Intercomparison of modal and sectional aerosol microphysics representations within the same 3-D global chemical transport model

G. W. Mann¹², K. S. Carslaw², D. A. Ridley³, D. V. Spracklen², K. J. Pringle², J. Merikanto⁴, H. Korhonen⁵, J. P. Schwarz⁶, L. A. Lee², P. T. Manktelow², M. T. Woodhouse², A. Schmidt², T. J. Breider⁷, K. M. Emmerson², C. L. Reddington², M. P. Chipperfield², and S. J. Pickering²

¹National Centre for Atmospheric Science, University of Leeds, Leeds, UK
²School of Earth and Environment, University of Leeds, Leeds, UK
³Dept of Atmospheric Science, Colorado State University, Fort Collins, Colorado, USA
⁴Department of Physics, University of Helsinki, Helsinki, Finland
⁵Finnish Meteorological Institute, Kuopio Unit, Kuopio, Finland
⁶Cooperative Institute for Research in Environmental Sciences, University of Colorado, Boulder, Colorado, USA
⁷School of Engineering and Applied Sciences, Harvard University, Cambridge, MA, USA
3-D global intercomparison of bin versus mode

G. W. Mann et al.

* now at: Halcrow Group Ltd, Headingley, Leeds, UK
* now at: CSIRO Marine and Atmospheric Research, Aspendale, Victoria 3195, Australia

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Correspondence to: G. W. Mann (g.w.mann@leeds.ac.uk)
Abstract

A global modal aerosol microphysics module (GLOMAP-mode) is evaluated and improved by comparing against a sectional version (GLOMAP-bin) and observations in the same 3-D global offline chemistry transport model. With both schemes, the model captures the main features of the global particle size distribution, with sub-micron aerosol approximately unimodal in continental regions and bi-modal in marine regions. Initial bin-mode comparisons showed that various size distribution parameter settings (mode widths and inter-modal separation sizes) resulted in clear biases compared to the sectional scheme. By adjusting these parameters in the modal scheme, much better agreement is achieved against the bin scheme and observations. Surface mass of sulphate, sea-salt, black carbon (BC) and organic carbon (OC) are, on the annual mean, within 25% in the two schemes in nearly all regions. On the annual mean, surface level concentrations of condensation nuclei (CN), cloud condensation nuclei (CCN), surface area density and condensation sink also compare within 25% in most regions. However, marine CCN concentrations between 30°N and 30°S are systematically higher in the modal scheme, by 25–60%, which we attribute to differences in size-resolved particle growth or cloud-processing. Larger differences also exist in regions or seasons dominated by biomass burning and in free-troposphere and high-latitude regions. Indeed, in the free-troposphere, GLOMAP-mode BC is a factor 2–4 higher than GLOMAP-bin, likely due to differences in size-resolved scavenging. Nevertheless, in most parts of the atmosphere, we conclude that bin-mode differences are much less than model-observation differences, although some processes are missing in these runs which may pose a bigger challenge to modal schemes (e.g. boundary layer nucleation, ultra-fine sea-spray). The findings here underline the need for a spectrum of complexity in global models, with size-resolved aerosol properties predicted by modal schemes needing to be continually benchmarked and improved against freely evolving sectional schemes and observations.
1 Introduction

Aerosol particles affect the radiative budget of the Earth’s atmosphere by scattering and absorbing solar and terrestrial radiation and by modifying the albedo and lifetime of clouds, referred to as the direct and indirect aerosol radiative effects (e.g., Seinfeld and Pandis, 1998). A better understanding of how the atmospheric aerosol has changed since the pre-industrial era is vital to improve the robustness of model predictions of anthropogenic climate change. Successive climate assessment reports (Schimel et al., 1996; Penner et al., 2001; Forster et al., 2007) have continued to classify aerosol radiative forcings as having a low level of scientific understanding and larger uncertainty than forcing from changes in long-lived greenhouse gases. However, Kulmala et al. (2011) show that in recent years, improved representations of aerosol properties and sources is leading to a substantial narrowing of the aerosol forcing uncertainty range.

In-situ observations and process modelling have led to major advances in the understanding of key aerosol processes and how they determine the evolution of the particle size distribution, and hence concentrations of cloud condensation nuclei (CCN). For instance, observations of marine aerosol size distributions (e.g., Hoppel et al., 1994) have shown a clear separation between Aitken and accumulation modes at around 100 nm dry diameter, most likely due to growth of activated particles via in-cloud aqueous sulphate production. Observations of high number concentrations in the upper troposphere in both marine (e.g., Clarke, 1993) and continental regions (e.g., Hofmann, 1993) combined with modelling of atmospheric transport (e.g., Raes, 1995) have led to an understanding that new particle formation in the free troposphere and subsequent entrainment and mixing into the boundary layer represents a major source of marine CCN (e.g., Raes et al., 2000; Merikanto et al., 2009). More recently, a wide range of observations (e.g., Kulmala et al., 2004) have shown that, in addition to nucleation in the free troposphere, new particle formation occurs very frequently in the boundary layer in a variety of continental environments.
The majority of general circulation models used to provide estimates of aerosol radiative forcings in the fourth IPCC climate assessment report (Forster et al., 2007) used relatively simple aerosol schemes whereby the main components (sulphate, sea-salt, carbonaceous aerosol and dust) are represented by distinct externally-mixed types with only the mass of each transported, and the particle size distribution prescribed at globally uniform values (e.g., Jones et al., 2001; Reddy et al., 2005). Fixing the size distribution means that any growth process included in the models (for instance aqueous sulphate production in clouds) will increase particle number when it increases mass, potentially causing artefacts in simulated aerosol-cloud interactions.

Since the early 1980s, detailed aerosol dynamics schemes, developed initially in box models, emerged to capture aerosol microphysical processes such as new particle formation, and growth by coagulation and condensation. For instance Gelbard et al. (1980) were among the first to produce a sectional aerosol model whereby the general dynamics equation (GDE) over the continuous size spectrum is reformulated according to a discrete set of size sections or bins. Whitby (1981) developed the computationally cheaper “modal” approach, whereby the continuous GDE is given in terms of integral moments of the size distribution within log-normal modes covering different parts of the particle size range.

The growing realisation of the importance of aerosol microphysical processes in determining aerosol properties led to the implementation of these more complex aerosol schemes into global models. For instance Adams and Seinfeld (2002) developed a dual moment sectional global aerosol microphysics model and Ghan et al. (2001) and Wilson et al. (2001) implemented two-moment modal schemes into global models. Following these pioneering studies, a new generation of global aerosol microphysics models have now been developed with both sectional (e.g., Spracklen et al., 2005, 2008; Kokkola et al., 2008; Luo and Yu, 2011) and modal schemes (Easter et al., 2004; Stier et al., 2005; Liu et al., 2005; Lauer et al., 2005; Bauer et al., 2008; Mann et al., 2010; Pringle et al., 2010; Vignati et al., 2010). Furthermore, Ghan and Schwarz (2007) explain that international climate modelling groups performing co-ordinated experiments...
for IPCC assessment reports are developing new climate model versions that include more sophisticated aerosol schemes to improve the fidelity of simulated climate forcings.

Modal approaches continue to be favoured over sectional schemes in global models due to lower computational costs. However, simplifications in the parametrized modal approach (e.g. fixing the standard deviation) can cause biases in simulated process rates and size distributions (e.g., Seigneur et al., 1986; Zhang et al., 1999). In box model studies, Herzog et al. (2004) found that number concentrations and surface area density simulated by a modal scheme are, on average, within 20% of a similar bin-resolved version, but found differences of 50% on average for simulated accumulation-mode number concentrations. Kokkola et al. (2009) found larger differences between sectional and modal schemes in volcanically perturbed stratospheric conditions, and explored ways to modify the modal scheme to reduce such biases.

Although these box model studies have demonstrated differences in certain conditions, the extent to which the use of modal schemes in 3-D global models leads to systematic biases in simulated aerosol properties has not yet been assessed. This study, for the first time in a 3-D global model, compares modal and sectional aerosol schemes, sharing the same process representations. Specifically, we compare the two-moment sectional (Spracklen et al., 2005, 2008) and two-moment modal (Mann et al., 2010) versions of the GLObal Model of Aerosol Processes (GLOMAP), known as GLOMAP-bin and GLOMAP-mode, respectively. Trivitayanurak et al. (2008) compared GLOMAP-bin against another two-moment sectional scheme (TOMAS, Adams and Seinfeld, 2002) in different 3-D global models, and found major inter-model differences and discrepancies to observations, but the schemes had different process representations, emission inventories, size assumptions, oxidant fields, clouds and transport. By contrast, here, we have the bin and mode schemes in the same 3-D global offline chemistry transport model (Chipperfield, 2006) with the same process representations and parameter settings, and ensure the intercomparison has the same meteorology, oxidants, clouds and aerosol precursor chemistry.
By examining particle size distributions simulated with the two schemes, we also aim to provide constraints for the choices of parameter values in the modal scheme to improve the comparison to the more sophisticated sectional scheme. We quantify differences in a wide range of simulated integral particle properties such as total and size-resolved particle number, speciated mass, CCN concentrations, surface area density and condensation sink. We also compare both models to benchmark observational datasets which constrain the skill of the simulated particle size distribution in marine and continental regions, both at the surface and in the vertical profile.

The GLOMAP-mode scheme has also been implemented (Johnson et al., 2010) in the HadGEM3-UKCA composition-climate model (Morgenstern et al., 2009; Telford et al., 2009). By making aerosol properties simulated by the modal scheme compare better to the sectional scheme in the offline transport model, we aim to increase the robustness of simulated aerosol radiative forcings, making simulations in the composition-climate model more reliable.

2 Model description

GLOMAP-bin and GLOMAP-mode are comprehensively described in Spracklen et al. (2005) and Mann et al. (2010), respectively. The GLOMAP-bin model used here is the multi-component version (v1a as in Merikanto et al., 2009), and differs from that used in the first papers (e.g., Spracklen et al., 2005; Spracklen et al., 2006) which did not discriminate between different aerosol components (e.g. sulphate, sea-salt, black carbon).

The GLOMAP-mode model here (v6), differs slightly from v5 in Mann et al. (2010), having been updated to more closely follow process formulations in GLOMAP-bin. These changes modify the routines for vapour condensation, nucleation scavenging and aqueous sulphate production, and are described in Appendix A.

Both GLOMAP schemes are run within the TOMCAT chemistry transport model (Chipperfield, 2006) using the approach described in Spracklen et al. (2005). Oxidant
concentrations are prescribed according to temporal interpolation between monthly-mean 3-D fields (at 00:00, 06:00, 12:00, 18:00 UT) from a full-chemistry simulation (Arnold et al., 2005). Wind and temperature fields in the model are also prescribed via interpolation between 6-hourly meteorological re-analysis fields from the European Centre for Medium-range Weather Forecasts (ECMWF).

2.1 Host model settings for the model runs

The GLOMAP-bin and GLOMAP-mode runs are at \( \approx 2.8^\circ \times 2.8^\circ \) horizontal resolution on 31 vertical hybrid sigma-pressure levels from the surface to 10 hPa. Monthly-means from January to December 2000 are used following a spin-up of 3 months from zero initial aerosol. Gas phase and aqueous chemistry are identical in the runs following that described in Mann et al. (2010). Briefly, gas phase reactions of DMS, SO\(_2\) and monoterpenes with OH, NO\(_3\) and O\(_3\) are included with a small source of SO\(_2\) from COS and CS\(_2\). Aqueous phase oxidation of dissolved SO\(_2\) is simulated via reaction with H\(_2\)O\(_2\), which is treated semi-prognostically, being transported, depleted via SO\(_2\) and replenished by gas-phase HO\(_2\) self-reaction up to a background value given by the prescribed 3-D oxidant fields. The ASAD chemical integration software (Carver et al., 1997) is used with identical rate files and gas-phase deposition settings, with tendencies for the transported gas-phase species given by the IMPACT algorithm (Carver and Stott, 2000).

The model runs were driven by identical offline fields of meteorology (6-hourly ECMWF ERA-40 re-analyses), cloud fields (monthly climatological low-cloud from ISCCP, Rossow and Schiffer, 1999) and oxidants (6-hourly monthly-means from a full-chemistry TOMCAT run, Arnold et al., 2005). Gaseous and primary particulate emissions for both models are as described in Mann et al. (2010) including DMS, SO\(_2\), COS, CS\(_2\), monoterpenes, primary sulphate, carbonaceous aerosol (speciated to BC and organic carbon, OC) and sea-salt, mostly following the AEROCOM recommended sources as in Dentener et al. (2006). The non-local closure boundary layer mixing scheme from Holtslag and Boville, (1993) and the convection parameterization from
Tiedtke (1989) is used, with identical settings for bin and mode runs. This set-up ensures that atmospheric transport, primary aerosol sources from direct emission and secondary sources from gas phase precursor oxidation were equivalent in all simulations. Note however, that differences (up to 20% in some marine regions) did arise between the bin and mode simulated gas phase precursors (see Table 6) which we attribute to different host transport model versions being used.

3 Comparison of simulated particle size distributions

Figure 1 compares simulated surface-level particle size distributions at three marine and three continental locations for GLOMAP-bin (black) and two GLOMAP-mode runs in June (a–f) and December (g–l). The first GLOMAP-mode run (blue line) has modal parameter settings as in Mann et al. (2010) with a standard deviation (σ) of 2.0 for the coarse mode and 1.59 for all other modes, and mode-edge radii at 5, 50 and 500 nm as separating nucleation-Aitken, Aitken-accumulation and accumulation-coarse modes, respectively. Note that these mode-edge dry-radii determine how primary emissions are mapped onto the modes, and is also the size at which particles are transferred to the adjacent larger mode by mode-merging. The second run (red line) has revised modal settings, to improve comparison with the bin scheme, whereby the soluble accumulation mode is set narrower with σ = 1.40 and the accumulation-coarse mode-edge dry-radius (r_{3,4}) is reduced to 250 nm.

In the three ocean locations (Fig. 1a–c and g–i), all three runs capture the main features of the marine boundary layer size distribution. During summer (Fig. 1a, b, i) the sub-µm aerosol is bi-modal (Aitken and accumulation) with a third coarse mode from sea-spray. During winter, (Fig. 1c, g, h), all three runs also show an additional distinct nucleation-mode below 5 nm dry radius that is not present during summer, or has merged with the Aitken-mode following growth.

In the three marine locations, and in both seasons, the model runs all show a “Hoppe-l gap” at about 25–40 nm dry radius, created by growth of activated particles via
in-cloud sulphate production (e.g., Raes, 1995). However, with the original modal settings ($\sigma_{\text{acc}} = 1.59$ and $r_{3,4} = 500\,\text{nm}$), the minimum in $dN/d\log r$ between Aitken and accumulation modes is, compared to bin, biased high in summer in all regions (Fig. 1a, b, i). For this run, the accumulation mode radius is also biased substantially high in winter against the peak in the bin scheme (Fig. 1c, g, h). With these original modal settings, the accumulation mode, is also too wide compared to bin in all 3 locations. By contrast, when $\sigma_{\text{acc}}$ is set to 1.4, and $r_{3,4}$ also reduced to 250 nm, GLOMAP-mode performs much better, with the size of the accumulation-mode peak, and its shift from summer to winter, matching well to the sectional scheme.

Another bias in GLOMAP-mode (with the original settings) is that, in all three marine locations, the coarse mode number is much less than in GLOMAP-bin (by factor 2–3) in both seasons. With the revised settings however, GLOMAP-mode compares well, due to the sea-spray emissions then being mapped more coherently onto the log-normal modes. In the sectional scheme, the simulated coarse mode begins at about 300 nm dry-radius with a peak at about 500–600 nm (Fig. 1). In GLOMAP-mode, the coarse-soluble mode only receives emitted sea-spray larger than $r_{3,4}$, so with this set to 500 nm the modal approach effectively splits the sea-spray mode in two where there should be a peak, leading to the low bias in coarse mode number. In the revised modal configuration (red line), the split occurs at 250 nm, matching that seen in the size distribution for the freely-evolving sectional scheme. The accumulation mode is also compromised in GLOMAP-mode when $r_{3,4} = 500\,\text{nm}$. Since the modal approach has to construct a single log-normal distribution for all particles in a mode, it must combine any sea-spray emitted into the mode with finer sulphate particles. In the $r_{3,4} = 500\,\text{nm}$ run, this effect pulls the mode radius to erroneously large sizes. whereas with $r_{3,4} = 250\,\text{nm}$, the bias from the effect is greatly reduced since sea-spray particles are then mostly emitted into the soluble coarse mode.

Figure 2 further examines accumulation mode size, showing the latitudinal variation of Aitken and accumulation mode dry diameter from the two GLOMAP-mode runs against observed values derived from a 30-yr compilation of marine size distribution.
measurements (Heintzenberg et al., 2000). The observed mean dry diameter values are from 3- and 4-mode log-normal fits to the ship-borne mobility and aerodynamic particle sizer measured dry size distributions (below 40% relative humidity). To compare against these observations, model marine-zonal-mean dry diameter values for each mode were obtained via number-weighted means over all ocean gridboxes on each latitude grid-point, and averaging up to the 15 degree grid in the observations. There is no significant difference in simulated Aitken-mode size or number between the two GLOMAP-mode runs, and good agreement between model and observations, with larger particle size in the tropics compared to mid-latitudes. However, in the accumulation mode, the original GLOMAP-mode settings give too large size in the Southern Ocean (due to the sea-spray effects described above), whereas much better agreement is seen with the revised settings.

In the three continental locations (all in the Northern Hemisphere), sub-µm size distributions in winter (Fig. 1j–l) are uni-modal in all three runs, with a peak at about 25 to 35 nm dry radius. A pronounced accumulation-mode shoulder is evident in summer (Fig. 1d–f) up to about 60–90 nm dry radius. However, in both GLOMAP-mode runs, the Aitken-mode number peak is substantially biased low compared to bin, particularly in winter, and is also too narrow. The low $dN/d\log r$ may indicate deficiencies in the modal treatment of coagulation where rates based on the mode mean radii could be too high. Another possible cause is that primary carbonaceous emissions are sized here to have $\sigma = 1.8$ following Dentener et al., 2006). Whereas the size sections in GLOMAP-bin can adapt freely to this prescribed shape, in GLOMAP-mode, the Aitken mode is forced to be narrower, held fixed at 1.59.

In the accumulation-mode, GLOMAP-mode compares better to bin than in the Aitken sizes. The size distribution for all three runs shows a similar pronounced shoulder during summer but, again, the original configuration of GLOMAP-mode has the accumulation mode too wide in the large-end tail, whereas much better agreement is achieved with the narrower $\sigma$. 
Figure 3 compares against a climatological observed size distribution from Raes et al. (2000) for the marine boundary layer at Tennerife in July, which combines differential mobility analyzer (sub-µm) and aerodynamic particle sizer (super-µm) measurements. In marine locations, the observed size distribution is tri-modal (as seen in the model in Fig. 1) with distinct Aitken and accumulation mode separated by a minimum at about 50 nm dry radius, and a distinct coarse mode at dry-radii larger than 300 nm. Figure 3 shows in black lines the July monthly-mean from GLOMAP-bin (solid), GLOMAP-mode as configured originally (dashed) and with the revised modal-settings (dot-dashed). The figure confirms that an accumulation mode $\sigma$ of 1.59 is too wide, with the $\sigma = 1.4$ run agreeing much better with the observed tail of the accumulation-mode. The observed coarse mode is also improved with the revised GLOMAP-mode settings, showing that the better agreement with bin also reduces bias against the measurements. The blue/red lines in Fig. 3 are minima/maxima over the 12 monthly-means and illustrate that revising the modal settings also improves simulated intra-annual variability in size distribution compared to bin.

4 Sub-micron mode widths

In Sect. 3, we found that size distributions simulated by the sectional scheme, which can evolve freely in response to the processes (without parametric constraints) result in a large-end tail of the accumulation mode consistent with $\sigma = 1.4$. The original GLOMAP-mode settings for the standard deviation were taken from the literature, matching those in the M7/HAM models (Vignati et al., 2004; Stier et al., 2005) with $\sigma = 1.59$ for all three sub-µm modes. The value of 1.59 originates from Wilson et al., (2001), and represents a compromise between values suggested by self-preserving theory (1.45) and from observations (1.4–2.0). In the light of the findings in the previous section, we revisit this issue and review values from observations in the literature.

Whitby (1978) presented a synthesis of observed size distributions from a range of environments, which suggested $\sigma$ should be between 1.8 and 2.2. Heintzenberg
et al. (2000) compiled 30 yr of marine dry size distribution observations and presented a global variation of Aitken and accumulation mode log-normal fit parameters on a 15 degree latitude grid. Their reported observed values of the accumulation mode standard deviation $\sigma_{\text{acc}}$ are between 1.4 and 1.6 with most values at 1.4. Birmili et al. (2001) follow a similar approach from log-normal fits to size distribution measurements at a continental site in Germany. They find the Aitken mode is generally wider than the accumulation mode with $\sigma_{\text{Ait}}$ tending to vary between 1.55 and 1.73, and $\sigma_{\text{acc}}$ between 1.41 and 1.57. However, Pirjola et al. (1999) find the opposite from measurements at the Hyytiala boreal forest site in Finland (Makela et al., 1997), finding $\sigma_{\text{Ait}} = 1.5$ and $\sigma_{\text{acc}} = 1.7$. Petzold et al. (2002) applied tri-modal log-normal fits to size distributions measured over Germany by aircraft-borne optical particle counters and found $\sigma$ for the accumulation mode to vary between 1.28 and 1.6 with the majority of values between 1.3 and 1.4. Asmi et al. (2011) present a synthesis of sub-µm size distribution measurements from 24 sites in Europe and applied bi-modal log-normal fits to the median size distributions in the 20–500 nm dry-diameter range. They find large spatial variation in $\sigma$, with values for the Aitken and accumulation modes being in the range 1.47–2.49 and 1.27–2.44, respectively.

Overall, although the observations suggest $\sigma$ varies substantially between different sites and environments, our revised values of 1.59 and 1.40 for the constant standard deviations for Aitken and accumulation soluble modes are reasonably consistent with findings from size distributions measured in both marine and continental regions. We also note that Pirjola et al. (1999) found that the self-preserving distribution for modal schemes which allow $\sigma$ to evolve in the model gave values in the range 1.36–1.45, giving additional evidence in support of reducing $\sigma$ for the accumulation mode (the longest lived of the modes) from 1.59 to 1.40.
5 Comparison of regional CN and CCN concentration

In this section, we quantify the impact of the changes to the mode-edge radius and standard deviations on simulated CN and CCN. Table 1 shows regional mean concentrations of CN (all particles with dry-diameter \(D_p > 3\) nm), \(\text{CCN}_{50}\) and \(\text{CCN}_{150}\) (soluble particles with \(D_p > 50\) and 150 nm) for the regions used in Merikanto et al. (2009).

The revisions to \(\sigma_{\text{acc}}\) and \(r_{3,4}\) have only a minor effect on GLOMAP-mode simulated CN and \(\text{CCN}_{50}\), although in each of the regions, the 10–20% high bias compared to bin is reduced slightly. However, simulated \(\text{CCN}_{150}\) are substantially too low with the original modal settings, with the low bias compared to bin in the range 33–45% in Europe, North America, North Asia, South East Asia and Oceania. On the global continental average, the modal scheme, as originally configured is 40% lower than bin, reflecting the biases in particles with \(D_p > 100\) nm seen in Fig. 1d–f, j–l. With the revisions to the modal settings, \(\text{CCN}_{150}\) agrees better with the bin scheme, with the low bias reduced in all regions, although still at 23% on the global continental average.

6 Comparison of model global burden and budgets

In this section, we examine the global aerosol lifecycle simulated by the bin and mode schemes, with the GLOMAP-mode run using the revised modal settings. Table 2 compares annual-mean global column-integrated mass burdens for each aerosol component (sulphate, sea-salt, BC and particulate organic matter, POM), along with their source mass fluxes from primary emissions and secondary production. Simulated lifetimes and percentage removal by wet deposition are also shown to aid the analysis.

Compared to GLOMAP-bin, simulated sulphate and POM burdens are slightly lower in GLOMAP-mode (−12 and −3%) whereas BC and sea-salt are slightly higher (+4 and +9%). These are reflected in the slightly shorter lifetimes for GLOMAP-mode simulated sulphate and POM, while BC and sea-salt are longer-lived with the modal approach. However, for each species, the global burdens compare well, with GLOMAP-mode...
within about 10 % of GLOMAP-bin. To set these differences in context, we note the findings in Textor et al. (2006), who examined diversity in simulated lifetimes among the AEROCOM models, finding standard deviations among the models of 58, 43, 18, 33 and 27 % for sea-salt, dust, sulphate, BC and POM, respectively. Thus, inter-modal diversity is much larger than the difference introduced by the simplified model treatment of the evolving size distribution. The percentage removal by wet deposition illustrates that wet removal is the dominant removal process for sulphate, BC and POM, which reside mainly in sub-µm particle sizes, whereas the coarser sea-salt aerosol is influenced strongly by sedimentation. While the bin and mode schemes predict similar wet removal for sulphate, BC and POM, there is a substantial difference for sea-salt, with 27.1 % of mass removal by wet deposition in GLOMAP-mode compared to 47.1 % in GLOMAP-bin. This suggests that the wet removal is acting on a larger proportion of the sea-salt particles in GLOMAP-bin than GLOMAP-mode, likely due to better treatment with the differential removal timescales of each size bin in the sectional scheme.

### 7 Comparison of global distributions of integral aerosol properties

In this section, we compare maps of the global surface-level distribution of a range of integral aerosol properties. The GLOMAP-mode run uses the revised modal settings. First, we compare surface mass concentrations of sulphate, sea-salt, BC and OC. We then quantify differences in aerosol microphysical properties, considering condensation nuclei (CN, \(D_p > 3 \text{ nm}\)) and CCN concentrations for \(D_p > 50\) and 70 nm. We also compare surface global maps of the first and second integral moments of the size distribution (in dry-radius space), which correspond to surface area concentration and condensation sink (in the continuum regime). To help understand differences in secondary (nucleated) CN and CCN, we also compare gas phase \(\text{H}_2\text{SO}_4\) in the two model runs. In each of the following figures, the global map simulated by GLOMAP-mode is shown on the left, with the bias relative to GLOMAP-bin shown on the right.
7.1 Speciated particle mass

Figure 4 compares surface-level mass concentrations of sulphate (a, b) and sea-salt (c, d). Over the vast majority of the domain, GLOMAP-mode sulphate is within 25% of GLOMAP-bin (Fig. 4b), with a weak low bias. In the Arctic and parts of the Southern Ocean, Central Africa and South America, the low bias is larger, but never exceeds a factor of two. The regions with lower sulphate in GLOMAP-mode also show a similar magnitude SO$_2$ low bias compared to GLOMAP-bin (not shown), which suggests that the different CTM version used (see Sect. 2.1) may be causing some of this bin-mode difference. In high altitude surface regions (e.g. the Himalayas, Canadian Rocky Mountains) however, GLOMAP-mode sulphate is biased low compared to GLOMAP-bin whereas SO$_2$ is not. In the tropics and sub-tropics, GLOMAP-mode simulated sulphate in the free troposphere is lower than GLOMAP-bin (not shown). The low bias is largest between 3 and 6km where rainout has a dominant influence on aerosol properties, suggesting that different size-resolved scavenging may be the cause. Rasch et al. (2000) showed that sulphate mixing ratios in the free troposphere vary by a factor 2–5 due to differences in convective transport and removal processes.

Simulated surface sea-salt mass is compared in Fig. 4d. In most marine regions the two schemes compare within 25%, although in the Southern Ocean and off the west coast of South America and South Africa, GLOMAP-bin sea-salt mass is systemically higher than in GLOMAP-mode by up to 50%. By contrast, in the Inter-Tropical Convergence Zone, where wet removal dominates, and also in some continental regions, sea-salt is higher in GLOMAP-mode.

In Sect. 6, we found that wet deposition accounts for a much larger fraction of sea-salt removal in GLOMAP-bin, which is consistent with higher sea-salt concentrations for GLOMAP-mode in regions dominated by wet deposition. The global sea-salt burden was slightly higher in GLOMAP-mode, which is consistent with the boundary layer being deeper between 30°S and 30°N (where GLOMAP-mode has higher sea-salt) than at mid-latitudes (where the modal scheme is lower).
Simulated surface BC and OC are compared in Fig. 5. Both components of the carbonaceous aerosol show very similar distributions in bin and mode, with concentrations being within 25%/−20% in most regions, although a larger low bias for GLOMAP-mode compared to GLOMAP-bin is evident in equatorial parts of Africa and South America. These regions also show similar magnitude differences in SO$_2$ and sulphate (see Table 6), suggesting that the different CTM version may be responsible for much of this discrepancy.

Comparing the simulated zonal-mean BC and OC against latitude and altitude shows that although in the lowest few km the two schemes compare well, above 3–4 km, GLOMAP-mode BC and OC both become substantially higher than GLOMAP-bin, by up to a factor 5 in some places. One can even see evidence of this at the surface in Fig. 5b and d with GLOMAP-mode systematically higher in marine regions between 30° N and 30° S, where free tropospheric air is entrained into the boundary layer due to the descending parts of the Hadley and Walker circulations (e.g., Raes et al., 2000).

Figure 6 compares GLOMAP-mode and GLOMAP-bin simulated remote BC profiles against a January 2009 multi-flight climatology (Schwarz et al., 2010) of aircraft measurements with the SP-2 instrument (Schwarz et al., 2008) from the HIPPO campaign. The model January-mean was averaged over the latitude and longitude range of the observations, which cover an altitude range of 300 m–14 km and span the latitudes 67° S–80° N. In the tropics, in common with the AEROCOM models, both schemes over-estimate free troposphere BC by a factor 20–100 compared to the observations, most likely due to poor treatment of convective scavenging. In mid- and high-latitudes however, whereas GLOMAP-bin represents the free-tropospheric BC concentrations rather well, GLOMAP-mode is biased high by a factor 2–3 in the Northern Hemisphere and by a factor 3–4 in the Southern Hemisphere. The bin-mode differences are about half the magnitude of the differences between the AEROCOM 15th and 85th percentile profiles, so here the simplification of the size distribution does seem to be a substantial source of bias. These bin-mode differences likely result mainly from size-resolved wet removal being more effective in the bin scheme.
7.2 Surface sulphuric acid vapour and CN concentrations

Figure 7a, b show the comparison of surface $H_2SO_4$ concentrations, which is a key factor determining simulated nucleation rates. In general, lower $H_2SO_4$ vapour concentrations are found in GLOMAP-mode, but the two schemes compare quite well, being mostly within $\pm 25\%$. Differences are larger in some regions however, up to a factor of two in the most polluted regions of China. Removal of $H_2SO_4$ occurs almost exclusively by condensation onto existing aerosol, and the lack of a bias suggests that, although it is not included in these runs, boundary layer nucleation rates (parameterized generally as a function of sulphuric acid concentration, e.g. Spracklen et al., 2010) may not be greatly affected by the simplifications in the modal scheme. However, we note that simpler size-resolved growth in the parameterized modal approach could lead to different condensation sink and cause subsequent biases when boundary layer nucleation is included.

Figure 7c, d compares surface CN concentrations. In continental regions, CN compare extremely well between GLOMAP-mode and GLOMAP-bin, being within 25% almost everywhere. A larger high bias (up to a factor of 2) in the modal scheme is seen in Antarctica, the Himalayas and Greenland, where the surface layer in the model is at high altitude, suggesting larger differences may be present in the free troposphere. This is investigated as part of Sect. 7.3.

7.3 CCN concentrations and vertical extent of biases

Comparing CCN in the bin and mode schemes gives an indication of how much the deficiencies in simulated size distribution seen in Sect. 3 are likely to propagate into errors in simulated aerosol-cloud interactions. In Fig. 8, we compare CCN concentrations based on dry-diameter thresholds of 50 nm (panels a and b) and 70 nm (panels c and d). These threshold sizes correspond to supersaturations of 0.35 and 0.22 %, respectively, representing values typical for marine stratocumulus, which have the largest spatial coverage and thus dominate aerosol indirect effects globally. Note that model
CCN are counted as particles in the soluble modes/distribution larger than the stated dry diameter threshold.

In all continental regions, where primary particles dominate CCN (Merikanto et al., 2009), bin and mode CCN concentrations compare well (within 25% on the annual-mean) at both supersaturations. This level of agreement extends to mid- and high-latitude marine regions, but in the 30° S–30° N marine regions, a systematic bias is apparent, with GLOMAP-mode CCN higher by up to 60%.

Figure 9 indicates the vertical extent of these differences, showing zonal-mean CN and CCN ($D_p > 50$ nm, CCN$_{50}$) against latitude and altitude. Figure 9a shows the expected vertical profile in CN concentrations with a clear nucleation layer between 12 and 17 km between 30° S and 30° N. The layer occurs in the upper troposphere and descends to lower altitudes at mid- and high-latitudes. This shape of the CCN$_{50}$ plot reflects the lifting of primary aerosol in the tropics and transport to higher altitudes in the free troposphere, with mixing of secondary, nucleated aerosol.

One potential source of bias for the parameterized modal aerosol dynamics identified in previous studies (e.g., Zhang et al., 1999) is in coagulation rates predicted when the standard deviation of each mode is held fixed. Coagulation acts to reduce the number concentration of the finest particles in the high-CN nucleation layer. Rates of coagulation depend strongly on particle diameter, and the monodisperse modal treatment in GLOMAP-mode could introduce biases by using the geometric-mean diameter as a representative size for the mode in this process. Such biases would affect growth of CN to larger sizes which could lead to CCN biases in the model. However, Fig. 9b shows that the CN bias is actually below 25% throughout most of the lowest 3 km (although larger in the mid and upper troposphere), suggesting the bias caused by this artefact may not be so important.

Figure 9d shows that the surface high bias in GLOMAP-mode marine CCN concentrations between 30° S and 30° N extends at a similar magnitude and latitudinal extent into the free troposphere, with the bias increasing slightly above 3 km. The spatial pattern of the GLOMAP-mode CCN high bias seen in Figs. 8b and 8d matches where
growth by cloud-processing is occurring most in the model. The modal treatment of cloud processing, involves particles in the large-end of the Aitken-soluble mode being transferred over to the accumulation-soluble mode, with the two modes subsequently re-constructed to give a log-normal shape. It is possible that the CCN high bias may be partly caused by this simplification, although it may just be reflecting differences in size-resolved growth via condensation or coagulation.

In summary, for CCN at the surface, the improved modal scheme is within 25% of the sectional scheme everywhere except in 30° S and 30° N marine regions. There, where photochemistry is strongest, GLOMAP-mode is high biased by up to 50%, likely resulting from differences in size-resolved growth or from the simplified treatment of cloud processing. Whereas bin-mode differences in mass are larger in the free troposphere than the boundary layer, for CN and CCN, relative differences in these two parts of the atmosphere are similar.

7.4 Surface area density and condensation sink

Figure 10 shows surface global maps of GLOMAP-mode simulated (dry) surface-area density (a) and continuum-regime condensation sink (b). These two quantities represent 2nd and 1st moment integrals across the size distribution, respectively, and are relevant as they influence rates of heterogenous chemistry and nucleation, respectively. Almost everywhere in the surface model domain, the modal scheme is within 25% of bin. The differences in surface-area density show a similar pattern as for CCN (Figs. 8b and 8d) with a maximum in the same 30° S to 30° N marine regions, but with a high bias weaker by a factor 2.

7.5 All measures at different levels

We use Taylor diagrams (Taylor et al., 2001) to summarise the bin-mode comparison for all the quantities shown in Figs. 4, 5, 7, 8, 10. Taylor diagrams combine statistical measures of the relative-variance and skewness into a single point in polar co-ordinates.
In Fig. 11, the distance of each point from the origin is the ratio of the standard deviations ($\text{sd}_{\text{rat}}$) between the two schemes, and the angle to the horizontal axis is the inverse cosine of the Pearson correlation coefficient ($\cos^{-1}(R)$). The figure illustrates $\text{sd}_{\text{rat}}$ and $\cos^{-1}(R)$ values based on the bin and mode simulated zonal-means over all latitude grid-points and on model levels between (a) 0–1 km, (b) 1–4 km and (c) 4–8 km altitude.

In the lowest km (Fig. 11a), the points for all of the variables are close to the “perfect comparison” point at $y = 0$ and $x = 1$, illustrating the general good agreement between bin and mode seen in the figures. DMS, $\text{SO}_2$ and sulphate compare the best, with standard deviation in GLOMAP-mode lower by only 8, 5 and 3%, respectively, and correlation coefficient very close to 1.0. Surface area and condensation sink are also close to the “perfect-comparison-point” with slightly higher variance in GLOMAP-mode (about 2% and 10% higher standard deviation, respectively). CN and CCN concentrations both have 20% higher standard deviation in GLOMAP-mode in the lowest km, whilst sea-salt, OC and $\text{H}_2\text{SO}_4$ have standard deviation about 20%, 15% and 22% lower. Simulated BC concentrations have the largest discrepancy with standard deviation 40% higher in GLOMAP-mode and only moderate correlation with GLOMAP-bin.

Between 1 and 4 km (Fig. 11b), there is also quite good agreement between the bin and mode schemes, although biases are higher than in the lowest km, particularly for CN and BC. In the 4–8 km altitude range (Fig. 11c), the differences get wider, but all points except CN, BC and NaCl are still between the 0.7 to 1.3 range for the ratio of standard deviations, suggesting the modal scheme is performing well in the free troposphere. The bias in CN, BC and NaCl are much larger however, with the mode-to-bin standard deviation ratio around 1.8, 2.5 and 3.5, respectively, indicative of large high biases in GLOMAP-mode in the free-troposphere compared to GLOMAP-bin (see Figs. 6 and 9d).
Comparison against benchmark observational datasets

Here we provide mean normalised bias ($b$) and Pearson correlation coefficient ($R$) for the GLOMAP-mode and GLOMAP-bin runs against each of the benchmark observational datasets compiled in Mann et al. (2010). The purpose of the paper is to quantify differences in predicted aerosol properties between sectional and modal aerosol microphysics models, and biases with observations can sometimes be misleading due to compensating errors or missing processes. Nevertheless, by comparing to the observations, we provide some context for the differences presented in the previous sections. The GLOMAP-mode simulation here is the improved version with the narrower standard deviation finer mode-edge radius for accumulation mode, with the run with the original settings also shown in Tables 3–5 to illustrate the sensitivity.

8.1 Aerosol precursor gases

Table 3 evaluates simulated DMS and SO$_2$ in the bin and mode runs against observations through the annual cycle at three remote Southern Hemisphere sites (Amsterdam Island, Cape Grim and Dumont D'Urville) and against winter and summer SO$_2$ observations across monitoring sites in Europe (EMEP, Loevblad et al., 2004) and North America (CASTNET, Holland et al., 1999). Bin-mode differences are here due to the different CTM version used, but we include them for completeness and to provide context for the biases against aerosol observations in the next sub-section.

For DMS, the models have good agreement with the observed temporal variability over the annual cycle at the three sites ($R = 0.62–0.72$), although a low bias is seen at Amsterdam Island and Dumont D'Urville. The annual variation of SO$_2$ at the remote sites is also well captured by the models ($R = 0.45–0.6$). While the magnitude of SO$_2$ compares within a factor of 2 on average at Amsterdam Island ($b = (-0.48)–(-0.39)$), there is a strong high bias ($b = 4.22–4.85$) at the Cape Grim site, likely due to the observations (Ayers et al., 1991) representing only clean air-masses and the coarse horizontal grid in the model run.
The spatial variability in simulated continental SO$_2$ compares well to the observations in both Europe ($R = 0.57–0.62$) and North America ($R = 0.79–0.89$). However, there is a strong high bias in SO$_2$ in Europe in summer ($b = 1.47–2.19$) and winter ($b = 2.33–3.04$), whereas in North America, the magnitude compares quite well in summer ($b = 0.25–0.55$) and very well in winter ($b = −0.05–0.05$). The high bias in modeled SO$_2$ over Europe is consistent with other large scale models and may be due to uncertainties in vertical mixing, emission heights (De Meij et al., 2006) or wet scavenging (Rasch et al., 2000).

### 8.2 Speciated particle masses

In Table 4 we present $b$ and $R$ values for the two schemes, evaluating simulated surface aerosol mass of sulphate, sea-salt, BC and OC. For Europe and North America, we compare winter and summer model values against year-2000 filter measurements in Europe (EMEP, Loevblad et al., 2004) and North America (IMPROVE, Malm et al., 2002). We also evaluate marine simulated sulphate and sea-salt by comparing to annual-mean observations over several years from monitoring stations in the University of Miami network (values from Stier et al., 2005).

In the previous section, we saw that simulated surface sulphate in the bin and mode schemes compares within 25% everywhere except in the Arctic and free-troposphere. Here, we examine how close they compare to the observations. The bin and mode runs represent well the observed spatial variability in Europe (each have $R = 0.63$ in winter and 0.60 in summer) and particularly well in North America (bin, mode $R = 0.76$, 0.72 in winter and 0.93, 0.94 in summer). However, in both schemes, sulphate has a winter low bias in both regions (bin, mode $b = −0.55$, −0.63 for Europe and −0.23, −0.29 in North America) and a summer high bias ($b = 0.83$, 0.68 for Europe and 0.26, 0.17 for North America). Comparing annual-mean sulphate at the University of Miami remote marine sites, both schemes are similar, representing the observed spatial variability very well ($R = 0.98$) with a moderate high bias ($b = 0.60$, 0.54 for bin, mode). The European sulphate winter low bias is likely mainly caused by the SO$_2$ low bias seen in Sect. 8.1,
although the omission of in-cloud sulphate production via ozone may also be a factor. Other sulphate production mechanisms not included here could also be important, including via heterogeneous chemistry on the surface of dust particles (e.g., Bauer and Koch, 2005) or other reactions with transition metals (Alexander et al., 2009).

Sea-salt in GLOMAP-bin is slightly longer lived than in GLOMAP-mode (see Table 2) leading to a slightly higher burden. At the surface (Fig. 4), GLOMAP-mode sea-salt was slightly lower over the Southern Ocean and higher in the tropics, where wet deposition dominates removal processes. Against the annual-mean observations, simulated sea-salt in both versions is only weakly spatially correlated against the University of Miami sites \((R = 0.13 \text{ for mode }, 0.02 \text{ for bin})\) although the magnitude compares fairly well on average \((b = -0.23, -0.30 \text{ in bin, mode})\). Note that this low bias in GLOMAP-mode sea-salt is considerably worse \((-0.44\) in the original run with the coarser accumulation-coarse mode edge-radius underlining the benefit gained from the revision to the modal settings.

Both schemes have quite good correlation with BC at the North American sites with \(R = 0.44, 0.41\) for bin and mode in winter and \(R = 0.69\) for both in summer. However, there is a moderate low bias in both winter (bin, mode \(b = -0.24, -0.31\)) and summer \((-0.36, -0.43\)) with mode slightly more low-biased in each case. Against the observations of organic carbon, both schemes have a very strong low bias during winter \((b = -0.73, -0.76 \text{ for bin, mode})\) and moderate low bias during summer \((b = -0.36, -0.45 \text{ for bin, mode})\). The correlation scores are almost identical for bin and mode and it is notable that \(R\) for both schemes is higher for OC than BC during summer but similar during winter. There is also a worse winter low bias in OC than BC for both schemes which suggests that simulation of winter organic aerosol is poorly simulated here.

8.3 Size-resolved particle number concentrations

Table 5 shows \(b\) and \(R\) values for simulated size-resolved particle number concentrations against the datasets compiled in Mann et al. (2010). The climatology of aerosol
properties from 30 years of marine particle size distribution measurements (mainly from cruises in field campaigns) compiled by Heintzenberg et al. (2000) is used to constrain simulated total, Aitken and accumulation mode number concentrations in the marine boundary layer. The vertical profile of total particle concentrations in marine regions is tested based on the aircraft observations compiled in Clarke and Kapustin (2002). Continental profiles of size-resolved number concentrations are evaluated against aircraft observations over Germany from the LACE campaign (Petzold et al., 2002) based on the 1km-resolution profiles compiled by Lauer et al. (2005). Several years of condensation particle counter measurements at Global Atmospheric Watch (GAW) stations covering free troposphere (FT), marine boundary layer (MBL) and continental boundary layer (CBL) environments (see Mann et al., 2010 for the range of years covered) is used as the observational benchmark for simulated total particle concentrations across the annual cycle. Finally, the CCN database compiled by Spracklen et al. (2011) is used to assess simulated CCN concentrations covering a range of supersaturations and environments.

We first describe the methods used to derive model values to compare against these datasets. For comparison to the Aitken and accumulation mode number concentrations (dataset 9), model values are zonal means (for gridboxes over ocean) of particle concentrations in the dry-diameter range 10–100 and 100–1000 nm, respectively. When comparing to dataset 11 (CN observations at the GAW sites, details in Mann et al., 2010), model values are concentrations of particles larger than the instrument cut-off dry-diameter (3 nm for Cape Grim and Hohenpeissenberg, 10 nm for all other sites). For the CN comparisons in datasets 9, 10 and 12, and the size-resolved concentrations in datasets 9 and 11, both soluble and insoluble modes/distributions are included. In comparing with dataset 13 (the CCN compilation from Spracklen et al., 2011), only particles in the soluble modes/distribution are included. The model CCN concentration is for particles larger than a minimum cut-off dry diameter, calculated from the supersaturation of the measurement, based on Kohler theory, and assuming sulphuric acid composition. The monthly-mean CCN corresponding to the month of the observation
is used. The simulated CN concentrations to compare at the GAW sites are from the
model level corresponding closest to the altitude of the measurement site.

The variation of marine CN concentrations in both schemes correlates only weakly
with the Heintzenberg et al. (2000) observations ($R = 0.12$ for bin, 0.13 for mode) al-
though the average bias is low ($b = -0.09$ for bin, 0.02 for mode). For particles in the
Aitken mode, both schemes fail to correlate with the observations ($R = -0.23, -0.26$ for
bin, mode), although GLOMAP-bin compares slightly better in terms of bias ($b = -0.19$
compared to $-0.27$ for mode). The negative correlation reflects substantial underes-
timation in the Southern Hemisphere and overestimation in the Northern Hemisphere
(see Fig. 12a), and the difference between bin and mode is not a significant source of
bias here. However, simulated accumulation mode number concentrations (Fig. 12b)
correlate very well with the observations in both schemes ($R = 0.73$ for bin, 0.77 for
mode) and also both have low bias ($b = -0.04$ for bin, 0.04 for mode). Note that al-
though the revision to the modal settings does not greatly improve the GLOMAP-mode
simulated number concentration in the accumulation mode, Fig. 2 clearly showed much
better agreement in simulated size against the observations.

The bin and modal schemes show good agreement to the aircraft profile CN ob-
servations over the Pacific from Clarke and Kapustin (2002), see Fig. 13. In all three
latitude ranges GLOMAP-mode is less low-biased ($b = -0.07, -0.44$ and $-0.15$) and
better correlated ($R = 0.85, 0.70$ and 0.87) than GLOMAP-bin ($b = -0.26, -0.56$ and
$-0.44; R = 0.84, 0.73$ and $0.84$) against the observations. We also note the improved
bias at GLOMAP-mode v6 compared to GLOMAP-mode v5 in Mann et al. (2010) which
had $b = -0.28$, $-0.57$ and $-0.15$.

Figure 14 shows a comparison to aircraft profiles over Germany of particle concen-
trations larger than 5, 15 and 120 nm dry-diameter ($N_{5}$, $N_{15}$ and $N_{120}$). Here the bin
and mode schemes differ more, with $N_{5}$ in GLOMAP-mode moderately high-biased
($b = 0.67$) albeit with good correlation ($R = 0.90$), whereas GLOMAP-bin $N_{5}$ has only
small high bias ($b = 0.19$) and is better correlated ($R = 0.95$). However, GLOMAP-
mode $N_{15}$ compares extremely well ($b = 0.0$, $R = 0.93$) suggesting it is only in the very
649 smallest particles (dry-diameter < 15 nm) where the bias occurs. Comparing to the observed N_{120} profiles, both schemes have moderate low bias (b = −0.28, −0.31 for bin, mode) and both correlate extremely well with the observations (R = 0.99).

At the free troposphere GAW sites, the bin and mode schemes are in very close agreement, and compare rather well against the observations. At Mauna Loa and Jungfraujoch, the bin and mode schemes have only a weak to moderate low bias in all months, (b = −0.12, −0.13 and −0.37, −0.38, respectively). At South Pole the seasonal cycle is also well captured (R = 0.75, 0.65), although the summer maximum is a factor 2–4 too high and the winter minimum underestimated (not shown). At marine GAW sites, bin and mode again have similar skill against the observations, both generally underpredicting either moderately (b = −0.38 at Samoa) or more substantially at the coastal mid-latitude (b = −0.64, −0.62 at Cape Grim and −0.53, −0.52 at Mace Head) and high-latitude sites (b = −0.67, −0.72 at Neumayer and −0.81, −0.78 at Barrow). At the continental GAW sites, the two schemes have a similar level of agreement with the observations, with weak to moderate low biases and poor correlation over the seasonal cycle. This poor correlation is consistent with Spracklen et al. (2010) who found that simulations with binary nucleation and primary emissions (the processes included here) could not explain the continental seasonal CN cycle, whereas considerably improved correlation was found when boundary layer nucleation was included.

Against the Spracklen et al. (2011) compilation of CCN measurements, both schemes have a strong high bias (b = 2.10 for mode, 1.82 for bin), although the spatial correlation is good (R = 0.74 for bin, 0.68 for mode), see Fig. 15. The CCN high bias in the runs here does not imply that the models always perform poorly against these observations. Indeed, we note specifically that the bias against those same observations was much lower (b = 0.49) in Mann et al. (2010) for GLOMAP-mode v5 and also in Spracklen et al. (2011) for GLOMAP-bin which had normalised mean bias of 0.37 for the “small CCA” run which matches the emissions sizes used here. Clearly, the aerosol parameter settings used in the runs here are not optimum for best agreement with these observations, but the bin-mode difference in bias and correlation is
much lower than differences to the observations, even considering the skill scores from Spracklen et al. (2011).

In Fig. 16, we compare the bin and mode CCN annual cycle to observations at Mace Head (Reade et al., 2006) and Cape Grim (Ayers and Gras, 1991). The high CCN bias is clearly evident at Cape Grim, and is slightly worse in GLOMAP-mode ($b = 0.97$, $1.90$ for $S=1.2$, $0.23\%$) than GLOMAP-bin ($b = 0.90–1.40$) and similarly at Mace Head ($b = 0.90$ for bin, $1.10$ for mode). However, GLOMAP-mode reproduces the seasonal variability better at Mace Head ($R = 0.42$ c.f. $0.30$) whereas GLOMAP-bin is better at Cape Grim ($R = 0.67–0.70$ c.f. $0.29–0.58$). Note that one might expect the model CCN to be higher than observed at Cape Grim since it covers all wind directions, whereas the measurements are for marine air masses only.

9 Conclusions

In this study we have, for the first time in a 3-D global model, carried out a thorough intercomparison of integral particle properties simulated by two-moment sectional and modal aerosol dynamics schemes. The assessment was carried out using the same offline chemistry transport model with equivalent meteorological, oxidant and cloud fields to help isolate simulated differences to the sophistication of the aerosol scheme.

We have compared size distributions simulated by the two schemes to constrain the choice of parameters in the modal scheme, and have quantified biases in size-resolved particle concentrations. We find that a value of $1.59$ for the accumulation mode standard deviation ($\sigma_{acc}$) is too wide, and reducing this and the separation-dry-radius between the accumulation and coarse modes ($r_{3,4}$), we achieve much better agreement against the bin scheme. With $\sigma_{acc}$ set to $1.59$, the particles in the large-end tail of the accumulation mode were over-estimated, whereas a value of $1.40$ fits much better the size distribution in the sectional scheme and, in general, against observations. With $r_{3,4}$ at $500$ nm, coarse particle concentrations were biased low with the bin scheme, but reducing $r_{3,4}$ to $250$ nm led to excellent agreement, avoiding problems in the previ-
ous configuration whereby the lower-end of the coarse mode size-resolved sea-spray emissions flux was emitting in the model accumulation-soluble mode.

We have shown that these revisions of the size-settings in the modal scheme lead to improved agreement against the sectional scheme in terms of simulated sea-salt, CN, CCN\textsubscript{50} and CCN\textsubscript{150}. Whereas the original configuration of GLOMAP-mode showed low-biased CCN\textsubscript{150} up to a factor two compared to GLOMAP-bin in some regions, the revised configuration approximately halved this bias in almost all regions (see Table 1).

With the improved configuration, globally and vertically integrated burdens (and hence lifetimes) of sulphate, sea-salt, BC and POM in the two schemes compare well, with GLOMAP-mode within about 10% of GLOMAP-bin.

At the surface, speciated sub-\(\mu\)m aerosol mass (sulphate, BC and OC) in the bin and mode schemes compared very well in general (within 25%) in both marine and continental regions. GLOMAP-mode sulphate mass has a moderate low bias compared to GLOMAP-bin in the Arctic however, and in regions with strong biomass burning emissions moderate biases were also seen. Generally, biases were slightly larger between bin and mode for mass in the coarse mode (sea-salt), although good general agreement was still found.

In the free troposphere, bin-mode differences in simulated mass are larger than at the surface, with the modal scheme higher there, which we attribute to the coarser size-resolution in the treatment of wet removal. In the sectional scheme, highly size-resolved nucleation-scavenging rates are possible, whereas the modal scheme cannot differentiate between particles within one size class.
With the improved settings, the modal scheme performs well against the bin scheme with differences in simulated CN, surface area density, and condensation sink less than 25% in almost all regions at the surface. Differences in simulated surface CCN are everywhere less than 25%, except in 30° S to 30° N marine regions, where the modal scheme is high-biased by ≈ 50%, likely caused by too effective growth and the simplified cloud-processing approach.

Although there are substantial bin-mode differences in mass in the free troposphere and remote locations, differences are generally less for aerosol microphysical integral-properties such as CN, CCN, surface area and condensation sink. This suggests that the simplification from bins to modes affects processes which influence the accumulation and coarse parts of the size distribution (where most of the mass resides) more than those at sub-100 nm sizes (which contain most of the number). We infer from this, that although growth by coagulation or condensation is better treated by the sectional scheme, the biases arising from the simplification to the modal scheme are larger on processes such as cloud processing, sedimentation and scavenging, which more strongly affect the accumulation and coarse size range.

In previous studies, we have assembled benchmark observational datasets against which to evaluate global aerosol microphysics models and to better understand the processes controlling the evolution of the particle size distribution in the atmosphere. Here, we have used these datasets to give context for the differences between the parametrized modal and more sophisticated sectional versions of the 3-D offline global aerosol microphysics model.

In this paper, we have refined the chosen values for the parameters inherent in a modal aerosol microphysics scheme to better compare with a sophisticated sectional scheme. By benchmarking the modal aerosol dynamics scheme against the bin scheme, we therefore reduce biases in simulated size-resolved number concentrations and CCN. In so-doing, we aim to better constrain modal-parameter settings and help improve predictions of aerosol properties and radiative forcings with two-moment modal schemes.
The bin and mode schemes perform similarly against observed CN and size-resolved particle concentration datasets in free troposphere, marine and continental regions. The simplification from bins to modes is found to be only a minor factor in determining the skill of the model against observations. We conclude therefore that the limitations in size-resolved growth and removal in the modal approach cause only small biases in the model simulations, with the updated modal scheme general able to reproduce the global distribution of size-resolved particle concentrations only slightly worse than the sectional scheme.

In summary, the comparisons have shown that, in most parts of the atmosphere, bin-mode differences are less than model-observation differences, although some processes missing in these runs (e.g. boundary layer nucleation, ultra-fine sea-spray) may well decrease the biases against observations stated here. However, the biases seen in the size distributions underline the need for a spectrum of complexity in global models, with size-resolved aerosol properties predicted by modal schemes needing to be continually benchmarked and improved against freely evolving sectional schemes and observations.

Appendix A

Modifications to GLOMAP-mode compared to Mann et al. (2010)

A1 Vapour condensation

The vapour-condensation routine has been updated to use an improved representative size for the mode and to have revised calculation for the vapour diffusion coefficient ($D_s$). The “condensation sink radius” $r_{i,\text{cond}}$ (see Lehtinen et al., 2003) is used when calculating the condensation coefficient whereas the geometric mean radius $r_{i,\text{g}}$ was used in Mann et al. (2010). Lehtinen et al. (2003) explain that using $r_{i,\text{g}}$ in monodisperse-modal models (like GLOMAP-mode) will introduce biases which can be-
come substantial in regions with high vapour condensation. They recommend instead using $r_{\text{cond}}$, defined as the size giving the same condensation sink as the polydisperse distribution with corresponding $r_{g}$. To evaluate $r_{\text{cond}}$, one integrates the condensation sink expression across the log-normally distributed particles, being essentially the 1st-moment-mean-radius in the continuum regime (small Knudsen number, large radius) and the 2nd-moment-mean-radius in the molecular regime (large Knudsen number, small radius). For a log-normal mode with geometric standard deviation $\sigma$, the condensation sink-radius is given by:

$$r_{i,\text{cond}} = r_i \left( A^2 \log^2 \sigma_g \right)$$  \hspace{1cm} (A1)

where $A$ is the “growth exponent” defined in Lehtinen et al. (2003). The condensation sink radius is evaluated with $A$ set to be 2.0, 1.9, 1.5 and 1.1 for nucleation, Aitken, accumulation and coarse modes, respectively. The new expression for $D_s$ matches that implemented in GLOMAP-bin at v1a (e.g., Merikanto et al., 2009) following the approach of Fuller et al. (1966), using atomic diffusion-volumes and a dependence on pressure and temperature (e.g., Poling et al., 2001). A further difference is that a minor bug was found which caused simulated nucleation rates to be slightly too high since gas phase H$_2$SO$_4$ was not being depleted for the (small) sink due to new particle formation.

### A2 Nucleation scavenging

In the GLOMAP-mode approach in Mann et al. (2010), nucleation-scavenging is applied only to soluble-accumulation and soluble-coarse modes. In the revised code here, we follow GLOMAP-bin in applying nucleation-scavenging to soluble particles larger than a size threshold $r_{\text{scav}}$, the “scavenging-radius”. Also, in these runs, insoluble modes are scavenged in precipitating gridboxes where the temperature is below the ice frost point (taken to be 258 K) to simulate removal as ice nuclei. This approach matches GLOMAP-bin, and leads to slightly reduced black carbon (BC) in the free tro-
posphere compared to the simulations in Mann et al. (2010), where insoluble modes were not nucleation-scavenged.

A3 Aqueous sulphate production

In Mann et al. (2010), the rate of aqueous phase sulphate production was calculated via an effective Henry’s law approach. Here, we use a diffusion-limited approach, again to match GLOMAP-bin, whereby the sulphate production is limited by the rate of diffusion of SO$_2$ to the cloud droplets. The gas-to-cloud-droplet transfer is calculated for each aerosol size class assuming cloud droplet radius is proportional to that of the aerosol particles (e.g., Spracklen et al., 2005).

A4 Other differences in process settings

Several process settings in the benchmark GLOMAP-bin run (B1) were different to those in the GLOMAP-mode version 5 run in Mann et al. (2010) (v5$_{M10}$). To make the bin and mode simulations consistent, we have changed several parameter settings in the GLOMAP-mode v6 runs here (v6$_{M11}$) compared to v5$_{M10}$, and these are listed here for completeness.

- **Sea-salt emission dry-radius range**: v5$_{M10}$ emitted in the range 17.5 nm up to 14.4 µm, whereas B1 and v6$_{M11}$ emit in the range 17.5 nm–7.0 µm. This explains why the sea-spray emission mass flux in v6$_{M11}$ is a factor of three lower than at v5$_{M10}$.

- **Condensation-ageing rate**: v5$_{M10}$ used a 10-monolayer ageing rate where B1 and v6$_{M11}$ use 1-monolayer, leading to a shorter BC lifetime in v6$_{M11}$ than v5$_{M10}$.

- **Activation dry-diameter**: At v5$_{M10}$ this minimum dry-diameter for cloud-processing was set to 75 nm whereas B1 and v6$_{M11}$ use 50 nm.

- **Sticking efficiency for insoluble particles** ($S_{\text{ins}}$): At v5$_{M10}$ $S_{\text{ins}} = 0.3$, whereas B1 and v6$_{M11}$ have $S_{\text{ins}} = 1.0$, as for soluble particles.
– Size distribution for primary carbonaceous emissions: v5M10 used dry-diameters of 60 nm for fossil-fuels and 150 nm for bio-fuel/biomass burning sources and \(\sigma = 1.59\) (as in Stier et al., 2005) whereas v6M11 and B1 use 80 nm and 30 nm with \(\sigma = 1.8\) as recommended by AEROCOM (Dentener et al., 2006). Several papers (e.g., Spracklen et al., 2010) have shown that simulated particle concentrations are sensitive to the assumed size for primary emissions. Reddington et al. (2011) explore the sensitivity specifically around the Stier et al. (2005) and Dentener et al. (2006) values and find reduced bias against observations when the finer sizes are used.

Note also that where dust emissions and in-cloud sulphate production via \(\text{O}_3\) were included in v5M10, they are de-activated in v6M11 since they were not included in B1.

One difference between B1, and GLOMAP-bin runs in other papers, is that the nucleation-scavenging wet radius \(r_{\text{scav}} = 150\) nm, whereas it is usually set to 103 nm. The rationale for changing \(r_{\text{scav}}\) is from findings in Korhonen et al. (2008) where a low bias in GLOMAP-bin simulated aerosol optical depth against observations was remedied when \(r_{\text{scav}}\) was increased from 103 to 206 nm. We carried out three B1 simulations with \(r_{\text{scav}}\) at 103 nm (as usual), at 150 (as here) and at 200 nm (as in Korhonen et al., 2008). Note that in v6M11, \(r_{\text{scav}} = 103\) nm. The different \(r_{\text{scav}}\) values for bin and mode is considered appropriate since the size bins in GLOMAP-bin allow the process to cut off size distributions sharply at that size. By contrast, in GLOMAP-mode, the modal size classes are much wider, and follow a prescribed \(\sigma\), so size distributions at the large-end of the accumulation mode will tail-off gradually, according to the prescribed \(\sigma\) for the mode.

Appendix B  Summary of the bin-mode comparison in different regions

Here, to clarify the extent of the bin-mode differences (which are only shown in broad terms in the figures), we tabulate regional-mean values of the range of aerosol properties covered in the paper. Table 6 shows the values for the key precursor gases...
and each of the aerosol component masses, while Table 7 presents the numbers for CN, CCN and moments of the aerosol size distribution. The first number shown is that simulated by GLOMAP-mode, with the bias compared to GLOMAP-bin shown in parentheses.

Acknowledgements. This work is supported by the Natural Environment Research Council (NERC) through the National Centre for Atmospheric Science. We acknowledge funding from the EU Framework Program 6 (FP6) project European Aerosol Cloud Climate and Air Quality Interactions (EUCAARI) and the FP7 project Monitoring of Atmospheric Composition and Climate (MACC). We also acknowledge funding from NERC project AEROFORM (NE/D01395X/1), the NERC Directed Research Programme APPRAISE (Theme 3) and the NERC QUEST programme (Earth System Modelling). The development of UKCA is supported by both NERC and the DECC and Defra Integrated Climate Programme – DECC/Defra (GA01101). We also gratefully acknowledge AEROCOM and GEIA for providing emissions datasets, ISCCP for the monthly low cloud fields and ECMWF for the ERA-40 re-analysis fields. We thank the scientists and data providers associated with the EMEP, CASTNET, IMPROVE and University of Miami networks. We also thank the data PIs for the CN and CCN surface measurements at the WMO-GAW sites (J. Ogren, NOAA, ESRL; J. Gras, CSIRO; U. Baltensberger, PSI; U. Kaminski, DWD; G. Jennings, NUI Galway; R. Weller, AWI; Y. Viisanen, FMI) for making their data available via the World Data Centre for Aerosols website. We are also grateful to F. O’Connor (UK Met Office Hadley Centre) for providing the TOMCAT oxidant fields.
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Pirjola, L., Kulmala, M., Wilck, M., Bischoff, A., Stratmann, F. and Otto, E.: Formation of sul-


Trivitayanurak, W., Adams, P. J., Spracklen, D. V., and Carslaw, K. S.: Tropospheric aerosol microphysics simulation with assimilated meteorology: model description and intermodel


Table 1. Regional-mean annual-mean CN (all particles with $D_p > 3$ nm), CCN$_{50}$ and CCN$_{150}$ (soluble particles with $D_p > 50$ and 150 nm, respectively). GLOMAP-bin values are shown first with the ratio of mode/bin shown in parentheses (original modal settings followed by revised). Regions match those defined in Merikanto et al. (2009).

<table>
<thead>
<tr>
<th>Region</th>
<th>CN</th>
<th>CCN$_{50}$</th>
<th>CCN$_{150}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Global</td>
<td>719.0</td>
<td>452.1</td>
<td>159.3</td>
</tr>
<tr>
<td>Global Continental</td>
<td>1561.0</td>
<td>1014.9</td>
<td>308.2</td>
</tr>
<tr>
<td>Global Marine</td>
<td>392.0</td>
<td>233.4</td>
<td>101.5</td>
</tr>
<tr>
<td>Europe</td>
<td>3227.0</td>
<td>1657.4</td>
<td>457.7</td>
</tr>
<tr>
<td>Africa</td>
<td>1290.0</td>
<td>1050.7</td>
<td>387.3</td>
</tr>
<tr>
<td>N. America</td>
<td>1689.0</td>
<td>1074.0</td>
<td>363.2</td>
</tr>
<tr>
<td>S. America</td>
<td>1458.0</td>
<td>1224.8</td>
<td>309.9</td>
</tr>
<tr>
<td>N. Asia</td>
<td>820.0</td>
<td>497.4</td>
<td>163.0</td>
</tr>
<tr>
<td>S.E. Asia</td>
<td>4067.0</td>
<td>2364.9</td>
<td>631.1</td>
</tr>
<tr>
<td>Oceana</td>
<td>1078.0</td>
<td>885.3</td>
<td>272.1</td>
</tr>
<tr>
<td>W. of N. America</td>
<td>407.0</td>
<td>241.7</td>
<td>83.2</td>
</tr>
<tr>
<td>W. of S. America</td>
<td>244.0</td>
<td>127.0</td>
<td>60.3</td>
</tr>
<tr>
<td>W. of N. Africa</td>
<td>399.0</td>
<td>240.0</td>
<td>116.5</td>
</tr>
<tr>
<td>W. of S. Africa</td>
<td>405.0</td>
<td>314.1</td>
<td>163.1</td>
</tr>
<tr>
<td>E. of N.E. Asia</td>
<td>1282.0</td>
<td>798.5</td>
<td>269.7</td>
</tr>
</tbody>
</table>
Table 2. Annual mean global mass burden (Tg), emission fluxes, secondary production fluxes (both Tg yr$^{-1}$) and lifetime (days) for each simulated aerosol component. Also shown is the % removal by wet deposition for each component. The values for the mode and bin versions of GLOMAP are shown before and after the comma, respectively. Values in parentheses are the median values simulated by AEROCOM models as documented in Textor et al. (2006).

<table>
<thead>
<tr>
<th>Species</th>
<th>Burden</th>
<th>Primary emission</th>
<th>Production</th>
<th>Lifetime</th>
<th>% loss by wdep</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulphate</td>
<td>0.51, 0.58 (0.66)</td>
<td>1.74, 1.72 (59.6)</td>
<td>48.5, 46.3</td>
<td>3.71, 4.44 (4.1)</td>
<td>87.5, 86.0 (88.5)</td>
</tr>
<tr>
<td>Sea-salt</td>
<td>3.39, 3.11 (6.39)</td>
<td>2806, 2806 (6280)</td>
<td>0.0, 0.0</td>
<td>0.44, 0.40 (0.4)</td>
<td>27.1, 47.1 (30.3)</td>
</tr>
<tr>
<td>BC</td>
<td>0.100, 0.096 (0.21)</td>
<td>7.72, 7.78 (11.3)</td>
<td>0.0, 0.0</td>
<td>4.76, 4.51 (6.5)</td>
<td>79.5, 81.5 (79.5)</td>
</tr>
<tr>
<td>POM</td>
<td>0.87, 0.90 (1.21)</td>
<td>47.0, 47.3 (69.9)</td>
<td>26.0, 25.9</td>
<td>4.59, 4.70 (6.1)</td>
<td>84.1, 85.4 (78.9)</td>
</tr>
</tbody>
</table>
Table 3. Simulated gas phase DMS and SO$_2$ against surface observations for bin and mode, respectively (comma-separated). Mean normalised bias ($b$) and Pearson correlation coefficient ($R$) are shown for each dataset. References for the observations are 1: Nguyen et al. (1992), 2: Ayers et al. (1991), 3: Jourdain and Legrand (2001) 4: Loevblad et al. (2004) 5: Holland et al. (1999).

<table>
<thead>
<tr>
<th>Species</th>
<th>Site name</th>
<th>$b$</th>
<th>$R$</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>DMS (ann. cycle)</td>
<td>Amsterdam I.</td>
<td>$-0.43, -0.45$</td>
<td>$0.71, 0.72$</td>
<td>1</td>
</tr>
<tr>
<td>DMS (ann. cycle)</td>
<td>Cape Grim</td>
<td>$0.31, 0.28$</td>
<td>$0.65, 0.65$</td>
<td>2</td>
</tr>
<tr>
<td>DMS (ann. cycle)</td>
<td>D. D’Urville</td>
<td>$-0.52, -0.56$</td>
<td>$0.64, 0.62$</td>
<td>3</td>
</tr>
<tr>
<td>SO$_2$ (ann. cycle)</td>
<td>Amsterdam I.</td>
<td>$-0.43, -0.48$</td>
<td>$0.62, 0.64$</td>
<td>1</td>
</tr>
<tr>
<td>SO$_2$ (ann. cycle)</td>
<td>Cape Grim</td>
<td>$4.85, 4.27$</td>
<td>$0.45, 0.47$</td>
<td>2</td>
</tr>
<tr>
<td>SO$_2$ (Dec)</td>
<td>EMEP</td>
<td>$3.04, 2.72$</td>
<td>$0.61, 0.61$</td>
<td>4</td>
</tr>
<tr>
<td>SO$_2$ (Jun)</td>
<td>EMEP</td>
<td>$2.19, 1.53$</td>
<td>$0.59, 0.57$</td>
<td>4</td>
</tr>
<tr>
<td>SO$_2$ (Dec)</td>
<td>CASTNET</td>
<td>$-0.05, -0.03$</td>
<td>$0.89, 0.88$</td>
<td>5</td>
</tr>
<tr>
<td>SO$_2$ (Jun)</td>
<td>CASTNET</td>
<td>$0.55, 0.25$</td>
<td>$0.80, 0.79$</td>
<td>5</td>
</tr>
</tbody>
</table>
Table 4. Simulated aerosol mass of sulphate, sea-salt, BC and OC against surface observations for bin and mode, respectively (comma-separated). For each of the benchmark observational datasets, Mean normalised bias ($b$) and Pearson correlation coefficient ($R$) are shown. The GLOMAP-mode run with the original accumulation mode standard-deviation and mode-edge radius is shown in parentheses. References for the observations are: 6: Loevblad et al. (2004) 7: Malm et al. (2002) 8: from Stier et al. (2005).

<table>
<thead>
<tr>
<th>Component</th>
<th>Sites</th>
<th>$b$</th>
<th>$R$</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>SO$_4$ (Dec)</td>
<td>EMEP</td>
<td>$-0.55, -0.63 (-0.65)$</td>
<td>$0.63, 0.63 (0.62)$</td>
<td>6</td>
</tr>
<tr>
<td>SO$_4$ (Jun)</td>
<td>EMEP</td>
<td>$0.83, 0.68 (0.69)$</td>
<td>$0.60, 0.60 (0.60)$</td>
<td>6</td>
</tr>
<tr>
<td>SO$_4$ (Dec)</td>
<td>IMPROVE</td>
<td>$-0.23, -0.29 (-0.29)$</td>
<td>$0.76, 0.72 (0.72)$</td>
<td>7</td>
</tr>
<tr>
<td>SO$_4$ (Jun)</td>
<td>IMPROVE</td>
<td>$0.26, 0.17 (0.18)$</td>
<td>$0.93, 0.94 (0.94)$</td>
<td>7</td>
</tr>
<tr>
<td>SO$_4$ (annual)</td>
<td>Univ. Miami</td>
<td>$0.60, 0.52 (0.54)$</td>
<td>$0.98, 0.98 (0.98)$</td>
<td>8</td>
</tr>
<tr>
<td>NaCl (annual)</td>
<td>Univ. Miami</td>
<td>$-0.23, -0.30 (-0.42)$</td>
<td>$0.02, 0.13 (0.13)$</td>
<td>8</td>
</tr>
<tr>
<td>BC (Dec)</td>
<td>IMPROVE</td>
<td>$-0.24, -0.31 (-0.31)$</td>
<td>$0.44, 0.41 (0.41)$</td>
<td>7</td>
</tr>
<tr>
<td>BC (Jun)</td>
<td>IMPROVE</td>
<td>$-0.36, -0.43 (-0.43)$</td>
<td>$0.69, 0.69 (0.69)$</td>
<td>7</td>
</tr>
<tr>
<td>OC (Dec)</td>
<td>IMPROVE</td>
<td>$-0.73, -0.76 (-0.76)$</td>
<td>$0.46, 0.43 (0.43)$</td>
<td>7</td>
</tr>
<tr>
<td>OC (Jun)</td>
<td>IMPROVE</td>
<td>$-0.36, -0.45 (-0.45)$</td>
<td>$0.84, 0.84 (0.84)$</td>
<td>7</td>
</tr>
</tbody>
</table>
Table 5. Simulated CN, CCN and size-resolved number concentrations for bin and mode, respectively (comma-separated) against each of the benchmark observational datasets. Mean normalised bias ($b$) and Pearson correlation coefficient ($R$) are shown. The GLOMAP-mode run with the original accumulation mode standard-deviation and mode-edge radius is shown in parentheses. References for the observations are 9: Heintzenberg et al. (2000), 10: Clarke and Kapustin et al. (2002), 11: from Lauer et al. (2005), 12: World Data Centre for Aerosols webpage (http://wdca.jrc.ec.europa.eu/data/parameters/datacnc.html), 13: from Spracklen et al., 2011.

<table>
<thead>
<tr>
<th>Property</th>
<th>Location</th>
<th>$b$</th>
<th>$R$</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surf CN (ann. mean)</td>
<td>Global marine</td>
<td>$-0.09, 0.02$ (0.04)</td>
<td>$0.12, 0.13$ (0.14)</td>
<td>9</td>
</tr>
<tr>
<td>Surf $N_{\text{A}}$ (ann. mean)</td>
<td>Global marine</td>
<td>$-0.19, -0.27$ (−0.15)</td>
<td>$-0.23, -0.26$ (−0.28)</td>
<td>9</td>
</tr>
<tr>
<td>Surf $N_{\text{acc}}$ (ann. mean)</td>
<td>Global marine</td>
<td>$-0.04, 0.04$ (−0.07)</td>
<td>$0.73, 0.77$ (0.77)</td>
<td>9</td>
</tr>
<tr>
<td>Prof CN (ann. mean)</td>
<td>N.H. marine</td>
<td>$-0.26, -0.07$ (−0.05)</td>
<td>$0.84, 0.85$ (0.85)</td>
<td>10</td>
</tr>
<tr>
<td>Prof CN (ann. mean)</td>
<td>Trop’l marine</td>
<td>$-0.56, -0.44$ (−0.44)</td>
<td>$0.73, 0.70$ (0.70)</td>
<td>10</td>
</tr>
<tr>
<td>Prof CN (ann. mean)</td>
<td>S.H. marine</td>
<td>$-0.44, -0.15$ (−0.14)</td>
<td>$0.84, 0.87$ (0.87)</td>
<td>10</td>
</tr>
<tr>
<td>Prof $N_5$ (ann. mean)</td>
<td>Germany</td>
<td>$0.19, 0.67$ (0.69)</td>
<td>$0.95, 0.90$ (0.91)</td>
<td>11</td>
</tr>
<tr>
<td>Prof $N_{15}$ (ann. mean)</td>
<td>Germany</td>
<td>$0.08, 0.00$ (0.03)</td>
<td>$0.95, 0.93$ (0.93)</td>
<td>11</td>
</tr>
<tr>
<td>Prof $N_{120}$ (ann. mean)</td>
<td>Germany</td>
<td>$-0.28, -0.31$ (−0.38)</td>
<td>$0.99, 0.99$ (0.99)</td>
<td>11</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Jungfrau’ (FT)</td>
<td>$-0.37, -0.38$ (−0.35)</td>
<td>$0.14, 0.23$ (0.30)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Mauna Loa (FT)</td>
<td>$-0.12, -0.13$ (−0.12)</td>
<td>$-0.32, -0.44$ (−0.45)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>South Pole (FT)</td>
<td>$0.21, 0.09$ (0.09)</td>
<td>$0.75, 0.65$ (0.65)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Mace H’d (MBL)</td>
<td>$-0.53, -0.52$ (−0.51)</td>
<td>$0.18, 0.17$ (0.19)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Neum’r (MBL)</td>
<td>$-0.67, -0.72$ (−0.71)</td>
<td>$0.81, 0.78$ (0.78)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Barrow (MBL)</td>
<td>$-0.78, -0.80$ (−0.79)</td>
<td>$-0.35, -0.19$ (−0.14)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Samoa (MBL)</td>
<td>$-0.39, -0.38$ (−0.39)</td>
<td>$-0.47, -0.47$ (−0.45)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Trin’d H’d (MBL)</td>
<td>$-0.02, 0.00$ (0.02)</td>
<td>$0.30, 0.40$ (0.39)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Cape Grim (MBL)</td>
<td>$-0.64, -0.62$ (−0.61)</td>
<td>$0.34, 0.43$ (0.43)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>SG Plains (CBL)</td>
<td>$-0.48, -0.43$ (−0.43)</td>
<td>$0.38, 0.47$ (0.46)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Bondville (CBL)</td>
<td>$-0.43, -0.35$ (−0.34)</td>
<td>$0.04, -0.02$ (−0.02)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Pallas (CBL)</td>
<td>$-0.07, -0.04$ (−0.03)</td>
<td>$-0.51, -0.60$ (−0.54)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CN (ann. cycle)</td>
<td>Hoh’berg (CBL)</td>
<td>$-0.14, -0.05$ (−0.03)</td>
<td>$0.47, -0.02$ (−0.02)</td>
<td>12</td>
</tr>
<tr>
<td>Surf CCN (monthly)</td>
<td>Global</td>
<td>$1.82, 2.10$ (2.09)</td>
<td>$0.74, 0.68$ (0.69)</td>
<td>13</td>
</tr>
<tr>
<td>Surf CCN0.5 (ann. cycle)</td>
<td>Mace Head</td>
<td>$0.90, 1.10$ (1.10)</td>
<td>$0.30, 0.42$ (0.42)</td>
<td>13</td>
</tr>
<tr>
<td>Surf CCN0.23 (ann. cycle)</td>
<td>Cape Grim</td>
<td>$1.40, 1.90$ (1.90)</td>
<td>$0.67, 0.29$ (0.29)</td>
<td>13</td>
</tr>
<tr>
<td>Surf CCN1.2 (ann. cycle)</td>
<td>Cape Grim</td>
<td>$0.80, 0.97$ (0.97)</td>
<td>$0.70, 0.58$ (0.58)</td>
<td>13</td>
</tr>
</tbody>
</table>
Table A1. Summary of annual-mean concentrations of SO$_2$, H$_2$SO$_4$, and mass of sulphate, EC, POM and sea-salt simulated by GLMAP-mode (improved configuration) with the ratio to that simulated by GLMAP-bin shown in parentheses. Units for the GLMAP-mode simulated values are µg S m$^{-3}$ for H$_2$SO$_4$, µg S m$^{-3}$ for SO$_2$ and SO$_4$, µg C m$^{-3}$ for EC and POM and µg m$^{-3}$ for NaCl. Regions match those defined in Merikanto et al. (2009).

<table>
<thead>
<tr>
<th>Region</th>
<th>SO$_2$</th>
<th>H$_2$SO$_4$</th>
<th>SO$_4$</th>
<th>EC</th>
<th>POM</th>
<th>NaCl</th>
</tr>
</thead>
<tbody>
<tr>
<td>Global</td>
<td>0.17 (0.90)</td>
<td>0.025 (0.82)</td>
<td>0.30 (0.93)</td>
<td>0.07 (0.88)</td>
<td>0.42 (0.85)</td>
<td>6.48 (0.92)</td>
</tr>
<tr>
<td>GloCnt</td>
<td>0.40 (0.89)</td>
<td>0.032 (0.79)</td>
<td>0.47 (0.91)</td>
<td>0.18 (0.86)</td>
<td>1.07 (0.82)</td>
<td>0.45 (1.17)</td>
</tr>
<tr>
<td>GloMrn</td>
<td>0.08 (0.92)</td>
<td>0.027 (0.84)</td>
<td>0.23 (0.94)</td>
<td>0.03 (0.94)</td>
<td>0.17 (0.93)</td>
<td>8.82 (0.91)</td>
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<td>Europe</td>
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<td>0.053 (0.78)</td>
<td>0.93 (0.93)</td>
<td>0.27 (0.87)</td>
<td>0.65 (0.84)</td>
<td>0.91 (1.06)</td>
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<td>Africa</td>
<td>0.18 (0.92)</td>
<td>0.024 (0.79)</td>
<td>0.61 (0.92)</td>
<td>0.23 (0.85)</td>
<td>1.77 (0.83)</td>
<td>0.52 (1.09)</td>
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<tr>
<td>N. Amer</td>
<td>0.69 (0.88)</td>
<td>0.069 (0.74)</td>
<td>0.61 (0.96)</td>
<td>0.14 (0.87)</td>
<td>0.84 (0.84)</td>
<td>0.38 (1.30)</td>
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<td>S. Amer</td>
<td>0.07 (0.82)</td>
<td>0.021 (0.77)</td>
<td>0.19 (0.79)</td>
<td>0.17 (0.79)</td>
<td>1.95 (0.77)</td>
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<td>0.27 (0.83)</td>
<td>0.06 (0.87)</td>
<td>0.35 (0.84)</td>
<td>0.08 (1.18)</td>
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<td>SE. Asa</td>
<td>1.17 (0.88)</td>
<td>0.039 (0.69)</td>
<td>1.07 (0.95)</td>
<td>0.51 (0.89)</td>
<td>1.43 (0.87)</td>
<td>0.32 (1.40)</td>
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<td>Oceana</td>
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<td>0.036 (0.78)</td>
<td>0.26 (0.91)</td>
<td>0.13 (0.82)</td>
<td>1.11 (0.82)</td>
<td>1.23 (0.90)</td>
</tr>
<tr>
<td>WofNAm</td>
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<td>0.010 (0.78)</td>
<td>0.23 (0.93)</td>
<td>0.02 (1.15)</td>
<td>0.10 (1.09)</td>
<td>8.04 (0.85)</td>
</tr>
<tr>
<td>WofSAm</td>
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<td>0.013 (0.74)</td>
<td>0.19 (0.96)</td>
<td>0.01 (1.01)</td>
<td>0.06 (0.93)</td>
<td>8.41 (0.79)</td>
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<tr>
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<td>0.056 (0.88)</td>
<td>0.36 (0.94)</td>
<td>0.02 (1.00)</td>
<td>0.09 (0.99)</td>
<td>9.14 (0.89)</td>
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<tr>
<td>WofSAf</td>
<td>0.03 (0.96)</td>
<td>0.014 (0.84)</td>
<td>0.25 (0.89)</td>
<td>0.08 (0.88)</td>
<td>0.63 (0.87)</td>
<td>8.44 (0.81)</td>
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<tr>
<td>EoNEAs</td>
<td>0.37 (0.91)</td>
<td>0.034 (0.82)</td>
<td>0.73 (0.98)</td>
<td>0.11 (0.92)</td>
<td>0.24 (0.97)</td>
<td>6.77 (1.07)</td>
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Table A2. Summary of annual-mean CN (3 nm dry diameter), CCN (50 nm dry diameter), CCN (70 nm dry diameter), surface area density (dry) and condensation sink in the continuum region (dry) simulated by GLOMAP-mode (improved configuration) with the ratio to that simulated by GLOMAP-bin shown in parentheses. Units are cm$^{-3}$ for CN and CCN, µm$^2$ cm$^{-3}$ for surface area concentration and µm cm$^{-3}$ for condensation sink. Regions match those defined in Merikanto et al. (2009).

<table>
<thead>
<tr>
<th>Region</th>
<th>CN$_3$ (1.15)</th>
<th>CCN$_{50}$ (1.18)</th>
<th>CCN$_{70}$ (1.17)</th>
<th>sarea$_{dry}$ (1.09)</th>
<th>csink$_{cntm,dry}$ (1.08)</th>
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<td>534.1</td>
<td>450.2</td>
<td>494.2</td>
<td>5852</td>
</tr>
<tr>
<td>GloCnt</td>
<td>1778</td>
<td>1148.3</td>
<td>914.2</td>
<td>810.7</td>
<td>12141</td>
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<tr>
<td>GloMrn</td>
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<td>295.4</td>
<td>269.9</td>
<td>371.3</td>
<td>3408</td>
</tr>
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<td>1345.9</td>
<td>1136.5</td>
<td>20279</td>
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<td>803.2</td>
<td>12445</td>
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<tr>
<td>S.Amer</td>
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<td>1141.4</td>
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<td>468.5</td>
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<td>SE.Asia</td>
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<td>WofNAf</td>
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<td>899.4</td>
<td>775.7</td>
<td>754.5</td>
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Fig. 1. Surface size distributions simulated by GLOMAP-bin (black) and the standard (v6I) GLOMAP-mode run (blue) and improved (v6R) GLOMAP-mode run (red). Number size distributions (dN/dlogr) are shown for three marine and three continental locations in June (a–f) and December (g–l). The exact locations are as follows: Atlantic (40° W, 43° N), N. Pacific (163° W, 43° N), Southern Ocean (163° W, 38° S), E. USA (84° W, 43° N), Europe (6° E, 43° N) and China (101° E, 35° N).
Fig. 2. Simulated marine surface zonal-mean geometric mean particle diameters in the Aitken and accumulation mode compared to observed values from a compilation of 30-yr of measured size distribution data (asterisks) for the marine boundary layer (Heintzenberg et al., 2000). The black lines show simulated annual mean (averaged to the 15-degree grid) and the blue/red lines show minimum/maximum monthly-mean values. Results from two GLOMAP-mode v6 runs are shown: as configured originally (solid lines) and after the revisions to the modal aerosol settings (dashed lines). The error-bars around the asterisks show the observed standard-deviation in each latitude band. Values of mean normalised bias and correlation coefficient are shown in Table 5 from model annual-means (ocean grid boxes only) and by averaging up to the 15 degree grid to match the observations.
Fig. 3. Surface number size distributions at (30–35° W, 40–45° N) as simulated by GLOMAP-bin (solid), the standard (v6I) GLOMAP-mode run (dashed) and improved (v6R) GLOMAP-mode run (dot-dashed). The observations (asterisks) represent the climatological size distribution reported for clean marine air masses in Raes et al. (2000). Model lines are means over the range 30–35° W, 40–45° N. The blue/red lines show minimum/maximum monthly-mean values from each model run.
Fig. 4. Global surface maps showing (a) sulphate (SO$_4$) and (c) sea-salt (NaCl) aerosol mass simulated by GLOMAP-mode (v6R) on the annual-mean. The ratio of GLOMAP-mode (v6R) to GLOMAP-bin simulated SO$_4$ and NaCl are shown in panels (b, d), respectively. Regions coloured yellow, orange and red in the relative bias maps indicate where GLOMAP-mode is higher than bin by 25–50, 50–100 and 100–200 %, respectively, whilst light-blue, dark-blue and black indicate where bin is higher by the same proportions. Regions where the two schemes are within 25 % are colored white.
Fig. 5. Global surface maps showing (a) BC and (c) OC aerosol mass simulated by GLOMAP-mode (v6R) on the annual-mean. The ratio of GLOMAP-mode (v6R) to GLOMAP-bin simulated BC and OC are shown in panels (b, d), respectively. Regions coloured yellow, orange and red in the relative bias maps indicate where GLOMAP-mode is higher than bin by 25–50, 50–100 and 100–200 %, respectively, whilst light-blue, dark-blue and black indicate where bin is higher by the same proportions. Regions where the two schemes are within 25 % are colored white.
Fig. 6. Vertical profiles of simulated BC mass mixing ratio for GLOMAP-bin (solid) and GLOMAP-mode v6R (dashed) against aircraft SP2 observations (asterisks) from Schwarz et al. (2010). The whiskers on the observations show the standard deviation over the measurements in each 1-km bin. The blue and red lines show the 25th and 75th percentiles from models participating in AEROCOM phase 1.
Fig. 7. Global surface maps showing (a) gas phase $\text{H}_2\text{SO}_4$ and (c) CN (dry-diameter $> 3$ nm) concentrations simulated by GLOMAP-mode (v6R) on the annual-mean. The ratio of GLOMAP-mode (v6R) to GLOMAP-bin simulated $\text{H}_2\text{SO}_4$ and CN are shown in panels (b, d), respectively. Regions coloured yellow, orange and red in the relative bias maps indicate where GLOMAP-mode is higher than bin by 25–50, 50–100 and 100–200 %, respectively, whilst light-blue, dark-blue and black indicate where bin is higher by the same proportions. Regions where the two schemes are within 25 % are colored white.

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Fig. 8. Global surface maps showing CCN concentrations for (a) dry diameter > 50 nm (CCN$_{50}$) and (c) dry diameter > 70 nm (CCN$_{70}$), as simulated by GLOMAP-mode (v6R) on the annual-mean. The ratio of GLOMAP-mode (v6R) to GLOMAP-bin simulated CCN$_{50}$ and CCN$_{70}$ are shown in panels (b, d), respectively. Regions coloured yellow, orange and red in the relative bias maps indicate where GLOMAP-mode is higher than bin by 25–50, 50–100 and 100–200 %, respectively, whilst light-blue, dark-blue and black indicate where bin is higher by the same proportions. Regions where the two schemes are within 25 % are colored white.
Fig. 9. Latitude-altitude plots of zonal-mean concentrations of (a) CN (all particles with dry diameter > 3 nm) and (c) CCN$_{50}$ (all soluble particles with dry diameter > 50 nm), as simulated by GLOMAP-mode (v6R) on the annual-mean. The ratio of GLOMAP-mode (v6R) to GLOMAP-bin simulated CN and CCN$_{50}$ are shown in panels (b, d), respectively. Regions coloured yellow, orange and red in the relative bias maps indicate where GLOMAP-mode is higher than bin by 25–50, 50–100 and 100–200 %, respectively, whilst light-blue, dark-blue and black indicate where bin is higher by the same proportions. Regions where the two schemes are within 25 % are colored white.
Fig. 10. Global surface maps showing (a) aerosol dry surface area density (SAD\textsubscript{dry}) and (c) dry condensation sink (CS\textsubscript{dry}) in the continuum regime as simulated by GLOMAP-mode (v6R) on the annual-mean. The ratio of GLOMAP-mode (v6R) to GLOMAP-bin simulated SAD\textsubscript{dry} and CS\textsubscript{dry} are shown in panels (b, d), respectively. Regions coloured yellow, orange and red in the relative bias maps indicate where GLOMAP-mode is higher than bin by 25–50, 50–100 and 100–200 %, respectively, whilst light-blue, dark-blue and black indicate where bin is higher by the same proportions. Regions where the two schemes are within 25 % are colored white.
Fig. 11. Taylor diagrams showing how well GLOMAP-mode (v6R) compares against GLOMAP-bin for a range of metrics in different altitude ranges of (a) 0–1 km, (b) 1–4 km, (c) 4–8 km. The distance to the perfect-model position indicating a measure of skill which combines the Pearson correlation coefficient and the relative variance for the two models runs.
Fig. 12. Simulated marine surface zonal-mean size-resolved number concentration in (a) Aitken mode and (b) accumulation mode, compared to those in the observed climatology (asterisks) for the marine boundary layer (Heintzenberg et al., 2000). The black lines show simulated annual mean values (averaged to the 15-degree grid) and the blue/red lines show minimum/maximum monthly-mean values. In (a) and (b) the solid lines are for GLOMAP-bin with the dashed lines for GLOMAP-mode (v6R). The error-bars around the asterisks show the observed standard-deviation in each latitude band. Values of $b$ and $R$ are shown in Table 5 from model annual-means (ocean grid boxes only) and by averaging up to the 15 degree grid to match the observations.
Fig. 13. Simulated CN profiles (lines) over the Pacific and Southern Oceans compared to aircraft observations (asterisks) from Clarke and Kapustin (2002) in the latitude ranges (a) 20° S–20° N, (b) 20° N–70° N and (c) 20° S–70° S. Simulated CN concentrations are at standard temperature and pressure and for particles larger than 3 nm dry diameter from means over the longitude ranges (a) 135–180° E, (b) 175–270° E and (c) 200–240° E respectively following the approach in Spracklen et al. (2005). Values of $b$ and $R$ are calculated for each region (see Table 5) from model values interpolated to a 1 km grid to match the observations. The error-bars around the asterisks show the observed standard-deviation in each latitude band. The black line shows simulated annual mean profile and the blue/red lines show minimum/maximum monthly-mean values for GLOMAP-bin (solid) and GLOMAP-mode v6R (dashed).
Fig. 14. Simulated vertical profiles of size-resolved number concentration compared to aircraft observations (Petzold et al., 2002) over NE Germany (13.5–14.5° E, 51.5–52.7° N) for particles larger than (a) 5 nm, (b) 15 nm and (c) 120 nm dry diameter. Values of $b$ and $R$ are shown in Table 5 for each size-range from model values interpolated to a 1km grid to match the observations. The error-bars around the asterisks show the observed 25th and 75th percentiles in each 1 km altitude range. The black lines shows simulated annual mean profiles and the blue/red lines show minimum/maximum monthly-mean values for GLOMAP-bin (solid) and GLOMAP-mode v6R (dashed).
Fig. 15. Scatter plots showing comparison of simulated CCN concentrations against observations at the range of sites compiled in Spracklen et al. (2011) for (a) GLOMAP-mode (v6R), (b) GLOMAP-bin. A scatter plot of GLOMAP-mode against GLOMAP-mode simulated CCN is shown in panel (c). Each model CCN is calculated based on the stated supersaturation for the corresponding measurement, using the monthly-mean for the month the observation was made.
Fig. 16. The simulated annual cycle of CCN concentrations (solid lines) against observations (asterisks) at Mace Head and Cape Grim. The measurements at Mace Head (a) are with 0.5% superstaturation (Reade et al., 2006) whilst the comparison at Cape Grim is shown for measured CCN concentrations at supersaturations of (b) 1.2% and (c) 0.23%, see Ayers and Gras (1991). Vertical bars around the Cape Grim observations show the observed range from 1981–1989. Solid line shows GLOMAP-bin, dashed line is GLOMAP-mode (v6R).