Interactive comments on “Regional-scale Simulations of Fungal Spore Aerosols Using an Emission Parameterization Adapted to Local Measurements of Fluorescent Biological Aerosol Particles” by M. Hummel et al.

M. Hummel et al.

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Anonymous Referee #1

We thank the reviewer for his/her comments which have improved our manuscript. The comments are listed below in italics.

In general, the FBAP measurements measure all particles in the range of 2-4µm, however, it the authors assume these are fungal spores. Bacterial agglomerate, as well as giant bacteria and agglomerates of free proteins and amino acids could contribute to this fraction. In this case, the simulations, referring only to fungal spores would not represent this data well. This point should be further addressed and quantified, as the introduction generalizes this issue to the entire PBAP population.

It is not entirely clear what kind of biological particles dominate FBAP number concentrations. Several field and lab studies show that fungal spores and bacteria cells or bacteria agglomerates can contribute to FBAP concentrations in a size range of 2-4 µm (Pöhlker et al., 2012; Healy et al., 2014; Huffman et al., 2010; Gabey et al., 2013; Robinson et al., 2013). Taking previous studies into account, we conclude that FBAP concentrations shown in this study are dominated by fungal spores, but might also include to some extent contributions of other PBAP. A summary of recent WIBS and UV-APS measurements focusing on this issue is already included in section 2.4.

For readers not familiar with the measurement techniques, online measurements alone weaken the robustness of the study. If there are previously published works that validate this measurement with more traditional ways such as filters and genomics, please add them to show that this technique is robust and validated. If not – such validation is needed.

Additional references to previous studies that validate FBAP measurements with traditional techniques are included in section 2.4.

Text added: “At the costal site of Killarney, results of fluorescence and optical microscopy of impacted biological particles reveal that some PBAP, e.g. Spores of Cladosporium spp., which have been frequently observed in many environments, were not correlated to the FBAP concentration (Healy et al., 2014). However, particle size modes of WIBS channel FL2_280 correlate with the concentration of airborne fungal spores commonly observed at the sampling site (Healy et al., 2014).”
In general, the authors should expend the experimental section for instrumentation and validations. For instance, what is the sensitivity of the instruments? This could lead to underestimation of biological particle detection.

Most FBAP measurement data is also shown elsewhere (Schumacher et al., 2013; Toprak and Schnaiter, 2013; Gabey et al., 2011) and some include information about the sensitivity of the instruments. This manuscript is already rather long and its focus is on the comparison of FBAP data to model results. Additional information about the experimental part is contained in other studies (Gabey et al., 2010; Huffman et al., 2010; Pöhlker et al., 2012; Robinson et al., 2013).

P. 10, line 8: this sentence is inaccurate, as not all proteins will contain fluorophore-containing amino acids.

This issue is now taken into account and the sentence has been corrected as follows: “The region of excitation near 270 nm includes certain amino acids (e.g. tryptophan) contained in most proteins.”

The Authors assume that there is no contribution from dust in this size range. Could the authors supply evidence (using back trajectory analysis for example) support this assumption?

Mineral dust also shows slight fluorescence signals which can result in a detected fluorescence fraction in WIBS instruments between <1% and 5.8% (Gallagher, 2014; Toprak and Schnaiter, 2013). It may therefore interfere with the FBAP signal (Pöhlker et al., 2012; Gabey et al., 2011). Contribution of mineral dust to FBAP concentration can only occur during dust events. For the investigated locations and time periods however, no contribution of desert dust is confirmed by using back trajectories (Figure 1 to Figure 3) by using HYSPLIT (Draxler and Rolph, 2013). All locations mostly show local contributions for the selected test cases. Contribution of soil dust may be part of particles from local sources. Soil dust can be internally mixed with fungal spores or other organic components.

Figure 1. Back trajectories for the locations Karlsruhe and Hyytiälä at the simulation case during July 2010
Figure 2. Back trajectories for the locations Karlsruhe and Hyytiälä at the simulation case during October 2010

Figure 3. Back trajectories for the locations Karlsruhe, Hyytiälä, Manchester, and Killarney at the simulation case during August 2010
The two previously published models are based on mannitol concentration, which can also indicate other types of particles, such as vegetation in addition to fungal spores. This may lead to overestimation of bio aerosol loads. This should be mentioned and discussed as one of the factors influencing the difference between models and measurements.

Heald and Spracklen (2009) use mannitol concentrations that are compiled in Elbert et al. (2007) in order to estimate fungal spore emissions which best represent mean observed concentrations. Additional information about PBAP which are also containing mannitol (i.e. bacteria, algae, lichens, and plant fragments) is now added to the manuscript.

The authors correct the spores suspension time in the atmosphere to be 4 ¾ [hours]. This significant claim needs to be validated before stated.

In the updated manuscript version, this statement has been clarified. In order to calculate eq. 7, we need to assume a constant boundary layer residence time, previously called atmospheric lifetime. For a first shot, we used an approximation for atmospheric lifetime found in literature. A first model run with the new emission parameterization (calculated by using $\tau = 1$ day) revealed that the simulated fungal spore concentrations still underestimates the measured FBAP concentrations, although they are adapted to the measurements. We now calculated a corrected boundary layer residence time which is supposed to close the gap between simulated and measured concentrations. This was found to be $\tau = 4 \frac{3}{4}$ hours and is not directly comparable to an atmospheric suspension time or lifetime. Following sentences were also added to the manuscript: “All discrepancies between this boundary layer mixing time and a typical atmospheric lifetime are caused by assumptions which are done for eq. 7. This difference may be caused by deviations from a well-mixed constant concentration profile within the boundary layer (Figure 4), because source and removal processes in the simulation are not in equilibrium and fungal spores are continuously removed at the model boundaries.

However, using eq. 7 for calculating a potential FBAP emission flux is reasonable, because simulated fungal spore concentrations typically decrease rapidly near boundary layer height. This behavior is shown for an exemplary vertical profile in Figure 4.”

![Figure 4. Exemplary vertical profile of simulated fungal spore concentration within and above the planetary boundary layer for Karlsruhe at 28 Aug 2010 14 UTC.](image)
We thank the reviewer for his/her comments which have improved our manuscript. The comments are listed below in italics.

There is insufficient statistical support for the arguments put forth in this study. The authors need to fully quantify model performance for each scheme (R=correlation coefficient and normalized mean bias statistics for Figure 3-7). In doing so, it is critical that the authors distinguish between bias and skill. Does the new scheme actually add to the model skill (i.e. improve R2) or simply eliminate bias? Are there other potential causes for the bias (i.e. LAI, qv, constants used in the model)?

In order to improve the statistical support for this study’s arguments further information on model performance is included in this reply and will be added to the manuscript. The following content to this comment is also added to the manuscript. For each time series, a correlation coefficient (R²) and a normalized mean bias (NMB; Im et al., 2013) have been calculated by

\[ R^2 = 1 - \frac{\sum_{i=1}^{n}(M_i - \hat{O}_i)^2}{\sum_{i=1}^{n}(M_i - \bar{M})^2} \]

\[ NMB = \frac{\sum_{i=1}^{n}(M_i - O_i)}{\sum_{i=1}^{n}O_i} \times 100 \]

which gives an indication for a correlation between simulated fungal spore concentrations (M: N_{H&S}, N_{S&D}, and N_{FBAP}) and measured FBAP concentrations (O: N_{F,c}). Index i represents single elements of each time series. Elements calculated by a linear regression between simulated and measured concentrations are labeled with a hat, the mean value of a time series is labeled with an overbar. Results are listed in Table 1. Please note that the bottom line in Table 1 is not representing a mean value of the numbers above, but represents the result for a combination of all the time series. As noted by referee, different behaviors are found for R² and NMB and therefore it is important to distinguish between skill and bias in further discussion.

<table>
<thead>
<tr>
<th></th>
<th>N_{H&amp;S}</th>
<th></th>
<th>N_{S&amp;D}</th>
<th></th>
<th>N_{FBAP}</th>
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<tbody>
<tr>
<td></td>
<td>R²</td>
<td>NMB</td>
<td>R²</td>
<td>NMB</td>
<td>R²</td>
</tr>
<tr>
<td>Karlsruhe, Jul10</td>
<td>0.0047</td>
<td>-57.54</td>
<td>0.0127</td>
<td>-66.30</td>
<td>0.0048</td>
</tr>
<tr>
<td>Karlsruhe, Aug10</td>
<td>0.0125</td>
<td>-63.82</td>
<td>0.0002</td>
<td>20.14</td>
<td>0.0274</td>
</tr>
<tr>
<td>Karlsruhe, Oct10</td>
<td>0.0125</td>
<td>-35.71</td>
<td>0.0052</td>
<td>-40.08</td>
<td>0.0124</td>
</tr>
<tr>
<td>Hyytiälä, Jul10</td>
<td>0.3268</td>
<td>3.18</td>
<td>0.3358</td>
<td>-67.49</td>
<td>0.3255</td>
</tr>
<tr>
<td>Hyytiälä, Aug10</td>
<td>0.0002</td>
<td>-57.54</td>
<td>0.0099</td>
<td>35.43</td>
<td>0.0511</td>
</tr>
<tr>
<td>Hyytiälä, Oct10</td>
<td>0.0016</td>
<td>-61.74</td>
<td>0.0350</td>
<td>-75.56</td>
<td>0.0000</td>
</tr>
<tr>
<td>Manchester, Aug10</td>
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<td>2.74</td>
<td>0.4274</td>
<td>37.72</td>
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</tr>
<tr>
<td>Killarney, Aug10</td>
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<td>84.42</td>
<td>0.0998</td>
<td>200.51</td>
<td>0.1982</td>
</tr>
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<td>-44.04</td>
<td>0.0551</td>
<td>-28.74</td>
<td>0.1877</td>
</tr>
</tbody>
</table>
The statistical analysis of the results indicates that NMB improves more than $R^2$. Differences in $R^2$ are especially small between $N_{\text{H&S}}$ and $N_{\text{FBAP}}$, because both make use of emission rates as a function of almost the same parameters ($N_{\text{FBAP}}$ includes an additional T-dependence). Parameters $b_1$ and $b_2$ (in eq. 8 of the manuscript) are estimated to give fungal spore concentrations matching best with measured FBAP concentrations. At $T = 275.82$ K, $F_{\text{H&S}}$ is equal to $F_{\text{FBAP}}$, and temperatures above this threshold (as it is the case for almost all locations) shift $F_{\text{FBAP}}$ to give a larger emission flux. At meteorological conditions present for the selected cases, the second part of eq. 8 dominates over the first part by a factor of $\sim 4$ and therefore temperature changes have only a secondary influence on the emission flux. Hence, $R^2$ is similar for both emission parameterizations.

Possible causes for the bias of $F_{\text{H&S}}$ and $F_{\text{S&D}}$ may come from different assumptions made to determine the fungal spore concentrations in ambient air. The mass size distribution of mannitol, which is used as a chemical tracer for fungal spores by Heald and Spracklen (2009), peaks in their study at particle diameters of $\sim 5$ µm. Additionally to fungal spores, bacteria, algae, lichens, and plant fragments, can produce mannitol and some of these can contribute to PBAP concentrations at $\sim 5$ µm. Similar assumptions are made for this study by linking FBAP to fungal spores, but chemical tracers vary between both studies. Furthermore, both literature-based emission fluxes compare local measurements to concentrations simulated on a global scale. Additional biases may arise when using these fluxes on a regional scale.

The modeling in general is not very compelling and it seems like a missed opportunity to investigate/comment on the factors controlling the variability of PBAP. In particular, given the general skill of the model (e.g. Figure 3), it would be useful to try to separate in the model how meteorology and emissions contribute to variability (i.e. perform a simulation with constant emissions). This may be the first time that a model has been compared with high resolution PBAP observations. This is the primary unique direction of this work, but it is insufficiently explored in the current manuscript.

We will expand the discussion on the relative importance of the meteorological and emission-related variability in the manuscript. A time-independent (but spatially varying) emission flux has already been included in the simulations by $F_{\text{S&D}}$. Three locations consist of similar surface ecosystem properties and their emission fluxes are therefore very close to each other ($F_{\text{KA}} = 782$ m$^2$ s$^{-1}$; $F_{\text{Man}} = 837$ m$^2$ s$^{-1}$; $F_{\text{Kil}} = 844$ m$^2$ s$^{-1}$). The results show that diel cycles in the fungal spore concentrations also develop when using $F_{\text{S&D}}$ for emission. In the manuscript, the description is described as follows: “By using a time-independent (but spatially varying) emission flux $F_{\text{S&D}}$, every development in the local temporal pattern arises from meteorological influences. A similar cycle develops between constant ($F_{\text{S&D}}$) and time-independent ($F_{\text{H&S}}$ and $F_{\text{FBAP}}$) simulated fungal spore concentrations, but the order of magnitude differs to varying extend. Therefore and from by visually comparing to simulated boundary layer height, a diurnal cycle in the simulated fungal spore concentrations with a maximum between midnight and sunrise is probably influenced at least partly by boundary layer compression at night.”

1. Page 9907, lines 27-30: Regarding the role of PBAP as IN, the authors may also want to comment on the work of Hoose et al, 2010 and perhaps Creamean et al., 2013 which seem quite relevant here.

Further comments on the role of PBAP acting as IN are now included and additional references are taken into account. The part has changed to:
“These bio-IN may be important for ice nucleation in mixed-phase clouds at temperatures warmer than -15°C (DeMott and Prenni, 2010). In regimes colder than that, mineral dust particles and other ice nucleators are also active and the relative atmospheric abundance of PBAP is probably too small to contribute significantly to formation and evolution of these colder clouds. Previous modelling studies suggest, that bio-IN concentrations are several orders of magnitude lower than IN concentrations from mineral dust or soot and hence the influence of bio-IN on precipitation is limited on the global scale (Hoose et al., 2010; Sesaric et al., 2012; Spracklen and Heald, 2013). In-situ analyses of insoluble cloud ice and precipitation residuals meanwhile highlight the contribution of bio-IN to precipitation, and back trajectories indicate that they can be transported over large distances (Creamean et al., 2013).”

2. Page 9909, lines 1 and 10 seem to contradict each other, FBAP cannot be a lower limit for PBAP if it may be contaminated by other fine particles.

Uncertainties in FBAP measurements include both, a possible underestimation of PBAP as not all PBAP cause a fluorescence signal and some smaller PBAP are excluded by size selection (<1 µm), and a possible overestimation as some non-PBAP contaminate the FBAP signal to some extent. By comparison to traditional measurement techniques for PBAP, it has been found that the former aspect dominated and that FBAP can be considered a lower limit for PBAP (Pöschl et al., 2010).

3. Section 2.1: much of this is section (page 9909, line 25 through all of page 9910) is basic model treatment of aerosols that does not need to be included in a scientific manuscript. I suggest the authors trim this (equations are not necessary).

We understand the reviewer is suggestion to shorten the methodology section, but we think that a brief description of basic aerosol treatment in the model is important, especially as we are addressing readers from both the modelling and the observational communities. Descriptions of dry and wet fungal spore deposition are especially relevant for a sufficient understanding of issues about the atmospheric lifetime of fungal spores (discussed in section 3.2).

4. Page 9911, line 14: typo “gases at particulate” should be “gases and particulate”

Corrected

5. Page 9911, line 23: what are “Anthropogenic primary aerosols”?

Aerosol emissions in PM2.5 and PM10 size mode are disaggregated into chemical components using a split table from TNO (Kuenen et al., 2011). The category “other anthropogenic primary aerosols” represents the remaining, non-carbonaceous primary part, including e.g. minerals, metal oxides, and product emissions (Knote et al., 2011).

6. Section 2.2: line 1 and line 27 incorrectly suggest that the H&S parameterization is a constant emission rate. The authors have described how it is a dynamic function of q and LAI, therefore is by definition, not constant in time or location.

Heald and Spracklen (2009) also used a constant emission rate as a first guess and corrected it afterwards in order to give a better representation of mean mannitol concentrations, which is described in eq. 6 with a function giving $F_{H&S}$. In order to not confuse the reader in future, we decided to ignore this detail and remove the citation in the first place.
7. Page 9913, line 3: Please quantitatively compare the impact of your different size assumption (3 µm) here with Hoose et al., 2010a (5 µm) and Heald and Spracklen, 2009 (PM2.5). Are your totals for mass, number emissions comparable? How did you scale your constant (c) term to account for the smaller size range from Hoose et al.?

A constant $c = 4.6$ is applied to the number emission flux of 3 µm particles in this study in order to have consistent mass emission flux with Hoose et al. (2010) and Heald and Spracklen (2009). The former is obtained by:

$$M_{\text{this study}} = M_{\text{Hoose 2010}} = \frac{\pi}{6} (3 \, \mu m)^3 \rho_{\text{spore}} \left( \frac{5 \, \mu m}{3 \, \mu m} \right)^3 \frac{F_{\text{Hoose 2010}}}{c=4.6}$$

The latter gives a maximum fungal spore emission of $M_{\text{H&S}} \approx 1 \, \text{g/m}^2\text{yr}$ in regions with highest LAI and $q_v$. For the same conditions of LAI and $q_v$, number emission flux from Hoose et al. (2010) gives $F_{\text{H10}} = 500 \, \text{m}^{-2} \text{s}^{-2}$, which matches $M_{\text{H&S}}$ for $\rho_{\text{spore}} = 1 \, \text{g/cm}^3$ (see Figure 3a in Heald and Spracklen (2009)).

8. Page 9913, equation 6: This equation does not explicitly match the parameterization of H&S (unless one assumes that their constant=$c/(\text{LAI} \_\text{max} \, q_v \_\text{max})$. Why is the parameterization given this way here? Is there a physical justification?

We agree that the chosen formulation including an index “max” is confusing and changed it to giving the numbers directly inside the equation.

9. Page 9913, line 11: what is the time resolution for $q_v$ (i.e. the meteorological time step)?

All meteorological parameters (including $q_v$) are written out hourly. When using measured FBAP concentrations for the regression analysis, they are averaged for one hour in order to be consistent to the model output.

10. Page 9913, lines 13-18: This section is confusing. There are 3 sentences that mention the IC/BC for aerosols, and it’s not clear what the authors mean by “No initial and boundary concentrations are predefined for aerosols or gases.” Please clarify or simplify this text.

Simplified in text:

“The COSMO-ART mesoscale model system is driven by initial and boundary data for meteorological conditions. They are updated every six hours and result from interpolation of the coarse grid operational atmospheric model analysis of the ECMWF (European Centre for Medium-Range Weather Forecasts).”

11. Figures 3-6: missing statistical quantification of model performance

See description above.

12. Figures 3-6: all 3 schemes co-vary. To what degree does variability reflect meteorology (PBL height, mixing, and precipitation) rather than variability in emissions driver ($q$, LAI)? This can be diagnosed in the simulation (statement on page 9919 lines 15-16 isn’t quite true. Some of these effects can be deconvolved in the model). What is the correlation between FBAP and PBL height?
By comparing the simulated concentrations from both literature-based emission parameterizations (\( F_{\text{S&D}} = \text{const.} \)), influences of varying meteorological conditions and varying emission drivers are expressed. These effects cannot clearly be separated in the temporal pattern of the FBAP concentrations.

13. Page 9919, line 1: Can you show the observed precipitation from the sites in the Figures?

Exploring the correlation of FBAP concentrations to precipitation is beyond the scope of this study and is described elsewhere (e.g. Schumacher et al., 2013).

14. Page 9920, lines 7-9: Is there any evidence of this phenomenon in the observations you are exploring here?

Some of the FBAP concentrations used in this study are also included in the referenced studies (Huffman et al., 2013; Schumacher et al., 2013). An in-depth analysis of this phenomenon is beyond the scope of this study.

15. Page 9921, line 9-10: Figure 7a does not support the statement that the simulated concentrations are “systematically underestimated”. Nor does Figure 7b demonstrate the improvement suggested on page 9924, line 8. Please show statistics to support these claims.

These effects are difficult to be recognize on a log-scale, but a more detailed statistical analyses is now included.

16. Page 9922, line 3-4: what is the lifetime in the model simulations?

In the model simulation, a local estimate of the fungal spore lifetime calculated as follows:

\[
\tau_{\text{spore}} = \frac{\int_{z=0}^{z_{\text{top}}} N_{\text{spore}}(z) \, dz}{F_{\text{spore}}}
\]

The domain-average fungal spore lifetime is calculated by:

\[
\tau_{\text{spore}} = \frac{\iint F_{\text{spore}}(x,y) \, dx \, dy \, dz}{\iint N_{\text{spore}}(x,y,z) \, dx \, dy \, dz}
\]

\( N_{\text{spore}} \) and \( F_{\text{spore}} \) give the fungal spore number concentration in \( 1/\text{m}^3 \) and emissions flux in \( 1/\text{m}^2 \text{s} \), respectively, and refer to any of the three emission parameterizations. The integral runs from surface model layer (~10 m above ground) to the top-most model layer (at a height of 24 km). The atmospheric lifetime for each time period and location as temporal average are shown in Table 2.

<table>
<thead>
<tr>
<th></th>
<th>Hyttiälä</th>
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<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>July</td>
<td>October</td>
<td>August</td>
<td>July</td>
<td>October</td>
<td>August</td>
<td>July</td>
<td>August</td>
</tr>
<tr>
<td>Lifetime</td>
<td>1.425</td>
<td>1.388</td>
<td>0.748</td>
<td>3.731</td>
<td>6.081</td>
<td>2.801</td>
<td>1.961</td>
<td>1.578</td>
</tr>
</tbody>
</table>

These calculations reveal a strong effect of wet deposition on the simulated lifetime, which is highest for the October case in Karlsruhe (where nearly no rain occurred in the simulation) and shortest for the August case in Hyttiälä (with extensive rain in the simulation).
17. Page 9922, line 7: This lifetime is very short. What physical mechanism could justify such a rapid removal rate?

As indicated in the answers to the previous comments, the model simulates a very strong wet removal of the spores. In addition, calculating the atmospheric lifetime in this simple way assumes equilibrium between local removal and source processes. At the model boundaries, fungal spores are only removed from the domain without fungal spores being transported into the domain. This disagreement may cause an underestimation of the lifetime. Sedimentation is not a major sink for fungal spores, as the average sedimentation velocity for fungal spores is \( v_{\text{sed}} = 0.035 \text{ cm/s} \) (calculated after Helbig et al., 2004), which is typical for 3 \( \mu \text{m} \) particles.

18. Page 9923, equation 8: Are the variables used here independent? Have you verified that you are not over-fitting?

\( q \) is a diagnostic variable of the model and to some extent depends on \( T \), but an additional \( T \)-dependence has been reported for other FBAP-studies (Jones and Harrison, 2004; Di Filippo et al., 2013).

19. Figure 9: Can you overplot the observations using the same assumptions?

Figures 8 and 9 are modified such that an additional circle around each location gives the average measured FBAP concentration / FBAP emission flux colored with the same colorbar as the map.

References


Gabey, A. M., Stanley, W. R., Gallagher, M. W., and Kaye, P. H.: The fluorescence properties of aerosol larger than 0.8 \( \mu \text{m} \) in urban and tropical rainforest locations, Atmos. Chem. Phys., 11, 5491-5504, 10.5194/acp-11-5491-2011, 2011.


Regional-scale Simulations of Fungal Spore Aerosols
Using an Emission Parameterization Adapted to Local Measurements of Fluorescent Biological Aerosol Particles

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Abstract
Fungal spores as a prominent type of primary biological aerosol particles (PBAP) have been incorporated into the COSMO-ART regional atmospheric model, using and comparing three different emission parameterizations. Two literature-based emission rates derived from fungal spore colony counts and chemical tracer measurements were used as a parameterization baseline for this study. A third, new emission parameterization was adapted to field measurements of fluorescent biological aerosol particles (FBAP) from four locations across Northern Europe. FBAP concentrations can be regarded as a lower estimate of total PBAP
concentrations. Size distributions of FBAP often show a distinct mode at approx. 3 µm, corresponding to a diameter range characteristic for many fungal spores. Previous studies have suggested the majority of FBAP in several locations are dominated by fungal spores. Thus, we suggest that simulated fungal spore concentrations obtained from the emission parameterizations can be compared to the sum of total FBAP concentrations. A comparison reveals that parameterized estimates of fungal spore concentrations based on literature numbers underestimate measured FBAP concentrations. In agreement with measurement data, the model results show a diurnal cycle in simulated fungal spore concentrations, which may develop partially as a consequence of a varying boundary layer height between day and night. Measured FBAP and simulated fungal spore concentrations also correlate similarly with simulated temperature and humidity. These meteorological variables, together with leaf area index, were chosen to drive the new emission parameterization discussed here. Using the new emission parameterization on a model domain covering Western Europe, fungal spores in the lowest model layer comprise a fraction of 15% of the total aerosol mass over land and reach average number concentrations of 26 L⁻¹. The results confirm that fungal spores and biological particles may account for a major fraction of supermicron aerosol particle number and mass concentration over vegetated continental regions and should thus be explicitly considered in air quality and climate studies.

1. Introduction

Particles emitted from biological sources are a miscellaneous and omnipresent group of the Earth’s atmospheric aerosols (Elbert et al., 2007; Després et al., 2012). These primary biological aerosol particles (PBAP) can be transported over large distances and their impacts are studied by various fields of research, such as atmosphere science, agricultural research, biogeography and public health (Burrows et al., 2009). PBAP are solid airborne particles of biological origin and include microorganisms or reproductive units (e.g. bacteria, fungi, spores, pollen or viruses) as well as excretions and fragments of biological organisms (e.g. detritus, microbial fragments or leaf debris) (Després et al., 2012). Typical sizes range from < 0.3 µm for viruses to diameters of single bacteria (0.25 - 3 µm), bacteria agglomerates (3 - 8 µm), fungal spores (1 - 30 µm), and up to 10 - 100 µm for airborne pollen (Jones and Harrison, 2004; Shaffer and Lighthart, 1997; Després et al., 2012).
The share of atmospheric aerosol composition belonging to PBAP is large and possibly underestimated (Jaenicke et al., 2007), but is also very uncertain. Estimates of relative PBAP fraction from global models and local measurements reveal large differences between reports. On one hand, the calculated global number concentration of PBAP (zonal annual mean surface concentrations of $10^2 - 10^4$ cm$^{-3}$) is below mineral dust (65 cm$^{-3}$) or soot (1000 cm$^{-3}$) concentrations by several orders of magnitude (Hoose et al., 2010b). Modeling studies yielded global source strengths of ~10 Tg/yr (plant debris and fungal spores, Winiwarter et al., 2009), 56 Tg/yr (Penner, 1995), 78 Tg/yr (bacteria, fungal spores and pollen, Hoose et al., 2010a), 164 Tg/yr (Mahowald et al., 2008) and 312 Tg/yr (bacteria, fungal spores and pollen, Jacobson and Streets, 2009) for different PBAP components. On the other hand, measurements of continental boundary layer air in remote vegetated regions indicate that the mass fraction of PBAP in the coarse particle size range can be as high as ~30% (>0.2 µm, Siberia, Matthias-Maser et al., 2000) or 65-85% (>1 µm, Amazonia, Martin et al., 2010; Pöschl et al., 2010; Huffman et al., 2012).

Like all other aerosol particles, PBAP can influence the Earth’s climate by forcing the radiation budget directly (by absorbing or scattering radiation) and indirectly (by affecting cloud microphysics) (Forster et al., 2007). The direct PBAP effect on climate is difficult to estimate, because evaluations of the atmospheric PBAP concentration vary by several orders of magnitude when taking spatial and temporal divergences into account. Describing the radiative properties of PBAP is complicated, because their size ranges from fine to coarse (up to 100 µm in diameter) and in many cases their shapes are non-spherical and not accurately known. Hence, the applicability of Mie scattering theory is limited (Després et al., 2012). However, the direct PBAP effect on global and regional climate is generally assumed to be small due to low average concentrations, in contrast to the numbers of sub-micron absorbing and scattering aerosols. The indirect PBAP effect on climate is caused by PBAP that act as cloud condensation nuclei (CCN) and/or as ice nuclei (IN). Generally, changing aerosol populations by increasing nuclei concentrations or behavior can alter the microphysical properties of clouds, thus influencing the climate system (Forster et al., 2007). Most PBAP are assumed to be good CCN, because their surface area is large compared to most other aerosol species (Petters and Kreidenweis, 2007; Ariya et al., 2009) and thus may act as so-called giant CCN (Pöschl et al, 2010). Here, the Kelvin effect can be neglected when describing water vapor condensation, and thus activation and growth proceeds quickly (Pope, 2010). Some particles of biological origin (e.g. *P. syringae* bacteria and some fungal species)
have been found to efficiently nucleate ice growth at relatively high temperatures (Després et al., 2012; Murray et al., 2012; Hoose and Möhler, 2012; Morris et al., 2004; Morris et al., 2013; Haga et al., 2013). Biological particles have been observed ubiquitously in precipitation, fog, and snowpack (e.g. Christner et al., 2008), in clouds from airborne measurements (e.g. Prenni et al., 2009; DeLeon-Rodriguez et al., 2013) and have been shown to be important fractions of IN measured at ground level (e.g. Huffman et al., 2013; Prenni et al., 2013). These bio-IN may be important for ice nucleation in mixed-phase clouds at temperatures warmer than -15°C (DeMott and Prenni, 2010). In regimes colder than that, mineral dust particles and other ice nucleators are also active and the relative atmospheric abundance of PBAP is probably too small to contribute significantly to formation and evolution of these colder clouds. Previous modelling studies suggest, that bio-IN concentrations are several orders of magnitude lower than IN concentrations from mineral dust or soot and hence the influence of bio-IN on precipitation is limited on the global scale (Hoose et al., 2010a; Sesartic et al., 2012; Spracklen and Heald, 2013). In-situ analyses of insoluble cloud ice and precipitation residuals meanwhile highlight the contribution of bio-IN to precipitation, and back trajectories indicate that they can be transported over large distances (Creamean et al., 2013). These bio-IN may be important to ice nucleation in mixed-phase clouds at temperatures warmer than -15°C (DeMott and Prenni, 2010; Hoose et al., 2010a; Creamean et al., 2013), and therefore could influence atmospheric radiative properties on up to regional scales. In regimes colder than that, mineral dust particles and other ice nucleators are also active and the relative atmospheric abundance of PBAP is probably too small to contribute significantly to formation and evolution of these colder clouds.

The methods for identifying and detecting PBAP are challenging and many different PBAP can introduce significant detection biases. Particle diameter often plays heavily into PBAP detection and characterization, and it should be noted that large discrepancies can exist between physical and aerodynamic diameter measurements (Huffman et al., 2010; Reponen et al., 2001). PBAP concentrations can be obtained either by online techniques, in which samples are analyzed by advanced instrumentation in real-time, or by offline measurement techniques. If measured offline, samples of airborne biological particles are stored under refrigeration and common methods include analysis by microscopy (stained or unmodified), by cultivation of the sample on growth media, and by amplification and detection of genetic material by sequencing or electrophoretic separation. Chemical and optical properties of PBAP samples or their tracers can be monitored in real time by: chromatography, mass
spectrometry, fluorescence spectrophotometry, LIDAR, and flow cytometry. Short overviews of PBAP analysis techniques have been given by Caruana et al. (2011) and Després et al. (2012).

This paper focuses on the mesoscale simulation of atmospheric concentrations of fungal spores. The COSMO-ART limited-area model is used for the simulations and the setup includes a model domain covering most parts of Europe with a horizontal resolution of 14 km. Two different fungal spore emission parameterizations (Heald and Spracklen, 2009; Sesartic and Dallafior, 2011) are tested by comparing their number concentrations to online laser-induced fluorescence (LIF) measurements of airborne fluorescent biological particles. Additionally, a new emission parameterization adapted to these measurements is introduced. Field data used here comes from a real-time measurement technique that detects the intrinsic (i.e. unstained) fluorescence signal, after UV excitation, of fluorophores commonly present in most biological materials (e.g. free proteins, fungal spores, bacteria, and leaf fragments). Detected particles are categorized as fluorescent biological aerosol particles (FBAP), which may broadly be considered a lower limit for the abundance of PBAP (Huffman et al., 2010; Pöhlker et al., 2012; Healy et al., 2014). FBAP were measured at four different locations (Table 1) concurrently during three focus periods in summer 2010 and fall 2010. The resulting FBAP size distribution is usually dominated by particles in the range from 2 µm to 4 µm, which is consistent with the size of fungal spores (Huffman et al., 2010; Pöschl et al., 2010; Huffman et al., 2012; Healy et al., 2012a; Toprak and Schnaiter, 2013; Huffman et al., 2013). Further, the concentration of FBAP in a given air-mass is generally considered to underestimate PBAP concentration due to biological particles that exhibit very low levels of fluorescent emission (Huffman et al., 2012). To some extent, non-biological aerosol components can also be part of the fluorescence signal for fine particles (~1 µm) (Huffman et al., 2010; Toprak and Schnaiter, 2013). These factors contribute uncertainty to the parameterizations discussed here, however the overall ability of LIF techniques to provide real-time FBAP measurements allows first approximation measurements that can be enlightening.
2. Methodology

2.1. Model Description

The COSMO-ART atmospheric model system is based on the forecast model of the German weather service, combined with an online coupled module for simulating the spatial and temporal distribution of reactive gaseous and particulate components (Vogel et al., 2009). Additionally, fungal spores are incorporated as an independent, monodisperse particle class ($d_p = 3 \mu m$).

Parameterizations for emission, sedimentation, and washout, which were originally developed for pollen dispersal, are included for this particle class (Helbig et al., 2004). Fungal spores are treated independently, as no interactions with other aerosols or gases (coagulation or condensation) are considered. The temporal development of the fungal spore number concentration is calculated by:

$$\frac{d\Psi}{dt} = -\nabla \cdot \vec{F}_T - \frac{\partial}{\partial z} F_S - \lambda \Psi - \frac{1}{N} \frac{\partial}{\partial z} F_E$$  

with the number mixing ratio of fungal spores being

$$\Psi = \frac{N_f}{N},$$

and the number concentration of fungal spores $N_f$, the total number of particles and air molecules $N$ per $m^3$, the air density $\rho$, the turbulent flux $\vec{F}_T$, the sedimentation flux $F_S$, a washout coefficient $\lambda$ and a vertical emission flux $F_E$ (Vogel et al., 2008). The turbulent flux is calculated by $\vec{F}_T = \rho \Psi \Psi'$, incorporating the turbulent fluctuations of wind speed $v'$ and fungal spore number mixing ratio $\Psi'$. Fungal spore sedimentation is calculated by $F_S = \rho \Psi v_s$. The fungal spore settling velocity $v_s$ is calculated by applying the volume-equivalent particle diameter $d_e = 2 \sqrt[3]{a^2 b}$, with $a = 2.1 \mu m$ and $b = 5 \mu m$ (Yamamoto et al., 2012) being the major and minor radius of a prolate spheroid. This results in:

$$v_s^2 = \frac{4 \rho_p d_e g}{3 \rho c_d}$$

where $\rho_p = 1 \text{ g/cm}^3$ is the spore density (Trail et al., 2005; Gregory, 1961) and $c_d$ the drag coefficient (Aylor, 2002). The calculation of the washout coefficient is based on the assumption of raindrops being much larger than aerosol particles and having a much higher terminal fall velocity. It yields:
\begin{equation}
\lambda(d_p) = \int_0^{\infty} \frac{\pi}{4} D_D^2 v_t(D_D) E(d_p, D_D) n(D_D) \, dD_D
\end{equation}

(Rinke, 2008). $D_D$ and $d_p$ are the diameters of raindrops and particles, respectively, $v_t(D_D)$ is the terminal fall velocity, $E$ is a collision efficiency and $n(D_D)$ is the size distribution of the raindrop number concentration. For fungal spores with a spherical diameter of 3 µm, the collision efficiency $E$ with 0.1 mm and 1 mm droplets is approximately 0.085 and 0.3, respectively.

Adapting the model for simulations of fungal spores requires inclusion of an emission flux $F_E$ in the source term of eq. (1) by means of an emission parameterization which will be described in the next section.

Together with fungal spore simulations COSMO-ART is used to compute the mass concentration of major atmospheric aerosol components. Hence, the proportion of fungal spores with respect to the dry aerosol mass can be estimated (section 3.4). In addition to primary aerosol emissions, further gaseous emissions given by the EMPA emission dataset (section 2.3) are taken into account. Partitioning of inorganic aerosol components between the gases and particulate phase is simulated by the ISORROPIA II module (Fountoukis and Nenes, 2007). Condensation on fungal spore aerosols is not included. The contribution of secondary organic aerosols (SOA) to the particles is handled by condensation of oxidized volatile organic compounds as described by Schell et al. (2001). When soot aerosols are not involved as a solid nuclei enabling condensation, clusters build by gas-to-particle conversion via binary nucleation of sulfuric acid and water. They are computed as an individual particle mode. All aerosol particles including these chemical compounds are assumed to be internally mixed. A soot mode without mixing of other chemical compounds is included as particles that are emitted directly into the atmosphere. Anthropogenic primary aerosols (aPA) in the coarse size range (<10 µm) are treated as a separate mode. Detailed descriptions are given in Vogel et al. (2009). Furthermore, sea salt is included in the model simulation and its emission is related to sea water temperature and wind speed (Lundgren et al., 2013). No desert dust emissions are included, as the model domain does not cover the corresponding emission regions and no transport into the model domain is taken into account.
2.2. Emission Parameterization of Fungal Spores

In literature, a constant emission rate was used as input of a global chemical transport model to represent the magnitude and range of measured concentrations of mannitol as a molecular tracer for basidiospores (Elbert et al., 2007). Broad geographical differences can be included in the emission flux by distinguishing between ecosystems. While reviewing the measured data available on measured fungal spore concentrations, Sesartic and Dallafior (2011) calculated number fluxes of fungal spore emissions for six different ecosystems (defined by Olson et al., 2001). Four of these emission fluxes were included into COSMO-ART, and coupled to ecosystem definitions by the GLC2000 (Global Landcover 2000 Database) (forest and shrubs) and Ramankutty et al. (2008) (grassland and crops). The sum of these fluxes, as defined by Sesartic and Dallafior (2011), are emitted from the land area fraction of each ecosystem $i$ ($\sum_{i} E_i = 1$ for $n$ number of ecosystems), gives the total emission flux $F_E = F_{S&D}$ in m$^{-2}$s$^{-1}$ of eq. (1) for fungal spores:

$$F_{S&D} = 214 \text{ m}^{-2}\text{s}^{-1} E_{forest} + 1203 \text{ m}^{-2}\text{s}^{-1} E_{shrub} + 165 \text{ m}^{-2}\text{s}^{-1} E_{grassland} + 2509 \text{ m}^{-2}\text{s}^{-1} E_{crop}$$ (5)

Additionally, a second emission parameterization was tested, which varies as a function of meteorological and surface conditions. Jones and Harrison (2004) reviewed the relations determined when analyzing the observed fungal spore concentrations and atmospheric factors. Seasonal variations can be explained by changes in the leaf area index (LAI). This was verified by correlation to the observed mannitol concentrations. Among the drivers of day-to-day variations, specific humidity ($q_v$) correlates best with the mannitol concentrations (Heald and Spracklen, 2009). It was argued that though other atmospheric factors (e.g. temperature) may actually drive the correlation, this does not change correlation results and thus parameterizations can proceed without having information about the root drivers of fungal spore release. A constant emission rate was used here by is linearly scaled with LAI and $q_v$ in order to give global fungal spore concentrations matching the mean mannitol concentrations (Heald and Spracklen, 2009). In order to fit to the emission flux specified in Hoose et al. (2010a) for a spore diameter of 5 µm, a constant $c$ is set to $c = 2315 \text{ m}^2\text{s}^{-1}$ to be appropriate for fungal spores with 3 µm in diameter. Based on the emission flux in eq. (1), this gives an alternative source $F_E = F_{H&S}$ of fungal spores in m$^{-2}$s$^{-1}$:
\[ F_{H & S} = c \frac{LAI}{5 \, m \, m^{-2}} \frac{q_v}{1.5 \times 10^{-2} \, kg \, kg^{-1}} \]  \hspace{1cm} (6)

\( LAI \) is the leaf area index, \( q_v \) is the specific humidity at the surface, and their scaling factors adapted from tropical rain forest conditions are assumed to be \( LAI_{\text{max}} = 5 \, m^2 \, m^{-2} \) and \( q_v_{\text{max}} = 1.5 \times 10^{-2} \, kg \, kg^{-1} \). In the COSMO-ART simulations \( LAI \) is horizontally distributed according to GLC2000 containing monthly variation and \( q_v \) is provided by the model as a meteorological variable.

### 2.3. Model Domain and Input Data

The COSMO-ART mesoscale model system is driven by initial and boundary data for meteorological and aerosol and chemistry conditions. The meteorological conditions are updated every six hours and result from interpolation of the coarse grid operational atmospheric model analysis of the ECMWF (European Centre for Medium-Range Weather Forecasts). No initial and boundary concentrations are predefined for aerosols or gases. Therefore, all gaseous species are set to a climatological, homogeneously distributed initial concentration. The emission rates for chemical compounds included in the ART part are updated hourly. They are provided by EMPA (Swiss Federal Laboratories for Materials Science and Technology) based on the TNO/MACC (Monitoring Atmospheric Composition and Climate) inventory (Kuenen et al., 2011). The treatment of emissions for COSMO-ART can be found in Knote et al. (2011). Homogeneously distributed mass densities for each aerosol are used as initial conditions, together with the aerosol size distribution and particle density. Primary particle emissions are included as parameterizations based on meteorological and surface conditions. Land use data and constant surface properties are derived from the GLC2000 database (Bartholomé and Belward, 2005). All parameters are post-processed to the rotated spherical coordinate system of COSMO-ART (Doms and Schättler, 2002). For the purpose of this paper, the model domain covers most parts of Western Europe from mainland Portugal to northern Finland, the longitudinal extension being 2849 km the latitudinal extension being 3803 km with a horizontal spacing of 0.125° (\( \cong 14 \) km) on a rotated grid. In vertical direction the model reaches up to an altitude of about 24 km distributed over 40 terrain-following levels. The time stepping of the Runge-Kutta dynamical core is set to 30 s.
2.4. Auto-fluorescence Measurements

Ambient aerosols can be roughly classified as biological or not by interrogating particles at characteristic wavelengths of excitation and measuring the resultant emission in a process called ultraviolet light-induced fluorescence (UV-LIF) (e.g. Hairston et al., 1997; Pan et al., 1999). In particular, the region of fluorescent excitation near 360 nm is often used as characteristic of certain cell metabolites present in all living cells, including riboflavin and reduced pyridine nucleotides (e.g. NAD(P)H). The region of excitation near 270 nm includes certain amino acids (e.g. tryptophan) contained in all proteins. However, many other biological fluorophores exist and the relationship between the measured fluorescence of complex biological particles and fluorophore assignment is very complex (Pöhlker et al., 2012; Pöhlker et al., 2013).

Two instrument types were utilized at four locations for the comparison discussed in this paper. The ultraviolet aerodynamic particle sizer (UV-APS; TSI, Inc., Shoreview, MN, USA) measures particle size aerodynamically, excites individual particles using a single Nd:YAG laser pulse at 355 nm, and detects integrated fluorescent emission (non-dispersed) in a single wavelength region between 420 nm and 575 nm (Hairston et al., 1997; Brosseau et al., 2000; Huffman et al., 2010). The Waveband Integrated Bioaerosol Sensor (WIBS, versions 3 and 4; University of Hertfordshire, UK) measures particle size optically and excites individual particles via two sequential pulses from a Xe-flash lamp, at 280 nm and 370 nm, respectively (e.g. Kaye et al., 2005; Foot et al., 2008). Fluorescence for each particle is then measured in one of two wavelength regions, resulting in three measured fluorescence parameters for each WIBS instrument named FL1_280, FL2_280, and FL3_370. See Gabey et al. (2010) and Robinson et al. (2013) for more details, including slight differences in WIBS-3 and WIBS-4 models. The number concentration of FBAP can be written as $N_{F,e}$ with subscripts referring to fluorescent and coarse particle size. The differences in the pairs of wavelengths used for fluorescence, as well as the possible differences in sensitivity between instruments, suggest that the term “FBAP” as determined by each instrument is not rigorously interchangeable, and it is critical to understand the method of analysis when comparing datasets. For example, the ambient FBAP number concentration as determined by UV-APS has been shown to be qualitatively consistent with the number concentration of particles that fluorescence in the WIBS FL3_370 channel, while the $N_{F,e}$ comparison between UV-APS and WIBS FL1_280 channel is relatively poor (Healy et al., 2014). Here we use the term FBAP from WIBS data to
mean particles that exhibit fluorescence simultaneously in both channels FL1_280 and FL3_370.

Particle size can aide differentiation between biological particles classes observed, however the selectivity based on size alone is very uncertain. For example, and to a rough first approximation, it may be true that many FBAP $\sim 1 \, \mu m$ are single bacterial particles and that many FBAP 2 - 6 $\mu m$ may be fungal spores or bacterial agglomerates (Shaffer and Lighthart, 1997). However, biological species can vary widely, and other FBAP classes (e.g. fragments of larger PBAP, internal components of burst pollen, the presence of other biological species) confound the simple assignment of FBAP based on size (Després et al., 2012).

Further, at least a fraction of fluorescent, supermicron particles are likely to come from non-biological sources, and thus could be counted as FBAP. These non-biological process include anthropogenic sources (e.g. polycyclic aromatic hydrocarbon particles from combustion and cigarette smoke), present most often in submicron particles (Huffman et al., 2010), select oxidized organic aerosol particles (e.g. absorbing brown carbon particles) (Bones et al., 2010; Lee et al., 2013), and some humic-like substances (Gabey et al., 2013). For example, at the rural, elevated site of Puy de Dôme, France, WIBS-3 FBAP measurements were compared to results from fluorescence microscopy paired with staining of fungal spores and bacteria. These results suggest that the real-time UV-LIF measurements indeed track the diurnal cycle of the bacteria concentration, but that non-biological particles still contributed significantly to fluorescent particle number (Gabey et al., 2013).

Virtually every ambient measurement study performed with the WIBS or UV-APS to date has shown a dominant FBAP mode peaking at 2 - 4 $\mu m$ in size (Huffman et al., 2010; Huffman et al., 2012; Huffman et al., 2013; Gabey et al., 2010; Toprak and Schnaiter, 2013; Healy et al., 2014). For example, the FBAP size distributions measured at each of the four sampling locations discussed here is shown in Figure 1, highlighting the common presence of the 2 - 4 $\mu m$ peak. It has been proposed that fungal spores and bacteria agglomerates are the most dominant biological aerosols in this size range (Jones and Harrison, 2004; Després et al., 2012; Fang et al., 2008) and that the FBAP signal in this size range is typically dominated by fungal spores. This was corroborated in more detail for a remote Amazonian site using FBAP analysis along with fluorescence microscopy of stained filter samples (Huffman et al., 2012), but has not yet been rigorously tested in other environments. At the coastal site of Killarney, results of fluorescence and optical microscopy of impacted biological particles reveal that...
some PBAP, e.g. Spores of *Cladosporium* spp., which have been frequently observed in many environments, were not correlated to the FBAP concentration (Healy et al., 2014). However, particle size modes of WIBS channel FL2_280 correlate with the concentration of airborne fungal spores commonly observed at the sampling site (Healy et al., 2014). Other microscopy and DNA-based studies have suggested that fungal spores constitute the largest fraction of PBAP in the 2 – 4 µm size (e.g. Graham et al., 2003; Lin and Li, 1996; Burch and Levetin, 2002). Bauer et al. (2008) showed that fungal spores account for an average of 60% of the organic content in the particulate matter in a size range of 2 - 10 µm in rural and urban areas of Vienna, Austria.
3. Results

3.1. Comparison of Time Series of Measured FBAP and Simulated Fungal Spores

Fungal spore concentrations simulated using the emission flux given in eqs. (5) and (6) according to Heald and Spracklen (2009) and Sesartic and Dallafior (2011) were first compared to FBAP measurements without further adjustment. An overview of time series for all measurements and simulations discussed here is shown in Figure 2 by a box-and-whiskers plot. Time periods for each of three case studies (Table 1) were chosen as exemplary periods when UV-LIF instruments were operating simultaneously at a minimum of two locations, with no requirements applied with respect to environmental conditions. For the statistical analysis, FBAP measurements were averaged over one hour periods in order to be consistent to the model output time steps. For most time periods at Karlsruhe and Hyytiälä the simulated fungal spore concentrations are smaller than the measured FBAP concentrations (Figure 2). This difference is highest at Hyytiälä in August 2010. At Hyytiälä in July and at Manchester and Killarney in August, the Heald and Spracklen (2009) emission ($F_{H&S}$) gives median concentration values which agree reasonably well with the measurements. During October, the fungal spores number concentrations based on constant emission fluxes given by Sesartic and Dallafior (2011) ($F_{S&D}$) agree best with the measured FBAP concentrations. Long-term analysis of FBAP measurements, including periods at the Karlsruhe (Toprak and Schnaiter, 2013) and Hyytiälä site (Schumacher et al., 2013) discussed here, shows an annual cycle of average FBAP number concentrations peaking in summer and lowest in winter. Thus, a simulation based on a constant emission flux may not be appropriate to reproduce the FBAP concentrations.

Figures 3 to 6 show a series of one-week long case studies, each representing two measurement sites. The plots show comparisons between simulation and measurement time series for each station. The simulated fungal spore number concentration is given for the model grid point closest to the measuring site. Due to model spin-up, the first six hours of the simulated fungal spore concentrations are removed from the figures and are not included in the analysis. The total precipitation calculated by the model is shown by gray bars with the ordinate on the right hand side of the figure. The simulated boundary layer height is also included at the bottom of each panel in the figures.
Measured FBAP number concentrations often exhibit distinct diel (24-h) cycles with a maximum in the morning hours or around midnight and a minimum around noon. These features have been consistently reported by most studies discussing temporal behavior of FBAP (Gabey et al., 2010; Huffman et al., 2010; Huffman et al., 2012; Toprak and Schnaiter, 2013). Here, a similar diel cycle is frequently obtained from simulations, and the simulated fungal spore concentrations often anti-correlate with the simulated boundary layer height \(h_{PBL}\) (Figures 3 to 6). The measured FBAP concentrations often qualitatively track the general pattern of simulated \(h_{PBL}\), however the magnitude of concentration change and the timing is often not consistent. For example, on 24 and 25 July at the Karlsruhe site (Figure 3a) a boundary layer compression during the night leads to an increase in the simulated fungal spore concentrations by a factor of ~4, and during day the concentrations decreases as the boundary layer rises again. In this case, the measured FBAP concentrations are in relatively good agreement with the simulated fungal spore numbers, with \(N_{F,C}\) dropping slowly during the day on 24 and 27 July, and to a rate closer to the simulations on 25 July. This suggests that FBAP concentrations were likely influenced, at least partially, by the changing boundary layer height, though diel changes in biological emission are also likely to influence diel FBAP patterns. A similar temporal pattern in simulated fungal spore concentrations is shown in Figure 4a, where a maximum in \(h_{PBL}\) at 12 and 13 October occurs approximately coincident with a minimum in the simulated number concentration. In this case, however, the measured FBAP concentrations do not reflect the diel pattern of the simulations. On 31 August (Figure 5a), measured FBAP and simulated fungal spore number concentration increase simultaneously and parallel to the boundary layer compression, but the increase is more intense for FBAP measurements than for spore simulations. Additionally, at Manchester between 31 August and 1 September (Figure 6a) measured and simulated concentrations are in good agreement. Distinct minima and maxima clearly anti-correlate with the minima and maxima of the boundary layer height. In contrast, during the same time period in Killarney (Figure 6b), small changes in the boundary layer were simulated along with minor changes in fungal spore concentrations. In this case, measured FBAP concentrations qualitatively reflect the same temporal pattern of number concentration, but show poor trend consistency with \(h_{PBL}\). The magnitude of diel FBAP concentration change was similar throughout the week shown, whereas \(h_{PBL}\) showed large diel variations between 26 and 30 August and relatively no change in \(h_{PBL}\) from 30 August to 1 September. Figures 3b, 4b, and 5b also show diel FBAP concentration changes that correlate poorly with simulated \(h_{PBL}\). We conclude that (i)
simulated fungal spore concentrations are sensitive to changes in the simulated boundary layer height, by extension, that (ii) diel cycles of FBAP concentrations are likely to be partially influenced by diel cycles of boundary layer height, but that (iii) the development of the FBAP concentration is in addition influenced by daily cycles in biological emission processes, including those of fungal spores and other PBAP classes. These competing effects are impossible to separate by this analysis.

A comparison of measured FBAP and simulated fungal spore number concentrations for July 2010 is shown in Figure 3. At the measurement site of Karlsruhe, diel cycles were found in the simulated and measured time series, with constantly lower concentrations being obtained from simulations based on emission parameterizations given in literature. When precipitation occurs in the simulation, the simulated fungal spore concentrations decrease due to washout and the diel development of the concentration is interrupted. Afterwards, the simulated concentrations quickly return to the previous baseline. At Hyytiälä a strong decrease in simulated fungal spore concentration on 24 July precisely overlaps with the simulation of precipitation. After hitting a minimum value during simulated precipitation, the simulated fungal spore concentration increases steadily for two days as a result of a post-frontal shift in wind direction and decrease in wind speed. The increase is also reflected in the measured FBAP concentrations. However, the simulated precipitation values do not always coincide with precipitation at the site, as was the case in this instance. As a result of no rain falling at the site on 24 July, the measured FBAP concentration was not affected by the simulated rain.

While this example shows that uncertainty in local meteorology contributes uncertainty to the aerosol output of the model, washout from precipitation remains an important modeled process for estimating FBAP concentrations. Additionally, other dynamic processes are known to affect FBAP concentrations. For example, FBAP has been shown to increase dramatically during rainfall, a process reported recently for both a site in Colorado (Huffman et al., 2013) and also at the Hyytiälä site (Schumacher et al., 2013). The reasons for this FBAP increase are unclear, but are thought to be related to mechanical ejection from terrestrial surfaces as a result of rain droplet splash (Huffman et al., 2013). These effects are known to be dependent on the local geography and ecology, however, and are outside the scope of the presented emission parameterizations.

During the simulation period of October 2010, the simulated fungal spore number concentrations $F_{H&SS}$ are consistently below the measured FBAP concentrations at the sites of
Karlsruhe and Hyytiälä, whereas $F_{SD}$ matches the relative magnitude of the measurements more closely in both cases (Figure 4). At Karlsruhe, concentrations simulated by each emission parameterization follow a distinct diel cycle and increase slightly through the week, reaching concentration maxima on 15 October. The measured FBAP concentration develops differently, with only very weak diel cycle present from 11 to 14 October, and showing little relationship to the simulated $h_{PBL}$, as discussed above.

At the end of August 2010, four different measurement series were available for a comparison to fungal spore simulations (Figure 5 and 6). The measured time series of FBAP number concentrations generally exhibit diel cycles, as discussed. The absolute FBAP concentration at Hyytiälä was consistently highest, when comparing all four sites. This trend is even more obvious when comparing the median concentrations on a linear scale (Figure 2). As a result, concentrations simulated from the literature-based parameterizations under-predict measurements by the greatest margin at Hyytiälä. This under-prediction is likely a result of the persistent precipitation simulated by the model and is an indication that precipitation has a stronger influence on the simulated concentrations than changes in the boundary layer height. Measured rainfall during this period at Hyytiälä was less consistent than the model predicts, but occurred with episodic peaks. In all other August case studies, simulated fungal spore concentrations show relatively good agreement with FBAP measurements.

### 3.2. Development of a Fungal Spore Emission Parameterization by Adaptation to FBAP Measurements

Direct comparison between simulated fungal spores and measured FBAP reveals that in general the simulated concentrations systematically underestimate the measured concentrations (Figure 8a). This difference is most distinct at Hyytiälä during the August case study and at Karlsruhe in the July and October case study. Here we suggest an improved parameterization, including meteorological and surface parameters identified earlier as drivers of fungal spore emissions. Additionally, new parameters driving fungal spore emissions have been investigated. The emission flux depends on these parameters and their fitting coefficients obtained from a regression analysis of the FBAP measurements. The new parameterization for fungal spore emissions has been incorporated into COSMO-ART and the resulting concentrations are included in Figures 3 to 6.
The emission flux from the regression analysis is adjusted to an emission flux $F_{F,c}$ estimated from the FBAP number concentration. For this, it is assumed that particles are evenly distributed throughout the planetary boundary layer and that the simulated fungal spore concentration negatively correlates with $h_{PBL}$. Together with a steady-state condition and neglecting horizontal exchanges with the surrounding air, the balance holds between the number concentration ($N_f$) and the emission rate ($F_{F,c}$) together with the atmospheric lifetime of fungal spores ($\tau$):

$$N_f = \frac{F_{F,c} \tau}{h_{PBL}} \quad (7)$$

(Seinfeld and Pandis, 2006). The boundary layer height at the measurement site needs to be taken from the model simulation as it is not measured consistently. Here, the fungal spore lifetime represents a boundary layer mixing time and is not identical to an atmospheric residence time. For an initial test simulation, $\tau$ is assumed to be constant and is estimated with an initial value of one day, as given in literature for atmospheric lifetimes of aerosol particles with 3 µm in diameter (Jaenicke, 1978). In this case, the fungal spore concentrations with the initial value of atmospheric spore lifetime reveals an underestimation compared to the FBAP measurements. As a remedy, the fungal spore lifetime $\tau$ is corrected to $\tau = 4 \frac{3}{4}$ hours which can be understood as a mean time for boundary layer mixing of fungal spores. The deviation from a lifetime of 3 µm particles given in literature may be attributed to the assumption of a constant vertical distribution of fungal spores with increasing altitude until boundary layer height. However, a ratio of approximately 1.75 between surface-level concentrations and mean concentrations within the boundary layer is too small to explain the discrepancy and fungal spores above boundary layer may stay in atmosphere for longer times in lifetime. All discrepancies between this boundary layer mixing time and a typical atmospheric lifetime are caused by assumptions which are done for eq. (7). This difference may be caused by deviations from a well-mixed constant concentration profile within the boundary layer, because source and removal processes in the simulation are not in equilibrium and fungal spores are continuously removed at the model boundaries. However, using eq. (7) for calculating a potential FBAP emission flux is reasonable, because simulated fungal spore concentrations typically decrease rapidly near boundary layer height. This behavior is shown for an exemplary vertical profile in Figure 7.
Two types of instruments operating with different numbers of channels and detecting fluorescence at different wavelengths are used here for deriving an emission parameterization appropriate for fungal spores. The technical difference may lead to slightly deviating FBAP concentrations (Healy et al., 2014), because the WIBS instrument only counts particles as FBAP when a signal exceeds a threshold in both channels (Pöhlker et al., 2012; Gabey et al., 2010). Some fungal spores most abundant in the Earth’s atmosphere and very common for fungal spores of 2 - 4 µm (*Cladosporium sp.*, *Aspergillus versicolor*, *Penicillium solitum*) (Fröhlich-Nowoisky et al., 2012; Hameed and Khodr, 2001) only show a weak signal in the emission wavelength of 310 nm to 400 nm (Saari et al., 2013; Healy et al., 2014). This difference needs to be taken into account when comparing absolute concentrations of fungal spores and FBAP. During the time periods shown here, the WIBS indicate slightly lower FBAP concentrations than the UV-APS when comparing to the model results. In general, this feature is not always valid and detailed side-by-side comparisons between the two types of instruments are required to determine their behavior in terms of FBAP detection and estimation of the PBAP concentration. In the attempt to factor out the technical difference between the instruments, we assume that FBAP concentration can be multiplied by a constant factor for the concentration values to match each other. The amount of FBAP given by the UV-APS may be represented best by WIBS channel FL3_370 (section 2.4). The FBAP concentration given by the UV-APS is therefore reduced by a factor derived from the WIBS instrument as the mean ratio between channel FL3_370 and the total FBAP concentration $N_{F,c}$ (channels FL1_280 and FL3_370). The factor is estimated to be 2.2 and identical for WIBS data at Karlsruhe and Manchester. The difference is not taken into account in the comparison of the time series in section 3.1, but corrected before applying eq. (7).

Analyzing the meteorological and surface parameters of the model output, it was found that a better correlation with the measured FBAP concentrations is achieved for specific humidity rather than relative humidity, as it was reported for previous field measurements (Gabey et al., 2010; Toprak and Schnaiter, 2013; Di Filippo et al., 2013). During the time period in July 2010, the measured FBAP concentrations vary in a narrow range of specific humidity, which is not reproduced by the literature-based simulation. For this reason, the July case study was removed from the regression analysis. A dependence on the LAI is required in order to take the seasonal change into account and to distinguish among various regions. A combination of
LAI and specific humidity in the regression has the advantage of reducing the fitting parameters. The same relation was chosen by Heald and Spracklen (2009) for the previously discussed emission parameterization. Additionally, surface temperature dependence as suggested by Di Filippo et al. (2013) is indicated by the time series and factored in a regression analysis. The parameters ($b_1 = 20.426$ and $b_2 = 3.93 \times 10^4$) are estimated to be the smallest sum of all squared residuals and result in a multiple linear regression giving an emission flux $F_E = F_{FBAP}$ in $m^{-2} s^{-1}$ for fungal spores fitted to FBAP measurements:

$$F_{FBAP} = b_1 (T - 275.82 K) + b_2 q_v LAI$$

where $T$ is the surface temperature in K, $q_v$ the specific humidity in kg kg$^{-1}$, and $LAI$ the leaf area index in $m^2 m^{-2}$. The parameter inside the parentheses is related to an emission offset of the regression and covers unknown influences. The coefficient $b_2$ is approximately the same as the constants in the Heald and Spracklen (2009) emission for a particle diameter of $3 \mu m$ given in eq. (6). The additional temperature dependence in eq. (8) increases the fungal spore emission for temperatures above 275.8 K and lowers the emission for temperatures below this value.

The multiple linear regression yields a coefficient of determination of $R^2 = 0.4$. By comparing the simulated concentrations (based on $F_{FBAP}$) to the measured FBAP concentrations, it is found that they distribute more evenly along the 1:1 line (Figure 8b). The statistical overview (Figure 2) shows a better agreement between the median concentrations of simulation and measurement for the new emission parameterization than for the literature-based emissions, which is most obvious at Karlsruhe in August and at Hyytiälä in July. The new emission parameterization only slightly reduces the underestimations found for Hyytiälä during August.

Figure 9 shows the emission flux for late August 2010, following the new parameterization, horizontally distributed over a model domain covering Europe. Here, averaged over land areas of the domain, $F_{FBAP}$ gives $1.03 \times 10^3 m^{-2} s^{-1}$. During July and October, the average flux is shifted to $1.4 \times 10^3 m^{-2} s^{-1}$ and $0.4 \times 10^3 m^{-2} s^{-1}$, respectively, mainly as a result of seasonal changes of $LAI$ and $T$.

When analyzing the temporal development of the simulated fungal spore concentrations for each time series, $F_{FBAP}$ mostly results in a slightly higher number concentration than $F_{H&S}$ or $F_{S&D}$ (Figures 3 to 6). This is not the case for October 2010, where the $F_{FBAP}$-concentrations
are in the range of the literature-based concentrations. A sharp decrease on 15 October at Hyytiälä, which is not reflected by the literature-based simulation, is caused by a rapid temperature change. Comparison for the August case study show that simulated $F_{FBAP}$ concentrations agree well with measured FBAP concentrations without overestimating the measurement at Manchester and Killarney, where literature-based simulations and measurements already correspond to each other. Only a slight overestimation can be found at Manchester, which might be due to an urban measuring site that is not represented accurately by the model setup with its broad resolution.

### 3.3. Statistical Correlation between Measured FBAP and Simulated Fungal Spore Concentration

The statistical analysis of the results indicates that normalized mean bias NMB improves more than the correlation coefficient $R^2$ (Table 2). Differences in $R^2$ are especially small between $N_{H&S}$ and $N_{FBAP}$, because both make use of emission rates as a function of almost the same parameters ($N_{FBAP}$ includes an additional $T$-dependence). Parameters $b_1$ and $b_2$ in eq. (8) are estimated to give fungal spore concentrations matching best with measured FBAP concentrations. At $T = 275.82 \text{K}$, $F_{H&S}$ is equal to $F_{FBAP}$, and temperatures above this threshold (as it is the case for almost all locations) shift $F_{FBAP}$ to give a larger emission flux. At meteorological conditions present for the selected cases, the second part of eq. (8) dominates over the first part by a factor of ~4 and therefore temperature changes have only a secondary influence on the emission flux. Hence, $R^2$ is similar for both emission parameterizations.

Possible causes for the bias of $F_{H&S}$ and $F_{S&D}$ may come from different assumptions made to determine the fungal spore concentrations in ambient air. The mass size distribution of mannitol, which is used as a chemical tracer for fungal spores by Heald and Spracklen (2009), peaks in their study at particle diameters of ~5 $\mu\text{m}$. Additionally to fungal spores, bacteria, algae, lichens, and plant fragments, can produce mannitol and some of these can contribute to PBAP concentrations at ~5 $\mu\text{m}$. Similar assumptions are made for this study by linking FBAP to fungal spores, but chemical tracers vary between both studies. Furthermore, both literature-based emission fluxes compare local measurements to concentrations simulated on a global scale. Additional biases may arise when using these fluxes on a regional scale.
3.3.3.4. Contribution of Fungal Spore to Near-surface Aerosol Composition

For a comparison of simulated fungal spores to the dry aerosol chemical composition, the fungal spore mass concentration is calculated from the number concentration assuming monodisperse and spherical particles ($\rho_p = 1 \text{ g/cm}^3$; section 2.1). The horizontally distributed near-surface (approximately 10 m above ground) fungal spore number concentration using $F_{FBAP}$ is shown in Figure 10. Concentrations simulated at the measurement locations are considerably lower than the high surface concentrations in the southern part of the model domain.

The simulated mass concentrations of each chemical aerosol compound are averaged over the land areas of the model domain and the time period of late August 2010 (Figure 11). The total aerosol mass concentration is approximately 2.5 µg/m$^3$. Fungal spores distribute in the domain with an average number concentration of 26 L$^{-1}$ over land. This corresponds to an average mass concentration of fungal spores of 0.37 µg/m$^3$ which accounts for 15.4% of the total simulated aerosol mass. The total aerosol mass excludes mineral dust as one of the main contributors to the chemical aerosol composition, which might lower the fraction of fungal spore mass considerably. A list of mass concentrations of the simulated chemical aerosol compounds, including fungal spores occurring at the measurement site, is given in Table 3. The fraction of fungal spores simulated for these sites varies between 9% and 20%. FBAP mass concentrations calculated from the measured FBAP number concentrations are also listed in Table 3. Here, the same spherical particle diameter and particle density as for the fungal spore simulation is assumed. The FBAP number concentrations are averaged over the same time period as covered by the aerosol simulation. Their share of the aerosol mass ranges from 5% at Manchester up to 64% at Hyytiälä. For Karlsruhe and Killarney, the fractions calculated from the measurements are in good agreement with the fractions resulting from the simulated fungal spore mass concentrations.
4. Discussion and Conclusions

FBAP measurements from four locations in Northern Europe were compared with simulated fungal spore concentrations. Fluorescent particles in the diameter range of 2 - 4 µm are highest in number concentration of FBAP measurements at the rural site near Karlsruhe, Germany (Huffman et al., 2010; Pöschl et al., 2010; Huffman et al., 2012; Healy et al., 2012a; Toprak and Schnaiter, 2013; Huffman et al., 2013). The diameter range for peak FBAP concentration matches closely with the modal size of many species of fungal spores known to be present in airborne concentrations. Simulated fungal spores have been adjusted to match this diameter. Contrary to that, an increase in number concentration towards small particles has been reported for some FBAP measurement series, but only a small fraction of particles could be counted as bacteria cells (Gabey et al., 2011; Huffman et al., 2010).

Comparison of simulations and measurements at four locations and the correlation of FBAP concentrations to meteorological and surface conditions are expected to be most robust when applying identical methods and conditions at all locations. These conditions were not fulfilled in our study. On one hand, site characteristics vary between the stations, which may influence the sensitivity of PBAP emission to surrounding conditions. On the other hand, the measurements are made with different instruments. The measurement series at Karlsruhe, Germany, are done with a WIBS-4 instrument which includes technical improvements compared to the WIBS-3 used at Manchester, UK and Cork, Ireland (Gabey et al., 2010; Healy et al., 2012b). At Hyytiälä, Finland, and Killarney, Ireland, the UV-APS is used to determine the FBAP concentration. This variation may lead to different estimation of the FBAP concentration and within this case study WIBS may report FBAP at lower concentrations than UV-APS at different locations but similar meteorological conditions.

In this paper, fungal spore concentrations are calculated with the COSMO-ART atmospheric model by using literature-based emission parameterizations which adapt simulated global atmospheric concentration to mannitol measurements or spore colony counts (Heald and Spracklen, 2009; Sesartic and Dallafior, 2011). Although mannitol concentration can include contributions from other PBAP (e.g. insects, bacteria, and algae) and from lower plants, the association to fungal spore concentration is reasonable (Di Filippo et al., 2013). Some differences in the comparison may occur from the usage of FBAP concentrations as a representative for fungal spores. Overall, the temporal development of the literature-based
simulated fungal spore concentrations calculated by COSMO-ART approximately reproduces the measured FBAP concentrations.

By using a time-independent (but spatially varying) emission flux $F_{S&D}$, every development in the local temporal pattern arises from meteorological influences. A similar cycle develops between constant ($F_{S&D}$) and time-independent ($F_{H&S}$ and $F_{FBAP}$) simulated fungal spore concentrations, but the order of magnitude differs to varying extend. Therefore and from by visually comparing to simulated boundary layer height, a diurnal cycle in the simulated fungal spore concentrations with a maximum between midnight and sunrise is probably influenced at least partly by boundary layer compression at night. Measured FBAP concentrations are often not consistent to simulated $h_{PBL}$ which suggest that $N_{F,c}$ is additionally influenced by increases in biological emission at night.

The purpose of the work reported here was to develop a new emission parameterization for fungal spores, because literature-based emissions for fungal spores have been found to significantly underestimate measured FBAP concentrations. The parameterization is therefore adjusted to the FBAP concentrations (section 3.2). As was formulated by Heald and Spracklen (2009), it depends on the specific humidity and the leaf area index, but is extended by temperature. The resulting concentrations are in better agreement to the measured FBAP concentrations on the average, but variations in the measurements are not always captured by the simulation. Another long-term analysis of FBAP concentrations and surrounding conditions may result in a further adjustment of the parameters or reveal another parameter driving the emission.

Using the new emission parameterization on a model domain for Europe, fungal spore emission fluxes are extrapolated from northern parts of the domain, where UV-LIF measurements were located, also to Southern Europe. There, much higher emission fluxes occur in the simulation, partially caused by higher specific humidity, which is also the case for $F_{H&S}$, as well by temperature dependence in $F_{FBAP}$. This extrapolation is without local Southern European measurements, however, and thus further UV-LIF measurements are recommended for this region in Southern Europe where fungal spore emission fluxes are potentially greater.

As a result of the relatively low horizontal resolution, small-scale variations influencing fungal spore emission at the measurement sites may not be resolved. Influences on a small scale might be due to an increased amount of fungi for the given vegetation type. When
taking the leaf area index as a surrogate for the vegetation type, uncertainties may result from
an insufficient relation to the presence of fungi or additional surrounding factors favoring
fungi growth. Furthermore, variations in precipitation may not be captured by the model,
which then may lead to improper fungal spore concentrations. The same holds for small wind
gusts and convective cells which may have a strong influence on spore dispersion, but are not
captured well in the model. An increase in fungal spore concentration during or shortly after
rain events (Huffman et al., 2013) could not be reproduced by the simulations, as this effect is
not included in the emission parameterization to an adequate extent.

The module calculating the dispersion of fungal spores does not include all processes of
aerosol dynamics and cloud physics. Of the processes not included, only breaking up of
spores can enhance their number concentration. Coagulation is neglected, as in most cases the
fungus number concentration is low and, hence, their collision is highly improbable. A
coagulation of spores with other aerosol particles is more likely to happen, but not included in
the simulations. Not much is known about the role of fungal spores in clouds and their ability
to act as cloud condensation nuclei.

The simulations presented in this paper highlight the importance of PBAP to the composition
of atmospheric aerosol. Fungal spores, the focus of this paper, are among the main
contributors to PBAP and therefore exert significant influence on aerosol loading. In this
study, COSMO-ART is used up to simulate all major chemical aerosol compounds except for
mineral dust in a domain covering Western Europe. When averaging the mass concentration
horizontally across the land-covered part of the model domain and over all time steps of the
simulation, fungal spores are among the major mass components (Figure 11). However, the
fraction of fungal spores might be overestimated here, as another major aerosol component,
mineral dust, is not included, because the domain does not include any desert dust source
areas. At the selected cases, back trajectories with the HYSPLIT (Draxler and Rolph, 2013)
suggest that transport of Sahara dust into the domain is low. An additional difference in the
treatment of aerosol dynamics implies that spores in the simulation are assumed to be
monodisperse with a diameter of 3 µm without being subjected to sedimentation.

A FBAP mass concentration, estimated from measured FBAP number concentrations
\( (d_p = 3 \, \mu m; \rho_p = 1 \, g/cm^3) \), may reach up to 64% of simulated near-surface chemical aerosol
mass components in rural areas of Finland (Table 3). In comparison to relations of PBAP to
total aerosol concentrations given in literature, their volume fraction of particles larger than
0.2 µm during one year of measurements at a remote site in Siberia reaches 28% on the average and at Mainz, Germany the volume fraction amounts to 22% (Matthias-Maser et al., 2000). Both of these fractions agree well with simulated mass fractions of this study for comparable locations, but simulated concentrations given in this study are much lower than total number concentrations given in Matthias-Maser et al. (2000). In contrast, the number and mass fractions in the Amazonian basin are above 80% and therefore much higher than in the highlighted urban and remote areas (Pöschl et al., 2010), but here the absolute concentrations are less and therefore in the order of magnitude given by the simulation of this study.

PBAP and especially fungal spores might account for a major part of the aerosol loading. Locally, a correlation between increasing FBAP and ice nuclei number concentration (Tobo et al., 2013) shows that future model studies of PBAP impacts on clouds are needed to determine their relevance to atmospheric ice nucleation.

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Table 1. Overview of the measurement sites, including their geographical location and the types of instrument used (d<sub>p</sub> corresponds to the optical particle diameter and d<sub>a</sub> to the aerodynamic particle diameter). The sections below show the simulation periods and the availability of data at this site (filled dot). Mean values for the simulated meteorological and surface conditions used for the new emission parameterization (section 3.2) at the measurement site during the corresponding time periods are added to each section.

<table>
<thead>
<tr>
<th>location</th>
<th>Karlsruhe, Germany</th>
<th>Hyytiälä, Finland</th>
<th>Manchester, UK</th>
<th>Killarney, Ireland</th>
</tr>
</thead>
<tbody>
<tr>
<td>coordinates</td>
<td>49° 5' 43.6&quot; N 8° 25' 45.0&quot; E</td>
<td>61° 50' 41.0&quot; N 24° 17' 17.4&quot; E</td>
<td>53° 27' 57.0&quot; N 2° 13' 56.0&quot; W</td>
<td>52° 3' 28.0&quot; N 9° 30' 16.4&quot; W</td>
</tr>
<tr>
<td>altitude</td>
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<td>152 m a.s.l.</td>
<td>45 m a.s.l.</td>
<td>34 m a.s.l.</td>
</tr>
<tr>
<td>instrument</td>
<td>WIBS-4</td>
<td>UV-APS</td>
<td>WIBS-3</td>
<td>UV-APS</td>
</tr>
<tr>
<td>size range</td>
<td>0.8≤d&lt;sub&gt;p&lt;/sub&gt;≤16 µm</td>
<td>1&lt;d&lt;sub&gt;a&lt;/sub&gt;≤20 µm</td>
<td>0.8≤d&lt;sub&gt;p&lt;/sub&gt;≤20 µm</td>
<td>1&lt;d&lt;sub&gt;a&lt;/sub&gt;≤20 µm</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>LAI (m&lt;sup&gt;2&lt;/sup&gt;/m&lt;sup&gt;2&lt;/sup&gt;)</th>
<th>mean T (°C)</th>
<th>mean q&lt;sub&gt;v&lt;/sub&gt; (kg/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>22 July 2010 - 28 July 2010</td>
<td>3.18</td>
<td>17.3</td>
<td>0.0088</td>
</tr>
<tr>
<td>26 August 2010 - 01 September 2010</td>
<td>2.94</td>
<td>16.6</td>
<td>0.0099</td>
</tr>
<tr>
<td>11 October 2010 - 21 October 2010</td>
<td>1.49</td>
<td>6.5</td>
<td>0.0055</td>
</tr>
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</table>
Table 2. Correlation coefficient ($R^2$) and normalized mean bias (NMB) for correlations between fungal spore and FBAP concentrations at different locations and three different time periods

<table>
<thead>
<tr>
<th>Location, Month</th>
<th>$R^2$</th>
<th>NMB</th>
<th>$R^2$</th>
<th>NMB</th>
<th>$R^2$</th>
<th>NMB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Karlsruhe, Jul10</td>
<td>0.0047</td>
<td>-57.54</td>
<td>0.0127</td>
<td>-66.30</td>
<td>0.0048</td>
<td>-31.86</td>
</tr>
<tr>
<td>Karlsruhe, Aug10</td>
<td>0.0125</td>
<td>-63.82</td>
<td>0.0002</td>
<td>20.14</td>
<td>0.0274</td>
<td>-34.68</td>
</tr>
<tr>
<td>Karlsruhe, Oct10</td>
<td>0.0125</td>
<td>-35.71</td>
<td>0.0052</td>
<td>-40.08</td>
<td>0.0124</td>
<td>1.36</td>
</tr>
<tr>
<td>Hyytiälä, Jul10</td>
<td>0.3268</td>
<td>3.18</td>
<td>0.3358</td>
<td>-67.49</td>
<td>0.3255</td>
<td>50.50</td>
</tr>
<tr>
<td>Hyytiälä, Aug10</td>
<td>0.0002</td>
<td>-57.54</td>
<td>0.0099</td>
<td>35.43</td>
<td>0.0511</td>
<td>-59.04</td>
</tr>
<tr>
<td>Hyytiälä, Oct10</td>
<td>0.0016</td>
<td>-61.74</td>
<td>0.0350</td>
<td>-75.56</td>
<td>0.0000</td>
<td>-45.97</td>
</tr>
<tr>
<td>Manchester, Aug10</td>
<td>0.4558</td>
<td>2.74</td>
<td>0.4274</td>
<td>37.72</td>
<td>0.4409</td>
<td>63.26</td>
</tr>
<tr>
<td>Killarney, Aug10</td>
<td>0.2408</td>
<td>84.42</td>
<td>0.0998</td>
<td>200.51</td>
<td>0.1982</td>
<td>213.81</td>
</tr>
<tr>
<td>all</td>
<td>0.1834</td>
<td>-44.04</td>
<td>0.0551</td>
<td>-28.74</td>
<td>0.1877</td>
<td>-0.43</td>
</tr>
</tbody>
</table>
Table 3. Simulated aerosol mass concentrations for aerosol chemical components, including fungal spores, together with measured FBAP values in µg/m³ at the measuring sites as averages over the time period during August 2010

<table>
<thead>
<tr>
<th>Particle Mass (µg/m³)</th>
<th>Karlsruhe, Germany</th>
<th>Hyytiälä, Finland</th>
<th>Manchester, UK</th>
<th>Killarney, Ireland</th>
</tr>
</thead>
<tbody>
<tr>
<td>measured FBAP</td>
<td>0.46</td>
<td>0.81</td>
<td>0.19</td>
<td>0.19</td>
</tr>
<tr>
<td>simulated fungal spores</td>
<td>0.41</td>
<td>0.20</td>
<td>0.35</td>
<td>0.28</td>
</tr>
<tr>
<td>sea salt</td>
<td>0.44</td>
<td>0.01</td>
<td>1.62</td>
<td>1.11</td>
</tr>
<tr>
<td>soot</td>
<td>0.19</td>
<td>0.06</td>
<td>0.42</td>
<td>0.04</td>
</tr>
<tr>
<td>SO₄²⁻</td>
<td>0.18</td>
<td>0.01</td>
<td>0.11</td>
<td>0.05</td>
</tr>
<tr>
<td>NH₄⁺</td>
<td>0.44</td>
<td>0.01</td>
<td>0.14</td>
<td>0.07</td>
</tr>
<tr>
<td>NO₃⁻</td>
<td>1.29</td>
<td>0.01</td>
<td>0.34</td>
<td>0.18</td>
</tr>
<tr>
<td>SOA</td>
<td>0.41</td>
<td>0.24</td>
<td>0.14</td>
<td>0.04</td>
</tr>
<tr>
<td>aPOA</td>
<td>0.67</td>
<td>0.13</td>
<td>0.85</td>
<td>0.11</td>
</tr>
</tbody>
</table>
Figure 1. Average FBAP size distributions derived from UV-LIF measurements during case studies in August 2010 at Karlsruhe (Germany), Hyytiälä (Finland), Manchester (UK), and Killarney (Ireland).
Figure 2. Box-whisker plots of: measured hourly FBAP concentration (open boxes), simulated fungal spore concentration emitted by \( F_{H&B} \) (horizontally hatched boxes), \( F_{FBAP} \) (filled boxes), and \( F_{S&D} \) (vertically hatched boxes) for all case studies. The central mark of each box shows the median, its edges the 25\(^{th}\) and 75\(^{th}\) percentiles, and whiskers show 5\(^{th}\) and 95\(^{th}\) percentiles. Dotes above whisker show outliers (>95\(^{th}\) percentile).
Figure 3. Time series of measured FBAP and simulated fungal spore number concentrations in l/L together with simulated precipitation in mm/h (right axis) and simulated boundary layer height in km (right axis) during the case study from 22 July to 28 July 2010 at (a) Karlsruhe, Germany and (b) Hyytiälä, Finland. Simulations were performed with three different emission parameterizations: $F_{H&S}$ from Heald and Spracklen (2009); $F_{S&D}$ from Sesartic and Dallafior (2011); $F_{FBAP}$ from this study.
Figure 4. Time series of measured FBAP and simulated fungal spore number concentrations in 1/L together with simulated precipitation in mm/h (right axis) and simulated boundary layer height in km (right axis) during the case study from 11 October 2010 to 21 October 2010 at (a) Karlsruhe, Germany and (b) Hyytiälä, Finland. Simulations were performed with three different emission parameterizations: $F_{H&S}$ from Heald and Spracklen (2009); $F_{S&D}$ from Sesaric and Dallafior (2011); $F_{FBAP}$ from this study.
Figure 5. Time series of measured FBAP and simulated fungal spore number concentrations in 1/L together with simulated precipitation in mm/h (right axis) and simulated boundary layer height in km (right axis) during the case study from 26 August 2010 to 01 September 2010 at (a) Karlsruhe, Germany. (b) Hyytiälä, Finland. Simulations were performed with three different emission parameterizations: $F_{H&S}$ from Heald and Spracklen (2009); $F_{S&D}$ from Sesartic and Dallafior (2011); $F_{FBAP}$ from this study.
Figure 6. Time series of measured FBAP and simulated fungal spore number concentrations in 1/L together with simulated precipitation in mm/h (right axis) and simulated boundary layer height in km (right axis) during the case study from 26 August 2010 to 01 September 2010 at (a) Manchester, UK and (b) Killarney, Ireland. Simulations were performed with three different emission parameterizations: $F_{HS}$ from Heald and Spracklen (2009); $F_{SD}$ from Sésaric and Dallafior (2011); $F_{FBAP}$ from this study.
Figure 7. Exemplary vertical profile of simulated fungal spore concentration within and above the planetary boundary layer for Karlsruhe at 28 Aug 2010 14 UTC.
Figure 8. Comparison for all case studies: Measured FBAP number concentrations plotted versus simulated fungal spore number concentrations (a) based on Heald and Spracklen (2009) emission flux and (b) based on emission parameterization derived from a multiple linear regression to FBAP concentrations. Solid black lines represent the 1:1-line, dashed lines the 1:2-line and dotted lines the 1:10-line.
Figure 9. Average simulated fungal spore emission flux \( F_{FBAP} \) in \( m^2s^{-1} \) from 26 August to 01 September 2010, excluding a spin-up period of 6 hours. White circles indicate the locations of the different FBAP measurement time series and the color within the white circles represents the mean emission flux calculated from FBAP measurements at each location.
Figure 10. Horizontally distributed fungal spore concentration in $1/L$, emitted by $F_{FBAP}$, in the lowest model layer, averaged from 26 August to 01 September 2010 (excluding a spin-up period of 6 hours). White circles indicate the locations of the different FBAP measurement time series and the color within the white circles represents the mean FBAP number concentration measured at each location.
Figure 11. Near-surface chemical aerosol mass composition simulated by COSMO-ART, horizontally averaged over the land area in the model domain and temporally averaged from 26 August to 01 September 2010 (excluding a spin-up period of 6 hours)