The regional aerosol-climate model REMO-HAM

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Abstract. REMO-HAM is a new regional aerosol-climate model. It is based on the REMO regional climate model and includes most of the major aerosol processes. The structure for aerosol is similar to the global aerosol-climate model ECHAM5-HAM, for example the aerosol module HAM is coupled with a two-moment stratiform cloud scheme. On the other hand, REMO-HAM does not include an online coupled aerosol-radiation nor a secondary organic aerosol module. In this work, we evaluate the model and compare the results against ECHAM5-HAM and measurements. Four different measurement sites were chosen for the comparison of total number concentrations, size distributions and gas phase sulfur dioxide concentrations: Hyytiälä in Finland, Melpitz in Germany, Mace Head in Ireland and Jungfraujoch in Switzerland. REMO-HAM is run with two different resolutions: 50 × 50 km2 and 10 × 10 km2. Based on our simulations, REMO-HAM is in reasonable agreement with the measured values. The differences in the total number concentrations between REMO-HAM and ECHAM5-HAM can be mainly explained by the difference in the nucleation mode. Since we did not use activation nor kinetic nucleation for the boundary layer, the total number concentrations are somewhat underestimated. From the meteorological point of view, REMO-HAM represents the precipitation fields and 2 m temperature profile very well compared to measurement. Overall, we show that REMO-HAM is a functional aerosol-climate model, which will be used in further studies.

1 Introduction

Aerosol particles have an important role in the climate system. They have the ability to scatter and absorb both solar and thermal radiation. These processes are referred to as the direct effect of aerosols (Haywood and Boucher, 2000). Absorbing aerosol particles can also heat the surrounding air,
which can lead to evaporation of cloud droplets. This effect is known as the semi-direct effect (Hansen et al., 1997). Finally, the ability to act as cloud condensation nuclei (CCN) and ice nuclei (IN) is referred to as the indirect effect of aerosol particles (Lohmann and Feichter, 2005). The indirect effect is usually divided into two parts: the cloud albedo effect (an increase in aerosol concentrations leads to an increase in cloud droplets, which increases the albedo) and the cloud lifetime effect (increase in aerosol concentrations leads to smaller droplets, which reduces precipitation efficiency and prolong the cloud lifetime). Aerosol effects have been widely studied experimentally and with models. However, the uncertainties of aerosol-cloud interactions still remain quite high (IPCC, 2007).

Aerosol particles originate both from natural and anthropogenic sources (Seinfeld and Pandis, 1998). Tropospheric aerosol particles are either formed from the gas phase through nucleation and growth, or directly emitted from the surface of the Earth. Downward mixing of aerosol particles from the stratosphere has a minor contribution to tropospheric aerosol loads. Kulmala et al. (2004) showed that nucleation events occur frequently around the globe and that nucleation has a significant role in global and regional CCN concentrations. Major sources of emitted particles are oceans, arid and semi-arid regions, volcanoes, wildfires and combustion of fossil and biomass fuels (Dentener et al., 2006).

Many of the aerosol processes have strong regional characteristics. Sogacheva et al. (2005, 2007) showed that nucleation events occur more frequently in Hyyttiälä, Finland, if the incoming air masses come from the Arctic Ocean rather than from Central Europe. This is interesting, because nucleation is a frequent process in Central Europe, and it is not known what suppresses nucleation when air flows towards Scandinavia. Atmospheric dynamics influence the properties of aerosol particles, but on the other hand, local aerosol processes change the number concentrations and size distributions, which in turn influences the dynamics. Laaksonen et al. (2005) showed that in the Po Valley, Italy, nucleation may contribute up to a third to the regional CCN budget. Moreover, Raatikainen et al. (2010) reported that the aerosol particle volume increases with increasing population along the parcel trajectory. This shows the importance of emissions to atmospheric aerosol properties.

Aerosol effects on warm clouds have been considered in climate models for many years (Jones et al., 1994). The first climate models used sulfate aerosols as a surrogate for all anthropogenic aerosols. During the development of the climate models, the major global aerosol components (such as sulfate, particulate organic matter, black carbon, sea salt and mineral dust) have been included, e.g. in ECHAM by Stier et al. (2005). Different approaches to describe the aerosol distribution have been implemented, such as in ECHAM modal representation by Stier et al. (2005) and a sectional representation by Kokkola et al. (2008). Aerosol modules are used in global (Stier et al., 2005) and regional (Zubler et al., 2011a) climate models with a coupling to the models’ cloud scheme (Lohmann et al., 2007).

Recently, many modeling efforts have been made to study the aerosol-cloud interactions on different scales. Just to list a few, Stier et al. (2005) presented the global aerosol-climate model ECHAM5-HAM and showed how well the model performs on global scale. Spracklen et al. (2006) used an offline version of a global aerosol model to study the contribution of boundary layer nucleation to particle concentrations. Lohmann et al. (2007) coupled the ECHAM5-HAM aerosol model with a large scale cloud scheme and studied the anthropogenic aerosol effects on the global scale. More detailed process studies (nucleation and secondary organics) with ECHAM5-HAM were carried out by Makkonen et al. (2009) and O’Donnell et al. (2011). Using a global aerosol microphysics model GLOMAP, Spracklen et al. (2010) studied the aerosol concentrations from various sites around the world. They showed, among other things, that continental boundary layer nucleation is an important process when trying to model the measured concentration. Yu et al. (2012) used the global chemistry transport model GEOS-Chem and the regional weather forecasting and chemistry model WRF-Chem, both with an advanced particle microphysics (APM) model, to calculate and investigate global and regional distributions of aerosol optical properties. On a regional scale, coupled aerosol-chemistry interactions were studied with the REMOTE model by Langmann et al. (2008). They showed the importance of emission sources for the regional scale concentrations. Finally, Zubler et al. (2011a) made simulations with the COSMO-CLM regional model in which they implemented the aerosol module HAM. This study showed how useful the regional aerosol-climate model is for studies of mesoscale phenomena.

The heavy computational load of aerosol physics and chemistry remains a major problem when modeling global and regional aerosol characteristics. Aerosol models need many species to be included even if only the main species are treated (Stier et al., 2005; Spracklen et al., 2006). This still limits the resolution of the models, although some parts of the widely used aerosol and chemistry modules have been kept simple (Feichter et al., 1996; Reddington et al., 2011). For example, Weigum et al. (2012) showed that black carbon plumes over the Pacific Ocean typically occur at scales smaller than global climate model resolution. In order to capture the regional characteristics, regional models can be used as a framework (Langmann et al., 2008; Langmann, 2000; Zubler et al., 2011a). This approach allows us to see, for example, how well the implemented aerosol physics and chemistry modules perform on smaller spatial scales. In a way, this can be seen as an intermediate step between fine scale process models and global scale climate models. This makes regional aerosol-climate models a very interesting tool to study phenomena like nucleation, particle growth, coagulation, deposition etc., and to get more detailed information.
on (locally) important species, such as black carbon and sulfur. Moreover, we can also study how well the local emission sources are represented in the databases and how they effect every day life, for example in air quality studies (e.g. Teichmann, 2010).

In this work, we present the new regional aerosol-climate model REMO-HAM, which is suitable for studying aerosol-cloud-interactions on a regional scale. It is based on the REMO regional climate model by Jacob and Podzun (1996). REMO can be used down to $10 \times 10 \text{km}^2$ resolution due to the hydrostatic limitation of the dynamical core. In this paper, we evaluate the REMO-HAM model against measurements, and compare it against the global aerosol-climate model ECHAM5-HAM.

2 Model description

2.1 General Circulation Model ECHAM5-HAM

The global climate model ECHAM5-HAM (Roeckner et al., 2003; Stier et al., 2005) is used in this study to provide lateral aerosol boundary data for the regional model simulations, and to provide data for model comparison. ECHAM5-HAM includes the aerosol module HAM (Stier et al., 2005), which uses the aerosol microphysical module M7 (Vignati et al., 2004). The aerosol model is coupled with a double-moment cloud scheme (e.g. Lohmann et al., 2007). In the following sections, more detailed information about the different modules will be provided.

2.2 The REMO model

The regional model REMO is a hydrostatic, three-dimensional atmosphere model, that has been developed at the Max Planck Institute for Meteorology in Hamburg. It is based on the Europa Model, the former numerical weather prediction model of the German Weather Service (for more details, see Jacob and Podzun, 1996; Jacob, 2001). REMO can be used for spatial resolutions from $10 \times 10 \text{km}^2$ upwards. The limitation comes from the dynamical core, which is hydrostatic (REMO also has a non-hydrostatic extension to the hydrostatic core, but it was not used in this study). The physical core of REMO is based on the physical package of the global circulation model ECHAM4 (Roeckner et al., 1996). Prognostic variables are the horizontal wind components, surface pressure, temperature, specific humidity, cloud liquid water and ice. The vertical levels in REMO are represented in a hybrid sigma-pressure coordinate system. Hybrid coordinates follow the surface orography in the lower levels and become independent from surface orography at higher atmospheric model levels.

In REMO, the stratiform (large-scale) cloud scheme is based on the original ECHAM4 cloud scheme (for details see Roeckner et al., 1996) and has been updated following ECHAM5 by Pfeifer (2003). The scheme includes prognostic equations for cloud water, water vapor and cloud ice, and has an empirical cloud cover scheme by Sundqvist et al. (1989). The cloud droplet number concentration is parameterized separately for continental and maritime climate, and is a function of height (Roeckner et al., 1996).

The convective (sub-grid) cloud parameterization is based on the mass-flux scheme from Tiedke (1989) with modifications by Nordeng (1994). Pfeifer (2003) also included a new type of convection to the cloud scheme, the so called cold convection. This type of convection might occur in cold air outbreaks over sea in the extra tropical atmosphere.

REMO does not include an aerosol module. The information about aerosols, for example in the radiation scheme, is based on the climatology from Tanre et al. (1984). In this climatology, the spatial distributions of the optical thickness of land, sea, urban, and desert aerosols, and well mixed tropospheric and stratospheric background aerosols are represented. The climatology is based on a global T10 spectral distribution ($\approx 1300 \text{km}$, fixed in time) and the aerosols in this climatology have no direct influence on the clouds, that is, the indirect aerosol effects are not represented. The climatology is highly absorbing, mainly over Africa and in Southern and Eastern Europe (Zubler et al., 2011b). The reason for this is the unrealistic dust component, which dominates the aerosol optical depth over these areas. In addition, the low resolution of the climatology is problematic, especially on regional scales. This is a known deficiency of the current model version and a new radiation scheme (coupled with HAM aerosol module) will be implemented.

2.3 Aerosol physics and chemistry

HAM is an aerosol chemistry and physics model, which predicts the evolution of an ensemble of microphysically interacting internally- and externally-mixed aerosol populations as well as their size-distribution and composition. HAM uses the aerosol microphysics module M7 in which the size-distribution is represented by a superposition of seven log-normal modes. HAM itself has all the main aerosol processes: water uptake, chemistry (gas- and liquid phase), emissions, sedimentation, deposition (wet and dry) and cloud processes (scavenging). The M7 microphysical core includes the following processes: coagulation, condensation, nucleation, thermodynamic equilibrium with water vapor and inter modal transfer. A detailed description of the HAM model can be found in Stier et al. (2005), and for more information about M7, see Vignati et al. (2004). In the following, we will present the main features of the aerosol model.

The seven log-normal modes are: soluble and insoluble Aitken, accumulation, and coarse modes, and one soluble nucleation mode. Table 1 shows how HAM treats different species. The five aerosol components considered in HAM are sulfate, black carbon, organic carbon, sea salt and mineral dust. These components are divided into different modes as shown in Table 1. The aerosol dynamics, e.g. coagulation,
can change the composition of each internally mixed mode. HAM has two water uptake methods: calculating the equilibrium liquid water content of the aerosol using the ZSR method (Jacobson et al., 1996), and the second is based on Kappa-Köhler theory (Petters and Kreidenweis, 2005). In this work, the latter is used.

In the framework of this work, four different nucleation schemes are included. The binary sulfuric acid – water based nucleation methods are used by Vehkamäki et al. (2002) and Kazil and Lovejoy (2007). Furthermore, two nucleation methods are implemented for the forested boundary layer: nucleation based on cluster activation following Kulmala et al. (2006) and nucleation based on kinetic activation following Laakso et al. (2004). Newly formed particles (number and mass) are placed in the nucleation mode. Nucleation is restricted to happen only in the cloud-free portions of the model grid boxes. In cloudy portions, sulfuric acid is removed by condensation. For simulations presented in this work, we are only using the nucleation scheme by Kazil and Lovejoy (2007).

In our current setup, we use a sulfate aerosol chemistry module described by Feichter et al. (1996). In this approach, dimethyl sulfide (DMS), sulfur dioxide (SO$_2$) and sulfate (SO$_4^{2-}$) are treated as prognostic variables. We use three dimensional monthly mean oxidant fields for hydroxyl (OH), hydrogen peroxide (H$_2$O$_2$), ozone (O$_3$) and nitrogen dioxide (NO$_2$) (Stier et al., 2005). These fields originate from the calculations of the comprehensive MOZART chemical transport model (Horowitz et al., 2003). This decision was made, because in this work we want to minimize the setup differences for the analysis part. However, this approach lowers the accuracy of the higher resolution model in terms of the chemistry. Higher resolution oxidation fields should be used in the future, or the online chemical transport version of REMO by Teichmann (2010) in a coupled mode.

The chemical module in our current version calculates the oxidation in the gaseous and aqueous phase. In the gas phase, SO$_2$ and DMS are oxidized by OH during the daytime, and DMS reacts with the nitrate radical (NO$_3$) during nighttime. NO$_3$ is assumed to be in steady state with its production and loss terms, which both include reactions with NO$_2$. The reactions of O$_3$, SO$_2$ and H$_2$O$_2$ are considered in the aqueous phase. The gas phase sulfuric acid concentration is calculated with an improved time integration scheme by Kokkola et al. (2009). As the oxidant fields are on a monthly time resolution, an artificial diurnal cycle for OH has been used. In this approach, the OH concentrations follow a cosine peak between sunrise and sunset. The peak amplitude has been scaled with day length so that the original monthly mean values are preserved. The same approach is also used in ECHAM5-HAM.

At the moment, the aerosol information from HAM is not coupled with the radiation scheme in REMO. ECHAM5-HAM has a radiation model, which includes the necessary physics for aerosols. We will update the radiation module of REMO(-HAM) in the future so that aerosols are also included.

### 2.3.1 Emissions used in the model

The emission module is based on the AEROCOM emission inventory for the year 2000 excluding sea salt (SS) emissions and dimethyl sulfide (DMS) emissions (Dentener et al., 2006). The sea salt emissions are based on the approach by Stier et al. (2005). In this approach, the emissions of sea salt particles are based on a look-up table for wind speeds between 1 and 40 m s$^{-1}$. All the emitted species, except the sulfur compounds, are treated as primary emissions (emitted directly). In addition, the DMS emissions from the marine biosphere are calculated based on DMS sea water concentrations using an air–sea exchange rate. This is calculated using the model 10 m wind speed. Terrestrial biogenic emissions of DMS are prescribed (for details of both DMS emission schemes, see Stier et al., 2005).

The AEROCOM emissions are presented on a 1° x 1° global grid. This is not an ideal accuracy REMO-HAM, but
for the validation of the model, it has been used in this work. We do want to point out that higher resolution emission datasets are available on global and regional scales, and that they can be easily used with REMO-HAM. For all of the AEROCOM compounds, the data is preprocessed using a 1st order conservative remapping method. The data is remapped to the grid used in REMO and read in once per modeled month. We assume homogeneous mixing across the model grid box for all aerosol species. The emissions are injected to the lowest model layer, except for biomass burning emissions, explosive and continuous volcanoes, and emissions coming from industry, shipping and power plants. For all of the vertically dependent emissions, the levels described by Dentener et al. (2006) are used. For detailed information of different AEROCOM species, see Dentener et al. (2006) or visit http://nansen.ipsl.jussieu.fr/AEROCOM/.

In the following, we describe the aerosol emission in the model. Details on the sources can be found in Dentener et al. (2006). For the mean radius and the standard deviation of different emission modes we have used the same values as in Stier et al. (2005). Sulfur (S) is divided between gas phase (97.5 %) and particles (2.5 %) as primary SO$_4$ (sulfate). For industry, shipping and power plants emissions, 50 % of the sulfate emissions are released to the accumulation mode and 50 % to the coarse mode. The remaining sulfate emissions are divided between Aitken mode (50 %) and accumulation mode (50 %). Secondary Organic Aerosol (SOA) formation is based on prescribed biogenic monoterpenes emissions assuming a fixed SOA yield (15 %), and that SOA formation takes place in the emitting grid box. The primary organic matter (POM) emissions are also released to organic carbon. The mass ratio of organic mass to organic carbon is $1.4, 65$ % of SOA and POM emissions are assumed to be soluble and 35 % insoluble. The insoluble organic matter is released entirely to the Aitken mode. With the exception of the fire and fuel sectors, soluble organic matter is emitted as organic carbon and is divided between Aitken and accumulation modes (50 % in each). The fire POM emissions are released to the soluble accumulation mode and fuel emissions to the insoluble Aitken mode. All black carbon emissions are released to the insoluble Aitken mode. For dust emissions, we use prescribed AEROCOM mass fluxes. These fluxes are on a monthly scale and have separate fields for accumulation and coarse modes (the mode properties are the same as in M7 Vignati et al., 2004).

### 2.4 Cloud microphysics

The stratiform cloud scheme described in Sect. 2.2 calculates the number of cloud droplets separately for maritime and continental clouds. The calculation does not take into account the information about aerosols provided by the HAM aerosol module. One way to use the aerosol information is to use the approach by Lohmann and Roeckner (1996), where the cloud droplet concentration for continental and marine clouds is calculated from the aerosol sulfate mass. This approach corrects the values calculated by the normal scheme but still does not fully take into account the information provided by the aerosol module. In order to use the information in the stratiform clouds, we have implemented the double-moment cloud microphysics scheme by Lohmann et al. (2007). This scheme has prognostic equations for water vapor, cloud water and cloud ice. In addition, it has prognostic equations for cloud droplet number concentration (CDNC) and ice crystal number concentration (ICNC). For the cloud cover (fractional), we use the approach by Sundqvist et al. (1989), in which the relative humidity determines whether a cloud is formed in the grid box or not. The autoconversion of cloud droplets can be calculated with two different parameterizations: Beheng (1994) and Khairoutdinov and Kogan (2000). The cloud scheme also includes a cirrus parameterization from Lohmann and Kärcher (2002).

The new cloud scheme is fully coupled with the HAM aerosol module. To connect the information from the aerosol module to the CCN activation, the activation schemes from Lin and Leaitch (1997) and Abdul-Razzak and Ghan (2000) are implemented. In this study, the latter is used. To connect the calculated cloud droplet and ice crystal number concentrations to the radiation, the approach by Lohmann et al. (2007) for cloud droplets and the approach by Lohmann et al. (2008) for ice crystals are implemented. In these approaches, using the CDNC and ICNC information combined with the cloud liquid water/ice content, the effective radius of droplets/crystals is calculated and passed to the radiation.

### 2.5 Tracer transport

The model tracers undergo the following transport processes: transport in convective clouds, sedimentation, dry and wet deposition, vertical diffusion and vertical- and horizontal advection. The convective transport of tracers is based on the mass-flux scheme described in Sect. 2.2. The sedimentation velocity calculation is based on the Stokes velocity with the Cunningham slip-flow correction factor accounting for non-continuum effects (Stier et al., 2005). The dry deposition is based on the same size-dependent parameterization as in the ECHAM5-HAM model (Stier et al., 2005). Current version of REMO-HAM does not include information about canopy height or soil pH. For these, REMO-HAM uses the same prescribed fields as ECHAM-HAM. Other land surface parameters are directly from the model. For the wet deposition, we have implemented two different approaches for below-cloud aerosol scavenging. The first one, by Stier et al. (2005), can be used, but also the size dependent below-cloud scavenging by rain, or by rain and snow, by Croft et al. (2009) is introduced and used in this work. The vertical diffusion fluxes due to subgrid scale turbulence are calculated for the lowest layer (surface layer) according to Louis (1979). For the other layers, a second-order closure scheme by Mellor and Yamada (1974), is used. For the horizontal and vertical
advection, we have implemented the finite difference, anti-diffusive scheme proposed by Smolarkiewicz (1984, 1983). The scheme is mass conserving, positive definite and computationally efficient. The monotonicity is achieved by using higher order flux corrections. We also use two corrective steps (using anti-diffusion velocity) in order to decrease the numerical diffusion (for details, see Langmann, 2000; Smolarkiewicz, 1983, 1984).

ECHAM5-HAM and REMO-HAM included a slightly improved dry deposition scheme. The dry deposition scheme calculates dry deposition velocities separately for vegetated surface, wet skin, bare soil or snow, water and ice (Ganzeveld and Lelieveld, 1995; Ganzeveld et al., 1998). It is updated to use separate values of virtual potential temperature, roughness length and stability functions over land, water and ice rather than grid box averages of these quantities in the calculation of dry deposition velocities. (Actually the virtual potential temperature was erroneously taken from the value for land, which is assigned a default value over water or ice). This results in a significant strengthening of the dry deposition sink for some species, for example in ECHAM5-HAM, global dry deposition of SO$_2$ increased from 17.6 Tg(S) yr$^{-1}$ before the changes to 20.8 Tg(S) yr$^{-1}$ afterwards (cf. a total emission flux of 71 Tg(S) yr$^{-1}$). We have used this updated scheme on all of the simulations done in this work.

2.5.1 Lateral and upper boundary

For the meteorology, the model uses the relaxation scheme by Davies (1976) at the lateral boundary. In this scheme, the eight outermost grid boxes are adjusted with an exponentially decreasing function. The influence of the large-scale driving fields decreases when moving towards the domain center. For the aerosol part, the lateral boundary treatment of tracers is implemented after Pleim et al. (1991). Due to discontinuities of the advection flux fields and the concentration fields at the boundaries, the lateral boundaries would have upstream reflections near the outflow boundary. The approach by Pleim et al. (1991) minimizes these discontinuities by setting the flux divergence to zero in the grid boxes next to the boundary.

We used the European Centre for Medium-Range Weather Forecasts (ECMWF) operational data set ($1.125^\circ \times 1.125^\circ$) for the meteorological data at the model domain boundaries. The aerosol data is from ECHAM5-HAM which is run for the needed time period nudging (assimilation by linear relaxation) the model towards the ECMWF data. The ECHAM5-HAM global output is conservatively remapped to the REMO grid using the Climate Data Operators (CDO) program (https://code.zmaw.de/projects/cdo/). We have also done the remapping for the vertical levels due to the better representation of orography. In this work, we used 31 vertical levels for the models. In ECHAM5-HAM, the uppermost level reaches 10 hPa, and in REMO-HAM 20 hPa is reached.

Fig. 1. The orographies of the domains used (in meters) and the selected CREATE database stations: Hyytiälä, Melpitz, Mace Head and Jungfraujoch. Shown on the left hand side is the 0.44° resolution domain and on the right hand side the 0.088° resolution domain.

2.5.2 Dynamical downscaling

The resolution of ECHAM5-HAM used in this study was T63L31 with a time step of 720 s. This resolution corresponds to a resolution of 1.9° (horizontally 210 km). A jump from 1.9° resolution to the resolution used in this study, 0.088° (10 × 10 km$^2$) is too large. In order to overcome this problem, dynamical downscaling was used (also known as the double-nesting method). In this method, the model is run with an intermediate resolution and the results from this simulation are used as boundary data for the higher resolution simulation. We used the resolution of 0.44° (50 × 50 km$^2$) for the intermediate step.

Figure 1 shows the orographies for the domains used in the dynamical downscaling process. The left figure represents the domain for the 1st step of dynamical downscaling. Using the data from this simulation, the model was run for the domain shown on the right hand side in Fig. 1.

3 Measurement data

We compare our results to measurement data from CREATE (Construction, use and delivery of an European aerosol database)$^1$ and EBAS$^2$ databases. From this dataset, we chose 4 different measurement sites: Hyytiälä (Finland 62° N and 24° E), Melpitz (Germany 51° N and 13° E), Mace Head (Ireland 53° N and 10° W) and Jungfraujoch (Switzerland 47° N and 8° E). From these stations, only the Melpitz and Jungfraujoch sites are within the 10 km resolution domain.

From the Hyytiälä (Dal Maso et al., 2005) and Melpitz (Birmili and Wiedensohler, 2000) stations we have used differential mobility particle sizer (DMPS) data for the total number concentrations and size distributions. The condensation particle counter (CPC) data was used for total number.

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$^1$http://tarantula.nilu.no/projects/ccc/create/general_info.htm
$^2$http://ebas.nilu.no
concentration at Jungfraujoch (Collaud Coen et al., 2007; Collaud Coen and Weingartner, 2011) and at Mace Head (O’Connor et al., 2008). In addition, from Mace Head the aerosol size distributions were measured with a Scanning Mobility Particle Sizer (SMPS).

The gas phase SO$_2$ concentrations are also compared against measurements. For Mace Head, there are no measurements of SO$_2$. As a proxy for Mace Head, we use data from Valentia Observatory which is located approximately 150 km due southwest from the Mace Head measurement station (Bashir et al., 2006). The data from Valentia is daily filtered data and should represent the concentrations from Mace Head reasonably well. The time resolution of one day was also used in the gas phase measurements from Jungfraujoch. For Hyytiälä and Melpitz, the gas phase data has a 30 min time resolution.

4 Simulations and results

The ECHAM5-HAM model was run for the time period of 1 July 2004–31 December 2005. The data output frequency used was 6 h and instantaneous values of the tracers were written out. From these results, the lateral tracer boundary data for REMO-HAM were processed, as described in Sect. 2.5.1. The REMO model was run for the time period of 1 October 2004–31 December 2005 for the 0.44° (50 km) resolution with a time step of 240 s, and 1 December 2004–31 December 2005 for the 0.088° (10 km) resolution using a 50 s time step. The model was run with the normal version (REMO) and with the aerosol module version (REMO-HAM). The time periods were chosen so that ECHAM5-HAM has 3 months of spin up time for the tracers, REMO/REMO-HAM with 50 km resolution has 2 months of spin up time and REMO/REMO-HAM with 10 km resolution has 1 month.

Of the four measurement sites, Jungfraujoch is the only one located in a mountainous area (the Alps) as can be seen from Fig. 1. In the case of Hyytiälä, Melpitz and Mace Head, the results from the models are from the lowest model layer. The mountainous location of Jungfraujoch (station at 3580 m height) makes the analysis different. The model orography for the Alpine region is flatter than in reality, especially in the case of ECHAM5-HAM. This means that the lowest model layer does not describe the actual measurement station altitude and instead the closest matching pressure level is chosen. The level is calculated separately for each of the models due to the different orography.

4.1 Aerosol total number concentrations

Figure 2 presents the measured and modelled weekly aerosol total number concentrations ($N_{\text{tot}}$) for different measurement sites. We can see that for Hyytiälä ECHAM5-HAM is in reasonable agreement with the measured concentrations. The modelled values are slightly lower than measured, especially during late winter ($\Delta N_{\text{tot}} = 1000 \, \text{#/cm}^3$), early summer ($\Delta N_{\text{tot}} = 500 \, \text{#/cm}^3$) and autumn ($\Delta N_{\text{tot}} = 1500 \, \text{#/cm}^3$), but the yearly cycle is fairly well captured. REMO-HAM has a similar yearly cycle to the measurements, but the fluctuations in concentrations are higher. In some cases, the concentration peaks are higher than the measurement or results from ECHAM5-HAM, but overall REMO-HAM gives lower concentrations. In any case, the values from ECHAM5-HAM are fairly close to the measurements, whereas REMO-HAM has problems during late winter ($\Delta N_{\text{tot}} = 3000 \, \text{#/cm}^3$), late summer ($\Delta N_{\text{tot}} = 2000 \, \text{#/cm}^3$) and autumn ($\Delta N_{\text{tot}} = 2500 \, \text{#/cm}^3$).

Although two boundary layer (BL) nucleation schemes are implemented in the models (Sect. 2.3), these schemes were turned off for these simulations. For testing purposes, we have done simulation with activation and kinetic BL nucleation, but the results showed severe overestimation in total particle numbers (results not shown). This suggests that the absence of a BL nucleation scheme has contributed to the underestimation of particle numbers at Hyytiälä during spring and autumn, which are typical nucleation event times at that site (Dal Maso et al., 2005). This has been also shown with different modeling studies. For example, Spracklen et al. (2010) and Reddington et al. (2011) showed that activation and kinetic BL nucleation does improve the modelled aerosol concentrations. Moreover, Spracklen et al. (2006) showed that either BL nucleation or primary emissions cannot directly explain the measured concentrations at Hyytiälä, but that the sum of these is the best fit to the measurements. This means that the models would give higher concentrations during spring, summer and autumn, which can be also seen from the study by Spracklen et al. (2010). The reason for the high nucleation rates with both of the BL nucleation
schemes is the too high SO$_2$ concentrations in the models (this is shown in Fig. 7). Overall, the values from ECHAM5-HAM and REMO-HAM are similar as in Spracklen et al. (2010) and Reddington et al. (2011), although it must be said that in the latter study the simulated period is only one month long.

From Fig. 3 we can see that the modal number concentrations between the models do not differ much at Hyytiälä. The Aitken mode is smaller in REMO-HAM during the whole year, which can explain the differences in Fig. 2. For example, the large underestimate of $N_{\text{tot}}$ in REMO-HAM for the late winter comes from the Aitken mode. One possible explanation is the resolution of the models: based on our other simulations, ECHAM5-HAM spreads the emissions much wider than REMO-HAM. Also, if the resolution of emissions is too coarse (like in our case, coarser than the grid used in REMO-HAM), local emission sources might not be positioned completely correctly. If we look at the nucleation mode, the concentrations are a bit higher in REMO-HAM, but the difference is not very big.

The results from Melpitz show that the models give lower concentration than the measurements throughout the whole year ($\Delta N_{\text{tot}} = 1000–4000 \, \text{#/cm}^3$) and that the difference between the models is quite small. Similar behavior and concentrations can also be seen in Reddington et al. (2011) and Spracklen et al. (2010). In the latter work, it was shown that for Melpitz, BL nucleation and/or increased carbon emissions could explain the underestimation. When looking at the two different REMO-HAM resolutions, the difference is barely discernible. Unlike at Hyytiälä, the nucleation mode concentrations differ the most. On the other hand, the Aitken mode concentrations are much higher and dominate the total number concentrations (Fig. 3). This is why the difference in nucleation mode is not visible in the total number concentrations.

At Mace Head, ECHAM5-HAM reproduces the measured values fairly well during the first half of the year, but underestimates the concentrations during the second half ($\Delta N_{\text{tot}} = 1000 \, \text{#/cm}^3$). REMO-HAM has the same pattern, although during late winter and spring time the concentrations are lower than in ECHAM5-HAM and measurements ($\Delta N_{\text{tot}} = 2500 \, \text{#/cm}^3$). Once again, this same behavior and concentrations have been modelled before, and the BL nucleation can be the key factor (Reddington et al., 2011; Spracklen et al., 2010). Figure 3 shows that the difference between ECHAM5-HAM and REMO-HAM comes from the nucleation mode, which is lower in REMO-HAM during the first quarter of the year, and in summer and autumn. This result is in good agreement with the low total number concentration seen in Fig. 2.

As mentioned in Sect. 4, the Jungfraujoch measurement site is located in a mountainous region. These kind of areas are demanding for the model and the results have greater error than usual, which can be seen from Fig. 2. The concentrations given by the models are too high throughout the year. ECHAM5-HAM has somewhat higher concentrations ($\Delta N_{\text{tot}} = 4000 \, \text{#/cm}^3$) than REMO-HAM ($\Delta N_{\text{tot}} = 3000 \, \text{#/cm}^3$). The 10 km resolution simulation has slightly higher concentrations, but the difference between the resolutions is not very significant ($\Delta N_{\text{tot}} = 3500 \, \text{#/cm}^3$). This is not the case with Reddington et al. (2011) and Spracklen et al. (2010), who both modelled much lower concentrations ($N_{\text{tot}} \approx 1000 \, \text{#/cm}^3$). The difference between ECHAM5-HAM and REMO-HAM (50 km) in nucleation mode can be seen from Fig. 3 and the interesting point is that the
nucleation mode concentrations are very high compared to other measurement sites. Later on in Fig. 4, it is shown that at higher altitudes the concentrations increase. This explains why the concentrations are so high at Jungfraujoch. The reason for the high concentration on higher altitudes is explained in Sect. 4.1.2.

Figures 2 and 3 show the number concentrations only from the lowest model layer, except for Jungfraujoch where the model level is matched to the measurement by pressure. In this work, we have not included vertical measurement data, but we compare ECHAM5-HAM and REMO-HAM (50 km resolution) profiles. From Fig. 4 we see how the vertical distribution of particle number changes seasonally. ECHAM5-HAM has higher concentrations at all sites below 750 hPa. Above this, REMO-HAM has higher concentrations at Hyytiälä and Melpitz, excluding the highest pressure levels (close to the surface). Also noticeable is that total number concentrations get higher until around 300 hPa and then start to decrease. This kind of behavior has been previously observed in ECHAM5-HAM by Kazil et al. (2010). The reason for this is the high concentration in the nucleation mode, which can be seen from Fig. 5 (note: these values are annual means which smooths out differences when compared to seasonal values). We can see how the nucleation mode dominates the total number concentrations from 200 hPa to 900 hPa. REMO-HAM has somewhat higher concentrations in all of the modes (with some fluctuations), except the nucleation mode, which is higher in ECHAM5-HAM. At higher altitudes, the difference is no longer noticeable. Also interesting is the effect of coarser orography at Jungfraujoch with ECHAM5-HAM, which causes the pressure levels at Jungfraujoch to be higher than in REMO-HAM (see Fig. 5). The low horizontal resolution tends to smoothen the orographical differences which is why REMO-HAM has a better representation of the Alps (the coarse and accumulation mode minima are at higher altitudes than with ECHAM5-HAM). Figure 5 also shows why the nucleation mode concentration is so high at Jungfraujoch in Fig. 2. As mentioned, Jungfraujoch station is located at the Alps and the values in Fig. 2 were chosen based on the pressure levels (higher altitude). This leads to higher nucleation mode concentrations at Jungfraujoch station.

Kazil et al. (2010) showed that in ECHAM5-HAM, the increase in nucleation mode concentrations as a function of altitude is stronger than measurements indicate, at least in Pacific regions. Possible reasons for this include too high SO\textsubscript{4}^2 concentrations which can lead to too high H\textsubscript{2}SO\textsubscript{4} levels affecting the nucleation. Also, the oxidation processes may influence the H\textsubscript{2}SO\textsubscript{4} budget and the treatment of ultrafine particles in M7 in terms of the mode limits can have an effect on the concentrations. In addition, Kazil et al. (2010) showed that the ultrafine particle number concentrations are high in ECHAM5-HAM regardless of the nucleation method used. In this work, we use the binary sulfuric acid – water based nucleation scheme, which favors colder temperatures. For the BL values, activation or kinetic nucleation would smooth out the difference due the higher concentrations on higher altitudes, which would of course be more important in the European boundary layer compared with Pacific areas.

### 4.1.1 Aerosol size distributions

As mentioned in the previous section, the nucleation mode can explain the difference in total number concentrations between the models at Melpitz, Mace Head and Jungfraujoch.
The high concentrations in the nucleation mode can be also seen in Fig. 6, where the yearly mean size distributions from Hyytiälä, Melpitz and Mace Head as well as the modeled mean distributions are shown. Results from Hyytiälä show that the distributions from the models are quite similar to the measurements, except for small particles, which are overestimated. The width of the distribution is not quite captured, although ECHAM5-HAM has a wider distribution than with REMO-HAM. The peak is almost in the right place for both of the models and ECHAM5-HAM has the height of the main peak very well captured, whereas REMO-HAM underestimates the concentrations. At Mace Head, similar behavior can be seen, although both of the models underestimate the concentration of the larger particles. The noticeable difference is that ECHAM5-HAM has more particles in the smallest sizes, which is in agreement with the spring and summer time higher nucleation mode concentrations as seen in Fig. 3. At Melpitz, we can see features similar to the other sites, but the form of the distributions gives some clue as to why the models differ. It seems that in the case of ECHAM5-HAM, more gas phase sulfuric acid condenses on the particles and this leads to smaller nucleation mode and to the higher and wider distribution. In the next section we will explain this in more detail.

4.1.2 Nucleation mode and gas phase

In this work, neither the activation nor the kinetic nucleation schemes were used. However, the neutral and charged H2SO4/H2O nucleation scheme by Kazil and Lovejoy (2007) was used in both of the models during the simulation period. In this approach, the nucleation rate depends on the sulfuric acid concentration. The source for gas phase sulfate is the oxidation processes of SO2 and DMS, as described in Sect. 2.3. The concentrations of SO2 (especially in continental areas) are much higher than those of DMS, and thus it is the main proxy for gas phase sulfate production (Hamed et al., 2010). The difference in the nucleation mode between ECHAM5-HAM and REMO-HAM could be explained by higher nucleation rates, which would be the results of higher sulfate (sulfuric acid) concentrations due to the different SO2 concentrations.

Figure 7 shows the SO2 gas phase concentrations for all the measurement sites and models (here the results from Valentia, Ireland, instead of Mace Head are shown). It is quite clear that both of the models overestimate SO2 concentrations at all of the measurement sites. At Hyytiälä, REMO-HAM has higher values than measurements, especially during winter and early spring. ECHAM5-HAM has similar values for the beginning of the year, although in a few cases slightly lower than those in REMO-HAM. Later on, values in REMO-HAM decrease, but are still higher than the measurements. ECHAM5-HAM decrease, but it also has a decreasing slope, but the values start to increase during the summer and are too high especially during the late autumn and winter. At Melpitz, the values are quite high in both of the models. The total number concentrations are underestimated at Melpitz and the reason for this is the low concentration in the nucleation mode. Although SO2 concentrations are high at Melpitz, the nucleation mode number concentration is not, because there is not enough sulfuric acid to activate nucleation. The reason for this is the Aitken mode. As we can see from Fig. 3, the Aitken mode concentration is quite high at Melpitz. This leads to a larger condensation sink, which in turn leads to low sulfuric acid concentrations. This can be also seen from Fig. 6, where the nucleation peaks are low at Melpitz, but otherwise the distribution is high and wide. In case of REMO-HAM, the condensation sink is lower and there is more sulfuric acid to activate nucleation. This can be seen from Figs. 3 and 6. The reason for differences in the Aitken mode are most probably due to resolution, which affects aerosol transport of species (emissions spread wider with coarse resolution). Another factor can be the coarse resolution of emission. Some local important emission sources can be mislocated (in terms of grid boxes) when the emissions are remapped from the coarse resolution to the higher resolution.

As previously mentioned, the model versions used included some changes to dry deposition, which decrease the SO2 concentrations. If we look more closely at how the nucleated particles grow in M7, the only condensing substance (besides water) is sulfuric acid. It seems that when the SO2 concentrations decrease, the sulfuric acid concentrations decrease as well and the condensation is no longer high enough to grow the particles. This can be seen from Fig. 6, where the nucleation mode has high peaks on every measurement site.

The activation and kinetic boundary layer nucleation schemes were not used in this work, because the sulfuric acid concentrations were so high. The implemented BL
nucleation schemes are very sensitive to $\text{H}_2\text{SO}_4$ concentration and without condensation of organic matter (SOA model), the nucleation mode concentrations would be unrealistically high. We are currently working on the problem of too high $\text{SO}_2$ and $\text{H}_2\text{SO}_4$ concentrations in other projects. For example, the chemistry part of the models is being improved in terms of the oxidant fields.

Figure 8 shows the zonal and yearly means for Europe for sulfate production rates and sulfuric acid concentrations for both of the models. Liquid phase production comes from the oxidation of $\text{SO}_2$ by $\text{H}_2\text{O}_2$ and $\text{O}_3$, and gas phase production from the oxidation of DMS by $\text{OH}$ and $\text{SO}_2$ by $\text{OH}$. We see that the production rates are fairly similar in the liquid and gas phases. The biggest difference is that in ECHAM5-HAM, the production rates are higher at higher altitudes. This is connected to the $\text{SO}_2$ concentrations, which are also higher in ECHAM5-HAM. On the other hand, the low liquid phase production rates over the Sahara in REMO-HAM are a direct effect of lower cloudiness (and precipitation). The precipitation bias can be seen (later on) from Fig. 11. The sulfuric acid ($\text{H}_2\text{SO}_4$) concentrations are somewhat lower in REMO-HAM, but the pattern is similar to ECHAM5-HAM. In any case, based on Fig. 8, we can state that the main processes in REMO-HAM for $\text{H}_2\text{SO}_4$ are working as they should. Small differences, such as those seen in the nucleation mode concentration in Fig. 3, can be explained by normal fluctuations arising from the different model resolutions.

### 4.1.3 Spatial variability

We can see from Fig. 9 how the different resolution of the models changes the spatial distribution of the total number concentration. The values shown are the mean values over summer months (June, July and August) and for lowest model layer. The difference in details between ECHAM5-HAM and REMO-HAM (50 km) is considerable, although the main features of spatial distribution can be seen from both. The values are fairly similar for different areas, which was also shown in Fig. 2. If the two different REMO-HAM resolutions are compared, the difference is quite small. Some of the very local concentration peaks are different, but the main features of the spatial distribution are more or less identical. This is because of two main reasons: firstly, the higher resolution was driven by the lower resolution simulation and secondly, the emissions database was the same for both of the simulations and it had coarser resolution than either of the model versions. It is quite clear that if we want to use the full benefit of the high resolution, the emission database needs to be updated, at least in terms of the spatial resolution. The resolution difference between the AEROCOM emissions...
and the 10 km simulation is too high. A higher resolution database should be used, otherwise the coarse emissions will smoothen out the benefits of accurate grid. It should also be mentioned that higher resolution in time has the potential to bring further differences between different model resolutions (now most of the species use monthly fields). In any case, Fig. 9 shows how much the higher resolution of regional aerosol-climate model can improve the spatial representation of the aerosol concentrations, and how sensitive the models are for the resolution of the emissions.

**4.2 Meteorological variables**

We have compared the meteorological model results with the Climatic Research Unit data (University of East Anglia Climatic Research Unit, 2008). The data used is the CRU TS 3.0 global dataset with 0.5° resolution. The dataset only has measurement data over land so all sea areas are excluded. The comparison with the dataset is done for the precipitation and temperature from ECHAM5-HAM, REMO and REMO-HAM. The model data is remapped to the CRU grid. Using a conservative remapping for precipitation fields and bilinear remapping for temperature fields the data is matched with the measurement grid. The temperature fields are also
Fig. 11. Precipitation difference ((model-measurements)/measurements*100 %) for ECHAM5-HAM, REMO and REMO-HAM.

Figure 11 shows the precipitation difference ((model-measurements)/measurements) for each of the models. ECHAM5-HAM seems to have a wet bias over Eastern and Central Europe, and a dry bias over some parts of Western Europe, North Africa and the Middle East. REMO and REMO-HAM have the same pattern, but the biases are not as high as with ECHAM5-HAM. REMO is overall closer to the measurements than REMO-HAM, which has a wet bias, especially over Eastern Europe. The pattern of precipitation between the models is quite similar. REMO and REMO-HAM have systematic precipitation differences along the boundaries, which is an effect of the relaxation scheme (prognostic variables are adjusted within eight outermost grid boxes). The large-scale driving field influence decreases exponentially from the outermost boundary towards the center of the domain.

REMO-HAM currently uses the same autoconversion parameters as used in ECHAM5-HAM with a resolution scaling factor. These scaling factors change the autoconversion parameters according to the resolution and have an impact on the precipitation. The effect of the scaling parameters should be studied further; this work is ongoing.

5 Conclusions

We have performed simulations with the regional aerosol-climate regional model REMO-HAM with two different horizontal resolutions: 50 × 50 km² and 10 × 10 km². In addition, we carried out simulations with the global aerosol-climate model...
model ECHAM5-HAM. The simulations were for the year 2005 after a sufficiently long spin-up time. The aerosol related tracer data from the ECHAM5-HAM simulation were used as lateral boundary data for REMO-HAM. ECHAM5-HAM was nudged with ECMWF operational analysis data and REMO-HAM used the same data at the lateral boundaries of the model domain.

The results from both of the models were compared against aerosol measurements from four different measurement sites. We have shown that the REMO-HAM can represent the measured aerosol concentrations and size distributions fairly realistically. Overall the concentrations are somewhat low, but one has to keep in mind that this version of REMO-HAM does not include an online SOA-model, and that neither the activation nor kinetic nucleation scheme for the boundary layer were used. With ECHAM5-HAM, the total number concentrations shows similar patterns as REMO-HAM. The difference between the models can be mainly explained by the nucleation mode, which differs in time and space. The nucleation mode is linked to the sulfur dioxide concentrations through oxidation and nucleation. We have shown that the models tend to overestimate the sulfur dioxide concentrations and this leads to high nucleation rates. In some cases, like at Melpitz, the background concentration of aerosol was high enough to increase the condensation of sulfuric acid to particles and this led to lower nucleation rates. We implemented some improvements to the models in terms of dry deposition. These improvements led to lower sulfur dioxide and sulfuric acid levels, but raised a new problem: the condensation of sulfuric acid is no longer high enough to grow the smaller particles to larger sizes. One thing missing in REMO-HAM is the growth of particles by organic matter. ECHAM5-HAM has a secondary organic aerosol version (O’Donnell et al., 2011), but it was not used in this study. Since the aerosol module structure is similar in both of the models, we will study the option of implementing the SOA model by O’Donnell et al. (2011) into REMO-HAM.

Emissions are the dominant factor for number concentrations in the lowest model layers. In this study, we tested REMO-HAM with two different horizontal resolutions (0.44° and 0.088°). Differences in the aerosol concentrations are small between the two REMO-HAM simulations, as both use the same emission fields. This means that although REMO-HAM with 0.088° resolution is able to solve the circulation with much higher accuracy, the coarse emissions were a restricting factor for the aerosol concentrations. In order to use high resolution aerosol-climate models for detailed studies, a high resolution emissions dataset should be used.

Temperature and precipitation fields from REMO-HAM were evaluated by comparing the parameters to CRU observations. REMO-HAM captures the annual mean temperature reasonably well, but overestimates the precipitation amount in most regions. The meteorology of the model will be studied more thoroughly in forthcoming publications (for example, the effect of scaling parameter to the autoconversion parameters). Moreover, the difference in sulfur species will be studied in more detail in the future.

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