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Investigation of bacterial and fungal communities in indoor and outdoor air of elementary school classrooms by 16S rRNA gene and ITS region sequencing

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Abstract

The advent of high-throughput sequencing methods allowed researchers to fully characterize microbial community in environmental samples, which is crucial to better understand their health effects upon exposures. In our study, we investigated bacterial and fungal community in indoor and outdoor air of nine classrooms in three elementary schools in Seoul, Korea. The extracted bacterial 16S rRNA gene and fungal ITS regions were sequenced, and their taxa were identified. Quantitative polymerase chain reaction for total bacteria DNA was also performed. The bacterial community was richer in outdoor air than classroom air, whereas fungal diversity was similar indoors and outdoors. Bacteria such as *Enhydrobacter*, *Micrococcus*, and *Staphylococcus* that are generally found in human skin, mucous membrane, and intestine were found in great abundance. For fungi, *Cladosporium*, *Clitocybe*, and *Daedaleopsis* were the most abundant genera in classroom air and mostly related to outdoor plants. Bacterial community composition in classroom air was similar among all classrooms but differed from that in outdoor air. However, indoor and outdoor fungal community compositions were similar for the same school but different among schools. Our study indicated the main source of airborne bacteria in classrooms was likely human occupants; however, classroom airborne fungi most likely originated from outdoors.

KEYWORDS

classroom air, elementary school classrooms, microbial communities, next-generation sequencing

Practical Implications

- Bacterial community composition in classroom air was similar among all classrooms but different from outdoor air.
- Indoor and outdoor fungal community compositions were similar for the same school but different among the schools.
- Our study indicated that the main source of airborne bacteria in classrooms was likely human occupants whereas most fungi in classroom air likely originated from outdoors.

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- Our finding that the amount of total bacterial DNA in outdoor air was higher than indoors also underscores the importance of controlling infiltration of outdoor microbes into classrooms.

1 | INTRODUCTION

Most people in modern society spend more than 80% of their time indoors.^{1,2} Therefore, indoor environmental quality has a significant effect on health and quality of life, especially for groups disproportionately affected such as children, adolescents, the elderly, and people suffering from respiratory diseases.³ Children may also be exposed to airborne pollutants more than adults because their respiration rate per unit weight is higher.⁴ In addition, the high activity level of students in elementary schools may also contribute to increase classroom airborne concentrations of particulate matter (PM) through resuspension of settled surface dusts.⁵

Indoor PM consists of biological particles, organic compounds adsorbed onto particles, transition metals, ions, reactive gases, particle core of carbonaceous material, and minerals.^{6,7} Exposures to indoor bacteria and fungi have been associated with the occurrence of infectious diseases, and exacerbation of asthma and allergies.⁸ While reducing exposures to pathogens and bacterial toxins in the indoor environments is important,⁹ exposures to human-associated indoor microbiomes may also provide beneficial effects on occupants' health.^{10–12} Hygiene hypothesis suggests that infants need to get sufficiently exposed to symbiotic microorganisms to properly develop the immune system that will prevent development of allergic diseases.¹³ Thus, exposure to indoor microbes might have both beneficial and adverse effects in occupants.¹⁴

It has been challenging to study the microbial community with the traditional culture method because most microorganisms existing in the environment are not easily cultured with selected media.^{15–18} Nonetheless, the microbial management standard for indoor air in Republic of Korea is based on the colony-forming units using the culture method that is likely to neglect some important indoor microorganisms. As metagenome analysis using next-generation sequencing becomes more readily available, studying microbial communities existing in the environment draws more attention from researchers.¹⁹ Nevertheless, only a limited number of studies exist that characterize microbial communities in school classroom air and examine their associations with classroom environments. Furthermore, the management of microbial community in indoor air in multi-user facilities has become a major issue during the pandemic of coronavirus disease 2019.²⁰ In this study, we selected three elementary schools located near busy roads with high vehicular traffic in Seoul, Republic of Korea to: (1) characterize microbial communities in classroom air; and (2) compare microbiomes inside and outside classrooms by bacterial 16S rRNA gene and fungal ITS (internal transcribed spacer) region sequencing.

2 | MATERIALS AND METHOD

2.1 | Site selection and sample collection

Three elementary schools adjacent to major roads with an average daily traffic volume of over 100 000 vehicles in Seoul, Korea, were selected.²¹ The schools were, on average, 226 ± 49 m apart from the main roads. Air samples were collected from nine classrooms (one per classroom and two to four classrooms per school) with natural ventilation made through opening interior windows between the classroom and the corridor during October through December 2019. We selected 1st, 2nd, and 3rd graders' classrooms considering their higher classroom activity levels compared to the higher grades.

Indoor and outdoor air filter samples for total suspended particulate matter were simultaneously collected during school hours on weekdays and during the absence of students on holidays or weekends. Three indoor sampling pumps (Sensidyne) were placed on the lockers at the back of the classroom at a height of 130 cm, similar to the average height of elementary school students in Korea,²² and two outdoor sampling pumps at the school playground (Figure S1). The sampling pumps connected to 37-mm closed cassettes loaded with polyvinyl chloride filters (pore size: 5 μ m, SKC, Inc.) were operated at 2.5 L/min indoors and 4 L/min outdoors from 8:00 a.m to 3:30 p.m. To secure enough genomic DNA in samples for the 16S rRNA gene and ITS region sequencing, three indoor filters were combined for each classroom and two outdoor filters for the corresponding outdoor sample. All equipment, materials, and supplies were sterilized before use. After completion of sampling, the filters were placed in a sterile filter container and sealed with parafilm using a sterile technique. The sealed filters were then placed in a cooler with ice packs to maintain cool temperature and minimize potential changes in the microbial community during transportation to the laboratory where all filter samples were stored at -80°C until the analysis. Environmental conditions of the classrooms were recorded on the day when the air samples were collected and they included the number of students, temperature, relative humidity, natural ventilation time, and air cleaner operation time. Outdoor wind speed, precipitation, temperature, and relative humidity on the sampling day were also recorded.

2.2 | Genomic DNA extraction

Genomic DNA (gDNA) in sampled filters was extracted using High Pure PCR Template Preparation Kit (Roche). The filter was put in a 2 mL screw-cap microcentrifuge tube along with 300 mg of 212–300 μ m sterilized glass beads (Sigma) and 650 μ L of ultrapure

water. It was pulverized twice for 30 s at 1484 g using a homogenizer (Allsheng).^{23,24} After centrifugation at 20 000 g for 1 min, the supernatant was collected and centrifuged again at 20 000 g for 1 min. A 400 µL aliquot of the supernatant was collected and used as a pre-treatment sample. Then, the sample was eluted from the column as recommended by the kit manufacturer and stored at -20°C.

2.3 | Microbial diversity analysis

Each sample was prepared according to the Illumina 16S Metagenomic Sequencing Library protocols. The amount and purity of DNA were measured using the QuantiFluor dsDNA System (Promega) and VICTOR Nivo (PerkinElmer). The samples that did not meet minimum cDNA concentration (10 ng/µL) were excluded from the 16S rRNA gene and ITS region sequencing (Table S1). ITS3-ITS4 region of fungal DNA²⁵ and 16S V3-V4 region of bacterial DNA²⁶ were used for gene amplification with the primer sequences shown in Table 1. The amplification step was performed to add multiplexing indices and Illumina sequencing adapters. The final products were normalized and pooled using the PicoGreen, and the size of libraries was verified using the TapeStation DNA screentape D1000 (Agilent). And then sequencing was performed using the Illumina MiSeq™ platform (Illumina).

Sequence analysis was performed using Quantitative Insights Into Microbial Ecology2 (QIIME2, v.2017.6.0), such as Caporaso et al.²⁷ At the Demultiplex phase, each sequence was divided into samples using DADA2 (ver.1.1.1) and quality checked to eliminate low quality.²⁸ The subsequent process also utilized DADA2 to filter out noisy sequences, correct errors in ambient sequences, and correct amplicon errors through the process of removing chimeric sequences and singletons. The clustering process used the q2-vsearch (ver.2019.7) tool and proceeded with an open reference OTU picking method. Sequences with similarity greater than 97% were clustered into Operational Taxonomic Units (OTUs), and OTUs were analyzed through SILVA and UNITE.^{29,30}

2.4 | Quantitative analysis of bacteria

For the quantitative analysis of the total bacterial DNA, qPCR (quantitative polymerase chain reaction) was performed by using 518F and

800R (Table 1) among 16S rRNA region.³¹ The reaction mixture was prepared with 4 µL of 5 × HOT FIREPol EvaGreen qPCR Supermix (Solis BioDyne), 14.6 µL of Nuclease-Free Water (QIAGEN), and forward and reverse primer that were mixed with 0.2 and 1 µL of DNA template, respectively. Initial denaturation was carried out at 95°C for 12 min, and 40 cycles of three-step process were carried out—denaturation at 95°C for 15 s, annealing at 55°C for 20 s, and elongation at 72°C for 30 s.

For qPCR, a LineGene 9600 (BIOER) was used. Using the qPCR absolute quantification method,³² qPCR was performed with the diluted *Escherichia coli* (*E. coli*) sample. PCR efficiency ($R^2 = 0.993$; PCR efficiency = 102.2%) was verified through a standard curve, and specificity was confirmed through a melting curve. The amount of bacterial cDNA in sample was calculated according to the cycle threshold value (C_t value) of the diluted *E. coli* sample. To calculate the copy number in the bacterial sample, the C_t value of the diluted *E. coli* and the C_t value of the sample were compared. The OD600 value of the undiluted standard solution of *E. coli* sample used in the experiment was 0.7, and the corresponding DNA concentration measured after DNA extraction was 55.0 ng/µL. The *E. coli* standard sample solution was serially diluted to 10^{-5} times by a factor of 10.

2.5 | Statistical analysis

Alpha and beta diversity were calculated and visualized using q2-diversity (ver.2019.7) in the QIIME2 plugin. Unidentified genera were excluded from the analysis. A rarefaction curve was created for each sample to estimate bacterial and fungal richness as a function of sequencing depth at genus level.³³ To explore the abundance and uniformity of bacterial and fungal taxa in the samples, a Shannon-Wiener index (HI), which is a diversity index considering both abundance and diversity, and Pielou's evenness index (EI), which represents the evenness of a community, were calculated. Using principal coordinates analysis (PCoA), we evaluated similarity of the microbial community among schools and between different sample types (inside vs. outside classroom) with Bray-Curtis dissimilarity index. Spearman's correlation coefficients among the diversity indices and environmental factors were calculated by sample location (indoors and outdoors). For bacteria, the DNA copy numbers from the qPCR analysis were also included in the correlation analysis. All statistical analyses were conducted in R version 4.0 (CRAN)

TABLE 1 The primer sequences used for gene sequencing and qPCR experiments

	Primer		Sequence
Bacteria	V3-V4	Forward	5' CCTACGGGNGGCWGCAG 3'
		Reverse	5' GACTACHVGGGTATCTAATCC 3'
	518F	Forward	5' CCAGCAGCCGCGTAATACG 3'
		Reverse	5' TACCAGGTATCTAATCC 3'
Fungi	ITS3-ITS4	Forward	5' TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGGTCATGATGAAGAACGCAGC 3'
		Reverse	5' GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGTCTCCGCTATTGATATGC 3'

and statistical significance set at $p < 0.05$ and marginal significance at $p < 0.1$.

3 | RESULTS AND DISCUSSION

3.1 | Classroom characteristics and outdoor weather conditions

All nine classrooms in three schools had wood flooring, and the number of students per occupied classroom ranged from 21 to 26 with an average of 23. Classroom average temperature was $22.6 \pm 1.6^\circ\text{C}$, and the average relative humidity was $42.2 \pm 3.2\%$ (Table 2). The average time of natural ventilation made through opening interior windows between the classroom and the corridor was 47.8 ± 80.5 min/day, while the average operating time of a classroom air purifier was 151.7 ± 142.5 min/day. Outdoor temperature during the sampling campaign varied widely from -0.5 (in December) to 17.0°C (in October), but was significantly lower than classroom temperature ($p < 0.01$). The outdoor average relative humidity was $62.5 \pm 11.2\%$ and significantly higher than indoors ($p < 0.01$). Average wind speed during outdoor sampling was 7.7 ± 2.5 m/s. It rained during the sampling period for classrooms 1, 3, and 4 in school "b," and the average precipitation was 3.3 ± 6.7 mm.

3.2 | Qualitative analysis of microbial community in classroom and outdoor air

Richness of bacterial and fungal samples is presented using the rarefaction curves (Figure 1). As rarefaction curve reached the asymptote (represented by a straight line with a zero slope), sufficient sequencing depth was confirmed. For bacteria, the indoor average OTUs were 240.3 ± 70.7 , which were lower than outdoors (310.9 ± 81.4). For fungi, the average OTUs of classroom and outdoor samples were similar— 120.7 ± 27.1 and 122.2 ± 37.7 , respectively. This indicated that bacteria in outdoor air samples were more diverse and more variable in diversity than classroom air, while fungal diversity was similar indoors and outdoors (similar in average richness) but more variable in diversity in outdoor air samples. The number of bacterial OTUs was significantly ($p < 0.01$) higher than fungal OTUs, indicating that bacterial taxa in both indoor and outdoor air were more diverse than fungal taxa. This result was similar to the studies that found more diverse bacteria than fungi in soil, sugarcane (roots, shoots, and leaves), and floor dust of homes.^{34–36}

Diversity and evenness (uniformity) of bacteria and fungi in the classroom and outdoor air is presented in Figure 2. Shannon-Wiener index (HI) of classroom airborne bacteria was lower (marginally significant, $p = 0.09$) than outdoors, but Pielou's evenness indices (EI) indoors and outdoors were similar while HI for fungi was not different between indoors and outdoors even if significance level was set at 0.1 although fungal taxa were more evenly distributed in classroom air than outdoor air ($p = 0.09$). Kembel et al³⁷ reported that diversity in the outdoor bacterial community was significantly greater

than indoors, which was similar to our study finding. Comparison of bacterial and fungal diversities in our study indicated that both HI and EI for bacteria were significantly ($p < 0.01$) higher than fungi, which is consistent with the findings of Kettleson et al³⁴ report.

The top 30 bacterial and fungal genera in relative abundance are illustrated in the stacked bar graphs for the classroom and outdoor air samples (Figure 3). The average cumulative relative abundance of the top 15 genera accounted for $35.6 \pm 9.0\%$ of total abundance for classroom bacterial samples, $22.3 \pm 7.0\%$ for outdoor bacterial samples, $56.3 \pm 7.8\%$ for classroom fungal samples, and $62.7 \pm 8.0\%$ for outdoor fungal samples (Figure S2). For bacteria, *Enhydrobacter*, *Streptococcus*, and *Micrococcus* were the most abundant top three genera in classroom air while *Methylobacterium*, *Pseudomonas*, and *Deinococcus* were the most abundant top three genera in outdoor air (Figure 3A). The literature showed the main source of *Enhydrobacter*, *Micrococcus*, *Staphylococcus*, *Streptococcus*, *Neisseria*, and *Haemophilus* in indoor environments that were also found within the top 22 genera in our study was human occupants. These bacterial genera live in the human skin, mucous membrane, and the intestine.^{38–43} *Methylobacterium*, a genus living in soil and water and existing widely in the environment,⁴⁴ was found in both indoor and outdoor environments as we observed in our study. *Kocuria* inhabits the human skin, mucous membrane, and larynx and was also detected in indoor dust.^{45,46} *Acinetobacter* is widely distributed in the environment such as soils and water and also found in hospital settings because it can survive exposure to various disinfectants.⁴⁷ For fungi, *Cladosporium*, *Trametes*, *Aureobasidium*, *Sistotrema*, and *Alternaria* were most abundant in both indoor and outdoor samples (Figure 3B). *Cladosporium* was the most commonly found genus inside and outside the classrooms.⁴⁸ Fungal spores of some species of *Alternaria* are allergens that cause hypersensitivity reactions, allergic asthma, keratitis, and skin infection.^{49,50} Chronic exposure to *Aureobasidium pullulans* (a black and yeast-like fungus found in various environments such as soil, water, air, and limestone) from a contaminated humidifier or air conditioner may result in hypersensitivity pneumonitis (extrinsic allergic alveolitis).^{51,52} Of the top 30 fungal genera in our study, 20 were parasitic or symbiotic on plants. *Cladosporium*, *Trametes*, *Phlebia*, *Daedaleopsis*, *Clitocybe*, and *Irpex* that were found to be relatively abundant in our study were mostly related to trees.^{53–56} Our finding that more abundant bacterial genera indoors than outdoors were mostly related to humans is consistent with other study findings suggesting that students and teachers are likely to be a strong source of these airborne bacteria in classrooms; however, fungi in classroom air likely originate from outdoors.^{9,57,58}

To confirm the main source of microbes in classroom air, we examined the similarity of bacterial and fungal community compositions among schools, classrooms, and outdoor air samples using principal coordinates analysis. Our results showed that bacterial community compositions in classroom air in all three schools were generally similar (clustered together) but different from those in outdoor air that also varied widely by school (Figure 4A). On the other hand, fungal community compositions in the samples were generally clustered by school (different location) regardless of indoor or outdoor samples, indicating that fungal communities in indoor and

TABLE 2 Indoor and outdoor environmental factors and number of students on the day when the air samples were collected

Sample type	Sampled month	Classroom/ outdoor	Number of students	Time of natural ventilation (min)	Operation time of air cleaner (min)	Temp (°C)	RH (%)	Wind velocity (m/s)	Precipitation (mm)
OC-Sa1	October	Classroom 1	26	0	160	21.4	46	-	-
OUT-Sa1	October	Outdoor	-	-	-	15.2	61	4.7	0
OC-Sa2	October	Classroom 2	23	170	0	23.7	45	-	-
OUT-Sa2	October	Outdoor	-	-	-	18.8	60	6.8	0
OC-Sb1	November	Classroom 3	21	220	290	23.1	45	-	-
OUT-Sb1	November	Outdoor	-	-	-	11.2	72	8.6	1.1
OC-Sb2	November	Classroom 4	22	0	285	22.6	41	-	-
OUT-Sb2	November	Outdoor	-	-	-	9.3	64	5	0
OC-Sb3	November	Classroom 5	24	40	0	23.7	43	-	-
OUT-Sb3	November	Outdoor	-	-	-	7.2	72	10.4	7.5
UOC-Sb4	November	Classroom 3	0	0	0	21.5	45	-	-
OUT-Sb4	November	Outdoor	-	-	-	8.3	79	11.2	21.1
OC-Sc1	December	Classroom 6	25	0	330	25.2	40	-	-
OUT-Sc1	December	Outdoor	-	-	-	-0.1	48	11.2	0
OC-Sc2	December	Classroom 7	23	0	300	22.4	37	-	-
OUT-Sc2	December	Outdoor	-	-	-	2	42	6.5	0
UOC-Sc3	December	Classroom 7	0	0	0	19.4	38	-	-
OUT-Sc3	December	Outdoor	-	-	-	-0.9	65	5.0	0

Note: Sample type: Occupied classroom sample (OC); unoccupied classroom sample (UOC); outdoor sample (OUT); schools a, b, and c (Sa, Sb, and Sc); and different days (sequential number within the same school). Classroom ID (number in the column for the classroom/outdoor); Temperature (Temp); Relative Humidity (RH).

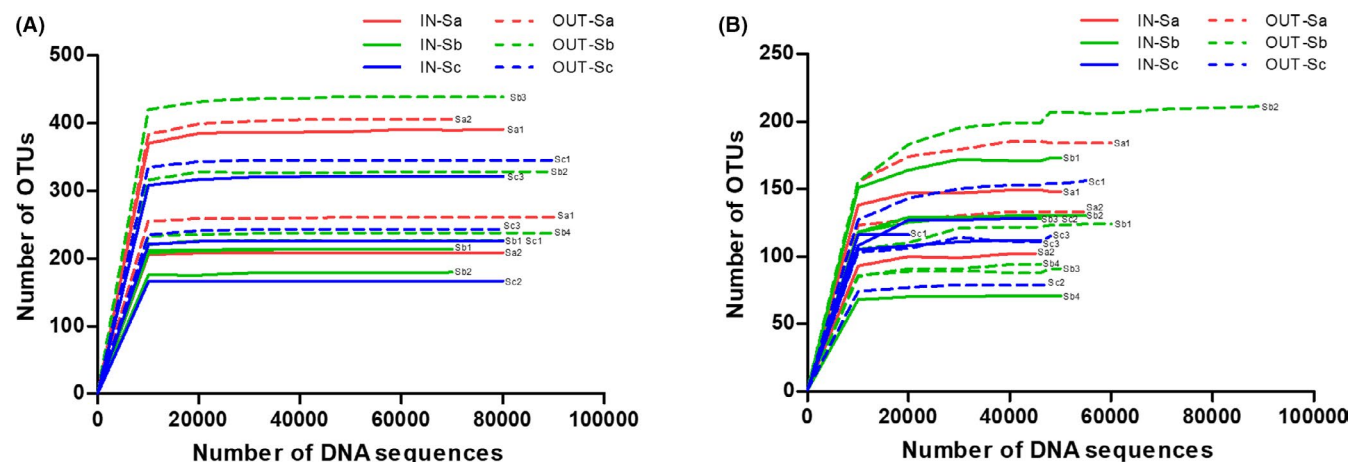


FIGURE 1 Rarefaction curves of (A) bacteria and (B) fungi in the samples. IN, classroom sample; OTUs, operational taxonomic units; OUT, outdoor sample; school ID: Sa, Sb, Sc. Two to four classrooms per school were sampled

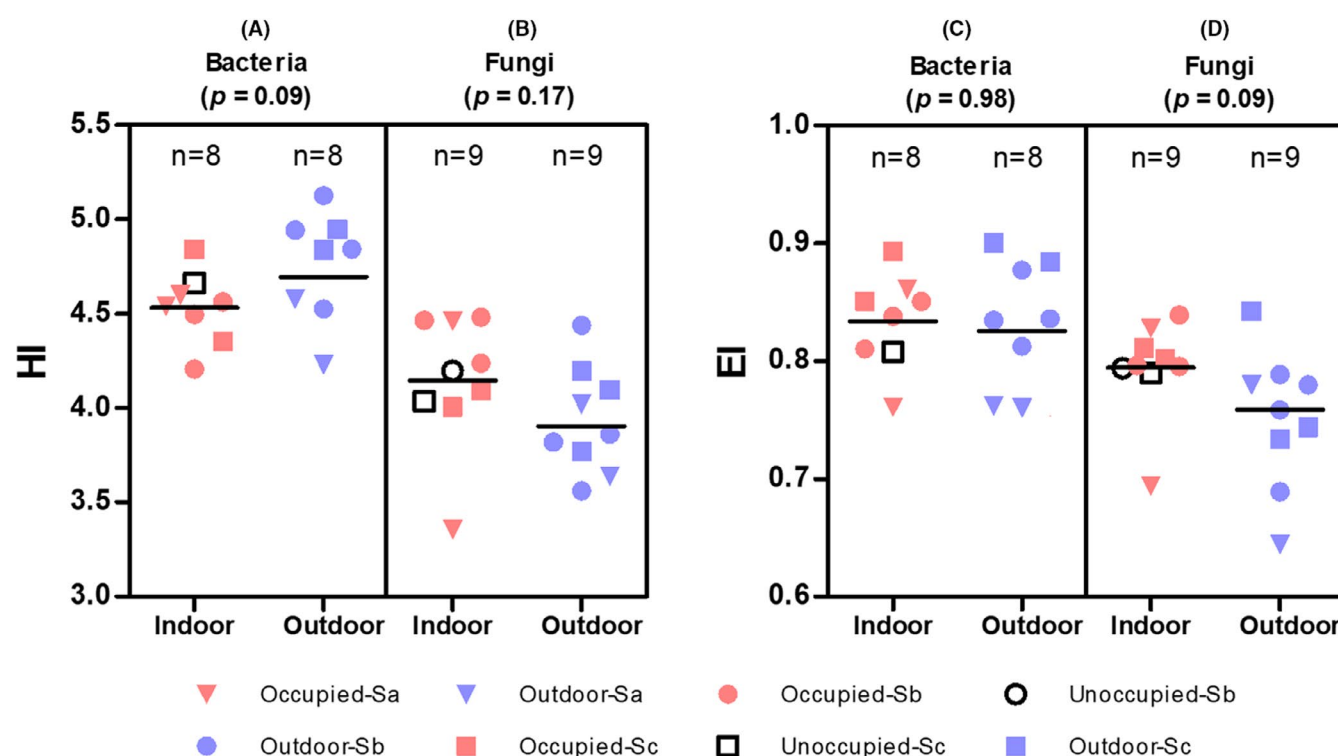


FIGURE 2 Shannon-Wiener index and Pielou's evenness index by sample type. HI: Shannon-Wiener index; EI: Pielou's evenness index; (A) Comparison of HI between indoor and outdoor bacterial samples; (B) comparison of HI between indoor and outdoor fungal samples; (C) comparison of EI between indoor and outdoor bacterial samples; (D) comparison of EI between indoor and outdoor fungal samples. OC, occupied classroom; OUT, outdoor sample; school ID: Sa, Sb, or Sc and classroom ID: 1, 2, or 3. The sample collected from "Unoccupied-Sa" was not sequenced because the amount of cDNA was too low

outdoor air were similar within the same school but different among schools (Figure 4B). However, we did not observe different community compositions among classrooms with or without operating air cleaner. These results of community composition confirmed that human occupants were likely to be the major factor determining bacterial community composition in classroom air whereas outdoor air was likely the main source contributing to fungal community composition in classroom air as described in Burge's report.⁵⁹

3.3 | Quantitative analysis of bacteria in classroom and outdoor air

The copy number of bacterial DNA in the occupied classroom air was lower than that in outdoor air samples on the same day for the most samples, except for the classroom "1" in school "a" (Sa1) and classroom "1" in school "c" (Sc1) (Figure 5). For the classroom in the school "a" (Sa1), the highest number of students (26) in the

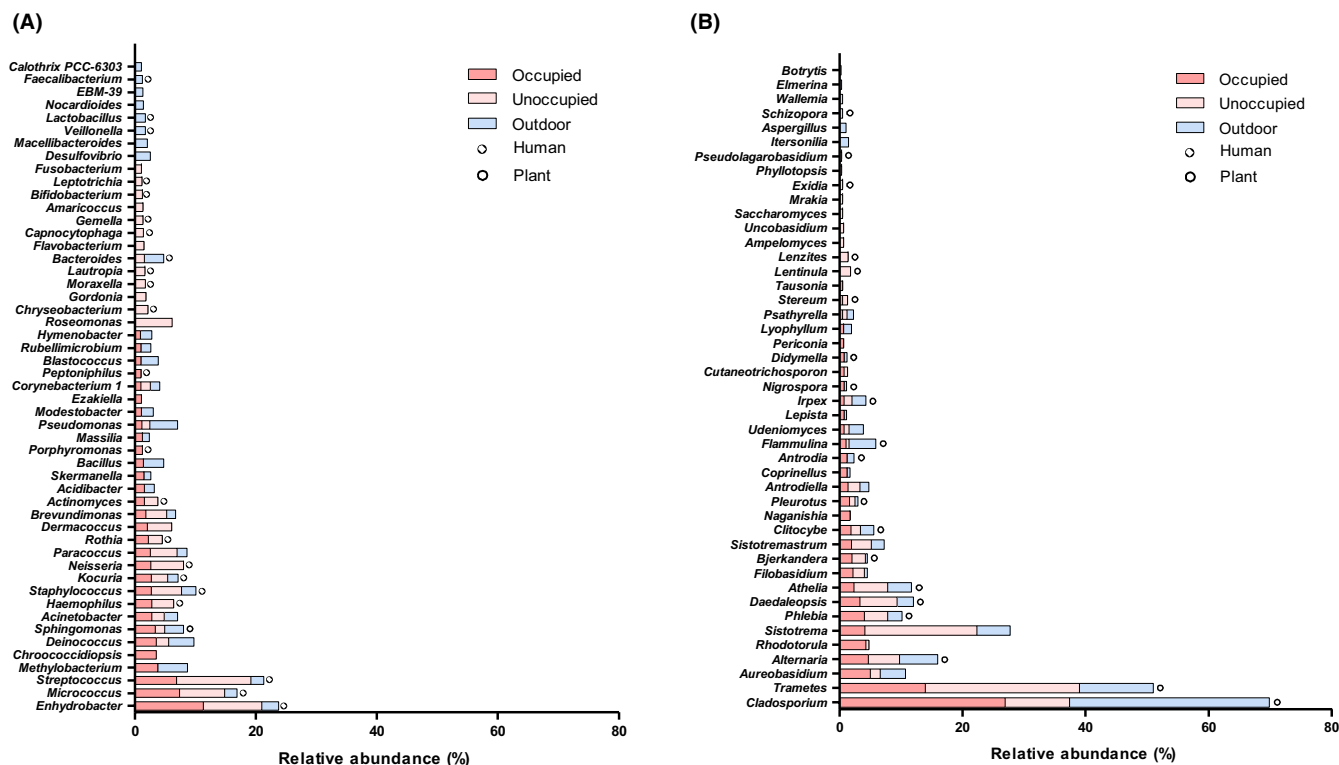


FIGURE 3 Relative abundance of bacteria (A) and fungi (B) at the genus level in occupied and unoccupied classroom and outdoor air. (●: human-related genus; ○: plant-related genus)

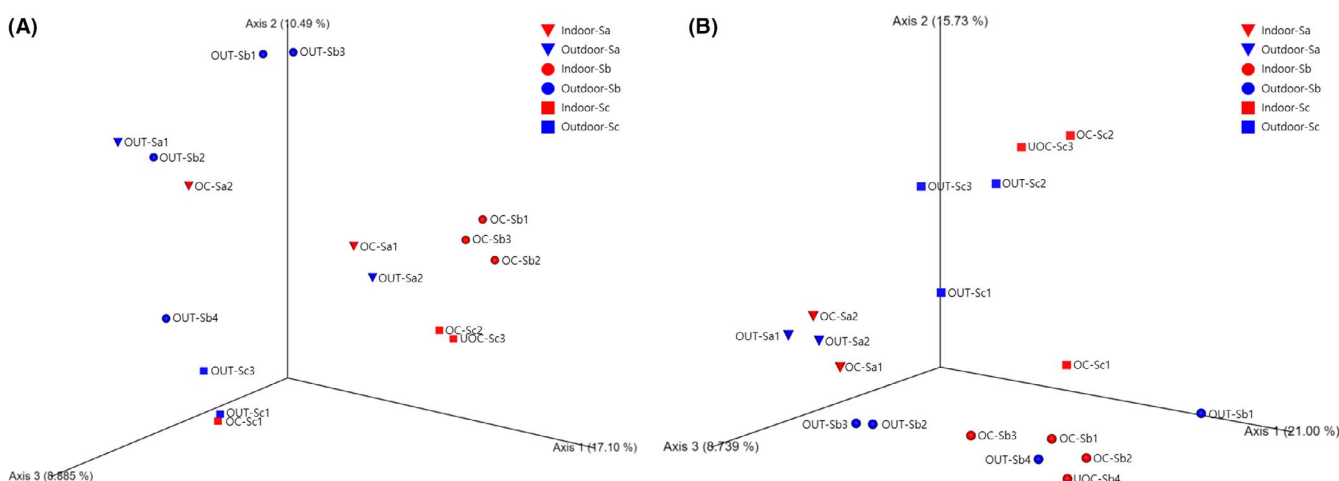


FIGURE 4 Principal coordinates analysis plot using Bray-Curtis dissimilarity index at the genus level for bacteria (A) and fungi (B). OC, occupied classroom; OUT, outdoor sample; School ID: Sa, Sb, or Sc and classroom ID: 1, 2, or 3

classroom might have influenced the concentration of airborne bacterial DNA in the classroom air. On the day of sampling at school “c,” outdoor temperature was below zero, and the sub-zero temperature might have influenced bacterial concentration in outdoor air. The copy number of bacterial DNA in the unoccupied classroom was similar to that in corresponding outdoor air. Our quantitative PCR results (amount of airborne bacterial DNA) were also consistent with the results of the semi-quantitative analyses of rarefaction curves and alpha diversity presented in the previous sections.

3.4 | Correlations between environmental factors and microbial community

Evenness index for bacteria in classroom air samples was significantly and positively correlated (correlation coefficient = 0.81, $p < 0.05$) with classroom temperature (Figure S3A). The number of students was positively correlated with the number of bacterial DNA copies but not statistically significant. For outdoor bacterial air samples, temperature was negatively correlated with evenness index (-0.83 , $p < 0.01$) but positively correlated with DNA copy

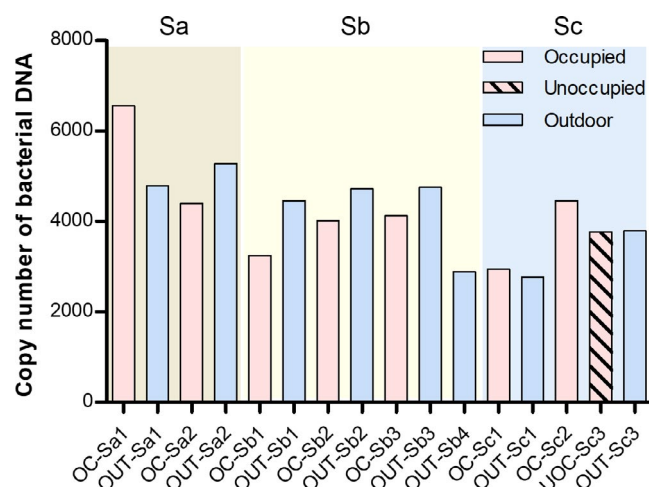


FIGURE 5 Copy number of bacterial DNA by indoor and outdoor sample for the same school and classroom. OC, occupied classroom; OUT, outdoor sample; School ID: Sa, Sb, or Sc and Classroom ID: 1, 2, or 3

numbers (0.74, $p < 0.05$) (Figure S3B). For fungi, the number of students indoors was positively correlated with evenness index (0.70, $p < 0.05$) (Figure S3C) whereas outdoor wind speed and evenness index of outdoor fungi had a positive correlation (0.77, $p < 0.05$) (Figure S3D). However, relative humidity, natural ventilation time, air cleaner operation time, and precipitation were not correlated with the microbial diversity indices both indoors and outdoors.

One of the limitations in our study was the small sample size, which might limit generalizability of the study findings. The small sample size also did not allow us to statistically evaluate the effects of environmental conditions on microbial community in classroom air. Future studies with a large number of schools and classrooms in different areas and seasons might provide more generalizable results. Nonetheless, our study findings provide useful information to better understand airborne bacterial and fungal microbiome including their major sources in occupied classrooms.

4 | CONCLUSIONS

From our study, we found that bacterial microbiota in occupied classroom air was likely associated with humans and different from that in outdoor air, indicating that human occupants may be the major source of indoor airborne bacteria. However, fungal microbial community in classroom air is similar to those in outdoor air, indicating that outdoor airborne fungi may be the major contributor to indoor airborne fungi. Our findings from a study of Korean school classrooms were in line with those from studies conducted in other countries. However, health effects of human occupant-originated bacteria in school classrooms are largely unknown and thus epidemiologic studies of classroom microbiomes and health in schoolteachers and students are warranted. Our findings of higher number of bacterial DNA copies in outdoors than indoors, and

outdoor airborne fungi as the main source of classroom fungi might indicate the importance of properly controlling infiltration of outdoor microbes into classrooms.

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CONFLICT OF INTEREST

The authors declare that they have no competing interest.

AUTHOR CONTRIBUTIONS

Bong Gu Lee involved in formal analysis (equal); writing-original draft (equal); data curation (equal); and methodology (equal). Jun IL Yang involved in data curation (equal); methodology (equal); and formal analysis (equal). Euna Kim and Sun Woo Geum involved in methodology (equal). Ju-Hyeong Park involved in conceptualization (equal); supervision (equal); and writing-review and editing (equal). Min-Kyeong Yeo involved in conceptualization (equal); funding acquisition (equal); supervision (equal); and writing-review and editing (equal).

DISCLAIMER

The findings and conclusions in this report are those of the authors and do not necessarily represent the official position of the National Institute for Occupational Safety and Health, Centers for Disease Control and Prevention.

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REFERENCES

- Klepeis NE, Nelson WC, Ott WR, et al. The National Human Activity Pattern Survey (NHAPS): a resource for assessing exposure to environmental pollutants. *J Exposure Sci Environ Epidemiol*. 2001;11:231-252.
- Yang J, Xu M, Zhang X, Hu Q, Sommerfeld M, Chen Y. Life-cycle analysis on biodiesel production from microalgae: water footprint and nutrients balance. *Biores Technol*. 2011;102:159-165.
- Cincinelli A, Martellini T. Indoor air quality and health. *Int J Environ Res Public Health*. 2017;14:1286.
- Yoon H, Seo J, Kim T, et al. Development of Korean exposure factors for children in Korea. *Korean J Environ Health Sci*. 2017;43:167-175.
- Ferro AR, Kopperud RJ, Hildemann LM. Source strengths for indoor human activities that resuspend particulate matter. *Environ Sci Technol*. 2004;38:1759-1764.
- Jones AP. Indoor air quality and health. *Atmos Environ*. 1999;33:4535-4564.
- Valavanidis A, Fiotakis K, Vlachogianni T. Airborne particulate matter and human health: toxicological assessment and importance of size and composition of particles for oxidative damage and carcinogenic mechanisms. *J Environ Sci Health C Environ Carcinog Ecotoxicol Rev*. 2008;26:339-362.

8. Douwes J, Thorne P, Pearce N, Heederik D. Bioaerosol health effects and exposure assessment: progress and prospects. *Ann Occup Hyg*. 2003;47:187-200.
9. Hospodsky D, Qian J, Nazaroff WW, et al. Human occupancy as a source of indoor airborne bacteria. *PLoS One*. 2012;7:e34867.
10. Blaser MJ, Falkow S. What are the consequences of the disappearing human microbiota? *Nat Rev Microbiol*. 2009;7:887-894.
11. Cho I, Blaser MJ. The human microbiome: at the interface of health and disease. *Nat Rev Genet*. 2012;13:260-270.
12. Weyrich LS, Dixit S, Farrer AG, Cooper AJ, Cooper AJ. The skin microbiome: associations between altered microbial communities and disease. *Australas J Dermatol*. 2015;56:268-274.
13. Yazdanbakhsh M, Kremsner PG, van Ree R. Allergy, parasites, and the hygiene hypothesis. *Science*. 2002;296:490-494.
14. Liddicoat C, Waycott M, Weinstein P. Environmental change and human health: can environmental proxies inform the biodiversity hypothesis for protective microbial-human contact? *Bioscience*. 2016;66:1023-1034.
15. Amann RI, Ludwig W, Schleifer K-H. Phylogenetic identification and in situ detection of individual microbial cells without cultivation. *Microbiol Rev*. 1995;59:143-169.
16. Bloomfield SF, Stewart GSAB, Dodd CER, Booth IR, Power EGM. The viable but non-culturable phenomenon explained? *Microbiology*. 1998;144:1-3.
17. Streit WR, Schmitz RA. Metagenomics – the key to the uncultured microbes. *Curr Opin Microbiol*. 2004;7:492-498.
18. Jünemann S, Kleinbölting N, Jaenicke S, et al. Bioinformatics for NGS-based metagenomics and the application to biogas research. *J Biotechnol*. 2017;261:10-23.
19. Shokralla S, Spall JL, Gibson JF, Hajibabaei M. Next-generation sequencing technologies for environmental DNA research. *Mol Ecol*. 2012;21:1794-1805.
20. Morawska L, Tang JW, Bahnfleth W, et al. How can airborne transmission of COVID-19 indoors be minimised? *Environ Int*. 2020;142(1):105832.
21. Seoul. Seoul Metropolitan Government Traffic Survey Data. Seoul; 2019.
22. MOE. *Results of 2018 Student Health Examination Sample Statistics Analysis*. Sejong: MOE; 2019.
23. Rittenour WR, Park J-H, Cox-Ganser JM, Beezhold DH, Green BJ. Comparison of DNA extraction methodologies used for assessing fungal diversity via ITS sequencing. *J Environ Monit*. 2012;14:766-774.
24. Lemons AR, Hogan MB, Gault RA, et al. Microbial rRNA sequencing analysis of evaporative cooler indoor environments located in the Great Basin Desert region of the United States. *Environ Sci Process Impacts*. 2017;19:101-110.
25. Kim D, Kim H, Seo K. Microbial composition of Korean kefir and antimicrobial activity of *Acetobacter fabarum* DH1801. *J Food Saf*. 2020;40:e12728.
26. Klindworth A, Pruesse E, Schweer T, et al. Evaluation of general 16S ribosomal RNA gene PCR primers for classical and next-generation sequencing-based diversity studies. *Nucleic Acids Res*. 2013;41:e1.
27. Caporaso JG, Kuczynski J, Stombaugh J, et al. QIIME allows analysis of high-throughput community sequencing data. *Nat Methods*. 2010;7:335-336.
28. Beloshapka AN, Forster GM. Fecal microbial communities of overweight and obese client-owned dogs fed cooked bean powders as assessed by 454-pyrosequencing. *J Vet Sci Technol*. 2016;7(1):2.
29. Abarenkov K, Henrik Nilsson R, Larsson K-H, et al. The UNITE database for molecular identification of fungi – recent updates and future perspectives. *New Phytol*. 2010;186:281-285.
30. Quast C, Pruesse E, Yilmaz P, et al. The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. *Nucleic Acids Res*. 2012;41:D590-D596.
31. Senthilraj R, Prasad GS, Janakiraman K. Sequence-based identification of microbial contaminants in non-parenteral products. *Braz J Pharm Sci*. 2016;52:329-336.
32. Larionov A, Krause A, Miller W. A standard curve based method for relative real time PCR data processing. *BMC Bioinformatics*. 2005;6(1):1-16.
33. Zaheer R, Noyes N, Ortega Polo R, et al. Impact of sequencing depth on the characterization of the microbiome and resistome. *Sci Rep*. 2018;8:1-11.
34. Kettleson EM, Adhikari A, Vesper S, Coombs K, Indugula R, Reponen T. Key determinants of the fungal and bacterial microbiomes in homes. *Environ Res*. 2015;138:130-135.
35. de Souza RSC, Okura VK, Armanhi JSL, et al. Unlocking the bacterial and fungal communities assemblages of sugarcane microbiome. *Sci Rep*. 2016;6:28774.
36. Wagg C, Schlaeppi K, Banerjee S, Kuramae EE, van der Heijden MGA. Fungal-bacterial diversity and microbiome complexity predict ecosystem functioning. *Nat Commun*. 2019;10:1-10.
37. Kembel SW, Jones E, Kline J, et al. Architectural design influences the diversity and structure of the built environment microbiome. *ISME J*. 2012;6:1469-1479.
38. Kloos WE, Musselwhite MS. Distribution and persistence of *Staphylococcus* and *Micrococcus* species and other aerobic bacteria on human skin. *Appl Microbiol*. 1975;30:381-395.
39. Rintala H, Pitkaranta M, Toivola M, Paulin L, Nevalainen A. Diversity and seasonal dynamics of bacterial community in indoor environment. *BMC Microbiol*. 2008;8:56.
40. Wilson M. *Bacteriology of Humans: An Ecological Perspective*. 1st ed. Hoboken: John Wiley & Sons; 2009.
41. Tortora GJ, Funke BR, Case CL, Johnson TR. *Microbiology: An Introduction*. 11th ed. San Francisco, CA: Benjamin Cummings; 2012.
42. Liu G, Tang CM, Exley RM. Non-pathogenic *Neisseria*: members of an abundant, multi-habitat, diverse genus. *Microbiology*. 2015;161:1297-1312.
43. Park J, Kim SJ, Lee J-A, Kim JW, Kim SB. Microbial forensic analysis of human-associated bacteria inhabiting hand surface. *Forensic Sci Int Genet Suppl Ser*. 2017;6:e510-e512.
44. Green PN. *Methylobacterium*. In: Whitman WB (ed.), *Bergey's Manual of Systematics of Archaea and Bacteria*. Hoboken: Wiley; 2015:1-8.
45. McManus CJ, Kelley ST. Molecular survey of aeroplane bacterial contamination. *J Appl Microbiol*. 2005;99:502-508.
46. Grice EA, Kong HH, Renaud G, et al. A diversity profile of the human skin microbiota. *Genome Res*. 2008;18:1043-1050.
47. Doughari HJ, Ndakidemi PA, Human IS, Benade S. The ecology, biology and pathogenesis of *Acinetobacter* spp.: an overview. *Microbes Environ*. 2011;26:101-112.
48. Bensch K, Braun U, Groenewald JZ, Crous PW. The genus *Cladosporium*. *Stud Mycol*. 2012;72:1-401.
49. Rotem J. *The Genus Alternaria: Biology, Epidemiology, and Pathogenicity*. St Paul, MN: American Phytopathological Society; 1994.
50. Meena M, Gupta SK, Swapnil P, Zehra A, Dubey MK, Upadhyay RS. *Alternaria* toxins: potential virulence factors and genes related to pathogenesis. *Front Microbiol*. 2017;8(1):1451.
51. Andrews JH, Spear RN, Nordheim EV. Population biology of *Aureobasidium pullulans* on apple leaf surfaces. *Can J Microbiol*. 2002;48:500-513.
52. Gostinčar C, Ohm RA, Kogej T, et al. Genome sequencing of four *Aureobasidium pullulans* varieties: biotechnological potential, stress tolerance, and description of new species. *BMC Genom*. 2014;15:549.
53. Kirk PM, Cannon PF, David JC, Stalpers JA. *Ainsworth and Bisby's Dictionary of the Fungi*. 9th ed. Egham: CABI Publishing; 2001.
54. Zalar P, de Hoog GS, Schroers H-J, Crous PW, Groenewald JZ, Gunde-Cimerman N. Phylogeny and ecology of the ubiquitous

saprobe *Cladosporium sphaerospermum*, with descriptions of seven new species from hypersaline environments. *Stud Mycol.* 2007;58:157-183.

55. Jakovlev J. Fungus gnats (*Diptera: Sciarioidea*) associated with dead wood and wood growing fungi: new rearing data from Finland and Russian Karelia and general analysis of known larval microhabitats in Europe. *Entomol Fenn.* 2011;22:157-189.
56. Li HJ, Si J, He SH. *Daedaleopsis hainanensis* sp. nov. (Polyporaceae, Basidiomycota) from tropical China based on morphological and molecular evidence. *Phytotaxa.* 2016;275:294-300.
57. Shelton BG, Kirkland KH, Flanders WD, Morris GK. Profiles of airborne fungi in buildings and outdoor environments in the United States. *Appl Environ Microbiol.* 2002;68:1743-1753.
58. Hewitt KM, Gerba CP, Maxwell SL, Kelley ST. Office space bacterial abundance and diversity in three metropolitan areas. *PLoS One.* 2012;7:e37849.
59. Burge HA. An update on pollen and fungal spore aerobiology. *J Allergy Clin Immunol.* 2002;110:544-552.

SUPPORTING INFORMATION

Additional Supporting Information may be found online in the Supporting Information section.

Fig S1-3

Table S1

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