS project (grant agreement no. 654109) and its provision of transnational access funds, and the Deutsche Forschungsgemeinschaft (DFG, FR3641/1-2, FOR 1525 INUIT).

The article processing charges for this open-access publication were covered by the Max Planck Society.

Review statement. This paper was edited by Koji Suzuki and reviewed by Alexandr Safatov and one anonymous referee.

References


Morrיס, C. E., Sands, D. C., Glaux, C., Samsatly, J., Asaad, S., Moukahel, A. R., Gonçalves, F. L. T., and Bigg, E. K.: Urediospores of rust fungi are ice nucleation active at −10°C and...


Yordanova, P.: Metagenomic analysis of the atmospheric microbial composition over Cyprus, Mgnify [data set], available at: https://www.ebi.ac.uk/metagenomics/, Study MGYS00001699, last access: 28 December 2021.
