Aerosol measurements during COPE: composition, size and sources of CCN and IN at the interface between marine and terrestrial influences

J. W. Taylor¹, T. W. Choularton¹, A. M. Blyth², M. J. Flynn¹, P. I. Williams¹,³, G. Young¹, K. N. Bower¹, J. Crosier¹,³, M. W. Gallagher¹, J. R. Dorsey¹,³, Z. Liu¹, P. D. Rosenberg⁴

¹ Centre for Atmospheric Science, School of Earth, Atmospheric and Environmental Sciences, University of Manchester, Manchester, UK
² National Centre for Atmospheric Science, University of Leeds, Leeds, UK
³ National Centre for Atmospheric Science, University of Manchester, Manchester, UK
⁴ School of Earth and Environment, University of Leeds, Leeds, UK

Correspondence to: Jonathan W. Taylor (jonathan.taylor@manchester.ac.uk)

Abstract. We present measurements of boundary layer aerosol concentration, size and composition from a series of research flights performed over the southwest peninsula of the UK during the COnvective Precipitation Experiment (COPE) of summer 2013. We place emphasis on periods of southwesterly winds, which locally are most conducive to convective cloud formation, when marine air from the Atlantic reached the peninsula. Accumulation mode mass loadings were typically 2 – 3 µg m⁻³, the majority of which was sulphuric acid over the sea, or ammonium sulphate inland, as terrestrial ammonia sources neutralised the aerosol. The cloud condensation nuclei (CCN) concentrations in these conditions were ~150 – 280 cm⁻³ at 0.1% and 400 – 500 cm⁻³ at 0.9% supersaturation (SST), which are in good agreement with previous Atlantic measurements, and the cloud drop concentrations at cloud base ranged from 100 – 500 cm⁻³. The concentration of CCN at 0.1% SST was well correlated with non-sea-salt sulphate, meaning marine sulphate formation was likely the main source of CCN. Marine organic aerosol (OA) had a similar mass spectrum to sea spray, and was poorly correlated with CCN.

In one case study that was significantly different to the rest, polluted anthropogenic emissions from the southern and central UK advected to the peninsula, with significant enhancements of OA, ammonium nitrate and sulphate, and black carbon. The CCN concentrations here were around six times higher than in the clean cases, and the cloud drop number concentrations were 3 – 4 times higher.

Sources of ice nuclei (IN) were assessed by comparing different parameterisations of the nucleation of ice, using measured aerosol concentrations as input. The parameterisations based on total aerosol produced IN concentrations that agreed within an order of magnitude with measured first ice concentrations at cloud temperatures as low as -12°C. Composition-specific parameterisations for mineral dust, fluorescent particles and sea spray OA were 3 – 4 orders of magnitude lower, meaning either a source of IN was present that was not characterised by our measurements, and/or one or more of the composition-specific parameterisations greatly underestimated IN in this environment.
1 Introduction

Flash flooding occurs when rain is heavy and persistent. The southwest peninsula of the UK is particularly vulnerable to flash flooding because convective clouds form along convergence lines that often persist for many hours, as was the case in the Boscastle flash flood of August 2004 (Golding et al., 2005). This type of event develops when the wind blows from the southwest and aligns with the peninsula. Convergence associated with sea breeze fronts from the north and south coasts generates quasi-stationary convective systems (Warren et al., 2014). In the Boscastle case, individual cells moved with the wind, but new cells were constantly generated at the upwind end, causing the system as a whole to remain in a quasi-stationary position. Each cell developed over a similar timeframe, and they therefore precipitate over the same region, causing highly localised flooding.

As convection is initiated in the newly developing cells, boundary layer aerosols are lifted up to provide the cloud condensation nuclei (CCN) necessary for cloud drop formation. Increased concentrations of larger and more soluble aerosols enhance CCN numbers and, consequently, enhanced cloud drop number concentration (CDNC). This inhibits the formation of warm precipitation by reducing average drop size (Fan et al., 2007; Huang et al., 2008). At lower temperatures, aerosols provide the ice nuclei (IN) required for primary ice formation, which may then initiate secondary ice processes (e.g. Huang et al., 2008; Phillips et al., 2001). Interactions between the warm rain and secondary ice processes are thought to be key to determining the timing and location of cold precipitation (Taylor et al., 2015). Aerosols may influence these processes by affecting cloud and drizzle drop size, and by determining the concentrations of primary ice forming at different cloud temperatures. Koren et al. (2005) found that convective clouds forming in polluted air may have a higher cloud fraction, lower ice fraction and lower droplet radius than those forming in clean Atlantic air.

In the prevailing west/southwesterly winds, Atlantic air influences the southwest peninsula, whereas in northerly or easterly winds the region receives anthropogenic pollution from the UK and Western Europe. Local emissions from farming and transport may also influence the aerosol over the peninsula.

The COnvective Precipitation Experiment (COPE) (Leon et al., 2015) took place in the southwest peninsula during July-August 2013. Multiple aircraft made in-situ measurements of aerosol and clouds, as well as airborne remote sensing measurements, on days predicted to be favourable for convective cloud formation. Additional ground-based aerosol and X-band radar measurements were performed throughout the duration of the experiment. The purpose of this paper is to describe the boundary layer aerosol measured during the aircraft campaign in terms of concentration, size, and composition, and to determine the main sources of CCN and IN, in order to inform modelling studies aiming to improve forecasts of convective precipitation in the region.
2. Experimental

2.1 Aircraft measurements

Most of the data used in this analysis were measured by instrumentation mounted on the UK BAe-146-301 Atmospheric Research Aircraft during six case study flights over the southwest peninsula, which took place between 05 July and 03 August 2013. Each flight had a designated flight number, which are listed in Table 1. Additional airborne measurements were performed with the University of Wyoming King Air 200T (UWKA) and UK Met Office Civil Contingency 208 Aircraft (MOCCA), but these are not discussed here. The BAe-146 was fitted with a variety of aerosol and cloud probes. Here we only discuss instruments relevant to our analysis; Leon et al (2015) provide a full list of instrumentation aboard the aircraft. All concentrations are corrected to standard temperature (273.15 K) and pressure (1013.25 hPa). A typical flight plan involved performing aerosol runs in the boundary layer at an altitude of ~500m, before making measurements of clouds at higher levels.

The concentration of condensation nuclei (CN, i.e. the aerosol number concentration) larger than 2.5 nm was measured using a model 3786-LP water-filled condensation particle counter (WCPC; Hering et al., 2005) manufactured by Aerosol Dynamics Inc., based on the TSI model 3786 but modified for use at low pressure. The absolute uncertainty in the CN concentration is ±12%.

A Droplet Measurement Technologies (DMT) dual-column cloud condensation nuclei counter (CCNC; Roberts and Nenes, 2005) measured the concentrations of CCN at nominal supersaturations (SST) of 0.1% and 0.9%, though on 18 July the higher SST was set to 0.6%. The relative uncertainty in the supersaturation measurements is ±10% (Roberts et al., 2010; Trembath, 2013), and the uncertainty associated with the flow calibration on the particle counter is typically ~10%.

The aerosol size distribution was measured with a custom-built inboard scanning mobility particle sizer (SMPS; Wang and Flagan, 1990), and wing-mounted Particle Measurement Systems passive cavity aerosol spectrometer probe (PCASP) -100X, with electronics upgraded by DMT, and DMT cloud droplet probe (CDP). The PCASP and CDP were size calibrated as described by Rosenberg et al. (2012). The SMPS measures the mobility diameter of particles 20 – 320 nm, while the PCASP and CDP measure the optical diameter of particles 0.12 – 3.4 μm and 2 – 50 μm respectively. The PCASP inlet has active drying, and the SMPS inlet line was dried to <45% RH by ram heating and the temperature increase as the sample line entered the cabin. The CDP measures aerosols and cloud drops at ambient humidity.

The composition of nonrefractory submicron aerosol was measured with an Aerodyne Research Inc. compact time-of-flight aerosol mass spectrometer (AMS; Canagaratna et al., 2007; Drewnick et al., 2005), which reports concentrations of organic aerosol (OA), non-sea-salt sulphate (nss-SO$_4^{2-}$), nitrate (NO$_3^-$), ammonium (NH$_4^+$) and non-sea salt chloride (nss-Chl$^-$) as standard. Additionally, we used the fragmentation table given by Langley et al (2010) to calculate concentrations of methanesulphonic acid (MSA), and adjusted the fragmentation tables of OA and nss-SO$_4$ accordingly.
The AMS was calibrated at the start and end of each flight using monodisperse ammonium nitrate to calculate the ionisation efficiency (IE) for NO$_3^-$, corrected as described by Morgan et al. (2014, Supplementary material). The ammonium nitrate calibrations were also used to calculate the relative ionisation efficiency (RIE) of NH$_4^+$, and a separate ammonium sulphate calibration was performed to calculate the RIE of SO$_4^{2-}$. The AMS IE calibration was stable within ±12% throughout the campaign, and the absolute accuracy is ~30% (Bahreini et al., 2009). The collection efficiency was calculated as recommended by Middlebrook et al. (2012), and the calculated aerosol volume showed qualitative agreement with the aerosol volume measured by the SMPS and PCASP.

Refractory black carbon (BC) concentrations were measured with a DMT single-particle soot photometer (SP2; Schwarz et al., 2010), which is the same instrument used in previous studies on the BAe-146 (e.g. Taylor et al., 2014), but upgraded to be functionally identical to the current SP2 model D. SP2 calibrations were carried out at the start of each flight, and in the laboratory after the campaign. The incandescence detectors, which measure BC mass on a single-particle basis, were calibrated using 300 nm mobility diameter Aquadag, corrected for the difference in response between the calibration particles and ambient soot (Baumgardner et al., 2012; Laborde et al., 2012b). Using this calibration, the uncertainty in the reported BC mass concentrations are ±10% when sampling diesel soot, and ~20% when sampling biomass burning (Laborde et al., 2012a, 2012b).

Bulk aerosol was collected on two 47-mm nuclepore filters, with 10µm and 1µm pore sizes, connected in serial. The filter sampling system is described by (Formenti et al., 2008). The size and composition of particles collected on the filters was determined offline after the campaign using scanning electron microscopy (Phillips FEI XL30 Environmental Scanning Electron Microscope with Field-Emission Gun (ESEM-FEG) , with an Energy-Dispersive X-Ray Spectroscopy (EDX) system (Hand et al., 2010)). Particles were classified by composition using the scheme described by Young et al. (2015). An example filter composition measurement is presented by Leon et al. (2015).

2.2 Ground site measurements

The COPE ground site hosted a suite of aerosol and meteorological instrumentation operating continuously throughout July and August 2013. Leon et al. (2015) provide a full list of instrumentation at the ground site. In this analysis, we only use the size-resolved measurements of fluorescent aerosol made by a DMT wideband integrated bioaerosol sensor (WIBS) 4, which we consider to be primary biological aerosol particles (PBAP). Although the WIBS may also detect other fluorescent particles, the fluorescent particle concentration is generally considered to be a lower limit on the PBAP concentration (Huffman et al., 2010).
3. Results and discussion

3.1 Airmass history

Figure 1 shows 5-day back trajectories for each of the case study flights, calculated using the Hybrid Single Particle Lagrangian Integrated Trajectory Model (HYSPLIT) (Draxler and Hess, 1998). Trajectories were initiated every 60 seconds from the aircraft flight track during the boundary layer aerosol runs. The trajectories were performed with 3-hourly 0.5° Global Data Assimilation System (GDAS) reanalysis data, using full vertical dynamics on 23 model levels. Although the turbulent mixing in the boundary layer means the accuracy of individual trajectories is uncertain, examining the general trend provides information on the history of the airmass, and possible changes in its cloud nucleating potential, which is determined by synoptic-scale winds.

Figure 2 shows the flight path the aircraft took during the below-cloud runs, and the boundary layer wind speed and direction measured during these runs. The wind data allow us to assess the accuracy and potential influence of the back trajectories’ final stages, and examine local wind fields such as sea breeze circulations that are not captured by the meteorology data used to initiate the trajectories.

Figures 1 and 2 panels (c), (e) and (f) show that the 25 and 29 July and 03 August case studies shared the synoptic features of the “Boscastle” type meteorology, i.e. winds blowing marine air from the North Atlantic over the peninsula from the southwest (hereafter referred to as southwesterly winds (SWW)) (Golding, 2005). The back trajectories showed that the airmasses had been over the North Atlantic for at least five days prior to landfall. They also showed sharp changes in direction over the ocean; comparison with the synoptic charts showed these were due to circulation around low-pressure systems as they moved east, and were associated with frontal precipitation, which is likely to have washed out any long-range transported aerosol. On 28 July, the situation was reasonably similar in that the airmass had spent time over the North Atlantic, but reached the peninsula from the south, and may have passed over the northwest tips of Spain and France. Local sea breeze circulations may still have caused convergence on these four days, but they had only a minor influence on the absolute wind speed and direction, as the synoptic-scale winds were so strong.

The meteorology on 05 and 18 July were markedly different to the other case studies. On these two days, high pressure systems located over the UK meant that synoptic-scale boundary layer winds were much weaker. The measured winds were generally only a few metres per second and the greater variation in wind direction suggest that the local sea-breeze circulations had more of an influence on absolute wind direction on these days. On 05 July, the back trajectories show that for the few days prior to being sampled, the airmass had passed over the Atlantic, though the easternmost trajectories passed over the southern tip of Ireland and the south coast of Wales. It is difficult to assess the accuracy of these parts of the trajectories when local winds were dominant, though there was some anthropogenic influence on this flight, as discussed in the next section. The trajectories for 18 July suggest the air over the peninsula had passed over many parts of England, Wales and Ireland, and are therefore likely to have also had a strong anthropogenic influence.
The colours in Fig. 2 divide sections along the flight track into “Northwest (NW) coast”, “Inland” and “Other” influenced regions. The “NW Coast” regions include areas over the sea, and the regions labelled “Other” are not used in this analysis. Depending on the wind directions, aerosol in these regions may be influenced by marine or terrestrial emissions, or both. The trajectories and wind data in Figs. 1 and 2 show that for flights on 05, 25 and 29 July, and 03 August, the sections on the NW coast were affected by marine air that had not been influenced by land, though this was not the case for the other two flights. Dividing the data into these sections allows us to investigate changes in aerosol chemistry and properties at the land-sea interface. This topic is discussed further in the next section.

### 3.2 Submicron aerosol composition

#### 3.2.1 Marine aerosol

Figure 3 shows the average aerosol composition measured by the AMS and SP2 in the NW coastal regions during the six case study flights. The data from Figure 3 are also listed in Table 1. In clean Atlantic air (NW coast on 05, 25 and 29 July and 03 August, as identified in the previous section) the composition was fairly consistent. Around 60 – 70% of submicron aerosol mass was sulphate, in the form of a mixture of sulphuric acid and ammonium bisulphate as there was insufficient ammonium for full neutralisation. This is in agreement with measurements of North Atlantic aerosol made by Ovadnevaite et al. (2014). In the marine airmasses, the ratio of MSA/nss-SO$_4$ was ~0.05, which is perhaps lower than average, but within the range typically observed in clean marine air in summer (e.g. Dall’Osto et al., 2010; Huebert et al., 1996; Ovadnevaite et al., 2014; Phinney et al., 2006). MSA and sulphate were well correlated ($R^2 = 0.66$), and the source of both is likely to be oxidation of biogenic dimethyl sulphide (DMS) emissions. OA was the only other major aerosol component, though the ratio of nss-SO$_4$/OA did show some variation. It is unlikely that anthropogenic pollution, either from shipping or long-range transported, made any significant contribution to the aerosol loadings; in clean Atlantic air, the OA showed no correlation with BC ($R^2 = 0.0$), while nss-SO$_4$ showed a weak negative correlation ($R^2 = 0.15$). These results are in agreement with a similar assessment of Atlantic aerosol made at Mace Head, Ireland (O’Dowd et al., 2014). The source of the marine OA is discussed further in Sect. 3.2.3.

The total mass loading measured by the AMS + SP2 varied from 2.4 – 3.6 µg m$^{-3}$ in these flights. Non-sea salt chloride and black carbon were minor components. Low, but non-zero levels of nitrate were measured, despite the fact that aerosol nitrate is not expected to form in acidic conditions as the sulphate displaces any nitrate to the gas phase. Interpretation of this apparent measured nitrate is discussed in Sect. 3.2.4.

#### 3.2.2 Inland aerosol

Submicron inland aerosol composition on flying days is shown in Figure 4. On flying days between 25 – 29 July and 03 August, inland aerosol had composition similar to the Atlantic marine air, but with the addition of ammonium. On 25 and 28 July, the ammonium was sufficient to fully neutralise the sulphate, but on 29 July and 03 August, the sulphate was a mixture
of ammonium sulphate and bisulphate. The total measured mass loadings were within 10% of the marine air for the corresponding flight.

Agriculture is the largest land use category in the southwest peninsula (Morton et al., 2011), including a mix of both arable and pastoral farming, predominantly cattle grazing on managed grass fields. Previous measurements have found elevated levels of ammonia in the southwest peninsula compared to those found in the Atlantic (Quinn et al., 1996; Sutton et al., 2001). The terrestrial ammonia emissions are therefore most likely to be of agricultural origin.

Compared to the last four flights, the inland aerosol on 05 July had a much larger OA fraction. The low wind speeds on this day mean local aerosol sources are likely to have had more of an effect. Inland on 05 July, OA had a correlation with BC of $R^2 = 0.56$, and with the aircraft’s CO measurement an $R^2 = 0.69$. Additionally, the BC mass loading was around a factor of 10 higher inland than in the Atlantic air. Taken together, this suggests that local fossil fuel emissions were at least partly responsible for the increased OA inland on 05 July.

18 July was somewhat of an outlier in terms of the COPE case studies in that the aerosol mass loadings were much higher than the other flying days. The back trajectories showed that the airmass had come from England, Wales and/or Ireland, so a strong anthropogenic influence would be expected. There was not much difference between the aerosol measured inland and on the NW coast. OA was the largest component, followed by similar amounts of ammonium nitrate and sulphate, which were both fully neutralised by ammonium.

### 3.2.3 OA composition and origin

Examining the ratios of different fragments in the OA mass spectrum can give an indication of OA sources as well as their level of ageing. AMS factor analysis most-commonly tends to pick out factors with differing levels of oxidation. Fresh fossil fuel emissions appear as hydrocarbon-like OA (HOA), while more aged aerosol including secondary OA (SOA) appears as oxygenated OA (OOA), which may be semi-volatile (SV-OOA) or more aged low-volatility (LV-OOA) (Jimenez et al., 2009).

OA measured in marine background air may be primary OA from sea spray (Ovadnevaite et al., 2011), SOA from marine VOC emissions (Decesari et al., 2011), or long-range transported pollution. Globally, primary marine OA is thought to dominate over marine SOA formation (Arnold et al., 2009; Gantt and Meskhidze, 2013). As discussed in Sect. 3.2.2, local anthropogenic emissions contribute to OA concentrations inland, while the marine OA would have less of an influence in areas further away from the sea.

The OA mass spectrum measured over the NW coast on 05 July is shown in Fig. 5. The spectrum is dominated by peaks at m/z 44 and 28, which are prescribed to be equal in the default fragmentation table. These peaks are much larger than m/z 43 as is typical in oxidised OA spectra. Table 2 presents correlations between NW coast OA spectra from 05, 25 and 29 July, and 03 August, with several spectra from literature. The literature spectra are HOA and two OOA factors (Morgan et al., 2010), sea spray (Ovadnevaite et al., 2011), marine biogenic SOA (Chang et al., 2011, which was well correlated with nss-
SO4), and the “Organic factor” measured in the Arctic ocean by Chang et al. (2011), which was thought most likely to be sea spray OA and was poorly correlated with sulphate. Though there are many previous measurements of different OA components in Europe, we choose those from Morgan et al (2010) because they were measured with the same instrument aboard the BAe-146, so any inter-instrument variability should be minimised. OOA-1 was more oxidised than OOA-2, but no assessment was made of their volatility. To ensure the comparison was fair, only m/z values where all spectra had data points were used. The marine OA spectra showed the best correlation with sea spray OA, indicating this is a likely source, though the correlations with the other oxidised factors were still reasonably good.

Figure 6 shows the fractions $f_{44}$ and $f_{43}$ (the fraction of OA measured at m/z 44 and 43 respectively) for all six case study flights, divided up into NW coastal and inland data. The literature values from Table 2 are also plotted. Ambient OOA measurements tend to fall within the diagonal dashed lines in Fig. 6 (Ng et al., 2010), and fossil fuel emissions may be expected to fall close to the HOA point from Morgan et al. (2010) upon emission and progress approximately through OOA-2 to OOA-1 with increased oxidation.

The four marine spectra from this study have $f_{44}$ indicative of a moderate level of oxidation but, on average, slightly low $f_{43}$. Of the selected literature values, the marine OA all fall closest to the sea spray OA, rather than the marine biogenic. If the inland OA was composed of a mix of marine and terrestrial OA, lines drawn from the marine data points through their corresponding inland points should intersect in a region representative of terrestrial OA. For the four flights with clean marine spectra, the lines would intersect in a region around (0.1, 0.1), close to the SV-OOA data point from Morgan et al (2010). This is consistent with the terrestrial aerosol source being similar on each flying day, and being moderately oxidised.

It is not fully clear what fraction of the terrestrial OA is anthropogenic or biogenic. It is difficult to pick out anthropogenic locally produced SOA from biogenic as the mass spectra look similar, and photochemical production is strongest for both during peak daylight hours. Most AMS factor analysis studies are only able to pick out regional background and locally produced OOA (Jimenez et al., 2009 and references therein). Slowik et al. (2010) identified a biogenic SOA factor by correlation with biogenic VOCs, but noted the mass spectrum was very similar to SV-OOA formed from anthropogenic pollution. As discussed in Sect. 3.2.2, the correlation between OA and CO is indicative of a fossil fuel component of the terrestrial OA, but we also cannot rule out a biogenic component.

### 3.2.4 Interpretation of nitrate measurements

The formation of nitrate aerosol is only possible when there is sufficient ammonium (or other cations) to fully neutralise any sulphate aerosol (Seinfeld and Pandis, 1998). However, our observations in Figure 3 and 4 show the apparent presence of aerosol nitrate under acidic conditions. The measurements are above the detection limit for nitrate (~0.03 µg/sm³ for a 30s averaging time), and are ubiquitous throughout the runs, meaning they are not an artefact of averaging two airmasses of differing composition.
Nitrate appears in the AMS almost entirely in peaks at m/z = 30 and 46 (Allan et al., 2003), for ions NO\(^+\) and NO\(_2^+\) respectively. When sampling nitrate species, the ratio of these two peaks is determined by the heater temperature and aerosol volatility. For a fixed heater temperature, less volatile nitrate species decompose further during their slower vaporisation, and have a higher 30/46 ratio when measured (Drewnick et al., 2015). The AMS may detect other nitrate species including salts such as NaNO\(_3\) and KNO\(_3\), as well as organic nitrates, as long as their boiling point is low enough.

Table 3 shows the ratios of m/z 36/40 measured on each flying day during ammonium nitrate calibrations and ambient sampling periods. For 18 July, the 30/46 ratio was the same during the calibration and coastal/inland aerosol. As the ammonium in these periods was sufficient to fully neutralise the aerosol, this gives us a high degree of confidence that the NO\(_3^-\) measured on this flight was ammonium nitrate. However, on all other flights the ambient 30/46 ratio was higher than the calibration values, particularly over the sea. Sea salt-based nitrates may be found in marine air, such as NaNO\(_3\) formed by reaction of nitric acid with NaCl. This would be found predominantly in coarse mode aerosol rather than accumulation mode (Harrison and Pio, 1983), but some particles may be detected by the AMS. Alfarra (2004) measured nebulised NaNO\(_3\) and recorded a 30/46 ratio of 29.2, which is significantly higher than those measured during COPE. Presumably, the same argument applies with these nitrate salts as with ammonium nitrate: the nitrate would be displaced by sulphate unless the aerosol was fully neutralised. NaNO\(_3\) would only be possible if the coarse mode aerosol was not in equilibrium with the accumulation mode. We must therefore investigate the possibility of organic compounds giving signals at the same m/z as the nitrate peaks.

Ovdanevaite et al. (2011) measured high-resolution mass spectra of primary marine OA, which allowed them to identify the specific ions measured at particular m/z values. Around 1.7% of primary marine OA was measured at m/z 30, most of which was CH\(_2\)O, and 0.2% was at m/z 46. If the measured nitrate concentrations in COPE were attributable to organic species, they would make up 6 – 12% of organic mass in marine air when corrected for relative ionization efficiency. Rollins et al. (2010) measured 30/46 ratios of 0.99 – 5.30 for various organonitrates, which is more in line with our measurements.

To summarise, we are not able to conclusively explain the reported nitrate measurements in acidic conditions. Several past studies have faced similar problems in various different regions e.g. (Allan et al., 2004, 2006, 2014). It is likely that they are partly due to interference from organic species, but the inland 36/40 ratios, where enhanced ammonium concentrations were measured, were closer to those expected from NaNO\(_3\). The measurements would need to be performed with a high-resolution mass spectrometer to confirm the organic fraction at the nitrate peaks. There may also be a contribution from sea salt-based nitrate species such as NaNO\(_3\), but these are also difficult to quantify with our measurements. On all flights other than 18 July, the total nitrate loadings were a relatively small fraction of the total mass measured by the AMS, as the nss-SO\(_4^2-\) and OA loadings were both several times higher.
3.3 Aerosol size distributions

Figure 7 shows the submicron aerosol size distributions measured in NW coastal regions during the six case study flights. The standard deviations of each in are plotted to show the spatial variation over the runs, and were much larger than the calculated standard errors for the size range shown. The SMPS data were normalised to the PCASP in the overlap region.

Three submicron aerosol modes are evident from the distributions: nucleation, Aitken, and accumulation modes, with varying magnitudes in the different flights.

All the measured distributions were open at the lower end, meaning nucleation mode aerosols were present smaller than the 20nm size of the smallest bin. In all flights, the concentrations of particles in all bins <40 nm in diameter was higher inland than over the sea. Additionally, the total CN concentration (listed in Table 1) was always higher and more variable inland, and the standard deviations of smallest particles measured were higher inland. These measurements suggest there was a source of small particles forming inland, which was not uniform in spatial location.

On most flights, the Aitken mode was centred on 30 – 60 nm, and the accumulation mode was centred around 150 – 220 nm. The gap in between was characteristic of the “Hoppel dip”, thought to be caused by cloud processing of marine aerosol (Hoppel et al., 1986). Similar size distributions have been measured in summertime North Atlantic aerosol (Asmi et al., 2011; Dall’Osto et al., 2011). The accumulation mode on 05 July was larger, with modal diameter ~275 nm over the sea, which may be due to increased photochemical aerosol formation in the clear weather. 18 July was also an exception, showing just one wide accumulation mode, and the size distribution was consistent with aged UK urban aerosol (e.g. Bohnenstengel et al., 2015). Other than these two flights, where fossil fuel emissions had a larger effect than the case studies, the accumulation mode aerosol size distribution was fairly consistent.

By replacing the H+ ions with NH4+, the neutralisation of the marine sulphuric acid inland should add to the inorganic mass, causing a marginal increase in average particle size. However, the corresponding increase in inorganics was balanced out by similar decrease in nss-SO4 inland, and no consistent differences were observed between the corresponding aerosol size distributions.

One clear feature of the terrestrial influence was the presence of nucleation mode aerosols inland. In all cases, the inland aerosol concentration was significantly higher than those measured over the sea, or typical marine concentrations in the absence of nucleation events (Dall’Osto et al., 2011; Jensen et al., 1996), and the aerosol size distributions showed concentrations of particles <60 nm were always higher inland. Coastal nucleation is unlikely to explain these measurements, as there was no bias in concentrations towards coastal regions.

It is possible that the formation of these particles is also linked to agricultural emissions. Although particle formation rates from H2SO4–NH3–H2O ternary nucleation are lower than those observed in ambient environments, ternary nucleation involving other agricultural emissions (such as amines) may explain our results (Almeida et al., 2013; Pirjola et al., 2000). The high standard deviation/mean ratio of the inland CN concentrations in SWW suggests the process was spatially variable,
rather than ubiquitous throughout the boundary layer. Again, however, it is also clear that the generation of these particles had little influence on the CCN concentrations, which were similar over the sea and inland.

It is also of interest to examine the concentrations of giant (dry diameter, $1 \leq D_p \leq 10$ µm) and ultragiant particles ($D_p > 10$ µm), as these may have an enhanced effect on precipitation formation via the warm rain process (Johnson, 1982). The number concentrations of giant ($N_{GA}$) and ultragiant ($N_{UGA}$) are listed in Table 1. $N_{GA}$ varied by around a factor of two between cases, with values of around one per cubic centimetre, but there was no obvious trend to explain the differences between the different days. For the SWW cases, the inland $N_{GA}$ was fairly proportional to the NW coastal values, but lower by 15 – 40%. These are likely to be sea spray aerosols, and indeed sea salt made up nearly 80% of supermicron aerosol volume on 3rd August (Leon et al., 2015). These measurements are consistent with dilution/deposition of sea salt particles inland being greater than any terrestrial coarse aerosol emissions such as PBAP or dust on these days.

The concentrations of ultragiant aerosol also showed around a factor of two variation between the different cases, and were 3 – 6 L$^{-1}$, other than the NW coastal measurement on 29 July, which was 11 L$^{-1}$. Again, it is difficult to pick out any clear trend in the values. On some days, $N_{UGA}$ was higher over the sea, whereas on others it was higher inland. In all cases, $N_{GA}$ and $N_{UGA}$ comprised a very small fraction of the total aerosol number concentration, including the concentration of particles large enough to affect CCN populations.

### 3.4. Aerosol influence on CCN and cloud drop concentrations

#### 3.4.1 Linking CCN and CDNC

CCN data were only available on 18, 25 and 29 July, and 03 August. Figure 8 shows the average boundary layer CCN concentrations measured on these flights, as well as a comparison to CDNC measured in updrafts near cloud base. By choosing only data in updrafts, as well as excluding data from cloud edges, we aim to minimise the influence of entrainment, though it is unlikely to be eliminated. The higher values of CDNC are those least influenced by entrainment, and those with the strongest updrafts when the cloud formed.

On 18 July, the peak CDNC was close to the CCN concentration at 0.1% SST. For the SWW flights, the peak CDNC was more comparable to the CCN concentrations measured at 0.9% SST, though the median CDNC was between the two CCN measurements, which is reasonable based on previous estimates of SST in cumulus clouds (e.g. Politovich and Cooper, 1988).

Comparing CCN measured inland and over the sea, the differences were generally <10%, with the largest being 30%. There was no clear systematic difference between land and sea. The addition of ammonium inland did not always increase inorganics, as the change was not always larger than the inherent variability in the nss-SO$_4$$_2$. On an airmass level, there appears to be an aerosol influence on cloud drop concentrations- the higher aerosol concentrations in the polluted air caused higher CCN concentrations. This difference was also manifested in the CDNC, which was several times higher in the polluted case.
Dall'Osto et al. (2010) reported measurements of aerosol composition and CCN at Mace Head, Ireland during late spring and early summer. Mace Head shares similarities with the southwest peninsula of the UK in that it experiences Atlantic air under the prevailing westerly winds but is also a receptor site for aged urban pollution during easterlies. The CCN concentrations on 18 July were around double those measured by Dall'Osto et al. (2010) in continental pollution, though the aerosol mass loadings here were also significantly higher. In marine airmasses, the CCN concentrations at Mace Head were \(200 \text{cm}^{-3}\) at 0.1% SST and \(400 \text{cm}^{-3}\) at 1.0% SST. These numbers are both in good agreement with our measurements in the SWW cases.

### 3.4.2 Aerosol chemistry and CCN

In marine air, CCN may be produced by sea spray or condensation of secondary aerosols, primarily nss-SO\(_4\). These processes can interact, for example primary sea spray subsequently coated with nss-SO\(_4\). Particles produced by sea spray may be composed of sea salt, OA, or a combination of the two, with OA more dominant in submicron particles (Cavalli et al., 2004). Sea salt is an effective CCN; OA from sea spray is likely to be insoluble (Gershey, 1983), but may be internally-mixed with sea salt and/or serve as a condensation nucleus for secondary organic or inorganic aerosol formation. During runs over the southwest peninsula, terrestrial emissions may also have influenced CCN concentrations, such as the addition of ammonia or advected aged urban emissions.

As an air parcel cools as it rises, there may be some co-condensation of semi-volatile species such as ammonium nitrate or SV-OOA, which may further enhance CCN concentrations (Topping et al., 2013). As the aerosol inlets on board the BAe-146 do not reliably sample aerosol during cloud penetrations, we are unable to assess how much of an impact this made, but we note that sulphuric acid / ammonium sulphate (the largest aerosol component on most case studies) is not semi-volatile at the temperatures sampled.

As the source functions of sea spray and secondary organic and inorganic marine and terrestrial aerosols all depend on different factors, they would not be expected to be correlated. Therefore, by comparing the measured CCN concentrations to different chemical species, we may gain information on the sources of the CCN.

Figure 9 shows the relationships between CCN concentrations measured aboard the BAe-146 at 0.1% SST and the PCASP number concentration, AMS total mass, non-sea salt inorganics, and OA. Although the validity of the nitrate measurement is unclear on some flights, the nitrate fraction on these days was low. The data in Figure 9 are split between the three SWW case study flights and the polluted case of 18 July. If we did not make this distinction, all the variation and correlation coefficients would be dominated by the difference between the marine and polluted airmasses, rather than variations in concentration within these airmasses.

The correlation between CCN and aerosol number concentration measured by the PCASP was good, and the data fell close to the 1:1 line, meaning the critical diameter was close to the PCASP's lower cutoff of \(120 \text{ nm}\). Good correlations (\(R^2 \geq 0.66\)) were found in both marine and inland environments comparing CCN with total AMS mass and inorganics. The
correlations with OA were lower, but non-zero, and the OA correlation in marine air was better than inland. These results are consistent with the main source of CCN being accumulation mode sulphate-containing particles in both marine and inland air. Liu et al. (1996) found a similar correlation with sulphate at 0.06% SST for cloud-processed aerosol.

Figure 10 shows correlations between CCN at 0.9% SST and the same variables as in Figure 9, but only for the SWW flights. The correlations with all four x-variables are lower, meaning CCN at the higher SST are less directly linked to accumulation mode aerosol. In Fig. 10 (a) the CCN are above the 1:1 line with the PCASP number concentration, meaning the critical diameter was below 120 nm. It is therefore not surprising that the correlations at this SST are lower, as the mass measurements are also biased towards larger accumulation mode particles. The species trends in correlation were the same as at 0.1% SST though, in that the correlation with inorganics and AMS total mass were higher than those with OA.

While our data suggest that the average CCN were composed primarily of nss-SO$_4^-$, it does not provide information on the origin of these particles. For example, marine nss-SO$_4^-$ may condense onto primary sea-spray OA particles, which are largely insoluble when initially formed (O’Dowd and de Leeuw, 2007). Pierce et al. (2007) showed that this type of process can significantly enhance CCN concentrations globally by providing a surface for sulphuric acid to condense on to. Particle emissions from sea spray are strongly linked to wind speed (O’Dowd et al., 1997). This would mean the marine particle concentrations on 29 July would likely be the highest due to the stronger wind speeds, as shown in Fig. 2. While the marine CN concentration on 29 July was higher than on 25 July, it is difficult to draw firm conclusions from such a limited comparison.

Shipping emissions may also serve as condensation nuclei for secondary aerosol (e.g. Langley et al., 2010), but the low BC concentrations in our case studies suggests this was of little importance. Alternatively, new particles may form through nucleation of gas-phase precursors, and open-ocean nucleation events are thought to be most likely to be related to biological activity (O’Dowd et al., 2010). We gain some insight into particle formation by comparing to the size distributions classified as “clean marine” and “open-ocean nucleation” by Dall’Osto et al. (2011), measured at Mace Head. In particular, the value of dN/dlogDp in our lowest size bin (20 nm) is a useful comparison for the smallest particles, which have experienced the least secondary growth. On 05 and 25 July, the values of the lowest size bin were ~100 cm$^{-3}$, which is consistent with the “clean marine” values (~80 – 150 cm$^{-3}$). The concentration on 29 July was higher at 985 cm$^{-3}$, which is closer to that in “open-ocean nucleation” airmasses (~800 – 2500 cm$^{-3}$), but the value on 03 August of 516 cm$^{-3}$ fell between the two categories. It is therefore not clear what contribution primary and secondary aerosol formation had overall to the aerosol number concentration in marine air, but the majority of CCN mass was composed of nss-SO$_4^-$, which must be secondary.

This leads us to the conclusions that in the SWW cases, the main factor determining CCN concentrations was the size and composition of marine aerosols reaching the peninsula. The majority of accumulation mode aerosol measured by the AMS in these cases was nss-SO$_4^-$. The measured CCN also exhibited a high degree of correlation with inorganics, which were predominantly nss-SO$_4^-$ and its cations. If this correlation were due to the mixing of two homogeneous airmasses, the OA would also show a similar degree of correlation with CCN; however, the R$^2$ values for OA were much lower. The data
therefore show that whatever process is responsible for the production of sulphate is also largely responsible for determining the concentration of CCN. Our results are consistent with the prevailing theory that the majority of marine CCN are composed of nss-SO$_4$ formed from DMS oxidation (Charlson et al., 1987).

3.5 Sources of IN

The concentrations of primary ice and the temperatures at which they develop are a large source of uncertainty in the timeline of cloud glaciation. During COPE, the lowest cloud top temperatures were ~15°C (Taylor et al., 2015). Reviews by Hoose and Möhler (2012) and Murray et al. (2012) identified the only species that have been shown to behave as ice nuclei in laboratory experiments at these temperatures as mineral dust; primary biological aerosol particles (PBAP) such as bacteria (Diehl et al., 2006), fungal spores and pollen (Pouleur et al., 1992); and black carbon, although the latter is relatively inefficient. Emissions data for bacteria and fungal spores is limited but global model studies on clouds and precipitation including these species acting as IN suggest a small but significant zonal influence on precipitation rates (Spracklen and Heald, 2014), but their concentrations at cloud formation levels are generally much lower than mineral dusts. Recently, attention has also focused on marine surface organics, which may be aerosolised as sea spray (Burrows et al., 2013).

Mineral dust concentration and size distribution were measured using the BAe-146 filter system, by combining the ‘silicates’ and ‘mixed silicate’ categories described by Young et al. (2015). BC and OA concentrations were measured online using the SP2 and AMS. There was no instrument capable of isolating PBAP aboard the aircraft, but the ground site hosted a WIBS, which made measurements of UV auto-fluorescent particles. The reader should see Sect. 2 for further details of the instrumentation.

In the absence of IN measurements, parameterisations must be used to calculate IN concentrations. These are empirical relationships derived by comparing measurements of IN and various chemical species. Figure 11 shows comparisons of several parameterisations, as well as measured first ice concentrations from COPE. The aerosol measurements used for Figure 11 are those from 03 August, as this had the best measurements of ice formation, as presented by (Taylor et al., 2015). The measured ice concentrations are from a series of penetrations through a cloud in the early stages of development, when secondary ice processes were thought to have at most a minor influence on ice concentrations (Taylor et al., 2015).

DeMott et al. (2010) based a parameterisation (hereafter D10) for IN concentrations on the total concentration of aerosol larger than 0.5 µm (D >0.5 µm), derived from ice nuclei measurements with continuous flow diffusion chambers in a variety of environments around the world. Tobo et al. (2013) formed a similar parameterisation (hereafter T13-T) using data measured in a North American pine forest. We used a D > 0.5 µm concentration of 4.5 cm$^{-3}$ derived from a combination of the PCASP and CDP measurements during the inland aerosol runs. For comparison, the concentration collected on the filters was 4 cm$^{-3}$, and calculations performed using this number were not discernibly different when plotted on Figure 11. The IN calculations based on D-10 and T13-T both fall within an order of magnitude of the measured first ice concentrations. The
T13-T trace is closer to the measured ice concentrations than D10, meaning the ice active fraction during COPE was more similar to that in a forest ecosystem than in the average of the environments studied in DeMott et al. (2010). Niemand et al. (2012) and DeMott et al. (2015) provide parameterisations (hereafter referred to as N12 and D15 respectively) base on mineral dusts. N12 uses the integrated mineral dust size distribution to calculate the number of ice surface active sites using measured size distributions, while D15 again simply uses the number concentration of dust particles larger than 0.5 µm, which in our case was 0.4 cm$^{-3}$ based on the filter measurements. Young et al. (2015) found that the concentrations of particles collected on the filters agreed well with in-situ probes for particles larger than 0.5 µm, but the filters’ collection efficiency was poor for smaller sizes. As D15 uses the dust concentration larger than this size, and N12 is based on the dust surface area distribution, which peaks at supermicron sizes, the filter measurements are expected to be a robust measurement for performing these calculations within the uncertainty of the parameterisations. Both mineral dust parameterisations give IN concentrations several orders of magnitude below the measured first ice concentrations.

As well as T13-T, Tobo et al. (2013) constructed a parameterisation (hereafter referred to as T13-F) based on measurements of fluorescent particles larger than 0.5 µm, which are likely to be PBAP. The measurements upon which this parameterisation was based were made using an ultraviolet aerodynamic particle sizer (UV-APS), which has a different instrument response to the WIBS. Healy et al. (2014) made concurrent measurements using a UV-APS and a WIBS, and found the WIBS channel 3 (which uses the same fluorophore as the UV-APS) was well correlated with the UV-APS fluorescent concentration, but a factor of 2.7 higher due to the instruments’ different size ranges.

Between 1100 – 1500 UTC on 03 August, the total concentration of fluorescent particles measured by the WIBS was 100 ± 40 L$^{-1}$. The fluorescent concentration measured using channel 3 was 50 ± 26 L$^{-1}$, which corresponds to a UV-APS equivalent concentration of 19 ± 10 L$^{-1}$. Using this concentration in the T13-F parameterisation generates IN concentrations that are also several orders of magnitude lower than the measured ice concentrations. The difference between the predicted IN concentrations based on T13-T and T13-F are due to the different coarse aerosol composition between the forest environment studied in Tobo et al. (2013) and the marine environment studied here, which is likely to have higher concentrations of sea salt.

As an additional estimate of biological IN, Mohler et al. (2008) performed chamber experiments using pseudomonas syringae bacteria, and found that the fraction of these bacteria that were active IN at -10C was 0.5%. Multiplying this fraction by the PBAP concentration measured by the WIBS gives an IN concentration of 0.5 L$^{-1}$ at this temperature. As pseudomonas syringae are amongst the most active biological IN studied thus far (Lorv et al., 2014), this value may be considered an upper estimate for biological IN at this temperature. Though it is possible to classify WIBS measurements into different PBAP types, even within simple metaclass discrimination (bacteria, fungal spores and pollen) the uncertainty in the derived concentrations can be large, e.g. up to a factor of 5 for bacteria, depending on the assumptions used in the analysis approach, as discussed by Crawford et al. (2015).
To date there are no IN parameterisations for soot in the relevant temperature range, and laboratory studies show mixed results, though it is likely to be a worse IN than mineral dust (Hoose and Möhler, 2012). The total mineral dust mass concentration measured was 4 µg m\(^{-3}\), whereas the total BC concentration was 30 ng m\(^{-3}\), so the IN concentrations from BC were likely to have been several orders of magnitude lower than mineral dust.

Wilson et al. (2015) recently published a mass-based parameterisation for marine surface organics measured in the European Arctic. To our knowledge, this is the only parameterisation published for this type of IN. Our results in Figure 11 show that aerosolised marine surface organics are likely to have produced IN in similar concentrations to mineral dust and PBAP.

In summary, both parameterisations based on total aerosol larger than 0.5 µm produced IN concentrations of a similar order of magnitude to the measured first ice concentrations. However, all the composition-specific parameterisations produced IN concentrations several orders of magnitude lower. This therefore leaves us with two possible conclusions: either there was a source of IN that was not characterised by our measurements (i.e. a species other than PBAP, mineral dust and sea spray OA that was significantly active IN at -5 > T > -15°C), and/or one or more of the composition-specific parameterisations underestimate IN concentrations when used in the southwest peninsula of the UK. The measured ice concentrations lie in the temperature range at the high end of most of the parameterisations tested. For example, the data used to generate the fit parameters for T13-F included just seven data points at -10 °C, and the majority of points were from temperatures ≤-20 °C. Mason et al. (2015) also noted the poor performance of IN parameterisations at a coastal site, and noted the good correlation of PBAP with IN between -15 and -25 °C in air from the Pacific Ocean. It is possible the sources of IN are similar between the two studies, but without further characterisation of IN active at temperatures higher than -15°C, we are limited to such speculation.

4. Summary and conclusions

We have presented measurements of boundary layer aerosol concentration, size distribution and composition from a series of research flights performed over the southwest peninsula of the UK during the COPE campaign of summer 2013. Several case studies showed marine air from the Atlantic Ocean advecting to the peninsula from the south/southwest. The CCN and submicron aerosol in these cases were predominantly sulphate-based, and typical for Atlantic marine air. During Easterly winds, polluted air from the UK brought urban pollution to the region; the submicron aerosol was composed of ammonium nitrate and organics, and CCN concentrations were several times higher than the marine cases. In one marine case, IN concentrations were estimated using by inputting field measurements into several size- and composition-specific parameterisations. Bulk parameterisations predicted IN concentrations within an order of magnitude to measured first ice concentrations, but attempts to identify the composition of the IN were largely inconclusive.

The periods of southwesterly winds, which brought marine air from the Atlantic to the peninsula, are of particular interest as they are the most conducive to convective cloud formation in the region. The CCN concentrations in these conditions were
~150 – 280 cm\(^{-3}\) at 0.1% SST, and 400 – 500 cm\(^{-3}\) at 0.9% SST, which are in good agreement with previous measurements in Atlantic air performed at Mace Head (Dall’Osto et al., 2010). The marine aerosol size distributions were also similar to those measured at Mace Head during “clean marine” conditions (Dall’Osto et al., 2011), though some flights had enhanced concentrations of nucleation mode particles, the source of which is not clear from our measurements.

The majority of accumulation mode mass was composed of nss-SO\(_4\) on these days, which is likely to be from biogenic DMS oxidation (Charlson et al., 1987). The high degree of correlation between CCN and nss-SO suggests the CCN were predominantly composed of nss-SO\(_4\), though the origin of these particles (i.e. coated primary or nucleated secondary particles) is again unclear. Accumulation mode OA had a similar mass spectrum to previous measurements of sea spray OA (Ovadnevaite et al., 2011), but a low correlation with CCN, meaning sea spray OA that was not internally-mixed with nss-SO\(_4\) was not a large source of CCN.

By comparing aerosol over the sea to that measured inland, we infer the influence terrestrial emissions had on the marine aerosol. The marine nss-SO\(_4\) was acidic over the sea, but was neutralised by terrestrial ammonia, which is likely to have been emitted by agricultural activities that are widespread throughout the peninsula. The increase in inorganics by this process was almost equally balanced out by a decrease in nss-SO\(_4\). Consequently, the differences between oceanic and inland CCN concentrations were less than the day-to-day variability in different marine cases. Additionally, large enhancements of nucleation mode particles were measured inland, which may also be related to agricultural emissions interacting with marine H\(_2\)SO\(_4\), but these particles were too small to affect CCN concentrations.

The largest difference in aerosol mass loadings and CCN were between the marine flights and a case study where polluted urban emissions advected to the site from the east. The CCN concentrations were ~6 times higher in this airmass, and the inorganics were composed of equal amounts of ammonium nitrate and sulphate. The difference in CCN also corresponded to differences in the cloud base CDNC. In the SWW cases the peak CDNCs were ~450 cm\(^{-3}\), which is in agreement with the CCN measured at 0.9% SST, though most values of CDNC at cloud base were closer to 300 cm\(^{-3}\), which suggests the SST at cloud base was typically between 0.1% and 0.9%. In the polluted case, the peak CDNC was ~1300 cm\(^{-3}\), which was in agreement with CCN measured at 0.1% SST in this case.

Possible sources of IN were assessed by comparing various different parameterisations, using our measured aerosol concentrations as input. We estimated IN concentrations using measurements of total aerosol larger than 0.5 µm (DeMott et al., 2010; Tobo et al., 2013), mineral dust (DeMott et al., 2015; Niemand et al., 2012), fluorescent aerosol (Möhler et al., 2008; Tobo et al., 2013) and sea spray OA (Wilson et al., 2015). The parameterisations based on total aerosol produced IN concentrations that agreed with measured first ice concentrations within an order of magnitude.

In contrast, the predicted IN concentrations derived using all of the composition-specific parameterisations were several orders of magnitude lower. Either there was a source of IN that was not characterised by our measurements, and/or one or more of the composition-specific parameterisations tested greatly underestimated IN concentrations when used in the southwest peninsula of the UK. The lowest cloud-top temperatures during COPE were -15 °C. Better characterisation of IN
at these relatively warm temperatures would be required to gain any further insight into the sources of IN during COPE. It is, however, very unlikely that the ice concentrations of up to several hundred per litre reported by Taylor et al. (2015) could be achieved without invoking the action of secondary ice processes.

The results presented in this paper serve as information for modellers simulating cloud-aerosol interactions for the COPE project, providing details on the aerosol size, composition, CCN and IN in the different case studies. Our data also provide characterisation of Atlantic aerosol, and are likely to be particularly applicable to regions where marine air makes landfall in the absence of strong fossil fuel emissions.

Acknowledgements

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References


Table 1 - Mean aerosol properties measured in NW Coast and inland on each of the case study flights. Where quoted, uncertainty values are standard deviations. All measurements are corrected to standard temperature (273.15 K) and pressure (1013.25 hPa). On 18 July, the CCN column 2 was set to 0.6% SST and 0.9% for all other flying days. Standard deviations are listed for the CCN and CN data, as they are representative of real variations that are much larger than the instrument uncertainty.

<table>
<thead>
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<th>Date</th>
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<th>Org ((\mu\text{g m}^{-3}))</th>
<th>MSA ((\mu\text{g m}^{-3}))</th>
<th>NO(_3) ((\mu\text{g m}^{-3}))</th>
<th>NH(_4) ((\mu\text{g m}^{-3}))</th>
<th>nss-Chl ((\mu\text{g m}^{-3}))</th>
<th>BC (ng m(^{-3}))</th>
<th>CCN 0.1% SST (cm(^3))</th>
<th>CCN Column 2 (cm(^3))</th>
<th>CN (cm(^3))</th>
<th>N(_{\text{GA}}) (cm(^{-1}))</th>
<th>N(_{\text{UGA}}) (L(^{-1}))</th>
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<td>-</td>
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Table 2 - Uncentred R (also known as normalised dot product) between marine OA measured on the SWW case studies and several literature spectra. m/z 28 was not used for the correlations due to gas-phase interference.

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<th>Date</th>
<th>Sea spray (Ovadnevaite et al., 2011)</th>
<th>Marine biogenic SOA (Chang et al., 2011)</th>
<th>Organic factor (Chang et al., 2011)</th>
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Table 3 – Comparison of the m/z 30/46 ratios measured in the AMS aboard the BAe-146.

<table>
<thead>
<tr>
<th>Date</th>
<th>(m/z 30) / (m/z 46) ratio</th>
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<tbody>
<tr>
<td></td>
<td>(NH₄)NO₃ calibration</td>
</tr>
<tr>
<td>05 July</td>
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</tr>
<tr>
<td>18 July</td>
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<tr>
<td>25 July</td>
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</tr>
<tr>
<td>28 July</td>
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<tr>
<td>29 July</td>
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<td>03 August</td>
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Figure 1 - 5-day HYSPLIT back trajectories initiated during the boundary layer aerosol runs. Panels (a) – (f) show the back trajectories on different flying days.
Figure 2 - Spatial location of the below-cloud aerosol runs on each case study flight. Panels (a) – (f) show the six case study flights. The arrow markers show the measured wind speed and direction. The black square shows the location of the Davidstow ground site.
Figure 3 - Aerosol composition measured over the NW coast during the six case study flights. In this plot only, nss-SO$_4$ and nss-Chl are simply referred to as ‘SO$_4$’ and ‘Chl’. The definitions of each chemical species are given in the text. The lower and middle dashed lines show the concentration of NH$_4$ required to neutralise the nss-SO$_4$ to ammonium bisulphate and sulphate respectively. The top dashed line shows the concentration of NH$_4$ required to fully neutralise all measured inorganic components. The reader should consult the text for discussion of the NO$_3$ measurements.

Figure 4 - As in Fig. 3 but for measurements taken inland over the southwest peninsula of the UK.
Figure 5 – Normalised mass spectrum of marine OA measured over the NW coast on 05 July.
Figure 6 - Relationship between the fraction of OA signal at (m/z) = 44 and 43, in coastal and inland areas. The diagonal dotted lines define the region where ambient OOA measurements tend to fall (Ng et al., 2010). The data points labelled “All marine” are the NW coast data from 05, 25 and 29 July and 03 August, whereas the points labelled “All inland” are from all six flights. The literature values are the same as those in Table 2.
Figure 7 – Submicron aerosol size distributions measured by the SMPS and PCASP probes aboard the BAe-146. Panels (a) – (f) show the inland and NW coast distributions for each of the case study flights, while panels (g) and (h) show comparisons of all the NW coast and inland distributions respectively. The error bars in panels (a) – (f) show the standard deviations over each measurement period.
Figure 8 – Comparison of CCN concentrations to CDNC measured in updrafts >2 m s⁻¹ within 500 m of cloud base. The boxes and whiskers show the 25/75th and the maximum/minimum. The triangles are the CCN concentration in marine air, and the squares are the CCN inland.
Figure 9 - Scatter plots of measured CCN concentration at 0.1% SST with concentrations of aerosol species. The data shown are from 18, 25 and 29 July, and 03 August, as CCN data were only available on these days. The blue data points were measured over the NW coast, and the red points were measured inland, as defined in Figure 2.
Figure 10 – As in Fig. 9 but at 0.9% SST. The data shown are from 25 and 29 July, and 03 August only. Data for 18 July are not included as the second CCN column was set to a different SST for this flight.
Figure 11 - Comparison of IN calculations using different parameterisations. The solid lines show the temperature ranges in which the IN parameterisations are deemed valid, while the dashed lines are extrapolations. The inland aerosol measurements were used as input for the parameterisations, other for Wilson et al. (2015) where the marine OA concentration was used. The blue and red markers show measured first ice concentrations from Taylor et al. (2015) at different cloud top temperatures.