



# 1 Impact of climate change on the production and transport of

- 2 sea salt aerosol on European seas
- 3
- 4 J. Soares<sup>1</sup>, M. Sofiev<sup>1</sup>, C.Geels<sup>2</sup>, J.H.Christensen<sup>2</sup>, C.Andersson<sup>3</sup>, S. Tsyro<sup>4</sup>,
- 5 J.Langner<sup>3</sup>
- 6 [1] Finnish Meteorological Institute, Helsinki, Finland
- 7 [2] Department of Environmental Science, Aarhus University, Roskilde, Denmark
- 8 [3] Swedish Meteorological and Hydrological Institute, Norrköping, Sweden
- 9 [4] EMEP MSC-W, Norwegian Meteorological Institute, Oslo, Norway
- 10 Correspondence to: J. Soares (joana.soares@fmil.fi)
- 11

# 12 Abstract

13 The impact of climate change on sea salt aerosol production, dispersion, and fate over the Europe is 14 studied using four offline regional chemistry transport models driven by the climate scenario SRES 15 A1B over two periods: 1990-2009 and 2040-2059. The study is focused mainly on European seas: 16 Baltic, Black, North and Mediterranean. The differences and similarities between predictions of the 17 individual models on the impact on sea salt emission, concentration and deposition due to changes in wind gusts and seawater temperature are analysed. The results show that the major driver for the 18 19 sea-salt flux changes will be the seawater temperature, as wind speed is projected to stay nearly the 20 same. There are, however, substantial differences between the model predictions and their 21 sensitivity to changing seawater temperature, which demonstrates substantial lack of current 22 understanding of the sea-salt flux predictions. Although seawater salinity changes are not evaluated 23 in this study, sensitivity of sea-salt aerosol production to salinity is similarly analysed, showing 24 once more the differences between the different models. An assessment on the impact of SSA to the 25 radiative balance is presented.





# 27 1 Introduction

The sea salt aerosol (SSA) affects the Earth radiation budget, atmospheric chemistry, cloud processes, and climate (O'Dowd et al., 1997; IPCC, 2013). Anthropogenic and natural aerosols have similar annual impacts on the global radiative balance, though being predominant in different locations (Textor et al., 2006). SSA dominates the particulate mass and it is the major contributor to aerosol optical depth (AOD) over the ocean (Quinn et al, 1998).

33 SSA originates from sea spray droplets resulting from waves breaking on the seawater surface, 34 forming whitecaps that cause the entrainment of air into the water. The two main mechanisms 35 responsible for sea spray formation are air bubble bursting during whitecap formation and decay, 36 and direct tearing of droplets from the top of breaking waves. Therefore, the formation of primary 37 SSA is mainly dependent on wind speed: the emission of SSA is generally considered to be 38 proportional to surface winds cubed (Monahan et al., 1986), suggesting that small changes in 39 surface winds can have a substantial impact on the emission of this natural aerosol. Further on, 40 studies on the marine aerosol size distribution (e.g. Covert et al., 1998; Russell and Heintzenberg, 41 2000; Bates et al., 2002; Huebert et al., 2003) suggest that for high wind speeds the production of 42 very coarse SSA (with particle diameter (Dp) > 20 mm) increases, contributing to a higher transfer of heat and water vapour from the ocean to the atmosphere (Andreas et al., 1995). These processes 43 44 have a strong impact on the climate forcing. Other parameters influencing the formation of primary 45 SSA have been identified, e.g., seawater temperature and salinity, atmospheric stability, and wave 46 height and steepness (O'Dowd and Smith, 1993; Gong et al., 1997; Gong, 2003; Mårtensson et al., 47 2003; Lewis and Schwartz, 2004; O'Dowd and de Leeuw, 2007; Witek et al., 2007a, 2007b; 48 Ovadnevaite et al., 2014). Laboratory studies by Mårtensson et al. (2003) and in situ measurements 49 by Nilsson et al. (2007) show that for nano-sized particles, the aerosol number emission decrease 50 with increasing seawater temperature, and for particles with Dp > 100 nm, the number SSA increase 51 with increasing seawater temperature; reflecting different sea spray formation processes. Seawater 52 salinity also affects the droplet formation, where formation of particles with  $Dp < 0.2 \mu m$  are not 53 affected by salinity, but for larger Dp's, salinity impact is substantial: higher salinity contributes to 54 higher production (Mårtensson et al., 2003). The SSA removal processes are scavenging by 55 precipitation and dry deposition (including gravitational settling). SSA has an effect on secondary 56 aerosols formed by gas-to-particulate conversion process such as condensation and nucleation 57 (binary homogeneous and heterogeneous) (Twomey, 1997). SSA serves as a sink for condensable 58 gases and smaller aerosol particles, and serves also as a medium for aqueous-phase reaction of





59 reactive gases, e.g. H<sub>2</sub>SO<sub>4</sub>. This can lead to nucleation suppression for other components of the 60 marine aerosol and consequently change their size distribution, creating a feedback on climate. 61 Furthermore, SSA formation results in a size spectrum ranging from 0.01 to 100 µm, which can 62 lead to cloud formation. With increasing concentrations of cloud condensation nuclei, the cloud microphysical properties change, i.e., the available water vapour is re-distributed over more 63 particles, on average resulting in smaller particle sizes, which in turn changes both cloud albedo and 64 65 precipitation (Latham et al., 2008, Lenton and Vaughan, 2009; Boyd, 2008; Korhonen et al., 2010, 66 Wang et al., 2011). With dry diameter lower than 1 µm, SSA can easily be transported for long 67 distances in the atmosphere, serving as a cloud seed outside of heavily clouded regions. The cloud drop number concentration can be spatially different, depending on the wind speed, atmospheric 68 69 transport and particle loss via dry and wet deposition (Korhonen et al., 2010).

70 Changes in atmospheric transport pathways, precipitation patterns, and sea ice cover influence 71 transport, removal and distribution of SSA. The main features of the regional and global SSA 72 distribution and the climate impact on SSA production due to these physical drivers have been 73 discussed in studies such as Liao et al. (2006), Pierce and Adams (2006), Manders et al. (2010), 74 Sofiev et al. (2011), Struthers et al. (2011), and Tsyro et al. (2011). The understanding of sea spray 75 emissions has increased substantially but process-based estimates of the total mass and size 76 distribution of emitted sea spray particles continue to have large uncertainties (de Leeuw et al., 77 2011). Chemical transport models (CTM) and general circulation models (GCM) estimates of sea 78 salt burden may vary over 2 orders of magnitude (Textor et al., 2006) and climate models disagree 79 about the balance of effects, ranging from little (Mahowald et al., 2006a) to a considerable 80 sensitivity to climate change (Bellouin et al., 2011). The difference between the available 81 estimations might be due to the wind speed predicted by the climate models, with little 82 understanding of how wind speed may change over the ocean in a warmer climate (IPCC, 2013).

The main goals of the current study are to assess the sensitivity of the production, surface concentrations and removal of SSA to climate change. A multi-model approach using four state-ofthe-art offline CTMs was taken to assess the uncertainty/robustness of model predictions over Europe. The sensitivity of simulated emission, concentration, and deposition of SSA to changes in climate was evaluated by comparing a past (1990-2009) and a future (2040-2049) period. This study is a follow-up to the climates studies of Langner et al. (2012) focusing on surface ozone and Simpson et al. (2014) focusing on nitrogen deposition.





# 91 2 Methods

92 This study uses the same modelling structure as in Langner et al. (2012) for ozone and in Simpson 93 et al (2014) for nitrogen. We focus on the comparison of SSA simulations from three offline 94 European-scale CTMs - EMEP MSC-W, MATCH and SILAM - and one offline hemispheric CTM, 95 DEHM. The models were run through a past (1990-2009) and a future (2040-2059) climate 96 scenarios and the results for the European seas (Baltic, North, Mediterranean, and Black Seas) were 97 compared. The climate meteorology data from a GCM were used in a regional climate model 98 (RCM) and the hemispheric model DEHM. The regional models where driven by the downscaled 99 meteorology from the RCM and the boundary conditions from DEHM. The horizontal grid for 100 DEHM is  $150 \times 150$  km<sup>2</sup> and for the regional CTMs identical to the RCM (ca.  $50 \times 50$  km<sup>2</sup>).

101 Throughout the paper, the SSA mass refers to the total mass of dry particles. Since the observations 102 measure sodium (Na<sup>+</sup>) concentrations rather than total SSA mass, it is assumed that Na<sup>+</sup> mass 103 fraction is ~30% (Seinfeld and Pandis, 2006). Particle sizes are also provided for dry conditions 104 and, unless otherwise stated, the dry diameter  $D_p$  ranges up to 10  $\mu$ m.

#### 105 2.1 Climate meteorology

106 Results of the global ECHAM5/MPIOM GCM (Roeckner et al., 2006), driven by emissions from 107 the SRES A1B scenario (Nakicenovic, 2000), were downscaled over Europe with the Rossby 108 Centre Regional Climate model, version 3 (RCA3) (Samuelsson et al., 2011; Kjellstrom et al., 109 2011). The global ECHAM5/MPIOM model is defined in spectral grid T63, which at mid-latitudes corresponds to a horizontal resolution of ca. 140×210 km<sup>2</sup>. The horizontal resolution of RCA3 was 110 111 0.44°×0.44° on a rotated latitude-longitude grid, and data were provided with 6-hourly resolution. 112 The climate, as downscaled by RCA3, reflects the broad features simulated by the parent GCM, but 113 from earlier studies with the current setup it is clear that the global ECHAM5/MPIOM model 114 projects a slightly warmer and wetter climate over Europe than the regional model RCA3 (Langner 115 et al., 2012; Simpson et al., 2014).

The wind speed is higher over the ocean and can be up to two times slower, in average, over the inner seas (Fig. 1, first panel on the left). Wind patterns are different between the Seas, with some areas over individual seas being more affected by wind gusts than others: e.g. in the Mediterranean, the wind speed is higher over the Levantine Sea than over other areas. For the wind speed, RCA3 predicts an stronger increase at the Norwegian Sea, Black Sea, Gulf of Bothnia (Baltic Sea) and Aegean Sea (Mediterranean Sea) and a stronger decrease between Italy and Tunisia and Libya





122 (Mediterranean Sea) in the future period (Fig. 1, first panel on the right). Nevertheless, the absolute

123 change is no more than 0.4 m/s. Trend analysis considering only marine grid cells for each sea (Fig.

124 S1 in supplementary material) shows that there is no significant trend between past and future

125 periods.

126 Typically, the surface water temperature is higher at southern latitudes. For the same latitude, the 127 Black and Mediterranean Seas have, in general, higher temperature than the Atlantic Ocean and the 128 Baltic Sea (Fig. 1, second panel on the left). RCA3 predicts a general increase of the water surface 129 temperature between the past and the future periods (Fig. 1, second panel on the left). The most 130 substantial changes are for the northern part of the Atlantic Ocean and for the Baltic Sea (maximum 131 1.17 °C). Trend analysis for the monthly mean temperature is significant for all the European inner-132 seas (Fig. S2 in supplementary material). The temperature is rising for all the seas with the highest 133 rise over the Black Sea and the lowest over the North Sea.

The precipitation tends to be higher over the ocean and lower over the inner seas. The lowest precipitation amount is seen over the Mediterranean Sea; on an annual level the difference from the ocean can be up-to two orders of magnitude (Fig. 1, third panel on the left). The climate model predicts that the precipitation will strongly decrease over the Mediterranean and increase over the Baltic and North Seas, whereas over different parts of Atlantic Ocean the opposite trends can coexist (Fig. 1, third panel on the right). Trend analysis shows that none of the trends is significant (Fig. S3 in supplementary material).

#### 141 **2.2 SSA boundary conditions**

Sea salt concentrations (as fine and coarse modes, see the description of DEHM below) provided by the hemispheric DEHM model, were used as lateral and top boundaries for the regional models. The boundary values taken from DEHM were updated every 6 h and interpolated from the DEHM grid to the respective geometry of each regional CTM. The DEHM model was driven by the global ECHAM5-r3 meteorology, without the RCA-3 downscaling.

#### 147 2.3 Chemical transport models

The models used in this study have been introduced in the previous studies: Langner et al. (2012) and Simpson et al. (2014). Below, we focus on their handling of the production and removal of SSA. All the SSA source functions in the current study are based on white-cap-area based parameterizations of Monahan et al. (1986), for formation of super-micron particles and follow





152 Mårtensson et al. (2003) for the sub-micron aerosols. The difference between the various source

153 functions is the dependence on temperature and salinity for the SSA generation (Table 1).

## 154 **2.3.1 DEHM**

155 In DEHM the production of SSA at the ocean surface is based on two parameterization schemes 156 describing the bubble-mediated sea spray production of smaller and larger aerosols. In each time 157 step the production is calculated for seven size bins and thereafter summed up to give an aggregated 158 production of fine (with dry diameters  $<1.3 \mu$ m) and coarse (with dry diameters ranging 1.3-6  $\mu$ m) 159 aerosols. For the fraction with dry diameters less than 1.25 µm a source function based on Mårtensson et al. (2003) is used, while for sizes larger than that the Monahan et al. (1986) source 160 function is applied. They both include an  $U_{10}^{3.41}$  dependency on wind speed and the production of 161 the smaller aerosols is also a function of the sea surface temperature. An ambient relative humidity 162 163 of 80% is assumed in the calculations and the size of the produced SSA is assumed to depend on the 164 salinity at the actual location. Here a monthly climatology of current day salinity on a 0.25°x0.25° 165 grid (Boyer et al., 2005) is applied for both time periods in focus in the current paper. Within the 166 atmosphere, the fine and coarse fraction of SSA is treated separately in terms of transport and 167 removal. Wet deposition includes in-cloud and below-cloud scavenging, while dry deposition 168 velocities are based on typical resistance methods for various land surface types (see Simpson et al., 169 2003; Emberson et al., 2000). The fine and coarse fractions in the DEHM model are in the current 170 paper assigned the dry diameters of 1 µm and 6 µm.

171 DEHM is continuously validated against available measurements from e.g. the EMEP network and 172 an evaluation of an earlier version of the sea salt routine in DEHM showed that the model gives

173 satisfactory results for sea salt over Europe (Brandt et al. 2012).

#### 174 2.3.2 EMEP MSC-W

175 The standard Unified EMEP model runs include sea salt particles with ambient diameters up to 176 about 10  $\mu$ m, which mainly originate from the bubble mediated sea spray (Tsyro et al, 2011). The 177 parameterisation scheme for calculating sea salt generation in the EMEP model makes use of two 178 source functions for bubble-mediated sea spray production. The first one is a source function for sea 179 spray droplets at 80% relative humidity from Monahan et al. (1986) and the second one is a source 180 function for sea salt particles from the work of Mårtensson et al. (2003), which is formulated for a 181 salinity of 33%. In the EMEP model, the SSA fluxes can be calculated for particle dry Dp ranging 182 from 0.02 to 12  $\mu$ m, whereas operationally and for this work SSA with Dp up to 6  $\mu$ m are included.





183 Mårtensson et al. (2003) parameterisation is applied for smaller size bins, while Monahan et al. 184 (1986) parameterisation is used for the coarser ones. From the fluxes of sea spray, the sea salt mass 185 is calculated assuming sea salt density of 2200 kg/m3. The total production rates of fine and coarse 186 sea salt are calculated by integrating the size resolved fluxes (7 in the fine and 3 in the coarse 187 fractions) over respective size intervals. In the model, generated SSA is assumed to be 188 instantaneously mixed within the model lowest layer at each time step. The transport and removal 189 of sea salt is described individually for the fine and coarse fractions in the EMEP model. Dry 190 deposition parameterisation for aerosols is calculated using a mass-conservative equation from 191 Venkatram and Pleim (1999). The dry deposition due to gravitational settling is size-dependent and 192 diameters of 0.33 and 4.8 µm are assumed for the fine and coarse SSA. . Wet scavenging is treated 193 with simple scavenging ratios, accounting for in-cloud and sub-cloud processes. The scavenging 194 ratios are assigned to crudely reflect the solubility of different aerosol components, and the size 195 differentiated collection efficiencies are used in sub-cloud aerosol washout.

The present sea salt parameterisation was shown to give the best overall results as compared to a number of other source functions within the EMEP model (Tsyro et al., 2011). The model SSA calculations are extensively evaluated against long-term observations (Tsyro et al., 2011; EMEP Reports http://www.emep.int).

#### 200 2.3.3 MATCH

201 The treatment of SSA production in MATCH is based on the parameterization of Mårtensson et al. 202 (2003) for dry particle sizes of up to  $0.4 \,\mu m$  aerodynamic radius, and on Monahan et al. (1986) for 203 larger particle sizes. The temperature correction following Sofiev et al. (2011) is applied to the 204 estimates from the Monahan scheme. The number of bins is flexible, but in this study four size bins 205 were used with Dp ranges 0.02–0.1  $\mu$ m, 0.1–1  $\mu$ m, 1–2.5  $\mu$ m, 2.5–10  $\mu$ m. The production of sea 206 salt droplets is calculated assuming an ambient relative humidity of 80% and a particle density of 207 1150 kg/m3 and is integrated over each size bin while dry removal rates are calculated using the 208 geometric mean size in each bin. Dry deposition over land is following Zhang et al. (2001) while a 209 separate parameterization accounting for bubble burst activity is used over sea (Pryor and 210 Barthelmie, 2000). Sea salt is assumed to 100% activated or scavenged by hydrometeors in-cloud 211 while below-cloud scavenging is handled following Dana and Hales (1976). The distribution of 212 salinity in sea water is taken from NOAA (2013). Further details and evaluation of MATCH sea salt 213 simulations using observed meteorology can be found in Foltescu et al. (2005) and Andersson et al. 214 (2014).





## 215 2.3.4 SILAM

216 The SSA production via bubble-mediated mechanism takes into account the effects of wind speed, 217 salinity, and water temperature and covers sea salt particles with dry diameter from 20 nm to 10 218  $\mu$ m. The observations from the Mårtensson et al. (2003) study for seawater surface temperature 298 219 K and sea water salinity 33 ‰ were used to extrapolate the scheme from Monahan et al. (1986) to 220 particle sizes down to 20 nm. To calculate SSA production for other water temperatures and 221 salinities, correction factors are applied which were derived based on the experimental data of 222 Mårtensson et al. (2003). The full description of the parameterisation in the SILAM model can be 223 found in Sofiev et al. (2011). The description of the temperature correction in Sofiev et al. (2011) 224 was changed. Currently, the water temperature reference for the unified shape function is 20 °C, 225 instead of 25 °C as referred in Sofiev et al. (2011). The shape function has been updated accordingly 226 and the new shape function  $(dF_0/dD_p)$  for particles with Dp ranging from 0.01 to 10 µm is described 227 below:

228 
$$\frac{dF_0}{dD_p} = (1+0.05*D_p)*\frac{\exp\left(\frac{-0.11}{D_p}\right)}{0.4+\exp\left(\frac{-0.2}{D_p}\right)}*\frac{6*10^5}{\left(1*10^{-4}*D_p^2+D_p\right)^3}*10^{1.19^8\exp\left(-\left(\frac{0.35-\log D_p}{0.8}\right)^2\right)}$$
(1)

229 For the current study the spume droplet formation based on Andreas (1998) was included, with 230 spume being supressed for 10m wind speed lower than 6 m/s. The production of sea salt droplets is 231 calculated assuming a dry particle density of 2200 kg/m<sup>3</sup>. The size distribution is described by 232 flexible bins. Production is integrated over each size bin while dry and wet removal rates are 233 calculated using mass-weighted mean diameter in each bin. Depending on particle size, mechanisms 234 of dry deposition vary from primarily turbulent diffusion driven removal of fine aerosols to 235 primarily gravitational settling of coarse particles (Kouznetsov and Sofiev, 2012). Wet deposition 236 distinguishes between sub- and in-cloud scavenging by both rain and snow (Sofiev et al., 2006; 237 Horn et al., 1987; Smith and Clark, 1989; Jylhä, 1991). Gravitational settling, dry deposition and 238 optical properties take into account the particle hygroscopic growth. For the simulations, five bins 239 with the Dp ranges of 0.01–0.1  $\mu$ m, 0.1–1.5  $\mu$ m, 1.5–6  $\mu$ m, 6–15  $\mu$ m; and 15-30  $\mu$ m were used. The 240 distribution of salinity in sea water is taken from NOAA (2013).

SILAM model has been evaluated against a wide range of observations and models utilizing the above described parameterization (Sofiev et al., 2011; Tsyro et al., 2011).





#### 243 2.4 Model evaluation

Sea water is the predominant source of Na<sup>+</sup> in the atmosphere, which can be used as its tracer in 244 245 most regions of Europe. Evaluation of the model predictions was performed via comparison with 246 observations available from the EMEP network (Co-operative Programme for monitoring and 247 evaluation of the long-range transmission of air pollutants in Europe, http://www.emep.int, Tørseth 248 et al. 2012) that perform regular measurements across Europe. The observations include Na<sup>+</sup> 249 concentration in aerosol and ion analysis of precipitation including Na<sup>+</sup>. Concentration 250 measurements are sampled daily by a filter pack sampler (cut-off at  $D_p = -10 \mu m$ ), at 2 m height; 251 the concentration in precipitation is mainly sampled by a "wet-only" sampler and, in a few places, 252 with bulk collectors. The wet deposition of Na<sup>+</sup> is obtained by multiplying the weighted mean 253 concentration by the total amount of precipitation in a daily basis. For more details about the 254 sampling the reader is referred to e.g., Hjellbrekke and Fjæra (2009). These sampling methods do 255 not distinguish if the sodium is originated from natural (e.g. mineral dust) or anthropogenic sources. 256 In some regions there might be certain amounts coming from combustion processes and industry, 257 but overall the contribution of anthropogenic sources to the sodium budget is low (van Loon et al., 258 2005).

259 The measurement data were averaged to monthly level with the minimum completeness 260 requirement of 75% temporal coverage per month and per year, between 1990 and 2009. The CTMs 261 predictions for the measurement sites satisfying the temporal criterion were averaged on a monthly 262 basis over the 20 years. Since the model computations were driven by climate model fields, no 263 temporal collocation was done. Therefore, the primary parameter considered was the monthly Na<sup>+</sup> 264 concentrations averaged over the past period. Modelled values were obtained from the model's 265 lowest layer mid-point, which is defined somewhat differently for each model (Table 1). No near-266 surface concentration profiling was made, with the exception of EMEP where concentrations are 267 corrected to 3 m height, largely due to unreliable stability estimates based on climate-model fields.

The model performance was evaluated by the following statistical measures: bias, spatial Pearson correlation coefficient (R), root mean square error (RMSE), bias and standard deviation (SD) ratio ( $SD_{model}/SD_{observations}$ ). The evaluation included Na<sup>+</sup> concentration in aerosols at 29 measurement sites and ion analysis of Na<sup>+</sup> wet deposition at 133 measurement sites, which we consider sufficient for computing the basic statistical scores and plotting scatter plots. The location of the measurement sites are shown in Fig. S4 in the supplementary material.





#### 274 2.5 Radiative transfer modelling

275 The radiative transfer modelling was completed offline with the libRadtran software package for 276 radiative transfer calculations (Mayer and Kylling, 2005). This tool calculates radiances, irradiances 277 and actinic fluxes for the given optical properties. The Earth radiative balance results from the 278 difference between the incoming (direct and diffusive-downwards) and outgoing (diffusive 279 upwards) radiation. The impact of SSA is assessed by the difference between an atmosphere with 280 SSA and without SSA, for the past and future periods. The calculations were defined at the top of 281 the atmosphere (TOA), with wavelength ranging from 0.2 to  $\sim 4 \mu m$ , in order to compute the 282 integrated shortwave irradiance. All the runs considered wet and icy clouds, with the cloud cover 283 taken from the climate model RCA3 and optical properties taken from MODIS observations (Pincus 284 et al. 2011). Monthly-basis observations from AOUA and TERRA obtained from 2002 to 2014 285 were averaged in order to have climatological cloud optical fields. These fields were the same for 286 both past and future period calculations. Earth albedo information is included in the calculations 287 and is obtained from the NASA model, GLDAS Noah Land Surface Model L4 (Rodell et al., 2004), 288 on a monthly basis for the period between 1990 and 2012. This dataset was averaged to obtain 289 climatological surface albedo fields, remaining the same for both past and future periods. The 290 calculations for an atmosphere with SSA included the AOD computed by SILAM: the AOD at 550 291 nm was computed for the full size-spectrum of the SSA described in Table 1. SILAM's optical 292 thickness predictions are based on size distribution and spectral refractive index of SSA (Prank, 293 2008). The AOD data was monthly-averaged for every hour in a day, for the past and future 294 periods. This allowed taking into consideration the length of the day, since solar zenith angle is 295 computed for every hour. The description of the runs and assumptions are provided in Table 2. This 296 setting was chosen in order to reflect an atmospheric state closer to reality, since there were no other 297 aerosols available for this study. Keeping the atmospheric and cloud conditions constant between 298 the past and the future, will allow pinpointing the impact of the SSA on the radiative balance.

299

#### 300 3 Results

#### 301 **3.1 Comparison with observations**

Figure 2 and Figure 3 show the performance of the CTMs estimating Na<sup>+</sup> surface concentrations and wet deposition, respectively, during the past period; Table 3 and Table 4 complete the statistical evaluation of the models for the surface concentrations and wet deposition, respectively. The





models showed similar performance with quite high correlation coefficients varying from 0.71 up to 305 306 0.85 for the concentrations but substantially lower for wet deposition (from 0.24 up to 0.41). The 307 difference between the model performances is quite small and varying for the different scores. The highest correlation with the concentration observations was shown by DEHM (0.85), which also 308 309 demonstrated the highest RMSE and bias originating from a stronger overestimation over the 310 regions with observed low concentrations. EMEP showed the lowest RMSE and bias, as well as one 311 of the best correlation factors. SILAM tends to overestimate the lowest observed values (positive 312 bias) whereas MATCH has a stronger underestimation of the highest values (negative bias). 313 Comparing the winter (December, January and February) and the summer (June, July and August) 314 seasons, one can notice that the models perform better in summer, with higher correlation and lower 315 bias. The observed winter time levels are likely harder to be reproduced due to stronger winds and 316 faster changing weather, which might not be captured by the climatological runs.

317 Comparison of Na<sup>+</sup> wet deposition with measurements shows low correlation and substantial under-318 prediction. This is particularly true for the high-deposition observations, which resulted in a strong 319 negative bias for all the models. The evaluation of modelled precipitation was presented in Simpson 320 et al. (2014), Table 4, and shows an overestimation of precipitation in the RCA3 model (reginal 321 CTMs) and underestimation in the precipitation used in DEHM. The overestimation leads to an 322 overestimation of the deposition of SSA close to the sources. Consequently, less SSA reaches the 323 shore and the measurement sites. The second major reason for discrepancy is that the observed wet 324 deposition does not cut-off the size of the particles, i.e. SSA coarser than 10 µm is accounted for, 325 including the SSA produced in the surf zone. This mostly explains the large negative bias of the 326 models, which reported PM<sub>10</sub> only, and, to some extent, the low correlation. This is demonstrated 327 when comparing SILAM scores taking into account the full size range available (Dp = [0.01-30]328  $\mu$ m): accounting for the coarser aerosols strongly reduced the bias, correlation strongly improved, 329 and RMSE became slightly smaller. In summer, the scores are slightly better than in winter, but the 330 absolute values and importance of this removal process is smaller in summer time.

In Simpson et al. (2014), it was shown that CTMs driven by RCM meteorology are likely to perform worse than they would with data from numerical weather prediction models. Nevertheless, the current comparison showed that CTMs can predict mean concentrations and depositions within ~30% uncertainty (for depositions, prediction of full size range is a pre-requisite), whereas the spatial distribution patterns are reproduced with correlation higher than 0.7 also when driven by climate model meteorology.





# 337 3.2 Current and future climate SSA emissions

338 The annual SSA emission in the reference period predicted by DEHM, MATCH and SILAM is 339 shown in Figure 4 (left panel). EMEP did not have this variable as an output. As expected, all 340 models predict the highest emissions over the Atlantic Ocean, with the Mediterranean Sea being the 341 second highest source. MATCH predicted, in average, 25% higher emissions over the 342 Mediterranean than SILAM. The emissions are mainly driven by the wind and typically expressed 343 by the white-cap produced by the surface-winds via the Monahan and O'Muircheartaigh (1980) 344 parameterisation. This empirical power-law is taken by all models participating in this study and 345 suggests emission (E) to be proportional to the 10m-wind speed (U<sub>10</sub>) to the power of 3.41: E  $\approx$ 346  $U_{10}^{3.41}$ , the so-called wind-forcing. Consequently, the SSA emissions (Fig. 4, left panel) clearly 347 correlate with the wind-forcing (Figure 5, left panel), in particular over the open ocean. However, 348 the use of the same functional dependence and input meteorology does not guarantee identical 349 emission, as it will be discussed further on. MATCH and SILAM seem more sensitive to the wind-350 forcing over the Mediterranean than DEHM, possibly due to the horizontal resolution difference 351 between the hemispheric and regional CTMs (e.g. the Mediterranean is not properly resolved by the 352 global climate model, the driver for DEHM). Apart from the wind forcing, laboratory studies have 353 shown the relation between the emissions of SSA and seawater surface temperature and salinity: 354 SSA mass will be higher at sea areas with higher surface water temperatures and salinity 355 (Mårtensson et al, 2003). The temperature and salinity dependencies are included in the 356 parameterizations, therefore, the models predict for the same wind forcing, higher emissions for 357 higher water temperatures: the Mediterranean and Black Seas (Fig. 1 and Fig. 4, left panel). The 358 effect of salinity is best seen in the Baltic Sea (salinity  $\sim 9$  ‰), which has comparable wind forcing 359 to some areas of the Mediterranean and the Atlantic (salinity ~33 ‰) but lower emission. SILAM 360 and MATCH show the highest difference between the inner-seas with at least 3 times lower 361 emissions over the Baltic Sea.

In absolute terms, the climate impact on SSA emissions (Fig. 4, right panel) is mainly positive according to the regional models whereas DEHM shows a general decrease. The exception goes for the Atlantic Ocean, in the west side of the domain, where all the models agree in a decrease of emissions. The difference between the past and future periods is only due to the wind forcing and temperature changes, since salinity was kept constant. Thus, this change (Fig. 4, right panel) highly correlates with the changes for wind-forcing (Fig. 5, right panel), adjusted by the changes in water temperature (Fig. 1, right panel). For example, the pronounced decrease of emission over western





369 Atlantic is mainly driven by the reduction of wind speed but the decrease is limited by the rising 370 temperature in the north and east: higher temperature leads to production of more SSA even for 371 somewhat slower wind speed.

372 The models demonstrated different sensitivity to seawater temperature: it seems to be less important 373 for DEHM than for other models, whereas SILAM is the most sensitive. For instance, MATCH and 374 SILAM showed an increase of emissions over the east of Iceland where temperature is predicted to 375 rise by almost 2 K. The increase of seawater temperature, supported by higher wind speed, over the 376 Black and Aegean Seas (Fig. 1, right panel), will lead to higher emissions. DEHM might not be so 377 sensitive to the local storms due to the coarse horizontal resolution. The absolute difference 378 between future and past is the smallest for the Baltic Sea, but in relative terms all the models show 379 an increase up to 20% in Gulf of Bothnia, which is actually higher than, e.g. 5-15% of increase 380 predicted for North Sea (minimum for DEHM and maximum for MATCH).

Trend analysis for the Baltic, Black, Mediterranean and North Seas (only sea cells are taken into consideration) is available as supplementary material: Fig. S5 for the Baltic, Fig. S6 for the Black, Fig. S7 for the Mediterranean, and Fig. S8 for the North Seas. The trend is only statistically significant (p < 0.001) for all the models for the Black Sea, with all models agreeing on an increase of concentration in the future.

386 Figure 6 (left panel) shows the SSA emission difference between the winter and summer for the 387 past period. The difference between seasons in terms of SSA production can be substantial: SSA 388 emission is up to 3 times higher in winter time. Seasonally, there are differences between the 389 driving processes for SSA production: the winter period has a larger SSA production, due to more 390 frequent and stronger storms; but the summer time shows pronounced maxima over specific areas 391 mostly influenced by the seawater temperature. The latter is mostly true for MATCH and SILAM, 392 since their temperature sensitivity is higher. SSA emission in winter will be accentuated in the 393 future for MATCH (more emphasized) and SILAM: Figure 6 (right panel) shows pronounced 394 maxima around Iceland and the British Isles; distinct differences in the SSA emission are also seen 395 in the Mediterranean. DEHM does not show much difference between the periods.

#### **396 3.3 Current and future climate SSA concentrations**

Concentration is a function of emission and transport of the SSA, that is dependent on ventilation of an area over inner seas (wind speed), and on removal processes largely controlled by precipitation and relative humidity (via settling). Generally, the pattern of SSA concentration follows the





400 emission areas with stronger winds and frequent storms. Concentrations are, therefore, higher at the 401 Atlantic Ocean and lower at the European inner-seas. All the models show lower concentrations for 402 the Baltic Sea, reaching up to 10 times difference from the ocean (Fig. 7, left panel). The 403 Mediterranean Sea is the inner sea with the highest concentrations. For the Baltic Sea, DEHM and 404 MATCH show the highest and the lowest concentrations, respectively, with a difference of a factor 405 of ~1.3 between each other. For the Black Sea, DEHM and EMEP show the highest concentrations 406 and a similar spatial distribution pattern, and SILAM the lowest; nonetheless the difference is not so 407 substantial. For the Mediterranean Sea, EMEP shows the lowest concentrations, MATCH being the 408 highest: with 30% difference. All models show pronounced maximums at the Balearic Sea and the 409 Levantine Sea. Transport over land is quite similar among the models, especially for the regional 410 CTMs. The biggest difference lies over the western-central Europe with MATCH showing lower 411 concentration over land. Transport of SSA inland is visible hundreds of km's inland; near the coast 412 line it can contribute up to 6  $\mu$ g/m<sup>3</sup> to PM<sub>10</sub>.

413 The models predict relatively similar pattern for the SSA spatial distribution for the past period but, 414 they seem to have different responses to the future climate, with MATCH and SILAM clearly being 415 the most sensitive and EMEP the least. Figure 7 (right panel) shows the difference between the past 416 and future periods for the different models. DEHM and EMEP foresee almost no change or a 417 decrease of SSA concentrations over the open sea, whereas MATCH and SILAM predict an 418 increase. These results were expected due to the predicted emissions (Sect. 3.2). All models agree in 419 an increase in SSA surface concentration over the north of Iceland, the Black Sea, and over land in 420 southern latitudes. The models agree somehow on an increase of the Mediterranean and Black Seas 421 SSA concentration but it is MATCH and SILAM that show the highest positive change in 422 concentrations. The impact over land is slightly positive for all the models in the Southern part of 423 the domain, while at more Northern latitudes DEHM and EMEP from one side, and MATCH and 424 SILAM models from another, disagree on the trend signal: a reduction of the SSA load over land is 425 predicted by the first two models and an increase by the latter pair.

Overall, EMEP is the least sensitive and MATCH the most sensitive model to a changing climate. SILAM is the most sensitive over the Norwegian Sea. The difference between the past and future period concentrations is more substantial than that of emissions: the factors seemingly having exacerbated this difference are the decrease of ventilation over the west-Mediterranean, changes in mixing patterns, etc.





Trend analysis (supplementary material: Fig. S9 for the Baltic, Fig. S10 for the Black, Fig. S11 for the Mediterranean, and Fig. S12 for the North Seas) suggest that trends are only significant (p < 0.001) for MATCH and S12 for the Mediterranean and Fig. S12 for the North Seas) suggest that trends are only significant (p < 0.001) for MATCH.

433 0.001) for MATCH and SILAM for both Mediterranean and Black seas, all with a positive signal.

434 Seasonally, the concentrations follow the same pattern as the emissions: higher in winter time. 435 When analysing the changes between winter and summer, the models can again be grouped into 436 DEHM-EMEP and MATCH-SILAM. In winter (Fig. 8, left panel), the first pair presents a larger 437 amount of SSA mass generally over sea and land surfaces. Conversely, MATCH and SILAM 438 predict a decrease of SSA surface concentration around the British Isles, Mediterranean and Black 439 Seas, though the coast lines have sharper peaks of SSA mass during winter. The difference between 440 the future and past periods (Fig. 8, right panel) is relatively similar for all the models over the open 441 sea: predictions show an increase of concentration around the British Isles and a decrease over the 442 Norwegian Sea, in the future. MATCH and SILAM show sharper increase or decrease along the 443 Mediterranean Sea. The changes predicted can be 3 times higher than the changes predicted for the 444 emissions (Figure 6, right panel). The changes can also have different signal, e.g. the Eastern-basin 445 of the Mediterranean where it is predicted an increase of emissions but a decrease of concentrations, 446 implicating that the ventilation over this area was quite effective.

#### 447 **3.4** Current and future climate SSA deposition

448 The deposition (wet+dry) patterns for SSA are depicted in Fig. 9 (left panel). Typically the 449 deposition is higher over the sources areas and close to the coastal areas. Over land, SILAM shows 450 less deposition and DEHM and EMEP predict the highest levels. There are different patterns over 451 the Atlantic, mostly attributable to the boundary conditions treatment by each model. DEHM 452 predicts quite high values over all the seas. Over the Black Sea, the deposition is more accentuated 453 in the predictions by EMEP and less by SILAM. MATCH also shows higher values for deposition 454 over the Mediterranean, and SILAM the lowest. Deposition is not substantial over the Baltic Sea, 455 with exception of DEHM, owing to low SSA mass released from its surface.

The impact of future climate conditions (Fig. 9, right panel) on deposition, in absolute levels, is small and mostly noticeable over the Atlantic Ocean. For all models, the most significant positive change in the deposition is seen around Iceland. This is expected according to the changes seen in precipitation between future and past periods (Fig. 1, third panel on the right). All regional CTMs show a strong signal on the west side of the domain, an artefact due to the boundary conditions. In





relative terms, Scandinavia, east of UK, central-western Europe and Mediterranean are the most
affected with 5-20% more deposition predicted by MATCH and SILAM.

463 Trend analysis (supplementary material: Fig. S13 for the Baltic, Fig. S14 for the Black, Fig. S15 for

- the Mediterranean, and Fig. S16 for the North Seas) suggests that none of models show a significanttrend.
- Seasonally, SSA deposition is higher in winter than in summer, due to the higher emissions and 466 467 frequent precipitation in winter months. This difference is mainly accentuated over the source areas: 468 MATCH and SILAM have the lowest difference over the Baltic and Black Seas, due to the lower 469 production; DEHM shows the highest at Mediterranean Sea. The difference of deposition between 470 winter and summer will also change in the future period (Fig. 10, right panel) with all models 471 showing a slight increase of the deposition in summer over the Mediterranean and along the coast of 472 Norway. An increase of deposition in winter was suggested around Iceland and British Isles, North 473 Sea and coastal areas of Mediterranean Sea.
- 474

#### 475 4 Impact of meteorology and seawater properties on the emission and fate of SSA

476 The multi-model comparison presented in Sect. 3 shows that there are significant difference 477 between the models in terms of emission and fate of the SSA. The latter is particularly true for the 478 inner seas. The differences between the models lead to a more uncertain answer about the impact of 479 the future climate on the production and transport of SSA and its possible feedback to climate. The 480 SSA emission in the models is driven by three parameters: wind speed, water temperature, and water salinity. All models use the same  $U_{10}^{3.41}$  dependence on wind speed; hence the differences in 481 482 emission have to be attributed to parameterization of temperature and salinity dependencies. 483 Formally, all models used the Monahan et al. (1986) and Mårtensson et al. (2003) parameterizations 484 or, at least, the available data for deriving the emission flux parametrizations (SILAM). Specifics of 485 the implementation, however, appeared significant. To understand the latter, box-model calculations 486 of the SSA mass flux as a function of temperature were made for seawater salinity 10 and 35 ‰, 487 representing Baltic Sea and Atlantic Ocean, respectively, and with wind-speed fixed at 15 m/s (Fig. 488 11, left-hand panel).

In general, all the models show an increase of mass flux of SSA with temperature and salinity, except EMEP that does not apply any correction for salinity. Both DEHM and EMEP mass flux show little difference between low and high temperatures; SILAM and MATCH show a substantial





492 dependency of the mass flux on temperature throughout the size ranges. This difference is explained 493 by the way dependency on seawater temperature is implemented: only for the fine mode in DEHM 494 and EMEP, based on the Mårtensson et al. (2003) source function, and for both fine and coarse 495 modes in SILAM and MATCH. In MATCH, the implementation of seawater temperature correction 496 is done by combining the temperature correction included in the Mårtensson et al. (2003) for size-497 range below  $Dp = 0.4 \ \mu m$  and the use of the temperature corrections from Sofiev et al. (2011) for 498 the coarser sizes. In SILAM the source function is scaled with Sofiev et al. (2011) size-dependent 499 temperature correction function. This explains why the results in Sect. 3 could be paired between 500 the models. EMEP is the model that shows the highest amount of SSA produced, with the exception 501 for seawater temperature higher than 15 °C and high salinity, with MATCH and SILAM predicting 502 the highest amount of SSA. For the lowest salinity, SILAM is the model that produces less SSA, 503 with DEHM being surpassed by MATCH around 17 °C. For the highest salinity, both MATCH and 504 SILAM start to predict higher SSA flux than DEHM around 9 °C. This is due to the temperature 505 correction factor described in Sofiev et al. (2011) that assumes that for low seawater temperature, 506 the production of coarse SSA, where the mass is significant, is very low. This analysis clarifies why 507 MATCH and SILAM tend to have higher emissions than DEHM where waters are warmer and 508 lower when colder (e.g. Baltic Sea), and why MATCH shows the highest values for the SSA mass 509 flux. Also explains the smaller difference between winter and summer predicted by DEHM, since 510 the changes in SSA mass flux depending on seawater temperature is very low.

511 Figure 11 (right panel) shows how the different models distribute the mass between the fine and 512 coarse modes, for the same wind and salinity conditions described above. Both DEHM and EMEP 513 assume that the contribution of the coarser mode is reduced with temperature, since more SSA is 514 produced with higher temperatures, for size ranges below 2.5 µm. EMEP has the highest 515 contribution for the coarse mode, independent of the temperature. For MATCH and SILAM, the 516 contribution to the coarser mode increases with temperature, though MATCH has a lower coarse 517 mode contribution than SILAM. The only agreement between the DEHM, MATCH and SILAM is 518 that for higher salinities, the coarse mode contribution is higher. The ratio between fine and coarse 519 mode is very relevant for the deposition processes, and it could explain why deposition is higher for 520 DEHM and EMEP (Fig. 9), though in this case, it is hard to evaluate the real impact due to different 521 deposition schemes implemented in the models.

It is pertinent to discuss the difference between DEHM, EMEP and MATCH, since these modelsapply the same parameterization for SSA number flux, though having different salinity fields and





salinity correction function. Mårtensson et al. (2003) defines very strict size ranges for the computation of the 6<sup>th</sup> order polynomial for particles between 0.02 to 2.8  $\mu$ m in dry diameter. In case the models define size ranges outside of the tabulated in that study, it can result in very different results. The linkage between the two parameterizations can also result in different outcomes: DEHM links the two parameterizations at dry diameter of 1.25  $\mu$ m, EMEP at 1.5  $\mu$ m and MATCH at 0.4  $\mu$ m. In the case of MATCH, an extrapolation of the Monahan et al. (1996) function is needed, in order to bring it to Mårtensson et al. (2003) range.

531

#### 532 5 SSA and climate change: production, fate and radiative impact

533 The regional-scale impact of SSA production and fate caused by a changing climate has been 534 shown in Sect. 3. We show that the change in SSA emission between the past and future periods is 535 not so large, arguably due to the small change in wind speed between the two time periods. 536 Climates studies such as Gregow et al. (2011) projected higher wind speed changes in periods 537 closer to the years 2100, in Scandinavia. Nevertheless, the available climate estimations of wind can 538 differ substantially given the little understanding of how wind speed may change over the ocean in a 539 warmer climate (IPCC, 2013). Studies such as Salisbury et al. (2013) suggest that other variables, in 540 addition to wind forcing, influence the whitecap fraction, such as the seawater temperature or the 541 sea state. New parameterization for whitecap fraction, based on satellite observations, claims that 542 the whitecap-area based parameterization used by all the models in this study is misrepresenting the absolute values. Albert et al. (2015) suggests that for higher latitudes the values are overestimated, 543 544 and underestimated for lower latitudes. If following that parameterization, the emission over the 545 Mediterranean is underestimated. This could mean that the changes in seawater temperature would 546 impact the SSA emission flux more substantially than suggested by this study.

547 The aerosol direct radiative effect (DRE) is defined as the difference between net radiative fluxes at 548 TOA in the presence and absence of SSA. The radiative forcing depends on the AOD of the aerosol 549 species in the atmosphere, the surface albedo and the vertical position of clouds. In this study, all-550 sky conditions were considered, i.e. clouds are included. Over the seawater surfaces, SSA directly 551 scatters solar radiation back to space, resulting in a cooling effect on the climate by decreasing the 552 amount of radiation absorbed by the water surface. Over land, there can be both cooling over the 553 low-reflectance surfaces, and warming over high-albedo surfaces (e.g., Haywood and Boucher, 554 2000). Adding only a low absorbing aerosol, such as SSA, and assuming the same atmospheric and





cloud conditions for the all the runs (with and without SSA), the upward scattering by SSA will bethe only radiation impact in this study.

557 Figure 12 shows the DRE due to SSA in the past (left panel) and the change in DRE due to the 558 changing climate (right panel). These calculations are based on the AOD predicted by SILAM for 559 the past and future. As expected, the past computations predict the highest cooling effect due to 560 SSA over the areas where concentrations (Fig. 7, left-lower panel) are the highest and where the 561 surface albedo is the lowest (seawater surfaces). The strongest effect is seen over the Mediterranean 562 Sea due to the lowest cloud cover and the largest number of hours of sunlight per year. Studies such 563 as Ma et al. (2008) and Lundgren et al. (2013), state that the impact of clouds can be substantial, 564 reducing the direct radiative impact of SSA. The lowest cooling effect is predicted over land where 565 the albedo is higher and SSA amount is the lowest. Conversely, warming is predicted where the albedo is high and the AOD is low, e.g. over the mountain tops in Norway and Italy. The current 566 study estimates the upward scattering by SSA, at TOA, to be up to 0.5 W m<sup>-2</sup> over seawater 567 568 surfaces. This value is within the estimates on upward scattering of radiation by SSA: ranging between 0.08 and 6 W m<sup>-2</sup>, at wavelengths in the range of 0.3-4 µm (Lewis and Schwartz, 2004). 569

570 Figure2, right panel, depicts the change in the DRE due to SSA between future and past. The results suggest overall cooling (negative change) in the future: North of Iceland, Norwegian and North 571 572 Seas are the areas where the cooling is more accentuated. The Mediterranean area seems to be again 573 the most sensitive area in our study: it is predicted an overall warming for this area, both over sea 574 and over land, but also cooling, in particular in the east of the eastern basin. DRE pattern for the 575 whole year is highly influenced by the summer period due to largest number of daylight hours. This 576 can be seen in Fig. 13, right panel, which shows the change between future and past but considering 577 only the summer months (JJA). This study predicts a substantial seasonal variation for the DRE in 578 the sea surface waters. This is expected due to the variation shown in Sect. 3.2 and 3.3. The upward scattering in the summer time can be up to 1.7 times higher than in winter, due to lower cloudiness 579 580 and lengthier daylight.

Figure 13 shows the change in winter (left panel) and summer (right) between the future and the past. The strongest impact in winter is seen over the Mediterranean area: negative over the sea surface and positive over land. In summer, the highest impact is over the seawater surfaces, predicting a cooling effect in the future, with exception over the western basin of the Mediterranean and the western side of the British Isles and France.





The results presented in this study for the present period are in accordance with the regional simulations for a summer month presented by Lundgren et al (2013) and the global simulations presented by, e.g. Grini et al. (2002) and Ma et al. (2008). The results are shown in Table 5.

589 The radiative forcing calculation is also sensitive to the SSA single scattering albedo. Thus, setting

590 the SSA's single scattering albedo as low as 0.95 (Russel et al, 2002), leads to a wide areas over

591 land where warming is substantial: essentially, over all surfaces with albedos exceeding 0.5 and low

592 (<0.03) aerosol load (not shown). We have chosen to show results for a more realistic SSA single

- scattering albedo of 0.99 (Lundgren et al., 2013).
- 594

#### 595 6 Conclusion

596 This study has compared predictions of SSA emissions, surface concentration and deposition from 597 four CTMs for both current condition and future scenarios, focusing on the European Seas: Baltic, 598 North, Mediterranean, and Black Seas. The three European-scale CTMs (EMEP, MATCH and 599 SILAM) were driven by the regional climate model (RCA3) meteorology and by the hemispheric 600 model (DEHM) boundary conditions. The hemispheric model was driven by the ECHMA5 601 meteorology. The impact of climate change on SSA production and fate, due to changes in wind 602 speed and seawater temperature, was analysed. Additionally, consideration about the impact of 603 seawater salinity on emissions was given.

604 The impact of climate change on SSA production and fate has different response from the models, 605 with the similar results between DEHM and EMEP, and between MATCH and SILAM. DEHM-606 EMEP show almost no difference between future and past periods, and MATCH-SILAM shows a 607 general increase of the emissions and surface concentrations with levels reaching 30% in change. The emissions increase is substantial in the Black Sea, Gulf of Bothnia (Baltic) and Levantine Sea 608 (Mediterranean), correlating well with the wind-forcing ( $\approx U_{10}^{3.41}$ ) computed with the changes 609 610 predicted between the same periods. Nevertheless, the major driver of the changes of the sea-salt 611 fluxes from the sea surface will be the changing seawater temperature, since near-surface wind 612 speed is projected to stay nearly the same in the climate scenario used, in absolute levels the wind 613 will change less than a meter per second, in average, between the two periods. The concentrations 614 are predominantly increasing in Black and Mediterranean Sea. The impact of climate change on 615 SSA on deposition is not really relevant; though an increase is projected around Iceland by all the 616 models. Boundary conditions impact on the predictions is substantial.





The discrepancies between the models raised additional question about the implementation of the SSA production formulations, since three of the models are based on the same parameterizations. This study shows that the way a given parameterizations is implemented in the models and the temperature and salinity correction functions play an important role for the final scaling of the SSA flux: size range prescription may play a substantial role on the SSA flux calculation.

622 Simple calculations with the libRadTran allowed understanding the impact of SSA on the direct 623 radiative forcing. According to this study the upward scattering by SSA, at TOA, can to be up to 0.5 624 W m<sup>-2</sup> over the seawater surfaces in the present period, predicting an overall cooling in the future. 625 The most affected areas by cooling will be North of Iceland, Norwegian and North Seas, and the 626 eastern basin of the Mediterranean; warming is predicted manly in Mediterranean Sea, including 627 over land.

628

#### 629 Acknowledgements

630 This study was supported by the Nordic Council of Ministers, EnsCLIM and CarboNord projects.

- 631 The authors also thank Antti Arola for his guidance in the radiative forcing calculations and
- 632 interpretation of the results.

633

#### 634 References

- Albert, M. F. M. A., Anguelova, M. D., Manders, A. M. M., Schaap, M., and de Leeuw, G.:
  Parameterization of oceanic whitecap fraction based on satellite observations, Atmos. Chem.
  Phys. Discuss., 15, 21219-21269, doi:10.5194/acpd-15-21219-2015, 2015.
- 638 Andersson, C., Bergström, R., Bennet, C., Robertson, L., Thomas, M., Korhonen, H., Lehtinen, K.
- E. J., and Kokkola, H.: MATCH-SALSA Multi-scale Atmospheric Transport and CHemistry
   model coupled to the SALSA aerosol microphysics model Part 1: Model description and
- 641 evaluation, Geosci. Model Dev., 8, 171-189, doi:10.5194/gmd-8-171-2015, 2015.
- Andreas, E. L.: A new sea spray generation function for wind speeds up to 32 m s-1, J. Phys.
  Oceanogr., 28, 2175–2184, doi:10.1175/1520-0485(1998)028<2175:ANSSGF>2.0.CO;2., 1998
- Andreas, E. L., Edson, J. B., Monahan, E. C., Rouault, M. P., and Smith, S. D.: The spray
  contribution to net evaporation from the sea: A review of recent progress, Boundary Layer
  Meteorol., 72(1–2), 3–52, doi:10.1007/BF00712389, 1995.





- Bates, T. S., Coffman, D. J., Covert, D. S., and Quinn, P. K.: Regional marine boundary layer
  aerosol size distributions in the Indian, Atlantic, and Pacific Oceans: A comparison of INDOEX
  measurements with ACE-1, ACE-2, and Aerosols99, J. Geophys. Res., 107(D18), 8026,
  doi:10.1029/2001JD001174, 2002.
- Bellouin, N., Rae, J., Jones, A., Johnson, C., Haywood, J., and Boucher, O.: Aerosol forcing in the
  Climate Model Intercomparison Project (CMIP5) simulations by HadGEM2-ES and the role of
  ammonium nitrate, J. Geophys. Res., 116, D20206, doi:10.1029/2011JD016074, 2011.
- Boyer, T. P., Levitus S., Garcia, H. E., Locarnini, R. A., Stephens, C., Antonov, J. I.; Objective
  analyses of annual, seasonal, and monthly temperature and salinity for the world ocean on a 0.25
  degrees grid. Int. J. Climatol. 25 (7), 931-945, 2005.
- Boyd, P. W.: Ranking geo-engineering schemes, Nat. Geosci., 1, 722–724, doi:10.1038/ngeo348,
  2008.
- 659 Brandt J, Brandt, J., Silver, J., Frohn, L. M., Geels, C., Gross, A., Hansen, A. B., Hansen, K. M.,
- 660 Hedegaard, G. B., Skjøth, C. A., Villadsen, H., Zare, A., and Christensen, J. H.: An integrated
- model study for Europe and North America using the Danish Eulerian Hemispheric Model with
- focus on intercontinental transport of air pollution. Atmos.Environ. 53:156-176, 2012.
- 663 Covert, D. S., Gras, J. L, Wiedensohler, A., and Stratmann, F.: Comparison of directly measured
- 664 CCN with CCN modeled from the number-size distribution in the marine boundary layer during
  665 ACE I at Cape Grim, Tasmania, J. Geophys. Res., 103, 16,597-16,608, 1998.
- de Leeuw, G., Andreas, E. L, Anguelova, M. D., Fairall, C. W., Lewis, E. R., O'Dowd, C., Schulz,
  M. and Schwartz, S. E.: Production flux of sea spray aerosol, Rev. Geophys., 49, RG2001,
- doi:10.1029/2010RG000349, 2011.
- Dana, M. T., and Hales, J. M.; Statistical aspects of the washout of polydisperse aerosols. Atmos.
  Environ. 10, 45-50, 1976.
- Emberson, L. D., Ashmore, M. R., Cambridge, H. M., Simpson, D., and Tuovinen, J. P.: Modelling
  stomatal ozone flux across Europe, Environ. Pollut., 109, 403–413, 2000.
- Foltescu, V. L., Pryor, S. C. and Bennet, C.: Sea salt generation, dispersionand removal on the
  regional scale. Atmospheric Environment 39, 2123-2133, 2005.





- Gregow, H., Ruosteenoja, K., Pimenoff, N., and Jylhä, K.: Changes in the mean and extreme
  geostrophic wind speeds in Northern Europe until 2100 based on nine global climate models. Int.
- 677 J. Climatol., 32, 1834–1846, 2011.
- Grini, A., Myhre, G., Sundet, J. K., and Isaksen, I. S. A.: Modeling the annual cycle of sea salt in
  the global 3D model OsloCTM2: Concentrations, fluxes, and radiative impact, J. Climate, 15,
  1717–1730, 2002.
- Gong, S. L.: A parameterization of sea-salt aerosol source function for sub- and super-micron
   particles, Global Biogeochem. Cycles, 17(4), 1097, doi:10.1029/2003GB002079, 2003.
- 683 Gong, S. L., Barrie, L. A., Prospero, J. M., Savoie, D. L., Ayers, G. P., Blanchet, J.-P., and Spacek,
- L.: Modelling sea-salt aerosols in the atmosphere: 2. Atmospheric concentrations and fluxes, J.
  Geophys. Res., 102(D3), 3819–3830, doi:10.1029/96JD03401, 1997.
- Haywood, J., and Boucher, O.: Estimates of the direct and indirect radiative forcing due to
  tropospheric aerosols: A review, Rev. Geophys., 38(4), 513–543, doi:10.1029/1999RG000078,
  2000.
- Haywood, J., Ramaswamy, V., and Soden, B.: Tropospheric aerosol climate forcing in clear-sky
  satellite observations over the oceans, Science, 283, 1299–1303, 1999.
- Hjellbrekke, A.-G. & Fjæra, A. M.: Data Report: Acidifying and eutrophying compounds and
  particulate matter. EMEP/CCC-Report 1/2011, 2009.
- Horn, H., Bonka, H., and Maqua, M.: Measured particle bound activity size-distribution, deposition
  velocity, 700 and activity concentration in rainwater after the Chernobyl accident, J. of Aerosol
  Science, 18, 681–684, 10.1016/0021-8502(87)90096-6, 1987.
- Huebert, B. J., Bates, T., Russell, P. B., Shi, G., Kim, Y. J., Kawamura, K., Carmichael, G., and
  Nakajima, T.: An overview of ACE-Asia: Strategies for quantifying the relationships between
  Asian aerosols and their climatic impacts, J. Geophys. Res., 108(D23), 8633,
  doi:10.1029/2003JD003550, 2003.
- 700 IPCC: Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the
- 701 Fifth Assessment Report of the Intergovernmental Panel on Climate Change [Stocker, T.F., D.
- 702 Qin, G.-K. Plattner, M. Tignor, S.K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex and P.M.
- 703 Midgley (eds.)]. Cambridge University Press, Cambridge, United Kingdom and New York, NY,
- 704 USA, 1535 pp, doi:10.1017/CBO9781107415324, 2013





- Jylhä, K.: Empirical scavenging coefficients of radioactive substances released from Chernobyl,
  Atmos. Environ. A., 25, 263–270, 10.1016/0960-1686(91)90297-K, 1991.
- Kjellström, E., Nikulin, G., Hansson, U., Strandberg, G., and Ullerstig, A.: 21st century changes in
  the European climate: uncertainties derived from an ensemble of regional climate model
  simulations, Tellus Series A-Dynamic Meteorology and Oceanography, 63, 24–40,
  10.1111/j.1600-0870.2010.00475.x, 2011.
- Kaskaoutis, D. G., Kambezidis, H. D., Hatzianastassiou, N., Kosmopoulos, P. G., and
  Badarinath, K. V. S.: Aerosol climatology: dependence of the Angstrom exponent on wavelength
  over four AERONET sites, Atmos. Chem. Phys. Discuss., 7, 7347-7397, doi:10.5194/acpd-77347-2007, 2007.
- Kusmierczyk-Michulec J. and van Eijk, A.M.J.: Atmospheric Optics: Models, Measurements, and
  Target-in-the-Loop Propagation III, 74630G; Stephen M. Hammel; Alexander M. J. van Eijk;
  Mikhail A. Vorontsov, Editor(s) doi: 10.1117/12.828394, 2009.
- Korhonen, H., K. Carslaw, S., and Romakkaniemi, S.: Enhancement of marine cloud albedo via
  controlled sea spray injections: A global model study of the influence of emission rates,
  microphysics and transport, Atmos. Chem. Phys., 10, 4133–4143, doi:10.5194/acp-10-41332010, 2010.
- Kouznetsov, R. and Sofiev, M.: A methodology for evaluation of vertical dispersion and dry
  deposition of atmospheric aerosols, J. Geophys. Res., 117, 10.1029/2011JD016366, 2012.
- Langner, J., Engardt, M., and Andersson, C.: European summer surface ozone 1990–2100, Atmos.
   Chem. Phys., 12, 10097-10105, doi:10.5194/acp-12-10097-2012, 2012.
- Latham, J., and Smith, M. H.: Effect on global warming of wind dependent aerosol generation at the
   ocean surface, Nature, 347, 372–373, doi:10.1038/347372a0, 1990.
- Lenton, T. M., and Vaughan, N. E.: The radiative forcing potential of different climate
  geoengineering options, Atmos. Chem. Phys., 9, 5539–5561, doi:10.5194/acp-9-5539-2009,
  2009.
- Lewis, E. R., and Schwartz, S. E.: Comment on "Size distribution of sea-salt emissions as a
  function of relative humidity," Atmos. Environ., 40, 588–590,
  doi:10.1016/j.atmosenv.2005.08.043, 2006.





- Liao, H., Chen, W. T., and Seinfeld, J. H.: Role of climate change in global predictions of future
  tropospheric ozone and aerosols. J. Geophys. Res., 111, D12304, 2006.
- T36 Lundgren, K., Vogel, B., Vogel, H., and Kottmeier, C.: Direct radiative effects of sea salt for the
- Mediterranean region under conditions of low to moderate wind speeds, J. Geophys. Res.
  Atmos., 118, 1906–1923, doi:10.1029/2012JD018629, 2013.
- Ma, X., von Salzen, K., and Li, J.: Modelling sea salt aerosol and its direct and indirect effects on
  climate, Atmos. Chem. Phys., 8, 1311-1327, doi:10.5194/acp-8-1311-2008, 2008.
- Mahowald, N. M., Lamarque, J.-F., Tie, X. X., and Wolff, E.: Sea-salt aerosol response to climate
  change: Last Glacial Maximum, preindustrial, and doubled carbon dioxide climates, J. Geophys.
- 743 Res., 111, D05303, doi:10.1029/2005JD006459, 2006.
- Manders, A. M. M., Schap, M., Querol, X., Albert, M. F., Vercauteren, J., Kuhlbusch, T. A. J., and
  Hoogerbrugge, R.: Sea salt concentrations across the European continent, Atmos. Environ., 44,
  2434–2442, doi:10.1016/j.atmosenv.2010.03.028, 2010.
- Mayer, B. and Kylling, A.: Technical note: The libRadtran software package for radiative transfer
  calculations description and examples of use, Atmos. Chem. Phys., 5, 1855-1877,
  doi:10.5194/acp-5-1855-2005, 2005.
- Mie, G.: Beiträge zur Optik trüber Medien, speziell kolloidaler Metallösungen [Contribution to the
  optics of turbid media, particularly of colloquial metal solutions Royal Aircraft Establishment],
  Ann. Phys.25(3), 377–445, 1908.
- Monahan, E. C., Spiel, D. E., and Davidson, K. L.: A model of marine aerosol generation via
  whitecaps and wave disruption, in Oceanic Whitecaps, edited by E. C. Monahan and G.
  MacNiochaill, 167–193, D. Reidel, Norwell, Mass, 1986.
- Mårtensson, E. M., Nilsson, E. D., de Leeuw, G., Cohen, L. H., and Hansson, H.-C.: Laboratory
  simulations and parameterization of the primary marine aerosol production, J. Geophys. Res.,
  108(D9), 4297, doi:10.1029/2002JD002263, 2003
- Nakićenović, N.: Global greenhouse gas emissions scenarios: Integrated modeling approaches,
  Tech. Forecasting & Social Change, 63, 105–109, 2000.
- Nilsson, E. D., Mårtensson, E. M., van Ekeren, J. S., de Leeuw, G., Moerman, M. M., and O'Dowd,
  C. D.: Primary marine aerosol emissions: Size resolved eddy covariance measurements with





- restimates of the sea salt and organic carbon fractions, Atmos. Chem. Phys. Discuss., 7, 13,345–
- 764 13,400, doi:10.5194/acpd-7-13345-2007, 2007.
- 765 NOAA: World Ocean Atlas 2013. http://www.nodc.noaa.gov/OC5/woa13/, 2013
- 766 O'Dowd, C., and de Leeuw, G.: Marine aerosol production: A review of the current knowledge,
- 767 Philos. Trans. R. Soc. A, 365, 1753–1774, doi:10.1098/rsta.2007.2043, 2007.
- O'Dowd, C. D., and Smith, M. H.: Physico-chemical properties of aerosol over the North East
  Atlantic: Evidence for wind speed related sub-micron sea-salt aerosol production, J. Geophys.
  Res., 98, 1137–1149, doi:10.1029/92JD02302, 1993.
- O'Dowd, C., Lowe, J., Smith, M., and Kaye, A.: The relative importance of non-seasalt sulphate
  and sea-salt aerosol to the marine cloud condensation nuclei population: An improved multicomponent aerosolcloud droplet parameterization, Q. J. R. Meteorol. Soc., 125, 1295–1313,
  1999.
- O'Dowd, C. D., Smith, M. H. Consterdine, I. E., and Lowe, J. A.: Marine aerosol, sea salt, and the
  marine sulphur cycle: A short review, Atmos. Environ., 31, 73–80, doi:10.1016/S13522310(96)00106-9, 1997
- Ovadnevaite, J., Manders, A., de Leeuw, G., Ceburnis, D., Monahan, C., Partanen, A.-I., Korhonen,
  H., and O'Dowd, C. D.: A sea spray aerosol flux parameterization encapsulating wave state,
  Atmos. Chem. Phys., 14, 1837-1852, doi:10.5194/acp-14-1837-2014, 2014.
- Pierce, J. R., and Adams, P. J.: Global evaluation of CCN formation by direct emission of sea salt
  and growth of ultrafine sea salt, J. Geophys. Res., 111, D06203, doi:10.1029/2005JD006186,
  2006.
- Pincus, R., Platnick, S., Ackerman, S. A., Hemler, R. S., and Hofmann, R. J. P.: Reconciling
  simulated and observed views of clouds: MODIS, ISCCP, and the limits of instrument
  simulators, Journal of Climate, 2011.
- Pryor, S. C., and Barthelmie, R. J.: Particle dry deposition to water surfaces: processes and
   consequences. Marine Pollution Bulletin 41, 220-231, 2000.
- 789 Prank, M.: Evaluation of atmospheric composition simulations via comparison with remote-sensing
- and in-situ observations, Master's thesis, 74 pp., Dep. of Sci. and Technol., Univ. of Tartu,
  Tartu, Estonia, 2008.





- Quinn, P. K., Coffman, D. J., Kapustin, V. N., Bates, T. S., Covert, D. S.: Aerosol optical properties
  in the marine boundary layer during the First Aerosol Characterization Experiment (ACE 1) and
  the underlying chemical and physical aerosol properties, J. Geophys. Res., 103, D13, 54716,563, 1998
- Roeckner, E., Brokopf, R., Esch, M., Giorgetta, M., Hagemann, S., Kornblueh, L., Manzini, E.,
  Schlese, U., and Schulzweida, U.: Sensitivity of simulated climate to horizontal and vertical
  resolution in the ECHAM5 atmosphere model, J. Clim., 19, 3771–3791, 10.1175/JCLI3824.1,
  2006.
- Rodell, M., Houser, P.R., Jambor, U., Gottschalck, J., Mitchell, K., Meng, C.-J., Arsenault, K.
  Cosgrove, B., Radakovich, J., Bosilovich, M., Entin, J.K., Walker, J.P., Lohmann, D., and Toll,
  D.: The Global Land Data Assimilation System, Bull. Amer. Meteor. Soc., 85(3), 381-394, 2004

Russel, P.B., and Heintzenberg, J.: An overview of the ACE-2 clear sky column closure experiment
(CLEARCOLUMN). Tellus B, [S.I.], 52(2), ISSN 1600-0889,
doi:http://dx.doi.org/10.3402/tellusb.v52i2.16173, 2011.

- Russell, P. B., Redemann, J., Schmid, B., Bergstrom., R. W., Livingston, J. M., McIntosh, D. M.,
  Ramirez, S. A., Hartley, S., Hobbs, P. V., Quinn, P. K., Carrico, C. M., Rood, M. J., Öström, E.,
  Noone, K. J., von Hoyningen-Huene, W., and Remer, L.: Comparison of Aerosol Single
  Scattering Albedos Derived by Diverse Techniques in Two North Atlantic Experiments. J.
- 810
   Atmos.
   Sci.,
   59,
   609–619,
   doi:
   http://dx.doi.org/10.1175/1520 

   811
   0469(2002)059<0609:COASSA>2.0.CO;2, 2002
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
   600
- Salisbury, D. J., Anguelova, M. D., and Brooks, I. M.: On the variability of whitecap fraction using
  satellite-based observations, J. Geophys. Res.-Oceans, 118, 6201–6222, 2013.
- Samuelsson, Jones, P., Willén, C. G., Ullerstig, U., Gollvik, A., Hansson, S., Jansson, U.,
  Kjellström, C., Nikulin, E., and Wyser, K.: The Rossby Centre Regional Climate model RCA3:
- 816 Model description and performance, Tellus A, 63, 4–23, doi:10.111/j.1600-0870.2010.00478.x,
- 817 2011.
- 818 Seinfield, J. H., and Pandis, S. N.: Atmospheric Chemistry and Physics: From Air Pollution to
  819 Climate Change, 2nd ed., 1232, John Wiley, Hoboken, N. J., 2006.
- 820 Simpson, D., Andersson, C., Christensen, J.H., Engardt, M., Geels, C., Nyiri, A., Posch, M., Soares,
- 821 J., Sofiev, M., Wind, P., and Langner, J.: Impacts of climate and emission changes on nitrogen





- deposition in Europe: a multi-model study, Atmos. Chem. Phys., 14, 6995-7017,
  doi:10.5194/acp-14-6995-2014, 2014.
- Simpson, D., Fagerli, H., Jonson, J. E., Tsyro, S., Wind, P., and Tuovinen, J.-P: Transboundary
  Acidification, Eutrophication and Ground Level Ozone in Europe, PART I, Unified EMEP
  Model Description, 1-104, 2003.
- Smith, F. B., and Clark, M. J.: The transport and deposition of radioactive debris from the
  Chernobyl nuclear power plant accident with special emphasis on consequences to the United
  Kingdom, Meteorological Office Scientific Paper no. 42, HMSO London, 1989.
- Sofiev, M., Siljamo, P., Valkama, I., Ilvonen, M., and Kukkonen, J.: A dispersion modelling system
  SILAM and its evaluation against ETEX data, Atmos. Environ., 40, 674–685,
  10.1016/j.atmosenv.2005.09.069, 2006.
- Sofiev, M., Soares, J., Prank, M., de Leeuw, G., and Kukkonen, J.: A regional-to-global model of
  emission and transport of sea salt particles in the atmosphere, J. Geophys. Res., 116, D21302,
  doi:10.1029/2010JD014713, 2011.
- Struthers, H., Ekman, A. M. L., Glantz, P., Iversen, T., Kirkevåg, A., Mårtensson, E. M., Seland,
  Ø., and Nilsson, E. D.: The effect of sea ice loss on sea salt aerosol concentrations and the
  radiative balance in the Arctic, Atmos. Chem. Phys., 11, 3459-3477, doi:10.5194/acp-11-34592011, 2011.
- Tsyro, S., Aas, W., Soares, J., Sofiev, M., Berge, H., and Spindler, G.: Modelling of sea salt
  pollution over Europe: key uncertainties and comparison with observations Atmos. Chem. Phys.,
  11, 10367-10388, doi:10.5194/acp-11-10367-2011, 2011.
- 843 Textor, C., Schulz, M., Guibert, S., Kinne, S., Balkanski, Y., Bauer, S., Berntsen, T., Berglen, T.,
- 844 Boucher, O., Chin, M., Dentener, F., Diehl, T., Easter, R., Feichter, H., Fillmore, D., Ghan, S.,
- Ginoux, P., Gong, S., Grini, A., Hendricks, J., Horowitz, L., Huang, P., Isaksen, I., Iversen, I.,
- 846 Kloster, S., Koch, D., Kirkevåg, A., Kristjansson, J. E., Krol, M., Lauer, A., Lamarque, J. F.,
- Liu, X., Montanaro, V., Myhre, G., Penner, J., Pitari, G., Reddy, S., Seland, Ø., Stier, P.,
- 848 Takemura, T., and Tie, X.: Analysis and quantification of the diversities of aerosol life cycles
- 849 within AeroCom, Atmos. Chem. Phys., 6, 1777-1813, doi:10.5194/acp-6-1777-2006, 2006.
- 850 Twomey S.: Atmospheric Aerosol, Elviser, Amesterdam, 1977.





- 851 Tørseth, K., Aas, W., Breivik, K., Fjæraa, A. M., Fiebig, M., Hjellbrekke, A. G., Lund Myhre, C.,
- 852 Solberg, S., and Yttri, K. E.: Introduction to the European Monitoring and Evaluation
- Programme (EMEP) and observed atmospheric composition change during 1972–2009, Atmos.
- Chem. Physics, 12, 5447–5481, 10.5194/acp-12-5447-2012, http://www.atmos-chemphys.net/12/5447/2012/, 2012.
- 856 Wang, M., Ghan, S., Ovchinnikov, M., Liu, X., Easter, R., Kassianov, E., Qian, Y., and Morrison,
- H.: Aerosol indirect effects in a multi-scale aerosol-climate model PNNL-MMF, Atmos. Chem.
  Phys., 11, 5431-5455, doi:10.5194/acp-11-5431-2011, 2011.
- Witek, M. L., P. J. Flatau, J. Teixeira, and K. M. Markowicz: Numerical investigation of sea salt
  aerosol size bin partitioning in global transport models: Implications for mass budget and optical
  depth, Aerosol Sci. Technol., 45, 401–414, doi:10.1080/02786826.2010.541957, 2011.
- Witek, M. L., Flatau, P. J., Quinn, P. K., and Westphal, D. L.: Global sea-salt modeling: Results
  and validation against multicampaign shipboard measurements, J. Geophys. Res., 112, D08215,
  doi:10.1029/2006JD007779, 2007a.
- Witek, M. L., Flatau, P. J., Teixeira, K., and Westphal, D. L: Coupling an ocean wave model with a
  global aerosol transport model: A sea salt aerosol parameterization perspective, Geophys. Res.
  Lett., 34, L14806, doi:10.1029/2007GL030106, 2007b.
- van Loon, M., Tarrasón, L., and Posch, M.: Modelling Base Cations in Europe, Technical Report
   MSC-W, Norwegian Meteorological Institute, ISSN 0804-2446, 2005
- Zhang, L., Gong, S., Padro, J., and Barrie, L.: A size-segregated particle dry deposition scheme for
  an atmospheric aerosol module. Atmospheric Environment 35, 549-560., 2001.
- 872
- 873





model	mode	Dp [µm]	source function	dependency	humidity	Lowest model layer thickness (m)
DEHM	fine	<1.3	MA02	T S	static	<i>c</i> 0
	coarse	[1.3-10]	MO86	S	(80%)	60
EMEP	fine	<2.5	MA02	Т	static	90
	coarse	[2.5-10]	MO86	-	(80%)	
МАТСН	fine	[0.02-0.1]		T S		60
		[0.1–1]	MA02			
		[1-2.5]			dynamic	
	coarse	[2.5–10]	MO86	T (SO11) S		
SILAM	fine	[0.01-0.1]	SO11	ΤS	dynamic	25
		[0.1 - 1.5]	5011	15		
	coarse	[1.5-6]		T S		
		[6–15]	SO11			
		[15-30]				

# 874 Table 1 Model characteristics for SSA computations.

875 T: temperature, S: salinity, MO86: Monahan et al. (1986); MA03: Mårtensson et al. (2003), SO11:

876 Sofiev et al. (2011). In bold, the modes not used for the  $PM_{10}$  analysis.

877

878





		nsfer modelling libRadTran2.0 for present and future.		
Clouds	cloud cover	monthly averaged RCA3 fields (1990-2009); same for both periods		
(icy and wet)	AOD	monthly averaged MODIS data (2002-2014) (Pincus e al. 2011); same for both periods		
	vertical profiles	wc.dat*; wc.dat*		
Atmospheric properties		subarctic winter, latitude over 60°: afglsw.dat*		
	vertical profiles	subarctic summer, latitude over 60°: afglss.dat*		
	ventical profiles	mid-latitude winter, latitude below 60°: afglmw.dat*		
		mid-latitude summer, latitude below 60°: afglms.dat*		
	altitude, pressure and	monthly averaged RCA3 fields (1990-2009); same for		
	temperature	both periods		
	vertical profile	aerosol_default*		
	AOD	dynamic: SILAM AOD calculations		
Aerosol	asymmetry factor	0.8 (Ma et al. 2008)		
properties	single scattering albedo	0.99 (Lundgren et al, 2013)		
	angstrom coefficient	0.2 (Kaskaoutis et al, 2007; Kusmierczyk-Michulec & van Eijk, 2009)		
solar zenith an	gle	dynamic: computed with libRadTran sza tool		
surface albedo		monthly averaged NOAA data (1990-2012) (Rodell e al., 2004); same for both periods		
RTE solver		DISORT		
integrated shortwave calculation scheme		KATO2 (wavelength ~[0.2, 4] $\mu$ m)		

880 Table 2 Assumption for the radiative transfer modelling libRadTran2.0 for present and future.

882





883Table 3 Statistical evaluation of model results for surface SSA concentration (Na<sup>+</sup> $\mu$ g m<sup>-3</sup>),884considering the whole year (annual), winter (December, January and February) and summer

periods (June, July and August), for 33 EMEP measuring sites, between 1990 and 2009.

	annual	winter	summer	annual	winter	summer
Obs	0.72	0.94	0.55			
DEHM	1.08	1.39	0.74			
EMEP	0.64	0.75	0.49			
MATCH	0.45	0.42	0.42			
SILAM	0.86	0.78	0.94			
	correlation			StdRatio		
DEHM	0.85	0.87	0.81	1.72	1.57	1.79
EMEP	0.82	0.84	0.80	0.69	0.54	0.85
MATCH	0.75	0.82	0.77	0.48	0.33	0.66
SILAM	0.71	0.77	0.75	1.05	0.75	1.59
	RMSE				Bias	
DEHM	0.97	1.11	0.70	0.36	0.45	0.18
EMEP	0.53	0.75	0.36	-0.08	-0.18	-0.06
MATCH	0.69	1.03	0.41	-0.27	-0.52	-0.14
SILAM	0.71	0.76	0.74	0.14	-0.16	0.38

886





- Table 4 Statistical evaluation of model results for SSA wet deposition (Na<sup>+</sup>mg m<sup>-2</sup>), considering
  the whole year (annual), winter (December, January and February) and summer periods (June, July
  and August) for 133 EMEP measurement sites, between 1990 and 2009. SILAM5m is the
- 891 evaluation if considering the whole possible size range for SSA  $Dp = [0.01-30] \mu m$ .

	annual	winter	summer	annual	winter	summer
obs	1.59E+06	6.88E+05	1.36E+05			
DEHM	1.41E+06	5.59E+05	1.40E+05			
EMEP	1.64E+06	6.44E+05	1.65E+05			
МАТСН	6.08E+05	1.77E+05	9.64E+04			
SILAM	8.42E+05	2.81E+05	1.25E+05			
SILAM5m	1.70E+06	6.70E+05	1.83E+05			
	correlation			StdRatio		
DEHM	0.55	0.53	0.41	0.36	0.31	0.55
EMEP	0.38	0.32	0.33	0.47	0.44	0.53
МАТСН	0.49	0.50	0.34	0.13	0.11	0.26
SILAM	0.49	0.45	0.38	0.22	0.19	0.41
SILAM5m	0.62	0.63	0.37	0.86	0.84	0.93
	RMSE			Bias		
DEHM	3477	5513	866	-114	-327	10
EMEP	3778	6006	912	34	-112	74
МАТСН	3879	6122	892	-634	-1304	-102
SILAM	3737	5945	871	-483	-1038	-29
SILAM5m	3335	5070	1032	73	-44	122

892





# 894 Table 5 Predicted direct radiative effect (W m<sup>-2</sup>) by SSA for the past period

	annual	winter	summer	
sea	-0.25±0.22	-0.077±0.053	-0.21±0.012	
land	-0.20±0.18	-0.073±0.0019	-0.083±0.0030	







896

Figure 1. Top: Sea surface temperature (K), middle: wind speed (m s<sup>-1</sup>), bottom: precipitation (mm). Left panel: mean value for the past period (1990-2009); right panel: absolute difference between the future (2040-

899 2059) and past periods.







1

Figure 2. Model-measurement comparison for Na<sup>+</sup> monthly mean concentration (µg m<sup>-3</sup>) for
29 EMEP measuring sites, between 1990 and 2009. The Person correlation (r), root mean
square error (rmse), bias, standard deviation ration (stdRatio), p-value (p), 1:1 (red solid), 1:2
(green), and 2:1 (green) lines are shown for each CTM.






Figure 3. Model-measurement comparison for Na<sup>+</sup> monthly wet deposition (µg m<sup>-2</sup>) for 133
EMEP measuring sites, between 1990 and 2009. The Person correlation (r), root mean square
error (rmse), bias, standard deviation ration (stdRatio), p-value (p), 1:1 (red solid), 1:5
(green), and 5:1 (green) lines are shown for each CTM.

6







Figure 4. Annual sea salt emission (mgPM<sub>10</sub> m<sup>-2</sup>) for DEHM, MATCH and SILAM models.
Left panel mean value for the past period (1990-2009); right panel: absolute difference
between the future (2040-2059) and past periods.







- 1 **Figure 5.** Wind forcing ( $\approx U_{10}^{3.41}$ ). Left panel: past period (1990-2009); right panel: absolute
- 2 difference between the future (2040-2059) and past periods.
- 3
- 4







Figure 6 Sea salt emission (mgPM<sub>10</sub> m<sup>-2</sup>) difference between winter (December, January and
February, DJF) and summer (June, July and August, JJA) for DEHM, MATCH and SILAM





- 1 models. Left panel: past period (1990-2009); right panel: absolute difference between the
- 2 future (2040-2059) and past periods.













- 2 Figure 7. Sea salt surface concentration ( $\mu g P M_{10} m^{-3}$ ) for DEHM, MATCH, EMEP and
- 3 SILAM models. Left panel: mean value for the past period (1990-2009); right panel: absolute
- 4 difference between the future (2040-2059) and past periods.
- 5







EMEP concentration (ugPM10/m<sup>3</sup>) 1990-2009 DJF-JJA



MATCH concentration (ugPM10/m<sup>3</sup>) 1990-2009 DJF-JJA





EMEP concentration (ugPM10/m<sup>3</sup>) change DJF-JJA



MATCH concentration (ugPM10/m<sup>3</sup>) change DJF-JJA



44







1

Figure 8 Sea salt concentration (μgPM<sub>10</sub> m<sup>-3</sup>) difference between winter (December, January
and February, DJF) and summer (June, July and August, JJA) for DEHM, MATCH and
SILAM models. Left panel: past period (1990-2009); right panel: absolute difference between
the future (2040-2059) and past periods.













- 1
- 2 Figure 9 Sea salt deposition (wet+dry) (mgPM<sub>10</sub> m<sup>-2</sup>) for DEHM, MATCH, EMEP and
- 3 SILAM models. Left panel: mean value for the past period (1990-2009); right panel: absolute
- 4 difference between the future (2040-2059) and past periods.
- 5













Figure 10 Sea salt annual deposition (gPM<sub>10</sub> m<sup>-2</sup>) difference between winter (December,
January and February, DJF) and summer (June, July and August, JJA) for DEHM, MATCH
and SILAM models. Left panel: past period (1990-2009); right panel: absolute difference
between future (2040-2059) and past periods.







Figure 11. SSA mass flux  $[gPM_{10} \text{ m}^{-2} \text{ s}^{-1})]$  box calculations (left) and coarse mode fraction of the mass flux (right): as a function of radius (dry for DEHM and SILAM and RH = 80 % for MATCH) and temperature, for wind speed 15 m s<sup>-1</sup> and salinities 10 ‰ and 35 ‰.







2

- 3 Figure 12. Radiative forcing by sea salt (W m<sup>-2</sup>). Left panel: past period (1990-2009); right
- 4 panel: absolute difference between future (2040-2059) and past periods.







2

Figure 13. Radiative forcing by sea salt (W m<sup>-2</sup>): difference between future (2040-2059) and
past periods. Left panel: winter (December, January and February); right panel: summer
(June, July and August)

6