Annual distribution of allergenic fungal spores in atmospheric particulate matter in the eastern Mediterranean; a comparative study between ergosterol and quantitative PCR analysis

N. Lang-Yona\textsuperscript{1}, K. Dannemiller\textsuperscript{2}, N. Yamamoto\textsuperscript{2}, N. Burshtein\textsuperscript{1}, J. Peccia\textsuperscript{2}, O. Yarden\textsuperscript{3}, and Y. Rudich\textsuperscript{1}

\textsuperscript{1}Department of Environmental Sciences, Weizmann Institute of Science, Rehovot, 76100, Israel
\textsuperscript{2}Department of Chemical and Environmental Engineering, Yale University, Mason Laboratory, 9 Hillhouse Avenue, New Haven, CT, 06520-8286, USA
\textsuperscript{3}Department of Plant Pathology and Microbiology, The R.H. Smith Faculty of Agriculture, Food and Environment, The Hebrew University of Jerusalem, Rehovot 76100, Israel

Received: 12 September 2011 – Accepted: 7 October 2011 – Published: 26 October 2011
Correspondence to: Y. Rudich (yinon.rudich@weizmann.ac.il)
Published by Copernicus Publications on behalf of the European Geosciences Union.
Abstract

Airborne fungal spores are an important fraction of atmospheric particulate matter and are major causative agents of allergenic and infectious diseases. Predicting the variability and species of allergy-causing fungal spores requires detailed and reliable methods for identification and quantification. There are diverse methods for their detection in the atmosphere and in the indoor environments; yet, it is important to optimize suitable methods for characterization of fungal spores in atmospheric samples. In this study we sampled and characterized total and specific airborne fungal spores from PM\textsubscript{10} samples collected in Rohovot, Israel over an entire year. The total fungal spore concentrations vary throughout the year although the species variability was nearly the same. Seasonal equivalent spore concentrations analyzed by real-time quantitative-PCR-based methods were fall \textgreater winter \textgreater spring \textgreater summer. Reported concentrations based on ergosterol analysis for the same samples were fall \textgreater spring \textgreater winter \textgreater summer. Correlation between the two analytical methods was found only for the spring season. These poor associations may be due to the per-spore ergosterol variations that arise from both varying production rates, as well as molecular degradation of ergosterol. While conversion of genome copies to spore concentration is not yet straightforward, the potential for improving this conversion and the ability of qPCR to identify groups of fungi or specific species makes this method preferable for environmental spore quantification. Identifying tools for establishing the relation between the presence of species and the actual ability to induce allergies is still needed in order to predict the effect on human health.

1 Introduction

Fungi are ubiquitous eukaryotic organisms that produce spores with the capability of long-range dispersal throughout the lower troposphere (Breitenbach, 2002). Understanding the diversity along with the spatial and temporal distribution of airborne fungal
spores is of increasing importance because fungal spores and other propagules can potentially act as sources of plant diseases, (Madelin, 1994) human allergens, (Breitenbach, 2002;Kurup, 2000) animal and human infectious agents, (Enoch et al., 2006) produce mycotoxins and other deleterious secondary metabolites (Garvey and Keller). The characteristics of an airborne fungal population may be related to different environmental factors, such as seasonality, location, meteorological parameters, relative humidity, and temperature (Burshteen et al., 2011; Fröhlich-Nowoisky et al., 2009; Oliveira, 2005). For example, elevated CO$_2$ concentrations (Kilronomos, 1997) or increased temperatures (Gange et al., 2007) have been shown to positively correlate with increases in fungal biomass. Some of these changes have been associated with global climate change and may impact human health and economics.

Several methods are commonly used to quantify the distribution of airborne fungal spores. Direct quantification can be performed either by conventional microbiological methods (e.g. spore counting or determining the number of colony forming units, CFUs), (Li, 2010; Takahashi, 1997) or by nucleic acid-based analyses (e.g. quantitative-Polymerase Chain Reaction (q-PCR) of extracted DNA) (Fröhlich-Nowoisky et al., 2009; Hospodsky et al., 2010; Yamamoto, 2011). Indirect methods of quantification utilize analytical chemistry for assessing the presence of fungal-specific biomarkers, (Bjurman, 1994; Burshtein et al., 2011; Lau et al., 2006; Mille-Lindblom et al., 2004; Miller, 1997; Robine et al., 2005) and include the analysis of ergosterol, a membrane sterol, found almost exclusively in fungi (Axelsson et al., 1995; Gessner and Chauvet, 1993; Lau et al., 2006; Mille-Lindblom et al., 2004; Pasanen, 1999). Ergosterol detection is typically performed by gas chromatography-mass spectrometry (GC-MS), (Burshtein et al., 2011; Lau et al., 2006) or high pressure liquid phase chromatography (HPLC) (Robine et al., 2005) and is reported as ergosterol mass per volume of air, or by using a conversion factor, as ergosterol mass per spore.

In order to determine if ergosterol content is consistently associated with the concentration of fungal spores, previous investigators have compared fungal concentrations based on culture enumeration with ergosterol mass concentration (Lau et al., 2006).
2006; Heid et al., 1996). However, spore counts based on culturability may not accurately reflect the true fungal content since nonviable species cannot be cultured, nor can fungal spores that require a different media or growth condition than provided in the culture assay. This limitation to culturing of environmental bacteria and fungi is well documented (Amann et al., 1995). In the case of airborne fungal spores, differences between total, intact spores and culturable counts as high as ten times have been documented (Hospodsky et al., 2010). In addition, sampling fungi onto agar plates is limited by desiccation and can only be performed for short (ca. ~15 min) sampling durations (Buttner, 1997).

Real-Time q-PCR analysis is based on direct detection of DNA extracted from sampled fungal spores and thus circumvents culturability limitations (Peccia and Hernandez, 2006; Alkan, 2004). In addition, bypassing the need for viable propagules enables sample collection onto filters and for longer periods of time, thus improving comparability by allowing fungal spore analysis to be performed on the same aerosol sample that is used for ergosterol.

In this study we analyzed the concentration of outdoor airborne fungal spores sampled throughout one year in Rehovot, Israel. We investigated the seasonal distribution of selected allergenic fungi species as indicators for the fungal community distribution, and compared total fungal concentrations (universal fungal qPCR primers) with ergosterol reported concentrations (Burshtein et al., 2011). Enumeration of airborne fungi by quantitative PCR is rapidly becoming the dominant method for exposure analysis in environmental and public health studies (Sacchi et al.; Haugland, 2002; Zhou et al., 2000; Mackay, 2007). It is important that these values be compared with other commonly used biomarker assays to ensure that these measures provide similar assessments of relative exposure.
2 Materials and methods

2.1 Sampling and storage

Airborne particles were collected using a high volume sampler (Ecotech High volume sampler model ECO-HVS3000 with PM$_{10}$ inlet, at a flow rate of 67.8 m$^3$ h$^{-1}$). Forty eight samples were collected over a period of one year (December 2008–December 2009), on the rooftop of a four store building in Rehovot, Israel (31.540200 N, 34.4803300 5 E). All sampling periods were 72 h, except for two cases (8/23/2009 and 9/22/2009) where the sampling period was 24 h. Sampled aerosols were collected on thermally pretreated (450°C) quartz fiber filters (20.3 cm x 25.4 cm Whatman). Immediately after sampling, filters were wrapped in aluminum foil and kept at −20°C until the sampling campaign was completed. Subsequently, all samples were archived at −80°C until analysis.

2.2 Universal and allergenic fungal spore identification and quantification by qPCR analysis

Quantitative PCR was performed in accordance with previously described standard fungal DNA preparation protocols, DNA extraction methods, and quality control measures (Haugland, 2002; Yamamoto, 2011; Hospodsky et al., 2010). Briefly, DNA was extracted directly from three different 3 cm x 3 cm subsamples filters for total fungi and 1 cm x 3 cm for specific species, using the PowerSoil DNA Isolation Kit (MO BIO Laboratories Inc.) according to the manufacturers protocol, with modifications for high yields, as previously described in detail for fungi collected on glass fiber filters (Hospodsky et al., 2010).

Quantitative PCR analyses for total fungi sampled (i.e. universal fungi) were performed with a Step One Plus Real-Time PCR (Applied Biosystems), using the FF2/FR1 assay (Table 1) (Zhou et al., 2000). A 10 µl reaction mixture consisted of 5 µl Fast SYBR Green Master Mix (Applied Biosystems), 1 µl extracted DNA, 0.5 µl of each
primer (0.5 µM), and 3 µl nuclease free water (NFW). The thermal cycling conditions consisted of an initial 20 s denaturation and enzyme activation at 95 °C, followed by 45 cycles of 3 s denaturation at 95 °C, and 30 s annealing and extension at 60 °C. Cycle thresholds were calculated using the auto function in Step One Plus. Negative controls, with NFW, were run on each qPCR 96-well plate, and all q-PCR measurements of each filter were replicated at least three times. Standard curves based on known quantities of *Aspergillus fumigatus* fungal spores versus cycle threshold values were established for each q-PCR 96-well plate analyzed. Results for universal primers are reported as *Aspergillus fumigatus* (ATCC 34506) equivalent fungal spores.

For 25 representative sampled filters, five medically important groups of allergenic fungi (*Aspergillus fumigatus*, *Alternaria alternata*, *Cladosporium cladosporioides*, *Epicoccum nigrum*, *Penicillium* spp. and the *Penicillium/Aspergillus* group) were chosen to be specifically analyzed (Table 1). In addition, a second set of biological replicates for universal fungi was repeated. DNA was extracted from filter samples as described above and real-time qPCR was performed. Specific allergens were quantified using the TaqMan method and employed the Afumi, Aaltr, Cclad2, Enigr, PenGrp3 and PenAsp1mgb assays (Table 1) reported by the U.S. EPA, (Haugland, 2002), and universal fungi detection was performed with SYBR green method (FastStart Universal SYBR Green Master (ROX); Roche Applied Science), using the same assay as described above (Table 1). A 25 µl reaction mixture containing the template DNA (1 µl sample extract), 1× TaqMan Universal PCR Master Mix (Applied Biosystems) for specific species or 12.5 µl of FastStart Universal SYBR Green Master (Roche Applied Science) for universal fungi, 1 µM of each primer, and 0.08 µM of the probe for the specific species assays. A real-time PCR system (ABI 7500 Fast Real-time PCR System; Applied Biosystems) was used with the following cycle condition: 50 °C for 2 min, 95 °C for 15 min of initial denaturation and 45 cycles at 95 °C for 15 s dissociation and 60 °C for 1 min of annealing and extension. All allergen specific qPCR measurements were performed in triplicate. Standard concentrations of genomic DNA for the five specific allergenic fungi were prepared (i.e. *Aspergillus fumigatus* (ATCC 34506), *Alternaria*...
Alternaria (PEM 01043), Cladosporium cladosporioides (ATCC 16022), Epicoccum nigrum (PEM 01043), Penicillium chrysogenum (CAES PC1)) and standard qPCR curves were based on spore concentration versus cycle threshold values. PenAsp1mgb standard curves (amplification of both Penicillium and Aspergillus species) were generated using the A. fumigatus (ATCC 34506) DNA standard.

To test for PCR inhibition, seven randomly selected extracts from the filter samples were added to subsets of diluted E. nigrum DNA, and standard curves were produced. No significant inhibition was observed in this study. Accuracy, precision, and method limit of detections where tested for these specific assays, and reported in Yamamoto et al. (2011).

2.3 Ergosterol analysis

Ergosterol extraction from the sampled filters used in this study was previously reported (Burshtein et al., 2011). In short, the analysis was performed on each of the 48 filters in triplicates. 3 × 3 cm² sampled filter pieces were hydrolyzed to release the ergosterol into methanol solution. Ergosterol was extracted using phase separation of deionized water and hexane, and an internal standard of 7-Dehydro cholesterol was added to each sample. Analysis was performed using GC-MS/MS (Varian GC-MS Saturn 2000 equipped with a Restek column).

2.4 Data analysis

Correlations between ergosterol mass concentration and equivalent fungal spore concentrations were analyzed by Pearson correlation test. Variance between fungal species distributions was analyzed by One-way ANOVA between groups. Analysis of significant differences between seasons was done by One-way ANOVA followed by Tukey SHD post-hoc test for each allergenic assay.
3 Results

Initially we followed the seasonal distribution of selected allergy-inducing fungal species (Fig. 1, Table 2). We found that *A. alternata* is the most abundant species out of all allergen-producing fungi tested, detected in three out of the four seasons (winter – 35%, spring – 34%, summer – 36%), whereas in the fall, *C. cladosporioides* was the most prevalent (38%) species. In the spring, a relatively high abundance of *E. nigrum* was detected (21% compared to 7%, 10% and 8% in winter, summer and fall respectively). *A. fumigatus*, *P. chrysogenum* and other groups of *Aspergillus* and *Penicillium* were relatively scarcer and observed throughout all seasons with a similar ratio. ANOVA test between groups confirmed the null hypothesis for significant similarity between the species seasonal distributions (*F*-value = 0.03, *P*-value = 1.00).

Figure 2 represents the values of both ergosterol (top graph) in ng m\(^{-3}\) air, and total fungi (bottom graph) in equivalent spore m\(^{-3}\). The two biological repeats using universal fungi primers for the q-PCR experiments are represented by “qPCR I” and “qPCR II” respectively, where the “qPCR II” analyses were done for a seasonally representative part of the filters. The two repeated fungal spore concentrations are statistically similar (tested by independent two sample *t*-test, *P*-value = 0.713). Both fungal spore and ergosterol concentrations indicate that fungal mean concentrations are lowest in the summer period (356.6 ± 265.7, 0.28 ± 0.23 respectively) and highest in the fall (2896.9 ± 2240.1, 0.94 ± 0.71 respectively, see Table 3). A local maximum at the spring period (between 14 and 18 May) also shows similarity between the two parameters. As can be seen, the inter-season and intra-season variability can span 1–2 orders of magnitude in concentration. No clear trend can be identified in our data. By averaging samples across individual seasons, the ranking of fungal spore concentrations is fall > winter > spring > summer. The seasonal ranking scheme of ergosterol concentrations is fall > spring > winter > summer. It is therefore important to understand the sources for these inconsistencies.
We tried to correlate ergosterol and fungal spore concentrations linearly, as detailed in Table 3 and Fig. 3. We did not find a yearly correlation between the two parameters \((r = 0.18 \pm 0.61, P\text{-value} = 0.21)\). When correlating seasonally, a positive correlation was observed only in the spring \((r = 0.61 \pm 0.63, P\text{-value} = 0.1\), double asterisk in Table 3). A semi-log correlation was performed as well, since spore concentration analyzed from qPCR data are log-normal distributed, (Maher, 1981; di Giorgio et al., 1996), however no improved correlation was detected.

In order to eliminate possible errors in qPCR analysis that may arise from genome-to total spore conversion, we further examined the prevalence of specific fungal species by utilizing qPCR analysis with species-specific assays (Table 1). Assuming that these species comprise a good representation of the total sampled fungal spores, we summed the analyzed species and compared this sum to the ergosterol concentrations (Fig. 4). Similarly to both ergosterol and qPCR analyses for equivalent spore concentration, a higher spore concentration of all species was detected in the fall, with the exception of *E. nigrum*, which had the highest values in the spring season (70.04 at spring compared to 62.45 spores m\(^{-3}\) at fall). However, only a weak yearly correlation was found between ergosterol and the sum of specific fungal spore concentrations \((r=0.35 \pm 0.96; P\text{-value} = 0.13)\). Seasonal correlations could not be calculated due to the low number of samples per season.

### 4 Discussion

The ability to quantify fungal spores concentration and species in atmospheric particulate matter is necessary for reducing or preventing exposure to infectious plant and human fungal agents and to fungal allergens (Madelin, 1994). Here we used DNA analysis to study the spore annual distribution of 5 allergy-causing fungal species in the Eastern Mediterranean. In addition, we conducted a comparative analysis of the airborne fungal spores and ergosterol concentrations and attempted to correlate the quality and reliability of conventional biomarker analyses to the emerging methods.
Based on DNA sequence analysis.

Fungal species distribution did not change significantly throughout the year. This might be attributed to local fungal sources, which did not experience extreme seasonal changes in the local climate in Israel, buffered by the Mediterranean Sea. In our previous published work (Burshtein et al., 2011) we calculated back trajectories to the sampled filters and found the highest ergosterol levels when back trajectories showed local terrestrial origin. Therefore, qualitatively we must conclude that the majority of the fungal spores most likely come from local origin. The only significant change in spore concentration was observed in the spring for *E. nigrum*. It might be that apart from the local source, this elevated value is attributed to additional sources. Further analysis is needed in order to understand the origin of different fungal species. For example, source apportionment analysis, or parallel sampling at different climatic region in the same area (such as the Negev desert in the south of Israel, or a rural Mediterranean forest in the northern Galilee) may lead to a better understanding of the species origin.

The observed species’ distributions are most important with respect to plant diseases and human allergy inducing, however we cannot deduce from these data on their potency, or the actual ability to induce allergies. Further investigation on the allergenic effects and potency is needed in order to clarify this point.

Ergosterol is considered a reliable proxy for fungal spore concentrations (Lau et al., 2006; Burshtein et al., 2011; Mille-Lindblom et al., 2004; Miller, 1997). We compared ergosterol concentrations with DNA-based analyses of airborne PM$_{10}$ fraction of fungal spores to test this notion. Although DNA-based analysis of environmental fungal content is now becoming routine, comparisons with chemical biomarker analysis have not been undertaken. Positive correlation was found between DNA-based analysis and ergosterol concentrations only in the spring season, with $P$-value higher, but close to the null limit of rejection, $\alpha$. Hence the observed correlation is weakened accordingly. Throughout the entire year, correlation for total fungi concentrations was not found. Ergosterol correlation to the sum of specific fungal species found to be weak, with low $P$-value.
The lack of correlations between ergosterol and equivalent fungal spore concentrations may be associated with the influence of sampling conditions, as well as of environmental factors on fungal growth and physiology, which lead to changes in ergosterol production, as investigated previously (Bjurman, 1994; West et al., 1987; Li et al., 2009; Nout et al., 1987). For example, as was shown by Y. Li et al. (2008), changes in growth temperature lead to changes in ergosterol content in *Penicillium roqueforti*, with an optimum content at 20 °C (Li et al., 2009).

Ergosterol degradation induced by potential environmental factors such as UV light, ozone, NO₅, high RH etc. may contribute to the observed differences as well. In general, using biomarkers for the determination of an organism’s concentrations holds an inherent uncertainty due to these factors, which could induce its degradation. DNA analyses (e.g. qPCR), on the other hand, would not be influenced by either growth conditions or environmental factors, and would reflect the true sampled fungi content.

Ergosterol mass concentration is often reported in equivalent spore concentrations (Pasanen et al., 1999; Lau et al., 2006) and references therein. This conversion, based on the assumption that ergosterol represents viable spores, (Mille-Lindblom et al., 2004) is calculated by comparing the parallel sampling of filters and viable sampling on agar plates. The enumerated viable fungal spores are correlated to the ergosterol mass concentration analyzed from the filters and a conversion factor is then determined. This conversion introduces errors into the reported data from both the different sampling time scale and the presence of non-viable or uncultivable spores that are not included in the viable measurement (Amann et al., 1995). Hence the uncertainty about the consistency of the conversion factor cannot be ignored.

DNA analysis is more reliable with respect to its molecular stability compared to ergosterol measurements. Although DNA molecules might degrade as well due to cell damages during spore transport through the air, the rates are expected to be low for spore genome (Desprès, 2007; Fröhlich-Nowoisky et al., 2009; Griffin, 2004a; Griffin and Kellogg, 2004b). Therefore, for atmospheric samples DNA-based analyses might be preferable in order to detect and quantify the presence of fungal spores rather than
biomarker molecules.

Fungal genome to spore conversion based on specific unicellular conidia (as was done in this study) may incorporate errors as well, as some fungal spores are multi-cellular, multi-nuclear, or both (Smith, 1975; Cummins, 2003). In addition, the genomes of fungal cells contain multiple copies of the target rRNA encoding genes, and the amount of copies may vary substantially between different species of fungi (DePriest, 1992; van Belkum et al., 1998; Gargas, 1996). Thus, while species level qPCR results can be reliably converted back to the number of genomes and ultimately whole cell concentrations, accurate whole cell concentrations cannot be derived when using universal qPCR primers due to the unknown mixture of species that are included in the universal qPCR assay. Amplifying segments of single copy genes (while also avoiding genes that belong to gene families) may be a good solution for calibrating DNA (Amnuaykanjanasasin and Epstein, 2003). A possible drawback in this approach is the low sensitivity due to low limit of detection, especially when studying environmental samples, since there is only one gene copy per cell. In addition, variations in the number of nuclei per conidia, common in fungi, as well as the fact that some fungi (e.g. A. alternata) produce conidia with several cells, will induce errors in this case as well.

Variations such as gene copy per genome or cell are typically inherent to the fungal species and are not dependent on growth conditions; therefore if the fungal population does not change from sample to sample, qPCR with universal primers should provide a consistent, relative measure of fungal spore content. In this study, the even distribution of species throughout the year (Fig. 1) suggests that the gene copy variations above did not contribute to the variability in the reported qPCR data. While this will not be the case for all environmental fungal samples, the consistent diversity throughout the year allows for the qPCR values presented here being a consistent representation of spore content. Therefore, we suggest that ergosterol variations detected between samples may be attributed to deviations per spore. These deviations are most likely caused by changes in fungal growth conditions in addition to different degradation rates of ergosterol and the fungal genome in the atmosphere and on the filter.
5 Summary

By analyzing airborne fungal spore concentration over a period of one year, we found that the variation between species is small in the studied region. A lack in correlation between ergosterol mass concentrations and fungal spore concentrations retrieved from qPCR analyses. This is most likely due to ergosterol production rates fluctuating with changing growth conditions, in addition to its degradation during spore transport in the atmosphere. Even though this was a one year experiment, the fact that no correlation was observed is sufficient to conclude that there are severe drawbacks to ergosterol-based quantification. This finding should be taken into consideration when monitoring fungal spore concentrations.

Acknowledgements. This research was supported by a research grant from the Angel Faivovich foundation for Ecological Studies. N. Lang-Yona was supported by a grant from the Environmental Health Fund, K. Dannemiller was supported by a U.S. National Science Foundation fellowship, and N. Yamamoto is supported by the Japan Society for the Promotion of Science. Y. R. acknowledges support by the Helen and Martin Kimmel Award for Innovative Investigation.

References


ACPD
11, 28689–28711, 2011

A comparative study between ergosterol and quantitative PCR analysis

N. Lang-Yona et al.


Miller, J. D. and Young, J. C.: The Use of Ergosterol to Measure Exposure to Fungal Propagules
**Table 1.** The qPCR assays as detailed in Zhou et al. (2000) and Haugland et al (2002).

<table>
<thead>
<tr>
<th>Assay name</th>
<th>Target fungal species</th>
<th>Primers and probes sequence 5'-3'</th>
</tr>
</thead>
<tbody>
<tr>
<td>FF2/FR1</td>
<td>Universal fungi</td>
<td>F: GGTTCTATTTTGTTGTTCTTA R: CTCTCAATCTGTCAATCCTATT P: GCCGGGCTGGAACCTC F: CCAATTACAAAGGTTATGTTGTCGTA P: TTACAGCCCTGCTGAATTATGCAACCCCTTTGCTATT</td>
</tr>
<tr>
<td>Aaltr</td>
<td>Alternaria alternata</td>
<td>Aaltr</td>
</tr>
<tr>
<td>Afumi</td>
<td>Aspergillus fumigatus, Neosartorya fischeri</td>
<td>Afumi</td>
</tr>
<tr>
<td>Cclad2</td>
<td>Cladosporium cladosporioides, svar.2</td>
<td>Cclad2</td>
</tr>
<tr>
<td>Enigr</td>
<td>Epicoccum nigrum</td>
<td>Enigr</td>
</tr>
<tr>
<td>PenGrp3</td>
<td>Penicillium chrysogenum/griseofulvum/ and glandicola/coprophilum/ expansum Eupenicillium</td>
<td>PenGrp3</td>
</tr>
<tr>
<td>PenAsp1mgb</td>
<td>Universal Penicillium, Aspergillus and Paecilomyces varioti</td>
<td>PenAsp1mgb</td>
</tr>
</tbody>
</table>

F-Forward primer; R-Reverse primer; P-Probe.
Table 2. Seasonally averaged values of fungal species concentrations (spore m$^{-1}$), analyzed by qPCR.

<table>
<thead>
<tr>
<th>Season</th>
<th>A.alternata (Aaltr)</th>
<th>A.fumigatus (Afumi)</th>
<th>C.cladosporidis (Cclad2)</th>
<th>E.nigrum (Enigr)</th>
<th>P.chrysogenum (PenGrp3)</th>
<th>Other* (PenAsp1mgb)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Winter</td>
<td>32.52 ± 26.95</td>
<td>2.33 ± 1.67</td>
<td>176.13 ± 101.73</td>
<td>8.67 ± 10.91</td>
<td>19.94 ± 38.79</td>
<td>106.05 ± 110.24</td>
</tr>
<tr>
<td>Spring</td>
<td>31.00 ± 25.90</td>
<td>2.15 ± 1.64</td>
<td>74.57 ± 105.44</td>
<td>70.04 ± 10.69</td>
<td>4.48 ± 37.85</td>
<td>52.97 ± 121.03</td>
</tr>
<tr>
<td>Summer</td>
<td>24.51 ± 30.85</td>
<td>2.15 ± 3.20</td>
<td>118.11 ± 103.36</td>
<td>51.98 ± 11.30</td>
<td>1.18 ± 37.33</td>
<td>131.82 ± 120.56</td>
</tr>
<tr>
<td>Fall</td>
<td>180.97 ± 26.44</td>
<td>10.99 ± 3.12</td>
<td>1354.72 ± 91.63</td>
<td>62.45 ± 27.61</td>
<td>20.92 ± 36.83</td>
<td>659.43 ± 116.66</td>
</tr>
</tbody>
</table>

*Other Penicillium and Aspergillus species, as detailed in Table 1.
Table 3. Seasonal correlation coefficients between Ergosterol and spore concentrations.\(^a\)

<table>
<thead>
<tr>
<th>Season</th>
<th>Ergosterol (\text{ng m}^{-3})</th>
<th>Spore (\text{m}^{-3})</th>
<th>Correlation coefficient, (r)</th>
<th>(P)-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Winter</td>
<td>0.42 ± 0.38</td>
<td>2044.6 ± 2772.5</td>
<td>0.03 ± 0.40</td>
<td>0.95</td>
</tr>
<tr>
<td>Spring</td>
<td>0.76 ± 0.74</td>
<td>1380.6 ± 2090.5</td>
<td>0.61 ± 0.63(^b)</td>
<td>0.10</td>
</tr>
<tr>
<td>Summer</td>
<td>0.28 ± 0.23</td>
<td>356.6 ± 265.7</td>
<td>0.06 ± 0.24</td>
<td>0.84</td>
</tr>
<tr>
<td>Fall</td>
<td>0.94 ± 0.71</td>
<td>2896.9 ± 2240.1</td>
<td>−0.15 ± 0.72</td>
<td>0.55</td>
</tr>
<tr>
<td>Whole year</td>
<td>0.61 ± 0.61</td>
<td>1873.8 ± 2195.0</td>
<td>0.18 ± 0.61</td>
<td>0.21</td>
</tr>
</tbody>
</table>

\(^a\) Ergosterol and spore values are presented by the season mean value.

\(^b\) Statistically significant.
Fig. 1. Seasonal distributions of specific fungi species detected in sampled filters by q-PCR analysis (represented by the specific detection assays). The values represent the relative abundance out of the sum of the specific fungi considered.
Fig. 2. Seasonal ergosterol concentrations (upper graph), (Burshtein et al., 2011) and fungal spore concentrations (bottom graph) from PM$_{10}$ filters. The qPCR I and qPCR II values refer to independent sample repeats. The error bars are standard deviations, $n = 48$ for ergosterol and qPCR I analysis of fungal spore, and $n = 24$ for qPCR II analysis of fungal spore.
Fig. 3. Seasonal correlation between equivalent total fungal spore concentrations analyzed from q-PCR measurements, and ergosterol mass concentrations (Burshtein et al., 2011). The error bars represent the standard deviation of ergosterol (vertical) and fungi spore sampling replicates (horizontal).
Fig. 4. Seasonal ergosterol concentrations (top right graph), (Burshtein et al., 2011) compared to the sum of specific fungal spore concentrations (bottom right graph) from PM$_{10}$ filters, with a correlation coefficient specified in the lower graph. The specific fungal species are specified on the left graphs (Alternaria alternata, Aaltr; Aspergillus fumigatus, Afumi; Cladosporium cladosporioides, Cclad2; Penicillum Chrysogenum, PenGrp3 Eppicocum nigrum, Enigr). The error bars are standard deviations, $n = 21$ samples for ergosterol, and $n = 25$ samples for fungal spore.