Chemical and physical characteristics of aerosol particles at a remote coastal location, Mace Head, Ireland, during NAMBLEX

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Abstract

Aerosol number concentrations and size distributions from 3 nm to 20 µm diameter were measured at the Mace Head Atmospheric Research Station, Co. Galway, Ireland, a coastal site on the eastern seaboard of the north Atlantic Ocean. Both on and offline size resolved aerosol composition measurements were also made using an Aerodyne Aerosol Mass Spectrometer (AMS) and ion chromatographic analysis of daily samples collected using a Micro-Orifice Uniform Deposit Impactor (MOUDI). Particle number concentrations, size distributions and AMS measurements were determined at 7 and 22 m above ground level to investigate local effects on the aerosol size distribution induced by the tidal zone. During periods of new particle formation ultrafine particle number concentrations are large and variable, however, outside these periods no variability in particle number was observed at any size, nor was the particle composition variable. Analysis of particle size distributions show that within each air mass observed particle number concentrations were very consistent. During anticyclonic periods and conditions of continental outflow Aitken and accumulation mode were enhanced by a factor of 5 compared to the marine sector, whilst coarse mode particles were enhanced during westerly conditions. Baseline marine conditions were rarely met at Mace Head during NAMBLEX and high wind speeds were observed for brief periods only.

Loss rates of gaseous species to aerosol surfaces were calculated for a range of uptake coefficients. Even when the accommodation coefficient is unity, lifetimes of less than 100 s were never observed and rarely were lifetimes less than 500 s. Diffusional limitation to mass transfer is important in most conditions as the coarse mode is always significant, we calculate a minimum overestimate of 50% in the loss rate if this is neglected and so it should always be considered when calculating loss rates of gaseous species to particle surfaces. \( \text{HO}_2 \) and HOI have accommodation coefficients of around 0.03 and hence we calculate lifetimes due to loss to particle surfaces of 2000 s or greater. Aerosol composition measurements using the AMS show accumulation mass modes of acidified sulphate and organic material, both of which have the same shape.
and are centred at around 350 nm vacuum aerodynamic diameter, implying an internal mixture. The organic and sulphate are approximately equally important, though the mass ratio varies considerably between air masses. Mass spectral fingerprints of the organic fraction in polluted conditions are similar to those observed at other locations that are characterised by aged continental aerosol. Even in marine conditions a background concentration of between 0.5 and 1 µg m⁻³ of both organic and sulphate was observed. Key differences in the mass spectra were observed during the few clean periods but were insufficient to ascertain whether these changes reflect differences in the source fingerprint of the organic aerosol. However, in an accompanying paper (Dall'Osto et al., 2005) periods of organic dominated aerosol particles were also observed and could be separated from the aged continental aerosol. The coarse mode was composed of sea salt and showed significant displacement of chloride by nitrate and to a lesser extent sulphate in polluted conditions.

1 Introduction

The standard picture of marine aerosols (Van Dingenen et al., 1995; Quinn et al., 1996) has been of a multi-modal distribution, with a fine (Aitken) mode from 20 to 100 nm and an accumulation mode between 100 and 500 nm. Particles greater than 250 nm in (dry) diameter have tended to be thought of as primarily sea salt (Clarke et al., 1997) while smaller aerosol have been thought to be mainly composed of sulphates. Recent work (Nilsson et al., 2001; Geever et al., 2005) shows sea salt aerosol present below 100 nm and fluxes of particles as small as 10 nm being significant depending on sea state. It is known that both sulphate aerosol and sea-salt aerosol play a role in determining the droplet number in marine stratocumulus clouds (O'Dowd et al., 1997; Bower et al., 1999) and this is the key to understanding the first and second indirect effects in these clouds. However, this picture is oversimplified as the role of organic material has not been considered. The presence of significant concentrations of organic material in marine aerosols has been appreciated for several decades (Hoffman and Duce, 1976). Only recently has this organic content been identified as being biogenically derived (O'Dowd et al., 2004). O'Dowd et al report that during phytoplankton blooms in the North Atlantic, the organic content of the submicrometre marine aerosols could constitute up to 63% of the total mass of the particles. By contrast, the winter contribution (low phytoplankton biomass) to aerosol mass was no more than 15%. These data strongly implicate phytoplankton as a critical determinant in the production of marine aerosols. The impact this material has on the cloud-forming properties of these aerosol remains highly uncertain but as the authors point out may be substantial and is likely to be largest for the smaller particles. It is the flux of dissolved organic matter (Larsson and Hågström, 1982) and exopolymer (Grossart, 1999) to particulate organic matter in the surface micro-layer, which is hypothesised to contribute significantly to the organic matter mass in marine aerosols. Dall'Osto et al. (2005) have observed organic rich sea salt particles at Mace Head that are correlated with Magnesium, postulated by Prather et al. (2005) to be derived from chlorophyll in the sea surface microlayer. Raes (1995) suggested that marine boundary layer sulphate may arise via the remote free troposphere where injections of marine sulphur gases have sufficient time to be oxidised at colder temperatures and particle surface losses will be much reduced. However, in the northern hemisphere anthropogenic sources of SO₂ dominate: these include oxidation via SO₂ emissions from shipping and sulphate advected from continental regions in the free troposphere and then mixed into the marine boundary layer (Dore et al., 2000). Anthropogenically derived aerosol contain significant organic material, which is most likely to be internally mixed with sulphate and may be transported significant distances when not lost through wet removal processes (Allan et al., 2004; de Gouw et al., 2005) and like the organic material in sea salt this will affect its cloud-forming properties.

In addition to increasing the albedo of the marine environment and influencing the cloud droplet number of marine stratocumulus clouds, aerosol particles provide signifi-

icant sinks for many trace gas and radicals. Bloss et al. (2005) used a photochemical box model to show that whilst halogen oxides provide an important loss of HO$_2$ in the marine boundary layer under low NO$_x$ conditions, aerosols are very significant. Haggerstone et al. (2005) showed that in clean marine conditions the loss of HO$_2$ is underestimated by nearly a factor of two if aerosol losses are not considered. Likewise, night-time radical losses are also controlled to a significant degree by aerosol particles. Carslaw et al. (1997) demonstrate that under moderate NO$_x$ concentrations the loss of N$_2$O$_5$ to aerosol surfaces can dominate the loss of NO$_2$ and provides a significant loss route for oxidised nitrogen from the gas phase.

This paper presents measurements of the physical and chemical properties of aerosol during the North Atlantic Marine Boundary Layer EXperiment (NAMBLEX). For an overview of the experiment the reader is referred to Heard et al. (2005). The influence of the coastline on the aerosol number and mass size distribution is examined using multi-height measurements as coastal stations are often seen as not being representative of the background marine environment. The measurements are used to identify loss rates of trace gases to particle surfaces as a function of accommodation coefficient and particle size. Both online and offline measurements of aerosol chemical composition are used to investigate the chemical composition of aerosol in the marine boundary layer on the western edge of Europe and explore the organic component of the aerosol in both polluted air transported over long distances and in clean air.

2 Measurements and instrumentation

Microphysical and chemical properties of aerosols were measured from 1 August 2002 until 31 August 2002 during the NAMBLEX experiment close to the shore laboratory of the Mace Head Atmospheric Research Station (53°19′ 34″ N, 9°53′ 14″ W). The site is described in detail in O’Dowd et al. (2002b) and the NAMBLEX experiment is described more fully in Heard et al. (2005). The shore laboratory is 5 m above mean sea level and is a distance of around 50 m from the shore in wind directions from around 180° to 330°. The terrain is mostly low lying and undulating, the soil is predominately peat covered in rough grasses with a significant amount of exposed granite rock.

Aerosol particle number size distributions from 3 nm to 20 µm diameter were measured using a range of differential mobility and optical probes. An in situ forward scattering spectrometer probe (FSSP), (Baumgardner et al., 1985), was used to measure the particle number size distributions from 1 to 20 µm diameter. The instrument was mounted 8 m above the ground at the shore laboratory and was rotated into wind. A large manifold, mounted to a tower was used to deliver ambient air to an air conditioned sea container that served as the main laboratory. The manifold sampled from one of two heights above the ground, 7 and 22 m, alternating on an hourly basis. An isokinetic sub-sampling pipe was fitted to the centre of the manifold at the height of the laboratory, distributing air to the instruments in the container. A particle counter (TSI 3025a) measured the total particle number concentration greater than 3 nm diameter and a Differential Mobility Particle Sizering system (DMPS) was used to measure the number size distribution of particles with diameters between 3 and 750 nm. The DMPS was built to the Vienna design (Winklmayr et al., 1991) and is described in Williams et al. (2000) and more fully in Williams (1999). The DMPS comprised two differential mobility analysers, the smaller of the two was run with an aerosol flow of 1.5 lpm and a sheath flow of 15 lpm, the larger DMA was run with flows of 1 and 10 lpm respectively. Both DMAs sampled dry aerosol and their sheath air lines were run as an open system with a relative humidity below 5%. These instruments were mounted close to the throat of the main sub sample pipe to minimise diffusional losses of small particles. Size distribution measurements presented in this paper are at ambient relative humidity, calculated from composition estimates using simple chemical volume mixing rule approximations.

The dimensions and flow rates of the sampling system were designed to minimise diffusional losses. A particle counter (TSI 3025) and a Scanning Mobility Particle Sizer (SMPS) were run adjacent to the throat of the 22 m manifold inlet for a day and found
to agree with the 3025a and the DMPS sampling through the inlet system to within the instrument uncertainties confirming the particle loss calculations. An optical particle counter (GRIMM 1.105) measured the particle number distribution for particle diameters between 450 nm and 6500 nm from the manifold.

An Aerodyne Aerosol Mass Spectrometer (AMS) system sampled through the main manifold and was used to determine the composition of the non refractory, non sea salt component of the submicron aerosol. The instrument is described in Jayne et al. (2000), whilst details of the data analysis procedures and error analysis are detailed in Jimenez et al. (2003) and Allan et al. (2003). The AMS sampled throughout the month of August 2002 during NAMBLEX. However, due to an instrument failure the data available from 13 August until 19 August, covering most of period 3, have considerably larger uncertainties.

An Aerosol Time Of Flight Mass Spectrometer (ATOFMS – TSI 3800) also sampled from the manifold. Data from this instrument is described in Dall’Osto et al. (2004) and a comparison between the AMS and the ATOFMS is described in Dall’Osto et al. (2005). An eight stage Micro-Orifice Uniform Deposit Impactor (MOUDI) was used to sample aerosol particles onto substrates for subsequent analysis by ion chromato- graphy for Na\(^+\), K\(^+\), Mg\(^{2+}\), NH\(_4\)\(^+\), SO\(_4^{2-}\), NO\(_3^-\), Cl\(^-\). Collections took place between 11 August 2002 and 22 August 2002, each sample period running from 11:00 for 24 h.

The National University of Ireland, Galway, continuously operate a Magee aethalometer model AE-9. This instrument, together with the wind direction, has often been used to identify the clean air sector at the research station (Jennings et al., 1997). The criteria used here are similar to those applied by O’Dowd et al. (2004) and are restricted to those periods when the aerosol absorption is less than 0.475 Mm\(^{-1}\), and the wind direction is to the west and south of 180\(^\circ\) and 330\(^\circ\), respectively. The absorption coefficient is equivalent to a black carbon concentration of 25 ng m\(^{-3}\), calculated using an instrumental attenuation cross section value of 19 m\(^2\) g\(^{-3}\).

3 Meteorological and physical properties of aerosol as a function of airmass

Norton et al. (2005) provided a detailed meteorological characterisation during the NAMBLEX experiment based on local wind data, radar, balloon ascents and ECMWF model products. They identified 5 main periods during August 2002. Statistics showing the main characteristics of the local wind speed, temperature, relative humidity and particle number concentration greater than 3 nm during each of these periods are presented in Table 1. Only a brief summary of that provided in Norton et al. (2005) is given here. Period 1 ran from 1 August to 5 August and could be identified by a semi-permanent decoupling of the surface layer from the mixed boundary layer above arising from a characteristic sea breeze development on most days, whilst the winds aloft were generally from the northeast. As Table 1 shows, the winds were very light and the temperatures were high and the air humid for much of the time. The particle counts show this period was comparatively polluted with a median particle concentration of 1600 cm\(^{-3}\). The air was identified as being from the clean sector only 2.5% of the time during this period.

Period 2 began after the passage of an occluded front on the 5 August and experienced westerly or north-westerly winds. There is some evidence of local influence on the wind profile during this period but the surface winds were similar to those aloft and Mace Head was exposed to the clean sector for 47% of the time. On the 8 August a depression tracked directly over the site and the winds shifted round 360\(^\circ\) though there was little evidence of land influences during the brief period of offshore wind.

During period 3 several frontal systems tracked over Mace Head delivering significant precipitation on most days. The wind direction remained westerly throughout this period both at the surface and aloft and the boundary layer appeared to be coupled for much of the time. This period gave rise to periods of enhanced wind speed, reaching a maximum of 20 m s\(^{-1}\) on the 17 August. Wind speeds on the 14 and 15 also exceeded 15 m s\(^{-1}\) for much of the day. Like periods 2 and 5, this period shows low particle number concentrations for much of the time. Clean air was observed 29% of the time,
mostly in the day, at night the surface air was often decoupled below strong inversions. An anticyclone developed during period 4 from 19 August to 27 August with light winds. Sea breezes were observed from the 18 to 20 August and again on the 25 August. Though the wind direction was from the marine sector for much of the time, the sea breeze circulation regularly introduced polluted air and the clean sector criteria were only met 30% of the time. In the final period between the 28 and 30 August the wind was predominately westerly and well coupled at all levels. The clean sector criteria were met 76% of the time during this period.

The Aitken and accumulation modes in the median number size distributions observed during the marine, predominately cyclonic, periods (3 and 5) were lower than those observed in the more polluted, anticyclonic periods by approximately a factor of 5 and the coarse mode particle number was enhanced by approximately the same ratio (see Fig. 1). In the same figure, the surface area distribution can be seen to have been approximately equally composed of sub and supermicron aerosol. The volume distribution was heavily skewed to the coarse mode as might be expected from a relative clean marine environment. Enhancements in the coarse aerosol mode were from sea spray generation when the wind speeds were significantly above 10 m s\(^{-1}\), which occurred only 14% of the time and only on the days 11, 14–15, 17 and 30 August 2002.

Attenuation coefficients measured by the aethelometer were used to separate aerosol sampled in clean and polluted periods once periods of new particle formation were separated from times when no ultrafine particles were observed in the lowest channels of the DMPS (Fig. 2). There is clear evidence of new particle formation and the existence of an ultrafine mode extending to around 20 nm in all conditions as has been seen previously, with the mode extending to slightly larger sizes in polluted conditions. Most likely this is due to the inter-tidal source of the precursors being further from the measuring site in easterly winds typical during polluted conditions rather than significant addition of anthropogenic secondary aerosol material being formed. There are differences in the accumulation and upper Aitken mode for attenuation coefficients below 1 Mm\(^{-1}\). Above 1 Mm\(^{-1}\) the air is considerably polluted and the accumulation mode size distribution is enhanced, broadens considerably, and displays a shape more characteristic of that of a rural continental aerosol. High winds were not observed during NAMBLEX and the clean sector identified by O’Dowd et al. (2004) was rarely sampled for continuous periods. As a result the very high coarse mode sea spray aerosol loadings observed by O’Dowd et al. (2001) were not seen and the loadings are highly variable and independent of the pollution conditions.

4 Coastal influences on aerosol particles at Mace Head

Coastal monitoring stations, such as Mace Head, provide extremely important sampling locations for investigating the marine atmosphere and long range transport of pollutants, and in the case of Mace Head offer important baseline information. However, the coastal zone can have a significant influence on the particle size distribution in the near shore surface layer. New particle formation has been observed regularly during low tide at Mace Head (O’Dowd et al., 1999) and has been linked to polymerisation reactions of oxidised iodine, the precursors of which appear to be emitted from the large exposed beds of macroalgae (McFiggans et al., 2004; O’Dowd et al., 2002a). The surf zone at the shore provides a large source of coarse mode particles, greatly enhancing the total surface area and volume of the particle population (de Leeuw et al., 2002; Kunz et al., 2002). Norton et al. (2005) have shown that the change in wind stress induced by the change in surface roughness at the shore propagates to a height of between 10 and 15 m at the sample site in all marine wind sectors and tidal states sampled. Hence all shore induced effects on the aerosol particle population at the sampling site will be restricted to heights below 15 m.

In this paper, the effect of the tidal zone on the particle population was studied using the switching inlet system. Figure 3 shows the correlation between the total particle number concentration of diameter greater than 3 nm, measured using the single CPC, sampled through the 7 and 22 m inlets. Minute average data were recorded and the inlet was switched hourly. The data in Fig. 3 are running averages calculated for each
inlet over three full cycles. There are clearly periods when the number concentrations sampled by the two inlets do not agree, when the concentrations are large these are during the nucleation bursts and the average data from the DMPS (not shown) at the ultrafine sizes also show similar variability. At very low concentrations the relative changes can be large though the increase in total number small, as there are few particles during these very clean periods the relative variability also increases at timescales less than an hour and the scatter in the correlation increased. However, for much of the time the correlation is very strong and the gradient of unity shows that there are no significant diffusion losses down the sampling inlet. The size resolved data show that there is little evidence of systematic enhancement of particles at the 7 m sampling level, though, a few local sources of pollution enhance the Aitken mode concentration for brief periods at the 7 m level but not at 22 m. Even particles in the 1 to 3 µm range showed no evidence of the surface layer perturbing the concentration of particles at 7 m above ground level. Figure 4 shows a scatter plot of the submicron sulphate and organic mass as measured by the AMS at the 7 and 22 m level. There is no systematic difference in either the sulphate or the organic concentration sampled at the two levels. A few periods show the enhancement of organic mass at 7 m, this correlates with the enhanced 50 nm particle number concentration. The mass spectra at the 7 m level, not shown, contain aliphatic fragmentation patterns similar to direct vehicle exhaust emissions and identify that these brief periods are due to local combustion sources. The aerosol population at Mace Head is little influenced by the coastal zone over much of its size range. The temporal and spatial variability of the ultrafine particles is known to be high (Hameri et al., 2002) and clearly the averaging methodology used here will not capture this. However, outside these periods the total particle number, the submicron mass loading and the Aitken and accumulation modes and the smaller submicron particles sampled at 7 m or higher remain unaffected by the coastal zone.

5 Diffusion limitations to mass transfer of trace gases to particle surfaces

Aerosol particles provide a sink for many important trace gas and radical species in the marine boundary layer. McFiggans et al. (2002) have shown the importance of the loss of HOI onto deliquesced sea salt particle as a recycling route for reactive iodine in the catalytic removal of ozone in the marine boundary layer. Nitric acid loss to aerosol in the remote marine boundary provides a significant loss of oxidised nitrogen from the gas phase. Accommodation proceeds rapidly, and often under laboratory conditions only a lower limit to the uptake coefficient can be determined as mass transfer limitations dominate. For example Abbatt and Waschewsky (1998) measured a lower limit of 0.2 for the uptake coefficient. Recently, Guimbaud et al. (2002) measured the loss of HNO3 to 70 nm diameter sea salt particles to overcome mass transfer limitations and calculated an uptake coefficient of 0.5, larger aerosol particles were shown to be diffusion limited. Clearly, as the accumulation and coarse modes provide the main surface area and the coarse mode the largest volume, mass transfer limitations of species with high surface uptake rates are likely to be important in the marine boundary layer.

The loss of a gas phase species with uptake coefficient γ. \( k_{\gamma} \) is often taken to be:

\[
k_{\gamma} = \frac{-\tilde{c}_g \gamma A}{4}
\]

(1)

where \( A \) is the particle surface area and \( \tilde{c}_g \) is the average velocity of the gas molecules, given by:

\[
\tilde{c}_g = \sqrt{\frac{8kT}{\pi M_w}}
\]

(2)

where \( M_w \) is the molecular weight of the gas, \( k \) is the Boltzmann constant and \( T \) is the temperature.
However, this expression is only valid in the molecular regime where the mass accommodation dominates the overall loss. If the particles are considerably larger than the mean free path of the gas molecules, this no longer holds and the mass transfer to the particle surfaces must also be considered. Schwartz (1986) gives an expression for $k_{iv}$ in the transition regime:

$$k_{iv} = \frac{4\pi}{3} \int_0^\infty \left( \frac{r^2}{3D_g} + \frac{4r}{3\sigma_g y} \right)^{-1} r^3 \frac{dN}{d\log r} \, dr$$  \hspace{1cm} (3)

where $r$ is the particle radius and $N$ is the number population of particles. The binary gaseous diffusion coefficient, $D_g$, is given by Maitland et al. (1981):

$$D_g = \frac{3}{16n_{air}} \left( \frac{2kT}{\pi M_w} \right)^{1/2} \frac{1}{d_{eff}^2} \hspace{1cm} (4)$$

where $n_{air}$ is the concentration of air molecules at sea level, and $d_{eff}$ is the effective collision diameter of the two molecules of the gas and air, given by the sum of their assumed radii. $M_w$ is the reduced molecular weight or the ratio of the product to the sum of the molecular weights of the diffusing gas and air.

Here we use measurements of the aerosol number size distribution presented in Sect. 3 to show the variability of $k_{iv}$ as a function of air mass history, windspeed, and aerosol size distribution and to show how the uptake is dependent on the uptake coefficient over a range of values that are typical of important molecules in the marine boundary layer. The effect of considering gaseous diffusion as well as accommodation is discussed.

Figure 5 shows time series of the uptake coefficient, $k_{iv}$, for the duration of the campaign with and without consideration of diffusional limitation to the particle surface and for two different values of $\gamma$, 0.03 and 1. The former value is typical of many molecules in the marine boundary layer. An example is HOI, important in the iodine cycle in the marine boundary layer and the molecular mass of HOI is used in the calculation of the binary diffusion coefficient. When $\gamma=1$ accommodation of the gas at the particle surface cannot be faster, providing the upper limit to surface transfer rate. Clearly, a large range of $k_{iv}$ values are observed during the course of the experimental period, considerably more than one order of magnitude variation is seen for either value of $\gamma$. This is typical of such an environment where the coarse mode particle loading is dictated to a large extent by the local windspeed and the pollution loadings of accumulation mode aerosol experienced are highly variable. In the accommodation limit, the lifetimes of the gas with respect to loss on particle surfaces is never less than 100 s and only in extreme wind conditions or during periods of significant aerosol loading does it fall below 200 s. For many gaseous species, such as HOI and HO$_2$, the loss to particle surfaces is relative slow and lifetimes of between $10^3$ and $10^4$ s are more typical (Fig. 5). These losses are often not significant compared to the gas phase cycling of such compounds. More detailed analyses are given in Bloss et al. (2005), and Haggerstone et al. (2005).

Figure 5 also shows the effects of not considering diffusional limitation of mass transfer to the surface. When $\gamma$ is of the order of 0.03, diffusional limitations lead to an increase in the lifetime of around 50% for most of the time during the NAMBLEX experiment. For higher values of $\gamma$ this increased so that in the mass accommodation limit the loss rate of a gas to the particle surface can be over-estimated by up to an order of magnitude.

The normalised size dependence of the uptake rate of a gaseous compound with an uptake coefficient of 0.03, in this example HOI, is shown in Fig. 6 during the clean and polluted conditions observed during NAMBLEX. The accumulation and coarse modes contribute approximately equally to the loss rate on average during the NAMBLEX experiment in both the heavily polluted and clean conditions. Also shown are the corresponding size dependent uptake rates assuming accommodation limitation at the surface only. Uptake to submicron aerosol is unaffected by diffusion limitation to the particle surface and a consideration of mass accommodation alone is sufficient to correctly estimate loss of gaseous material to these particles. However, without consideration of the diffusion limitation uptake rates are overestimated by a factor two for...
coarse mode aerosol in this case. It should be borne in mind that if gases with larger accommodation coefficients are considered these arguments extend to smaller sized aerosol particles. Given that considerably higher coarse mode particle concentrations have been observed in the marine boundary layer than were present during NAMBLEX (O’Dowd et al., 2001) then it is to be expected that uptake rates are dominated by the coarse mode in many clean marine environments.

6 The chemical composition of aerosol at Mace Head during NAMBLEX

Figure 7 shows the time trends of mass loadings of sulphate and organic material measured using the AMS during the NAMBLEX experiment. The five periods identified in Sect. 3 are marked. There is clearly much variability through the experiment with the highest loadings occurring during periods of decoupled (period 1) and anticyclonic (period 4) flow. Black carbon concentrations are also shown and highlight periods of polluted and clean air at the site. The organic and sulphate concentrations are in a broad sense correlated indicating that the organic material is derived from anthropogenic sources, though their relative abundances vary. However, during period 3 the organic concentration is larger than sulphate and not correlated with the latter. This may indicate a marine source, though the uncertainties during this period are large making interpretation during the baseline conditions difficult.

The average, median, quartile and 10th and 90th percentile values for sulphate, organic and ammonium are shown for each period in Fig. 8. Throughout the NAMBLEX experiment, the organic and sulphate mass loadings are greater than 0.5 µg m⁻³ for at least 75% of the time. As already discussed Mace Head experienced few periods of background conditions during NAMBLEX but these data serve to indicate the background level of pollution on the marine edge of western Europe. The exception to this is period 5 which was subject to significant precipitation and hence aerosol removal.

Average size distributions of organic, sulphate and nitrate aerosol mass are shown for polluted (c>1 Mm⁻¹) and clean (c<0.475 Mm⁻¹) periods over the campaign period.

The average size distributions in the clean and polluted periods are similar in shape, though the former is larger in magnitude. The extension of the organic mode to smaller sizes is due to brief periods of near source pollution adding a separate mode similar to that observed at urban locations (Allan et al., 2003) and are not representative of a background aerosol distribution. Nevertheless around 0.5 µg m⁻³ of both sulphate and organic material are present in clean air during NAMBLEX. The modal peaks of the sulphate and organic are similar, implying that the two components are internally mixed, externally mixed particles with different densities would have different vacuum aerodynamic diameters and hence different modal centres.

There is a significant change in the average mass spectrum collected during the clean period and that collected during the polluted period (Fig. 9). The polluted period is more similar to the average mass spectra taken at the Jungfraujoch at an altitude of 3500 m in the Swiss Alps. The latter is representative of mass spectra obtained from a wide range of sites across the northern hemisphere and represents an aged pollution aerosol that is highly oxygenated and is dominated by the CO₂⁻ ion at m/z 44. Zhang et al. (2005) have used a modified principle components analysis to show that the dominant component mass spectrum in many cases is very similar to that shown in Fig. 9. The mass spectrum obtained in clean conditions is similar to that obtained in polluted conditions but differs in a number of ways. Most notably the ratio m/z 44 to m/z 43 is much lower than in the aged pollution mass spectrum and a peak at m/z 56 is present. This may well indicate a different chemical signature and/or source. There is some evidence to suggest that the MS collected during the polluted period could be seen as a superposition of the Jungfraujoch MS and that from the clean period though there is insufficient data to statistically confirm this.

Coarse mode particulate composition was measured during periods 3 and 4 only. As is to be expected the supermicron aerosol was dominated by sea salt. During period 3, the only significant period of background marine air experienced during the impactor sampling period, no significant depletion of chloride was observed. However, under the more polluted conditions during period 4, the acid displacement of chloride can explain
the attainment of charge balance (Fig. 10). Nitrate appeared to be the primary cause of the chloride displacement, contributing at least 3 times more than sulphate on a molar basis (not shown). The impactor analyses also show that the sulphate dominates in the fine mode throughout both sample periods, being an order of magnitude greater than chloride at particle aerodynamic diameters less than 600 nm and only comparable with the sulphate loading at larger sizes. The impactor analyses also show that the sulphate mass dominates in the fine mode throughout both sample periods, being an order of magnitude greater than chloride at particle aerodynamic diameters less than 600 nm. The chloride loadings are only comparable with the sulphate loadings at larger sizes.

### 7 Discussion and conclusions

A characterisation of the physical and chemical composition was conducted at the Mace Head Atmospheric Observatory on the west coast of Ireland during August 2002 as part of the NAMBLEX experiment. Iodocarbon and ultrafine particle measurements have shown that this site is influenced by the littoral zone at low tide. Here multi-height measurements of aerosol number and composition show that Aitken and accumulation mode aerosol are not influenced by the coast under a wide rage of conditions.

An assessment of the diffusion limitation to mass transfer of trace gases to particles was conducted. Gaseous uptake to submicron particle surfaces is dominated by mass accommodation at the surface, even for gases with accommodation coefficients that approach unity. However, the uptake rates of gases to coarse mode particles are significantly suppressed by gaseous diffusion limitations. Under the conditions observed during NAMBLEX such reductions in uptake rate can be as large as a factor of 5 for high accommodation rates and even a factor of 2 for accommodation coefficients of 0.03, typical of a wide range of trace gases in marine boundary layers. Higher winds, generating significant sea spray, will most likely enhance this phenomenon.

The measurements of aerosol chemical composition show that there is often a significant semi ubiquitous background organic component of the submicron particle mass at Mace Head in a wide range of conditions with a mass loading of between 0.5 and 1 µg m$^{-3}$ and there is often evidence of significant organic and sulphate mass in air masses transported over considerable distances. The organic mass fraction appears to have a mass spectral signature that is similar to other sites sampled with an AMS in the northern mid latitudes (Allan et al., 2003, 2004; Alfarra et al., 2004) during polluted conditions. The shape of the mass distributions of organic and sulphate are similar suggesting that the components are often internally mixed and hence indicate long range transportation. However at times there is an increase in the organic mass at larger sizes that does not correlate with sulphate or other aged pollution markers. This is discussed further in Dall’Osto et al. (2005). There is little contribution of sea salt to the submicron aerosol, though it dominates the coarse mode. When exposed to polluted air the coarse mode aerosol shows evidence for chloride displacement that appears largely due to nitrate uptake.

O’Dowd et al. (2004) have shown that significant organic material is present in the submicron marine aerosol during clean conditions (identified by measured absorption coefficients below 0.475 Mm$^{-1}$ and wind directions between 180° and 330°). They link the organic in the particles to that found in the organic material in the sea surface microlayer and postulate that sea spray generation through bubble bursting when a surfactant layer is present is likely to lead to submicron particles with a high organic content and little sea salt. The particles were observed to extend in size down to a hundred nanometres or less. Dall’Osto et al. (2005) identified Magnesium rich particles at Mace Head during high wind speeds in the clean air sector using a laser desorption-ionisation single particle time of flight mass spectrometer and showed that organic material is present in the same particles. Prather et al. (2005)$^1$ were the first to identify Mg in particles combined with organic material and postulated that this arises from Magnesium contained in chlorophyll.

In this paper the mass spectral fingerprints of the organic fraction of the submicron aerosol as measured by the AMS show distinctly different fragmentation from the more polluted periods. These changes may well reflect the change in source, such as the pri-
mary surfactant organic formed from sea spray generation as postulated by O’Dowd et al. (2004). Dall’Osto et al. (2005) separated AMS size distributions that occurred during periods when Mg-rich particles were observed from those measured during other periods. Although most of the organic mass was shown to have a similar size distribution to that of sulphate, indicating long range transported pollution similar to that observed in other periods, a second mode was observed in particles above 1000 nm diameter, extending beyond the 50% upper size cut off of the inlet lens. Mass spectral analysis of the periods when Mg was present and when it was not, do not show statistically significant differences with the AMS. However, the larger mode appeared at sizes that are transmitted very inefficiently by the AMS lens and therefore material in such particles is undersampled and does not contribute to the total measured particle ensemble mass spectra. It is difficult to establish whether the organic material in the mode seen at large particle sizes is the same as that observed by O’Dowd et al. (2004) as the ultra-clean conditions observed by O’Dowd et al. (2004) were not observed during NAMBLEX. Although the measured size distributions were different to those observed in O’Dowd et al. (2004), the ATOFMS has a strongly size dependent efficiency and may simply not have observed smaller particles of a similar composition. In addition, the AMS only operated for a fraction of the available time in the clean conditions during NAMBLEX. Furthermore, the presence of a significant mixed sulphate-organic mass mode prevented causal differences being established from the AMS data set presented here as an unequivocal mass spectrum could not be obtained to identify a possible second source of organic material.

Though these results indicate that under clean conditions there may be a significant change in the mass spectral fingerprint of the organic fraction and this may give rise to size dependent changes in the organic aerosol size distribution there is at present insufficient data to draw causal relationships from the observations. Size dependent mass spectra collected over a longer time period in clean conditions would enable such relationships to be drawn. Such measurements are now possible using an AMS fitted with a Time of Flight Mass spectrometer (Drewnick et al., 2005) instead of the quadrupole. This instrument is capable of producing size resolved mass spectra with a much improved signal to noise ratio. The instrument is also capable of delivering complete mass spectra of individual particles, enabling the study of mixing states. This will in turn allow the direct observation of particles produced locally from the sea surface layer separately from the background aerosol. It is recommended that such a study is carried out to determine the variability and size distribution of the organic component of marine aerosol.

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Table 1. Statistics for wind speed, temperature, relative humidity and total particle number concentration greater than 3 nm diameter in the five sampling periods identified by Norton et al. (2005).

<table>
<thead>
<tr>
<th>Period</th>
<th>Parameter</th>
<th>10 Percentile</th>
<th>25 Percentile</th>
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<th>mean</th>
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Fig. 1. Median number (a), surface area (b) and volume distributions (c) for each of the five sampling periods. Aerosol data presented at ambient humidity.

Fig. 2. Average number size distributions separated as a function of attenuation coefficient, $c$. The value of 0.475 Mm$^{-1}$ represents the identified clean air sector threshold used by O'Dowd et al. (2004) and periods of nucleation around low tide are considered separately to the remainder of the data.
Fig. 3. Correlation between total particle number concentration greater than 3 nm diameter measured from the 22 m inlet and the 7 m inlet. The data were sampled alternately through each inlet on an hourly basis and each data point represents a running average over three cycles.

Fig. 4. Correlation between the submicron organic and sulphate mass loadings sampled through the 7 m and 22 m inlets. The data were sampled alternately through each inlet on an hourly basis and each data point represents a running average over three cycles.
Fig. 5. Time series of uptake coefficients calculated using the expression provided by Schwartz (1986) (Eq. 3), considering diffusional limitation to mass transfer of gas to the particle surfaces compared to those assuming only a mass accommodation limitation (Eq. 1). Two different mass accommodation coefficients are shown to represent the mass accommodation limit, $\gamma = 1$, and a value representative of a range of trace gases in the marine boundary layer, $\gamma = 0.03$. Values of molecular diameter and molecular mass for HOI have been used to in the calculation of the binary diffusion coefficient.

Fig. 6. Normalised size dependent uptake coefficients for clean and polluted conditions during NAMBLEX calculated considering diffusion limitation to the particle surface and compared with those assuming mass accommodation limitation only. An accommodation coefficient of 0.03 has been assumed and values of HOI were incorporated in the calculation of the binary diffusion coefficient.
Fig. 6. Normalised size dependent uptake coefficients for clean and polluted conditions during NAMBLEX calculated considering diffusion limitation to the particle surface and compared with those assuming mass accommodation limitation only. An accommodation coefficient of 0.03 has been assumed and values of HOI were incorporated in the calculation of the binary diffusion coefficient.

Fig. 7. Time series of the submicron organic and sulphate mass loadings as measured by the AMS during NAMBLEX. The five different meteorological periods are clearly marked. The bottom panel shows the black carbon mass concentrations derived from the aethelometer.

Fig. 8. The mean, median, and 10, 25, 75, and 90th percentiles of the submicron mass loadings of organic (a), sulphate (b) and ammonium (c) measured by the AMS during each of the five meteorological periods identified during the NAMBLEX experiment.
Fig. 9. Submicron average mass size distributions of organic, sulphate and nitrate measured using the AMS during polluted (c>1 Mm$^{-1}$) and clean (<0.475 Mm$^{-1}$) (panel a). Also shown are the averaged organic mass spectra for the same two conditions during the NAMBLEX experiment. A mass spectrum taken using the same AMS at the Jungfraujoch is shown for comparison (panel b).

Fig. 10. Mean (bows), median (horizontal line), 25th and 75th percentiles (grey box) and 10th and 90th percentiles (black vertical line) of the supermicron Na:Cl molar ratio for periods 3 and 4. Also shown are the corresponding charge equivalent ratios accounting for displacement of chloride by nitrate and non sea salt sulphate. It is worth noting that Na:Cl in sea salt is 0.86.