RESPONSES TO ANONYMOUS REFEREE #2

(1) Reviewer comments are in black text.
(2) Author responses are in blue text.
(3) Additions/modifications made to the manuscript.

We thank the reviewer for their detailed comments and helpful recommendations, which are addressed in the following responses.

The manuscript concerns the observations of bioaerosols using WIBS-3 and WIBS-4 and their main classification into types of bioaerosols using clustering, their typical patterns and potential sources, where the source analysis has been done with ArcGIS. The observational period cover 4 sites in the UK and the observations are Jan-March 2009, June-August 2013 (WIBS-4), Feb-June 2013 (WIBS-4) and August 2009 (WIBS-3). Please find below a numbered set of comments to the manuscript as well as one specific comment

1. The manuscript cover an area there is of relevance to ACP and an area where there is very few studies.

2. The study itself contains a large data set, about 9 months of data of bioaerosols obtained with WIBS instruments. However the data itself are not part of the manuscript, but only coarse numerical summaries.

We would like to re-affirm that we will provide access to the raw data as per Copernicus data sharing guidelines, but focus on the scientific elements of our analysis in this paper.

3. The conclusions in the manuscript are mainly related to clustering of data into 4 or 5 main clusters and there are some indication to potential source areas. These conclusions seem however to be indicative and qualitative instead of quantitative.

We would like to clarify that the cluster procedure provides and quantitative analysis of the UV-LIF spectral data but we have qualitatively assigned these clusters to potential bio-aerosol types using a standard comparison of fluorescence profiles. As we note in response to reviewer 1, the fluorescent signals of ambient derived clusters have been compared to laboratory data using trends in fluorescent channels in order to initially group the 18 clusters for further analysis. Specifically, the clusters from each field site were compared to existing (‘Dstl experiment 2014’) and new (‘Dstl experiment 2017’) laboratory data, depending upon the instrument used, in addition to published data e.g. Savage et al 2017. Laboratory data was available from a WIBS-3 for the 2014 Dstl dataset, and from a WIBS-4 for the 2017 Dstl dataset (the results from such are to be published in 2019). Prior to comparing the ambient clusters to the laboratory data according to broad fluorescent signature, the process for deriving these clusters was the same as used in all previous ambient studies. Additionally, data from published laboratory experiments (e.g. Savage et al 2017, Hernandez et al
2016) were used to provide some further support, and aid the initial grouping of these clusters into suspected particle type groups.

4. The scientific methods are valid and clearly described

5. The results and the methods are not described in depth to reach the current conclusion. This relates to both clustering and the mapping using ArcGIS. See issues below

We apologise if the reviewer feels this is the case, and we hope our detailed responses given here and the changes to the manuscript address this point.

5a. Issues on mapping: There is no exact geographical location of the sites. Please add this to the manuscript

We have added the geographic location of each site to Table 1 in section 2.3.

5b. Issues on mapping: I could identify the Weybourne observatory and compared Figure 1 with both google maps and land cover 2015 (Digimap). The land cover in Fig 1 shows large amounts of Coniferous woodland near Weyborne. However Googlemaps and Digimap shows that this area is improved grassland. Is this a simple mapping mistake when drawing figure 1 or is there a more systematic mistake in the manuscript, where the land cover has not been used correctly for all the sites?

It is correct that there is no coniferous woodland around the Weybourne site, however, the land cover map used shows that the area around the Weybourne site comprises improved grassland, not coniferous woodland. This may be unclear owing to the two shades of green used to identify improved grassland areas and coniferous woodland. To resolve this, a clearer distinction has been made between the two shades of green, with a much lighter green colour to represent improved grassland, and a darker shade of green for the coniferous woodland (Figure 1).
5c. Issues on mapping. Several times in the manuscript including the conclusion there is a connection between the observations and specific farming activities. However the manuscript does not contain any information about farm location. This connection can therefore not be made unless such data are present. Furthermore, why have those specific farms been attributed as source and not other farms in the area?

There are particular sampling sites which are in close proximity to a dairy factory (Davidstow) and a mushroom/composting facility (Chilbolton) which were observed during each experimental campaign within narrow wind sectors. Due to the nature of the land cover maps, details such as farm locations/nearby areas of interest are not included. As such, there was some consideration given to include a separate map of each site (either a simple OS basemap or Satellite imagery) outlining any potential influencing sources. However, adding such extra maps would add substantially to the size of the manuscript. As an alternative, altering the transparency of the land cover map and overlaying this upon an OS basemap has allowed for further information of the surrounding area at each site to be presented (Figure 2).

Additionally, ‘Potential point sources’ have been added to the maps from each site and have been split into ‘Farming sites’ and ‘Other potential influencing sources’ as identified during each campaign and with the use of online maps (e.g. Google Maps). Apart from the dairy factory at Davidstow, this site now features other potential point sources, including a dairy farm and other surrounding farms, a small garden centre, and a slaughterhouse which are in close proximity to the sampling site. Additionally, the presence of a livestock breeder near to the MST Capel Dewi site has been added. A description of these influencing sources has been added to the text.
With regards to the changes made in the manuscript, the land cover maps from each site have been overlaid on OS basemaps to provide geographical context to each site location. Additionally, potential influencing sources have been added to illustrate farming activity or any other potential sources of biological or interferent particle material, which can be seen in Figure 2 (Figure 1 in the manuscript).

A sentence has been added to Section 2.1, paragraph 3, to explain this change -

“To provide geographic context to each site, the LCM2015 has been overlaid on an OS basemap. Additionally, the presence of local farming activity and other potential influencing sources are illustrated in Figure 1”

Figure 2 – LCM layer overlaid on OS basemaps, potential point sources are split into farming sites and ‘others’, illustrated by black/white asterisks respectively.
5d. Issues on mapping. The chosen land cover map is probably among the best maps in the UK. However, it has some limitations. Smaller features such as smaller woodlands are not part of this map. The authors have not taken these limitations into account.

The chosen land cover map does have its limitations; however, the advantages do outweigh these, especially as this is an up-to-date land cover map. The method in which the land cover map is produced does mean that smaller features are excluded from the map, which is noticeable when comparing a basemap to the LCM. A sentence will be added to the manuscript to ascertain that there may be some smaller features which have been excluded.

The statement below has been added to Section 2.1, paragraph 3.

“Though the LCM2015 provides up-to-date data on the land cover characteristics of each site, it is acknowledged that smaller features are not identified and thereby not considered as potential sources.”

5e. Issues on clustering. The clustering uses an approach by Crawford et al. (2015). This requires use of dry materials that are aerosolised and added to the instrument in a laboratory. This calibration data is not present in this paper. We refer the readers to Crawford et al (2015) and noted data access procedures in the paper for the calibration data used in that study. Whilst the use of hierarchical agglomerative clustering (HAC) correctly attributed 98% of laboratory generated fluorescent test particle data, its limitation on more representative biological aerosol is noted in our latest study, and work on alternative methods is currently ongoing (e.g. Ruske et al 2018) following new laboratory studies before alternative recommendations can be made.

5f. Issues on clustering. The paper by Crawford et al. (2015) only describe pollen but not if other bioaerosols have been used. This is correct, the Crawford 2017 paper does not describe laboratory data other than the pollens used. When comparing the fluorescent signature of Cluster 1 and Cluster 3 from Weybourne, in which there is a greater fluorescent signature in channel FL3, the Crawford et al 2017 reference (Page 10, line 9) used is misleading as this infers that there was laboratory data to compare to.

Instead, in the Crawford 2017 paper, Cluster 2 is strongly fluorescent in FL3, similar to Cluster 1 and Cluster 3 from Weybourne (Table 1). However, when considering the size (7.7µm) and shape (Af, 20) of Cluster 2 from Crawford et al 2017, it was speculated that this may represent a bacterial aggregate or a larger dust particle containing uncharacterised bacteria. As per existing bioaerosol studies, cluster profiles were compared to results from Hernandez et al 2016 and Savage et al 2017, which both show a strong FL1 signature for bacteria.
Table 1 - Comparison between Weybourne Cluster 1 and Cluster 3, and Cluster 2 (Halley) from Crawford et al 2017.

<table>
<thead>
<tr>
<th>CL</th>
<th>FL1</th>
<th>FL2</th>
<th>FL3</th>
<th>D(µm)</th>
<th>AF</th>
<th>% Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weybourne 1</td>
<td>3.8 ± 25.2</td>
<td>12.5 ± 35.6</td>
<td>303.6 ± 295.0</td>
<td>5.0 ± 2.2</td>
<td>36.6 ± 15.9</td>
<td>33.7</td>
</tr>
<tr>
<td>Weybourne 3</td>
<td>5.1 ± 27.3</td>
<td>3.2 ± 15.8</td>
<td>192.6 ± 200.8</td>
<td>2.0 ± 1.1</td>
<td>17.2 ± 8.7</td>
<td>50.6</td>
</tr>
<tr>
<td>Halley 2</td>
<td>135.8 ± 227.4</td>
<td>172.1 ± 185.1</td>
<td>765.6 ± 535.9</td>
<td>7.7 ± 4.0</td>
<td>19.9 ± 9.2</td>
<td>2.1</td>
</tr>
</tbody>
</table>

To clarify, the Crawford et al 2017 reference has been removed from this sentence. Instead, a new sentence has been added to the manuscript (Section 3.2.1, paragraph 1) illustrating the fluorescent profile of Cluster 2 from the Antarctic site -

“.......Laboratory data collected using a WIBS-3 have shown that bacteria such as unwashed E-Coli and Bacillus atrophaeus (BG) spores exhibit higher fluorescence in channel FL3 (Dstl Experiment 2014). The presence of a highly fluorescent FL3 channel was found for Cluster 2 in Crawford et al 2017 which was used to infer a bacterial particle, or dust containing bacteria, as a result of the larger size and shape of the particles in this cluster. This is contrary to other studies which have found a strong FL1 signature for bacteria (Hernandez et al., 2016; Savage et al., 2017).”

.......Crawford et al. writes that the four pollen types are common in the UK. This is not correct. Two of the four allergens (paper mulberry and ragweed) are rare in the UK. The third in Crawford (birch) is common in the UK, typically with a season in April. This suggest that in this manuscript only Capel Dewi would have had a chance to detect this. The fourth pollen in Crawford et al (2015) is ryegrass......

We note that paper was a limited study. To clarify our results, of the total 18 clusters, following initial comparison of fluorescent signatures from laboratory data to fluorescent channel responses from each cluster, 3 clusters were considered to be pollen fragments (Section 3.2). These were Cluster 1 from Davidstow, Cluster 4 from Weybourne, and Cluster 4 from Chilbolton.

In the manuscript fluorescent signals from three different pollens sampled using a WIBS-3, during the Dstl 2014 experiment, comprising Ryegrass (as in Crawford et al 2017), Aspen, and Poplar pollen. In Crawford et al 2017 ‘four typical pollens’ birch, paper mulberry, ragweed, and ryegrass were used from the sample set.

Two different tree pollens were selected for comparison to the cluster data, these being Aspen pollen, a part of the poplar family, which was selected due to its status as a native tree species to the UK and parts of Europe. Poplar pollen was also selected as was the grass pollen, Ryegrass. This study did not include Paper Mulberry, Ragweed, or Birch as in Crawford et al (2017).
Grass pollens are common from mid-May to July (Met Office 2018) of which the sample period from Chilbolton falls too early (20\textsuperscript{th} January to 20\textsuperscript{th} March) and the sample period from Weybourne falls too late (17\textsuperscript{th} August to 25\textsuperscript{th} August). However, at Davidstow, data collection was conducted from 25\textsuperscript{th} June to 28\textsuperscript{th} August, which is well within the grass pollen season.

Tree pollens, in particular Poplar, are common from around mid-March to early-April (Met Office 2018). The sample period from Weybourne and Davidstow occurs too late in the year for it to be affected by this pollen type. However, at Chilbolton the sample period covers a small part of this period towards the end of the sampling campaign.

Birch pollen was not included in this manuscript, however, as a common UK tree species, it is common from late March to around the middle of May (Met Office 2018). The time period of such would most likely just miss the end of the sampling period for Chilbolton, and start before the sampling period at Davidstow and Weybourne.

….However, the pollen size is typically 30-40, which is above the typical detection limit of the WIBS.

In Appendix A of Crawford et al (2017) the 'four typical pollens' were clustered, and the cluster which accounted for ~70% of the fluorescent material was considered to be representative of the sampled pollens. The average size of this group of these pollen particles (11.8\textmu m) is within the detection range of the WIBS and provided confirmation that ambient Cluster 4 (8.1\textmu m) within the Halley dataset was a potential representative of pollen.

The size of the particles in the manuscript from Cluster 1 in Davidstow (16.5\textmu m), Cluster 4 from Weybourne (4.4\textmu m), and Cluster 4 from Chilbolton (2.4\textmu m) were considered to be pollen fragments owing to the size of these particles being smaller than that of intact pollens. Specifically, pollens such as Birch can range from 24\textmu m to 28\textmu m (Detweiler and Hurst 1930), which would not be detectable by a WIBS. Therefore any clusters with similar spectral signatures which were assumed to be pollen, were considered pollen fragments, as the WIBS would not be able to sample larger sized intact pollen grains, but would be able to detect fragments as described by Savage et al 2017.

Have the authors also calibrated with pollen and have they also used pollen that are less likely to be in the UK atmosphere and less likely to be detected by the WIBS?

As discussed above, the pollens used from the WIBS-3 data (Poplar, Aspen, and Ryegrass) were chosen following research into the different pollen type abundance in the UK, and were selected, as they are most likely to be in the UK atmosphere. These pollens would be likely detected by a WIBS, for example in Savage et al (2017) 14 intact pollens and 13 pollen fragments were sampled and displayed significant fluorescence above the set threshold when using a WIBS-4A.
5g. Issues on clustering. In the paper by Crawford et al (2015) the team has used dry pollen. Dry pollen from commercial samples will have a very different shape to fresh airborne pollen as pollen can take up and loose water. Using dry pollen will generally cause poor calibration of real-time instruments as the shape of dry pollen is very different compared to fresh pollen.

Dry pollen was used within this manuscript (referred to as Dstl Experiment 2014). The authors understand that commercial samples will not necessarily be representative of ambient pollens or other biological material within the atmosphere. Further work comparing the fluorescent signals of ambient and laboratory data of the same particle type is to be conducted in the future.

Secondly has there been any investigations if dry pollen will cause different excitation compared to fresh pollen?

Additional work has been recently conducted (cited as Dstl Experiment 2017 in the manuscript) using dry pollen to compare fluorescence profiles between multiple UV-LIF instruments. As we note in response to reviewer 1, we are planning on conducting a much more thorough evaluation of statistical methods once we have published and had this data appropriately peer reviewed.

6. The methods section are generally good if the issues in section 5 can be solved

We thank the reviewer for this supportive comment and hope that the above has addressed all aforementioned concerns.

7. The citations and reference list seems to be up-to-date with a good selections of citations to new and relevant literature. However the manuscript is not clear where the studies confirms existing knowledge and more importantly where it contributes with new knowledge by positioning the results against published literature

As the reviewer already notes, there are very few studies in this area. This manuscript has aimed to characterise biological particles following HAC analysis, whilst assessing sensitivity to a new recommended threshold, following Savage et al 2017. This paper highlights the method of characterising clusters by firstly comparing the fluorescent profiles to laboratory data, secondly, by assessing the clusters diurnal variation (following grouping as based on their fluorescent profiles), and thirdly, characterising these clusters as based on the response to temperature and relative humidity.

In particular, results were compared to the Hernandez et al 2016 laboratory data study in which different particle types were sampled by a WIBS-4, which was used to help classify the different clusters into groups, with an inferred cluster type. Those in Group 6 were considered to be interferent particles, given the higher FL2 fluorescent profile of these clusters (as found in Hernandez et al 2016). However, it was concluded that Group 6 was not consistent with
interferent particles (as illustrated by Figure 4 in the manuscript) in which of the three clusters one was determined to be bacteria, the other two wet discharged fungal spores.

8. The title of the paper reflects parts of the study, but not the part that try to associate the observed bioaerosols with potential sources (the ArcGIS part)

To represent the potential sources identification part of the paper, we have changed the title of paper to:
Characterisation and source identification of biofluorescent aerosol emissions over winter and summer periods in the United Kingdom

9. The abstract cover well the contents of the paper

10. The presentation is generally clear and well-structured but the conclusion might need some work (see point 13)

This was raised by Reviewer 1 and we fully address this issue in response to point 13 below.

11. The language is generally clear and fluent and does not need further improvements

12. The manuscript does not include mathematical formula. However the manuscript describes the use of a third order polynemia with R values between the observations and the polynemia (Table 5). The polynemia is not found anywhere in the manuscript (or in supplementary information) and the results (including low R values) will probably need a discussion.

This is commented on by Reviewer 1 also, and further discussion regarding these results would add to the already lengthy size of the manuscript. As a result, the statistical analysis section has been removed from the manuscript.

13. The conclusion is almost two pages and part of the conclusion seems to be a discussion (e.g. the section concerning difficulties in the clustering). Maybe the conclusion should be shortened to make it more sharp and part of the material should be moved to the discussion section.

The conclusion has been shortened to provide a clearer concluding message (page 23 – 25).
If the authors have used calibration of the instrument against known material, then this calibration needs to be described in more detail and in particular how well the instrument is able to identify test samples similar to the calibration material.

We have added an appendix section with a brief overview of Dstl data collection from 2014 and 2017, and references to published work showing the ability of other UV-LIF spectrometers to identify biological particulates –

“Appendix A: Laboratory characterisation of biological particulates
Laboratory data from a WIBS-3D were collected during a series of characterisation studies at the Defence, Science, and Technology Laboratory (Dstl). This data included bacteria comprising unwashed E-Coli and BG spores, and following research into pollen type abundance in the United Kingdom, Poplar, Aspen, and Ryegrass pollen. Additionally, fungal spore data (Alternaria and Cladosporium) collected using a WIBS-4M during an intensive chamber experiment conducted at Dstl, were also used for comparison to the ambient clusters. Further details of this experiment are to be published in 2019. Similar particle types included in this study have been sampled previously by other UV-LIF spectrometers e.g. Hernandez et al 2016, Savage et al 2017. Here, data from both experiments were clustered in the same way as the ambient data (as described in Section 2.3), with the dominant cluster compared to the ambient fluorescent profiles.”

14. There seem to be 60-65 references in the manuscript. This seems appropriate for this type of manuscript

15. There is no supplementary information. The authors might consider if adding supplementary information can improve transparency of the methods and the documentation.

As requested by Reviewer 1, supplementary information has been added to include plots which show the different meteorological variables and the clusters/total fluorescent particles, in addition to other supporting plots.

Specific comments: On page 25, line 4 onwards, the authors write that this is the first time ArcGIS has been used in relation to land cover mapping and bioaerosols to derive emission patterns etc. As far as I know there are many such studies (some of them are in fact in the reference list), but it is the first time it has been done in connection with the WIBS instrument.

This is a miswording and is meant to state that this is the first time using UV-LIF instrumentation in relation to land cover mapping.
This has been changed on page 25 from:

‘To our knowledge this is the first use of both ArcGIS land cover mapping, in association with airborne bioaerosol concentrations, to identify distinctive 5 emission patterns and factors.’

To:

‘To our knowledge this is the first use of both ArcGIS land cover mapping, in association with airborne bioaerosol concentrations collected using a UV-LIF spectrometer, to identify distinctive emission patterns and factors.’

References


