

Seasonal differences in observed versus modeled new particle formation at two European boreal stations

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Abstract. Realistic representation of atmospheric aerosol size distribution dynamics in large scale climate models is important for developing accurate descriptions of aerosol-cloud interactions. Despite the dynamic nature of the distributions, which have large seasonal and diurnal changes, model evaluations often focus on the annual median size distribution. Using more comprehensive monthly and diurnal model illustrations can be crucial for evaluating model performance and potential aerosol effects for short term variations. In this study, we assess the impact of a molecular model scheme for $\text{NH}_3\text{--H}_2\text{SO}_4$ nucleation integrated into the Earth System Model (ESM) EC-Earth3, across different seasons, months, and days within the boreal climate during the year 2018. Measured number size distributions from two in-situ boreal stations are used to evaluate and to study particle formation and growth representation in EC-Earth3 over 2018. Additionally, we utilize results from the ADCHEM model, a state of the art 1-D Lagrangian aerosol-chemistry model. This allows us to compare EC-Earth3 against results from highly detailed model description of aerosol formation and growth at the boreal stations. When comparing diurnal EC-Earth3 model results with ~~ADCHEM and observations~~ [in-situ observations at an hourly temporal resolution](#), we establish that using solely organic- H_2SO_4 nucleation parameterization will underestimate the aerosol number concentrations. The new added $\text{NH}_3\text{--H}_2\text{SO}_4$ nucleation parameterization in this study improves the resulting aerosol number concentrations and reproduction of particle formation events with EC-Earth3. However, from March to October, the EC-Earth3 still underestimates particle formation and growth.

1 Introduction

Aerosol particles suspended in the atmosphere are constantly forming and growing around us. *Secondary* aerosols are particulate matter formed in the atmosphere from precursor gases, either by the condensation of vapors onto existing particles or by the nucleation of new particles. Secondary aerosols have been shown to significantly impact the global climate and human health (Szopa et al., 2021; Nault et al., 2021; Pye et al., 2021). New-particle-formation (NPF) occurs when particles are formed from the clustering and condensation of various gases. The gases can originate from both natural and anthropogenic sources. A major gas compound driving new-particle formation is sulfuric acid (H_2SO_4). Atmospheric H_2SO_4 is formed in the gas-phase when SO_2 is oxidized by OH radicals, and when dimethyl sulfate (DMS) is oxidized by OH, Cl, and BrO radicals (see e.g.

Wollesen de Jonge et al. 2021). Sulfur dioxide is predominantly emitted from anthropogenic sources and volcanoes, while
25 DMS is mainly emitted by phytoplankton in the surface ocean. Secondary aerosols involving organic compounds (SOA) are
produced from volatile organic compounds (VOCs) in the atmosphere. 85% of the VOCs are estimated to originate from natural
sources (Lamarque et al., 2010; Guenther et al., 2012), labelled biogenic volatile organic compounds (BVOCs). The abundance
of these natural gas compounds can have strong fluctuations over the year, in relation to the growing season. Furthermore, gas-
phase ammonia ($\text{NH}_3(\text{g})$) has been shown to be a key species in the NPF processes by recent model and experimental studies
30 (Dunne et al., 2016; Roldin et al., 2019). Atmospheric NH_3 is mainly a by-product emission from agricultural and industrial
activities. Including NH_3 in the NPF parameterization has not yet been well established in most large-scale climate models, so
evaluating the impact of the recently introduced NH_3 -enhanced formation mechanism is a key focus of this study.

Quantification of aerosol formation mechanisms has advanced significantly in the past decades with the development of
computational atmospheric chemistry models. The essential prerequisites for achieving reasonable modelling of aerosols and
35 NPF is to define an accurate representation of the aerosols' physical and chemical processes. However, the price for increasing
the complexity in models has to be balanced with the model's computational capacity. All models are restricted within limited
computational process time. Generally, the complexity of model chemistry and physics is limited by the size of the spatial
and temporal scale it has to cover. Therefore, it is more common for global Earth system models (ESMs) to have more crude
parameterization for aerosols and chemistry compared to e.g. a 1-D model. Nonetheless, improving the representation of
40 aerosols in Earth system models is crucial for reducing the large uncertainties associated with aerosols regarding global climate
forcing (Forster et al., 2021).

Atmospheric aerosols tend to contribute to a net cooling of the global climate by scattering incoming short-wave solar
radiation, here referred to as the direct radiative effect (DRE) given in units of Wm^{-2} . Higher aerosol concentration can also
increase the number of cloud condensation nuclei (CCN), which can promote clouds to have higher cloud droplet number
45 concentrations (CDNC) and cloud albedo. This will increase the cloud's ability to effectively scatter short-wave radiation,
which promotes cooling of the climate as a consequence (Twomey, 1974; Albrecht, 1989). The changes in cloud radiative effect
(CRE) from aerosols are also mainly promoting net negative (cooling) climate forcings. The aerosol-cloud effect is considered
to produce a stronger net forcing than the direct scattering from aerosols (Forster et al., 2021). Nevertheless, aerosol-cloud-
climate effects hold some of the highest uncertainty of our known climate forcings. It is worth mentioning that recent studies
50 also suggests that increased NPF production can significantly decrease the CRE outcome. This can occur when a large portion
of the available vapors are consumed by strong NPF events, which limits the availability of vapors for existing particles to grow
into the CCN sizes (Sullivan et al., 2018; Roldin et al., 2019; Blichner et al., 2021; Patoulias et al., 2024).

Results from Svenhag et al. (2024) showed that the cloud radiative effect (CRE) in the Earth system model (ESM) EC-Earth3
exhibits high sensitivity to changes in the new particle formation scheme. Similarly, Sporre et al. (2020) showed the input SOA
55 yields and VOC precursors in EC-Earth3-AerChem resulted in high sensitivity in the CRE. The Sporre et al. (2020) study
also showed that different ESMs can produce opposite CRE outcomes from removing a VOC precursor in the models. This
further implies that the modelled NPF and sub-100 nm aerosol dynamics have a strong effect on the CRE, and it can vary
significantly between Earth system models. Furthermore, evaluation of ESMs and radiative effects commonly uses temporal

averaging to monthly mean or median outputs from the model (Mann et al., 2014; Sporre et al., 2020; Svenhag et al., 2024), and this could discount shorter extreme NPF events which could have a large impact on the environment and CRE locally.

The current default NPF parameterization scheme in EC-Earth3-AerChem include two nucleation pathways from: H_2SO_4 with water (relative humidity), and H_2SO_4 with extremely low-volatile organic compounds (ELVOCs) (Vehkamäki, 2002; Riccobono et al., 2014; Bergman et al., 2022). Using our new molecular clustering model scheme, here referred to as CLUST, incorporated through lookup table in EC-Earth3-AerChem, we introduce the $\text{H}_2\text{SO}_4\text{--NH}_3$ NPF mechanism in the EC-Earth3 aerosol module M7. The look-up table uses H_2SO_4 , NH_3 , temperature, ion-production, and cluster scavenging sink to derive the nucleation rate (Svenhag et al., 2024). A recent extensive study by Zhao et al. (2024) indicated the importance of including various NPF mechanisms in global models. They model and discuss how implementing 11 unique pathways had a significant effect on the vertical and horizontal aerosol size distributions.

A difficult task for all global model studies is to accurately validate the model, especially for aerosol formation and growth of sub-100 nm aerosols. This is partly due to the lack of available aerosol measurements globally. The global distribution of measurement stations is scarce, aerosol sizes below 10 nm in diameter are rarely measured, and observations of vertical profiles of aerosol distributions are lacking. Some aeronautical measurement campaigns from e.g. ATom (Brock et al., 2019) and CAFE-Brazil (Curtius et al., 2024) at various locations have made efforts to capture this, but they are not evaluated in this study. The coarse spatial resolution in the large-scale models also makes extrapolating single spatial points like the ground station measurement to a $2^\circ \times 3^\circ$ (latitude $\times$ longitude) grid-box likely non-representative. The modal aerosol size distribution representation in EC-Earth3, which is typical in ESMs, also hinders us to compare growth rates similar to what is seen in measured observations. Additionally, observations from satellite instrumentation can not confidently measure aerosols in the sub-100 nm diameter sizes, and aerial campaign measurements for vertical distributions are not sufficient yet.

For this study, we selected to compare and validate our EC-Earth simulations against two rural boreal stations in Sweden and Finland. The surrounding area is primarily spruce forest, with some anthropogenic and agricultural influences, particularly near the southern part of the station. This makes the extrapolation of the station observations to the model EC-Earth grid more accurate, as the areas are expected to share a more homogeneous air mass (Nieminen et al., 2014). At the Swedish site, we would expect higher NH_3 and H_2SO_4 levels compared to the Finnish site due to its proximity to more agricultural and urban regions.

The focus of this study is to determine the model's ability to predict NPF events and to investigate the timing of these events in the model. In addition to comparing the results to observations, we also compare EC-Earth simulations to results from the detailed chemistry process model ADCHEM. ADCHEM is operated as a Lagrangian 1-D column model along air mass trajectories, with similar nucleation and growth schemes as EC-Earth but including detailed aerosol size distribution dynamics and gas-phase and aerosol chemistry. This enables evaluation of the more simplified approaches used in ESMs against detailed modeling, which is essential for understanding the ESM performance. Comparisons to only field measurements of NPF do not give full information on the reasons for the differences between model and observations, for example, whether they originate from missing chemical components or from inadequate approximations used for included components. They also do not guarantee that a model is right for the right reasons. Therefore, we apply ADCHEM results to benchmark and

understand the aerosol formation representation in EC-Earth. This might reveal missing model features in EC-Earth3 that can
95 be addressed in future development.

2 Method

2.1 EC-Earth3.4.0

The Earth system model configuration used in this study is the EC-Earth3-AerChem (version EC-Earth3.4.0). This contains the
global circulation model Integrated Forecast System (IFS) cycle 36r4, which is coupled to the atmospheric chemistry model
100 Tracer Model 5 (TM5) version 1.2 with the carbon bond 5 (CB05) setting (Krol et al., 2005; van Noije et al., 2014). EC-Earth3
uses the model coupler OASIS3-MCT version 3.0 (Craig et al., 2017) where the information exchange between IFS and TM5-
MP is made every 6-hours, model time. The IFS model time step is 45 minutes and set to generate output every 3 hours on
a 0.7° spectral truncation grid. TM5 uses hourly time steps and is set to produce hourly model output with $2^\circ \times 3^\circ$ (latitude
 $\times$ longitude) resolution. The model is run as atmosphere-only, where EC-Earth3 is fed fixed sea-surface temperatures (SST)
105 and sea-ice content from the AMIP reader (van Noije et al., 2021). The vertical resolution in TM5 is represented by 34 hybrid
sigma pressure levels, and IFS have the same hybrid pressure levels, but extrapolated to 91 layers.

2.1.1 Aerosol module M7

The size-resolved aerosol micro-physics module M7 (Vignati et al., 2004), here used in TM5, has been developed for use in
large-scale transport models (e.g. Earth system models). Application of M7 within ESMs is motivated by the modal-based
110 size distribution that increase their computational efficiency. The size distribution in M7 is described by seven log-normal
modes with fixed standard deviation. This way, the model solely has to track the aerosol mass and the aerosol number of
each mode for every time-step. This is generally much more computationally efficient compared to a sectional scheme, which
requires significantly more tracers (Vignati et al., 2004). M7 uses four water-soluble (S) and three insoluble (I) size modes
with restricted diameter ranges as: nucleation mode (NUS) for $d < 10$ nm, Aitken modes (AIS,AII) for $10 < d < 100$ nm,
115 $100 < d < 1000$ nm, accumulation (ASC,ASI) $100 < d < 1000$ nm, and coarse modes (COS,COI) for $d > 1000$ nm. However, this modal system in M7 limits the size distribution appearance to fall within specific log-normal
modes at fixed sizes, and the transfer of aerosol mass and number between modes have some assumptions which are considered
unrealistic (Vignati et al., 2004). M7 includes seven aerosol species: sea salt (SS), dust (DU), black carbon (BC), sulfate (SO₄),
primary organic aerosol (POA), and secondary organic aerosols (SOA).

120 2.1.2 New particle formation and growth

The formation of new particles in M7 is modeled by calculating a rate (J) of newly formed 5 nm diameter aerosols every time
step in the units $\text{s}^{-1}\text{cm}^{-3}$. The default nucleation rate parameterization (Riccobono et al., 2014) in M7 used for EC-Earth3-

AerChem is:

$$J_{Riccobono} = K_m [H_2SO_4]^2 [ELVOC] \quad (1)$$

125 The equation includes a constant empirical factor ($K_m = 3.27 \times 10^{-21} \text{ cm}^6 \text{ s}^{-1}$) with the gas concentrations of H_2SO_4 and ELVOC. The $J_{Riccobono}$ in the default TM5-M7 parameterization is summed with the nucleation rate for binary homogenous nucleation (BHN) of H_2SO_4 and water (described by specific humidity) based on a classical nucleation theory approach (Vehkamäki, 2002). The nucleation rate J corresponds to particles of ca. 1 nm in diameter, and the growth to 5 nm is described using the Kerminen and Kulmala (KK) equation (Kerminen and Kulmala, 2002). The KK function in M7 determines
130 the particle formation rate using available gas phase ELVOCs and H_2SO_4 concentrations for estimated particle survival through condensational growth, ~~and the resulting particles then enter the explicitly modelled size distribution in the nucleation mode.~~ Only after this growth to 5 nm are the aerosols introduced into the modal size distribution (Bergman et al., 2022). The H_2SO_4 , ELVOCs, and semi-volatile organic compounds (SVOCs) can grow particles in the modal system through condensation. For the soluble accumulation mode, additional reactions of ammonia and nitric acid (HNO_3) can form particulate ammonium nitrate
135 (NH_4NO_3), and also methane sulfonic acid (MSA) can condense on existing ASC-mode particles (van Noije et al., 2014).

2.2 Detailed H_2SO_4 – NH_3 nucleation scheme

In accordance with Svenhag et al. (2024), this study also uses the implementation of a new approach for sulfuric acid and ammonia nucleation. Here, nucleation rates are obtained by molecular modeling and stored in lookup tables. The tables are generated and interpolated by the J-GAIN tool (Yazgi and Olenius, 2023b). The rates are calculated by applying benchmarked,
140 high-level quantum chemistry data to a molecular kinetics model, where the kinetic equations are solved by the state-of-the-art Aerosol Cluster and Dynamics Code (ACDC) (Olenius, 2021). The lookup table implemented in M7 gives the nucleation rate (J) as a function of the five variables (1) gas-phase H_2SO_4 , (2) gas-phase NH_3 , (3) temperature, (4) atmospheric ion production rate (IPR), and (5) cluster scavenging sink (Olenius et al., 2013). The lookup table contains nucleation rates for all combinations of the five variables that define the rate, and rates at specific conditions are determined by multivariate interpolation. This approach ensures accurate rates, avoiding the typical problem of nucleation rate parameterizations, namely that the rate may not be reproduced under all different conditions of the parameter space. The (5) molecular cluster scavenging sink in CLUST is calculated from sulfuric acid condensation sink which is scaled for different cluster sizes (Yazgi and Olenius, 2023b; Lehtinen et al., 2007). In the present EC-Earth3 setup, the input total condensation sink of sulfuric acid to CLUST is calculated from all 7 aerosol modes at every model time step. We test two H_2SO_4 – NH_3 schemes, created by different quantum
150 chemistry input data: and older data set (computed using the RICC method) and a newer dataset (computed using the DLPNO method), referred to as CLUST-High and CLUST-Low, respectively. The RICC2 (CLUST-High) version method is based on Olenius et al. (2013) and the newer DLPNO (CLUST-Low) version is based on Besel et al. (2020b). Further details on the EC-Earth3 implemented look-up table is described Svenhag et al. (2024). We study the ranges produced from the CLUST-High and CLUST-Low inputs, for assessing the model sensitivity to ammonia. An IPR lookup table ~~for~~ with global coverage of galactic
155 cosmic rays and soil radon is used (Yu et al., 2019). It reads model pressure (203 altitude levels), magnetic latitude (91 bands),

and the model grid land cover fraction (for ²²²Rn) to generate the ion production rates, analogous to the implementations in Svenhag et al. (2024) implementation in Svenhag et al. (2024).

2.3 EC-Earth3.4.0 simulations and emissions

We include five separate "atmosphere only" simulations for EC-Earth3.4.0-AerChem over the year 2018 with a 3-month spin-up period (i.e. from 2017-10-01 to 2017-12-31). Each simulation differs only in terms of the NPF parameterization. The five different simulation setup cases are shown in Table 1, as in Svenhag et al. (2024) with an additional "no NPF" case. All EC-Earth3 simulations have nudged meteorology from ERA-Interim for the IFS model (every 6 hours) for surface pressure, wind divergence, and vorticity to obtain more homogenous synoptic weather between cases.

Table 1. The 5 EC-Earth3 simulation cases setup.

EC-Earth3 case:	Nucleation Scheme:	Involved species:
Control	Equation. 1 (Riccobono et al., 2014) + BHN	H_2SO_4 , $ELVOC$, H_2O
CLUST-High	RICC2 generated Look-up table + BHN	H_2SO_4 , NH_3 , H_2O
CLUST-Low	DLPNO generated Look-up table + BHN	H_2SO_4 , NH_3 , H_2O
CLUST-Low + Riccobono	Look-up table and Equation. 1 + BHN	H_2SO_4 , $ELVOC$, NH_3 , H_2O
No NPF	No particle formation rates	-

All simulations are run with emissions in accordance with CMIP6 scenario SSP3-7.0, which includes monthly varying emissions for SO₂, BVOCs and NH₃ from 2015-2100. ELVOC and SVOC production is derived from monoterpene and isoprene reactions with OH and O₃ with specific rates and yields (Atkinson et al., 2006; Jokinen et al., 2015). Further descriptions for BVOCs in TM5 is stated in (Bergman et al., 2022). The isoprene and monoterpene emissions are based on the MEGAN-MACC inventory (Sindelarova et al., 2014b). The SO₂ and NH₃ anthropogenic primary emission sources include: agriculture, energy production, industrial, transportation, residential-commercial-other, solvents production and application, waste, international shipping, air, and open burning (Lamarque et al., 2010). The biogenic NH₃ emissions are from ocean and land with natural vegetation (Bouwman et al., 1997), and natural sources of SO₂ include volcanic emissions and oceanic dimethyl sulphide (DMS) that is oxidized by OH and NO₃. Further description of emissions for EC-Earth3-AerChem are given in van Noije et al. (2021).

2.4 ADCHEM simulation and emissions

The process-based chemistry transport model ADCHEM was used to provide detailed modeling of gas and aerosol species and NPF for comparisons with EC-Earth3 and field measurements obtained at the Hyytiälä and Hyltemossa research field stations during the year of 2018 (Hari and Kulmala, 2005; Neefjes et al., 2022). ADCHEM was operated as a one-dimensional Lagrangian column model, running along air-mass back trajectories generated by the Hybrid Single Particle Lagrangian Inte-

grated Trajectory Model (HYSPLIT) with input data on meteorology from the Global Data Assimilation System (GDAS) (Stein et al., 2015; Rolph et al., 2017). Back-trajectories were simulated seven days backwards in time ~~with the using the HYSPLIT default output interval of 1 hour for the~~ Hyytiälä and Hyltemossa field stations~~as receptor points, allowing the model to source~~ . The trajectory coordinates were then linearly interpolated to a temporal resolution of 10 minutes when calculating the emissions of gases and particles from the urban areas, ocean, and forested regions surrounding the stations. In this version of ADCHEM, the one-dimensional column model consists of 20 vertical layers spaced logarithmically, starting from the first
185 layer that representing the lowest 10 meters up to the 20th layer which represents the atmospheric layer between 1900 m and 2100 m above ground level. Emissions from these sources were obtained from the Copernicus Atmosphere Monitoring Service (CAMS), including emissions from the global ocean inventory (Lennartz et al., 2017; Ziska et al., 2013; Nightingale et al., 2000; Lana et al., 2011), the global anthropogenic inventory (Granier et al., 2019) and the global biogenic inventory (Sindelarova et al., 2014a). ADCHEM used a main model time step of 60 seconds when solving the atmospheric chemistry, aerosol
190 dynamics and vertical mixing.

New particle formation represented by an explicit coupling of molecular cluster and aerosol dynamics was obtained via the molecular cluster plugin (ClusterIn) (Olenius and Roldin, 2022). ClusterIn considered the ion-induced and neutral clustering of $\text{H}_2\text{SO}_4\text{--NH}_3$ and $\text{H}_2\text{SO}_4\text{--DMA}$ along with neutral clustering of $\text{HIO}_3\text{--HIO}_2$ and $\text{HIO}_3\text{--DMA}$. Data-sets for the $\text{H}_2\text{SO}_4\text{--NH}_3$ and $\text{H}_2\text{SO}_4\text{--DMA}$ mechanisms were computed at the DLPNO-CCSD(T)/aug-cc-pVTZ// $\omega\text{B97X-D/6-31++G(d,p)}$
195 quantum chemical level of theory. The $\text{HIO}_3\text{--HIO}_2$ data set was calculated at the DLPNO-CCSD(T)//M06-2X method as detailed by Zhang et al. (2022). The $\text{HIO}_3\text{--DMA}$ data was calculated applying the older RICC2 method (RI-CC2// $\omega\text{B97X-D}$), and can thus be considered likely an upper estimate for the pathway. (Ning et al., 2022; Besel et al., 2020a; Myllys et al., 2019). The subsequent growth of these particles was described by considering the condensation, dissolution and evaporation of 873 organic and inorganic gas-phase species to the aerosol particle population. Particles and gasses were mixed by use of the GDAS
200 meteorology throughout the 20 horizontal layers spaced logarithmically and spanning 2100 ~~meters~~-m into the atmosphere. ~~The ADCHEM model~~ For the gases, the model simulates the gas-phase and aqueous-phase chemistry of 5005 species via 13062 reactions. Strong inorganic acids (H_2SO_4 ,

The model solves the atmospheric diffusion equation in the vertical direction. The vertical diffusion coefficients (K_z) were calculated based on a slightly modified Grisogono scheme (??), where the K_z depends on the height above ground, the friction
205 velocity and the height of the atmospheric boundary layer, which is sourced from the GDAS meteorology. GDAS metrology is used in ADCHEM since the trajectory model HYSPLIT which calculates back-trajectories for the model is based on GDAS metrology. The ADCHEM model thereby attempts to reproduce the concentration of gasses and particles at different heights in accordance with the surface measurements made at both the Hyytiälä and Hyltemossa field stations. A more detailed description of the model along with specific cases where the model has been used can be found in Roldin et al. (2019) and
210 Wollesen de Jonge et al. (2024). The ADCHEM simulation for Hyytiälä were only generated for May to August 2018.

2.5 Hyltemossa and Hyytiälä stations

The 2018 measurement data set for aerosol particle number size distributions (PNSD) at the two forest stations are produced from Differential mobility Particle Sizer (DMPS) instruments, and retrieved from the online EBAS inventory (Tørseth et al., 2012). The instruments measure aerosol diameters down to ~ 3 nm, but with increasing uncertainty towards the lowest diam-
eters (Wiedensohler et al., 2012). Uncertainties in observational datasets obtained with the DMPS can also arise from factors including calibration and environmental influences, combined with diffusion and electrostatic losses for the smallest diameter aerosols. These uncertainties can potentially affect our interpretation of the PNSD at sub-100 nm. Each ramp (size distribution measurement) takes approximately 10 minutes with the DMPS. The sampled aerosol numbers are distributed into 52 (Hyytiälä) and 37 (Hyltemossa) size-bins and averaged to 1-hour arithmetic means with percentiles. Hyytiälä is the area and nearest small village where the SMEAR II station is located, and the coordinates are tabulated in Table 2.

Table 2. Station descriptions for the measurements used in this study (Tørseth et al., 2012).

Station Name	Location	Instrument	Data Time Period	Lat °N	Lon °E	Altitude	Setting
SMEAR II	Finland	DMPS	18/01/01-18/12/31	61.84	24.29	180 m	Forest/Rural
Hyltemossa	Sweden	DMPS	18/01/01-18/12/31	56.10	13.42	5 m	Forest/Rural

2.6 Model post-process methods

The ADCHEM and measured DMPS data sets of PNSD are presented in sectional bins. For EC-Earth3 TM5’s hourly output, the modal aerosol distribution is redistributed to 100 sectional bins from 1 nm to ~ 25 μ m diameter for each 1-hour time step. Applying this modal-to-sectional conversion for every 1-hour output (same as the TM5 time step) prevents the size distribution from being unrepresentative when using longer temporal averages, which can occur when the median modal-radius output from M7 for each mode is averaged over longer time-periods. ADCHEM is run along individual air mass trajectories that arrive at the stations every 3rd hour. As EC-Earth3 and the DMPS measurements are 1-hourly values, data for occurrences when the measured DMPS is missing over more than two time steps within a single 3-hour step for ADCHEM, is removed from the analysis.

3 Results and discussion

The differences between the modeled and measured PNSD are presented in this section, categorized by seasonal and weekly periods for the year 2018. The seasonal median PNSD for observations and models at the two stations are shown in Fig. 1. The measured PSND at the two stations, Hyytiälä and Hyltemossa, displays a comparable seasonal variability, with lower aerosol number concentrations during winter and peak concentrations in summer. At both stations, the five EC-Earth3 cases and ADCHEM underestimate aerosol number concentrations in summer (June to August), and overestimates in winter (December to

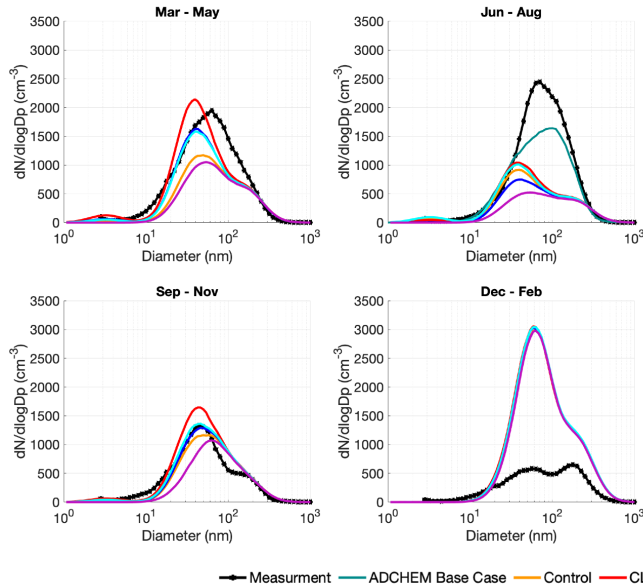
February). When comparing the seasonal PNSD from EC-Earth3 and ADCHEM with the measurements, the ADCHEM model shows substantially better performance. EC-Earth3's median aerosol number concentrations for winter are highly overestimated at both stations (shown in Fig. 1), which is driven by primary aerosol emissions, as is clear based on the overestimation also in the no-NPF simulation. Peculiarly, the overestimation by EC-Earth during winter is much stronger at Hyytiälä than at Hyltemossa. In springtime at both stations, the EC-Earth3 simulations have the highest disparity between model runs with the different NPF schemes. At such boreal conditions, springtime is favorable to nucleation as the period is photochemically active, but temperatures are not too high. The default ELVOC–H₂SO₄ (Riccobono) nucleation depend on biogenic emission from vegetation, and the NPF is significantly lower in the control case during spring before the initiated growing season. On the other hand, agricultural NH₃ emissions peak in spring due to application of fertilizers. The inclusion of NH₃–H₂SO₄ nucleation in spring, as modeled by the CLUST scheme cases, improves the agreement between the model and observations during this period. Shown in the hourly distributions and the seasonal median (Fig. 1 and Fig. 3), the model's ability to capture NPF-days is improved, and the results show more accurate magnitude of the nucleation events and the subsequent Aitken-mode number concentration.

The total aerosol concentrations during summer (Jun-Aug) is underestimated in EC-Earth3 shown for all model cases in Fig. 1, and similarly in an hour-to-hour comparisons during August and July shown in Fig.4. This insufficiency in NPF and growth could be attributed to low availability of condensable vapors, or a limitation for the model growth mechanisms in EC-Earth3, since ADCHEM can produce higher levels with similar nucleation schemes. The highest levels of available ELVOCs occurs in summer, and for these months the default ELVOC–H₂SO₄ will produce higher surface particle formation rates at the model surface level compared to the CLUST-High and CLUST-Low case (Fig. 4). However, the CLUST cases still have greater Aitken-mode aerosol number concentrations at the surface due to transport. As CLUST have substantially greater aerosol formation rates in the overlying grids (or possibly in neighboring grids) and these additional aerosols can then descend (or move laterally) to the surface grid representing the station.

3.1 Vertical distribution

For the 5 EC-Earth3 model variant cases, there is a great difference in the vertical particle formation rate profiles from the second model layer upward (approx at 900 hPa and above) when comparing the control case (ELVOC–H₂SO₄) and CLUST cases (NH₃–H₂SO₄) simulated rates, as shown in Fig. 2d. This occurs up to ~400 hPa, then the water-H₂SO₄ (BHN) nucleation prevails in the upper troposphere similarly for all cases. The cause of this model difference is related with the available precursor gases at different altitudes, as shown in Fig. 2a-c. The decline in particle formation rates in the control case is explained by the available modelled ELVOC gas concentration rapidly declining with altitude, and consequently the ELVOC–H₂SO₄ nucleation is limited to only near-surface NPF in the boundary layer. In contrast, the CLUST nucleation rate remains approximately unaffected by altitude because of compensating effects, as its formation is governed by temperature, ionization, and cluster scavenging sinks. Higher altitudes often offer more favorable conditions for these processes, supporting consistent nucleation rates (Svenhag et al., 2024). However, this advantage is likely counterbalanced by the decreasing NH₃ concentrations with altitude, which is why the formation rate remains fairly unchanged vertically. In Fig. A1d the difference

HYTTIÄLÄ



HYLTEMOSSA

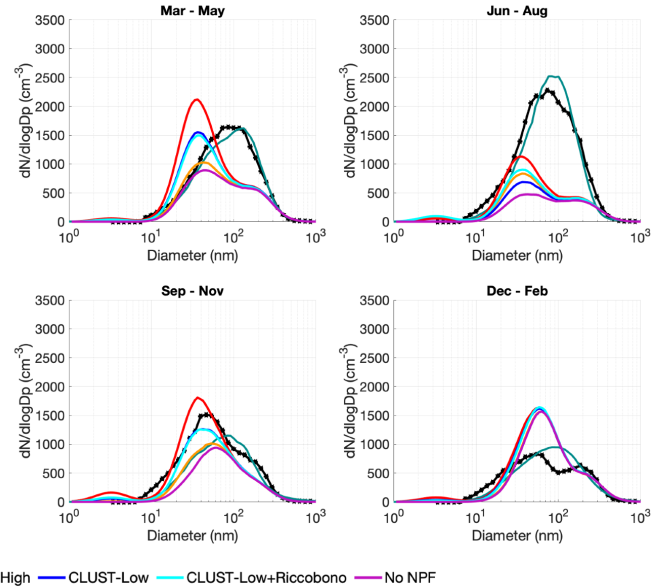


Figure 1. The seasonal median size distribution for the five EC-Earth3 variants, the ADCHEM simulations, and the observations. Note, the ADCHEM model only generated output from May to August for Hyytiälä and thus is only shown for Jun-Aug.

in gas concentrations and particle formation rates are shown at a lower altitude interval (800 hPa - 1000 hPa). The particle formation rate for the four EC-Earth3 cases is represented over the seasons (as a daily mean) at Hyytiälä is shown in Fig. A2. The largest case differences here between CLUST and the control case are found at altitudes between 800-400 hPa. There are occasions when the formation rates for the Control case are occurring at altitudes between 800-400 hPa, but these rates are almost entirely produced by the model from only BHN of water-H₂SO₄. During summer, nucleation throughout the troposphere seems to substantially decrease for all cases. The evaluated weekly cases for each season are made to additionally highlight the variations in EC-Earth3 between model level 1 (surface) and level 2 (~100 m) for aerosol formation rates, as well as ELVOC, H₂SO₄, and NH₃ concentrations (as presented in the following section).

3.2 Seasonal new particle formation

In this section, we evaluate hourly model output over separate weeks with observed NPF events from March to October, with most focus on the springtime. Firstly, two weekly spring cases in March and April are shown for the modelled aerosol size distributions and observed springtime new-particle formation events measured by the DMPS at the two stations (Fig. 3). As expected, the more detailed chemistry and sectional aerosol scheme in ADCHEM captures and reproduces these example events with better resemblance to observations compared to the EC-Earth3 model cases. In Fig. 3 (left panel) both observed or modelled NPF at Hyytiälä is initially (March 12th-13th) likely suppressed by a high concentration of background Aitken and

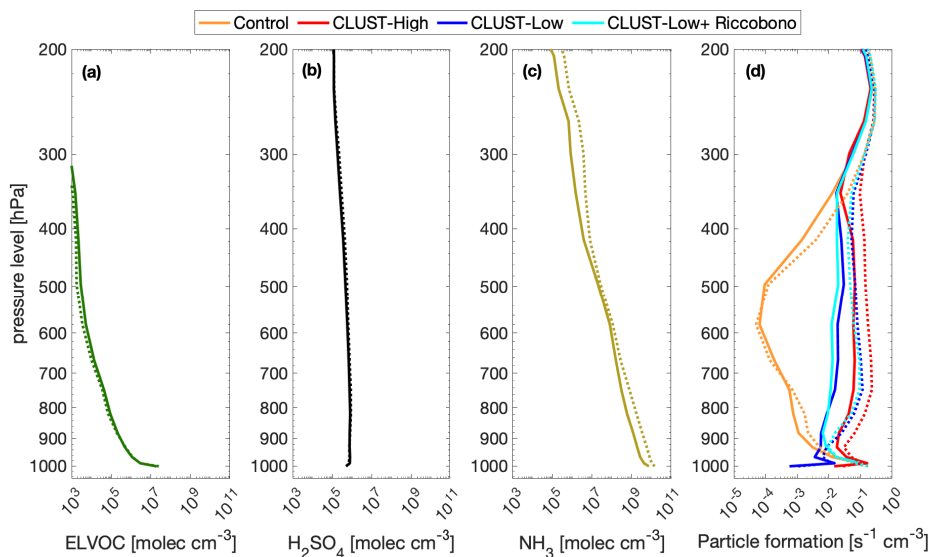


Figure 2. EC-Earth3's modelled annual mean vertical precursor (a) ELVOC, (b) H_2SO_4 (c) NH_3 concentrations, and (d) the 95th percentile of the particle formation rate for 2018. The bold line represent Hyytiälä and dotted is Hyltemossa station. The gas concentrations in a-c are from the No-NPF case, and the median differences between EC-Earth3 model cases would not be visible in this figure.

285 accumulation mode aerosols. EC-Earth3's mechanisms halts NPF on these days from the sinks produced by the larger mode aerosols, since the concentration of precursor gases are at similar levels to the following days when NPF is occurring, shown in Fig. 3. Afterward, four exemplary NPF and growth (banana) events at Hyytiälä are measured the succeeding days of March 14th to 17th. These four NPF events modelled by EC-Earth3 cannot be physically replicated (with a banana shape) due to the limitations of how aerosols grow and move between the modes. Nonetheless, regarding the resulting total concentration in
290 the nucleation and Aitken mode agrees better with observations when using the CLUST scheme compared to using only the default ELVOC– H_2SO_4 nucleation. Comparing the modelled "No NPF" case with the Control case for Hyytiälä in Fig. 3 (left panel) the weak visible nucleation in the control case gives almost no growth to the Aitken mode as the two cases have the same number size distributions shown in Fig. A8 a.

Fig. 3 (right-side panels) shows the modelled Hyltemossa station during seven days in April, and it presents three measured
295 typical nucleation (banana) events which are also captured differently by the model cases on April 6th–8th. The ADCHEM model captures these three events fairly well, but still show some deficiencies. From EC-Earth3, the resulting events are more irregular. The three events on April 6th–8th are captured, but underestimated in intensity and growth, with a continuously high NUS concentration throughout the day during April 6th. The EC-Earth3 CLUST cases gives aerosol number concentrations closer to the measured concentrations for the events on the 6th–8th compared to the default control case. This strong event
300 on April 6th shown in Fig. 3 i, j result in large differences for H_2SO_4 gas-phase concentrations between EC-Earth3 cases. Significantly more H_2SO_4 is consumed on this day in the CLUST scheme cases. The ELVOC concentrations are lower at this

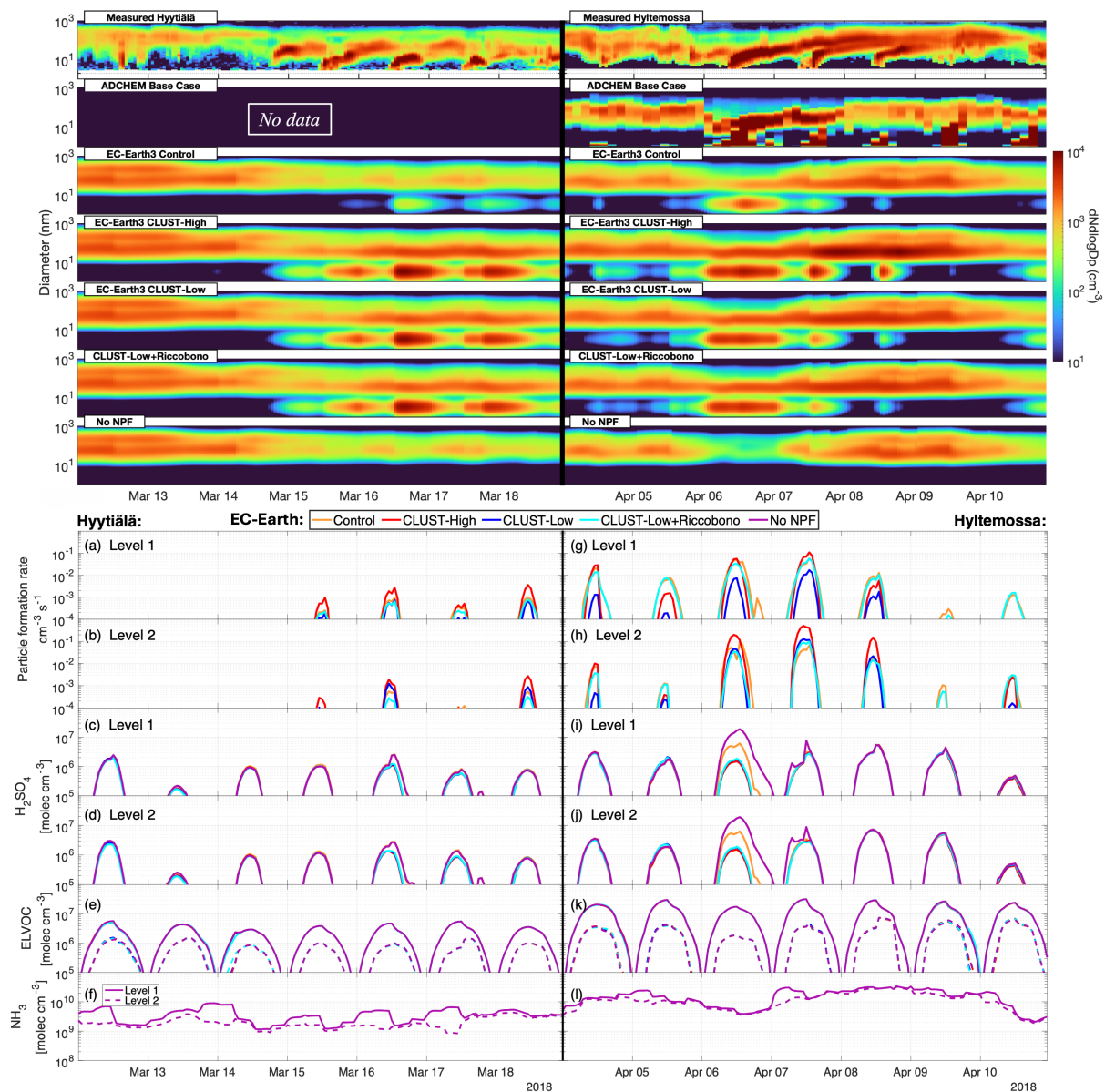


Figure 3. The top two figures shows show the surface aerosol number size distribution over the springtime for the 5 modelled EC-Earth3 cases, with the DMPS measured aerosols at Hyytiälä and Hyltemossa. ADCHEM simulations have no available hourly data outside the Summer months for Hyytiälä. The bottom section shows EC-Earth3 modelled layer 1 and layer 2 Hyytiälä and Hyltemossa springtime cases showing (a, b, g, h) daily maximum particle formation rate, (c, d, i, j) H_2SO_4 , (e, k) ELVOC, (f, l) and NH_3 gas concentration. The concentrations for ELVOC and NH_3 level 2 are shown as dotted lines in e, f, k, l. The missing particle formation rates below 10^{-4} in panel a, b, g, and h are considered practically zero.

earlier period of spring, shown in Fig. A6. A similarly strong NPF event is captured by EC-Earth3 at Hyltemossa on the 24th of October where the resulting aerosol and H₂SO₄ concentrations again vary between cases (shown in Fig. A7).

In this section, we include model results from the 2nd model height-level in TM5 to highlight the significant changes in formation rates and gas concentrations between model layers 1 and 2, with the differences for the nucleation schemes. Fig. 3 demonstrates the differences in the scheme's nucleation rates and gas concentrations for the same spring week as in Fig. 3. In the studied weekly cases, there are days when the different nucleation scheme experiment cases are diverging in the H₂SO₄ concentrations. The difference from the "No NPF" case shows how the different nucleation schemes are consuming H₂SO₄ to form and grow new particles, e.g. shown in Fig. 3 i, j on April 6th. On average, the H₂SO₄ concentrations are higher in the second model layer compared to the surface layer, shown in Fig. A1b. The ELVOC concentrations decrease by a factor of approximately 10 between levels 1 and 2 (Fig. 3 e, k), a pattern also observed for the SVOC concentrations (not shown). The springtime modelled ELVOC abundance at Hyltemossa compared to (more northern) Hyytiälä location drops by a factor of 10 at Hyytiälä, while contrarily, the measured total aerosol concentrations are slightly higher at Hyytiälä. This further suggest that we are lacking some model emissions or chemistry in EC-Earth3 around the station. Yet, underestimated summer PNSD at Hyytiälä is shown for both ADCHEM and EC-Earth3, which indicates some parameterizations and/or emissions for SOA might be missing at Hyytiälä in both models.

We evaluate an additional two weeks for the summer period at Hyltemossa and Hyytiälä (Fig. 4). It shows large underestimations in EC-Earth3 (all cases) for NPF magnitude and growth. The control case with ELVOC-H₂SO₄ results in higher particle formation rates (compared to CLUST) at the surface during this period at both locations, as shown in Fig. 4 a, g. Despite the higher surface formation rates for ELVOC-H₂SO₄ nucleation during these days, this case still yields a lower total surface aerosol number concentration compared to the CLUST-High case, shown in Fig. A8b, e. The particles produced in overlying and neighboring grids for CLUST-High is likely responsible for the resulting higher median summer surface concentrations here. Similar to the hourly results in Fig. 3, the CLUST annual median particle formation rates in Fig. A1 show significantly higher particle production in the second model layer compared to the surface layer.

3.3 Further discussion

Expanding on the vertical particle formation rates, the detailed ADCHEM model results suggest that the particle formation rate between the surface and 2000 m (800 hPa), shown in Fig. A3, exceeds what EC-Earth3's ELVOC-H₂SO₄ and water-H₂SO₄ nucleation (control case) produces ($< 10^{-3} \text{ s}^{-1} \text{ cm}^{-3}$). These results further support the inclusion of NH₃-H₂SO₄ NPF (like the CLUST table) as a nucleation pathway for the lower troposphere. Fig. 4 in Zhao et al. (2024) also shows NH₃-H₂SO₄ nucleation (neutral and ion-induced) as a key contributing pathway to particle formation in the upper and lower troposphere. However, while the newly-implemented ion-dependent NH₃-H₂SO₄ mechanism is an important NPF pathway, also other mechanisms and species may contribute. Studies have suggested the potential importance of amine and iodine enhanced nucleation (Wollesen de Jonge et al., 2024; Zhao et al., 2024), which are pathways not included in the EC-Earth3 model. Further, our current CLUST scheme does not include the effects of e.g. hydration or nitric acid (HNO₃) on NH₃-H₂SO₄ driven NPF, due to the lack of complete molecular data sets for calculating the formation rates, and testing these effects could be beneficial

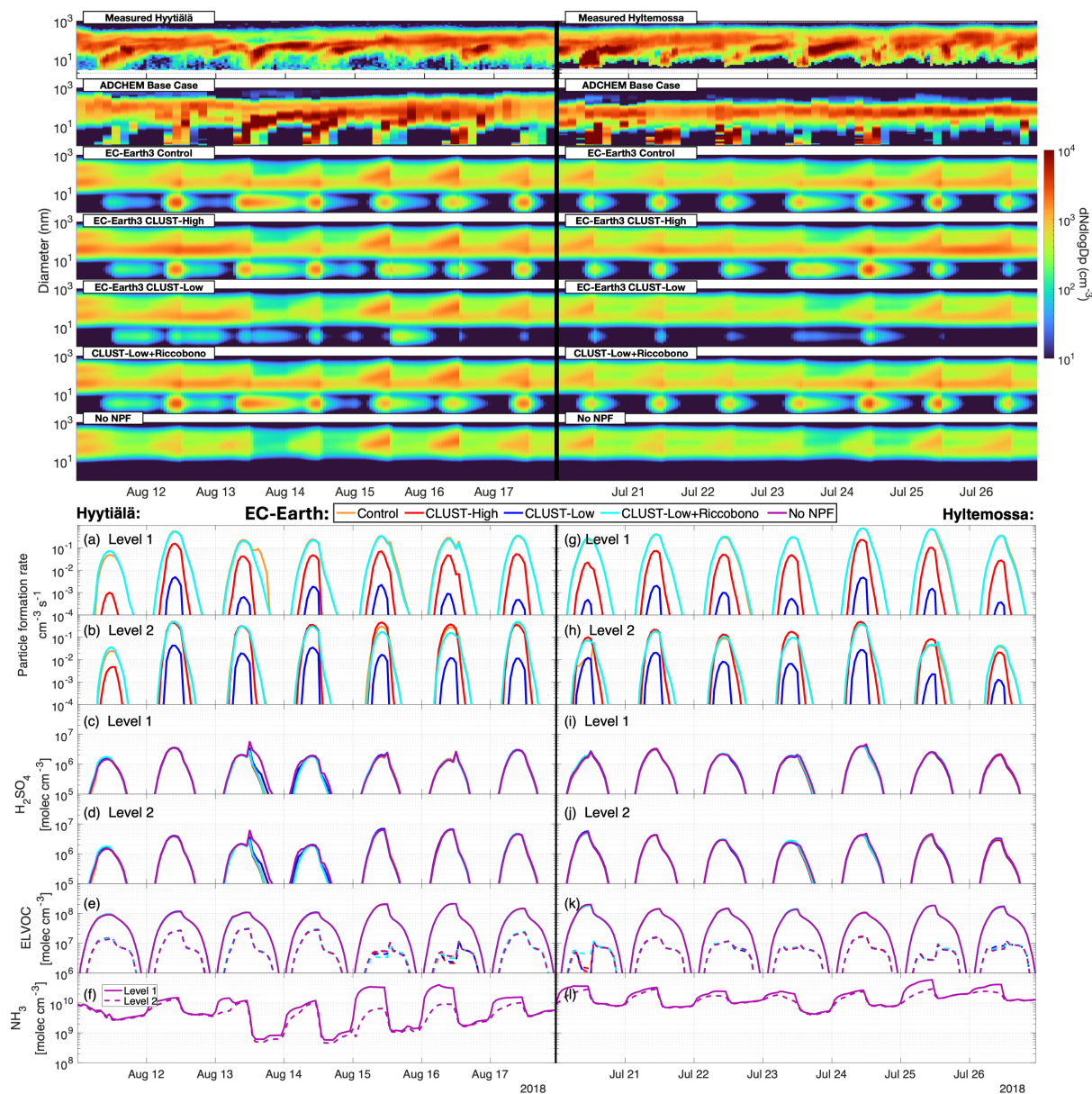


Figure 4. Same as Fig. 3 for the 2 weeks during summer.

(Zhao et al., 2024). Additionally, validating TM5's very low vertical distribution for the BVOCs concentrations above the surface model layer, could improve the boreal (and global) representation of aerosol in EC-Earth3. Further improvements could be achieved by incorporating detailed model comparisons to refine the fixed ELVOC and SVOC yields beyond the two categorical species currently used in TM5. Such enhancements could better capture VOC abundance and thereby improve predictions of particle formation rates and growth. Assuming the more sophisticated chemistry from ADCHEM would produce "significant

~~enough results" with respect to the increased computational burden~~The inclusion of some detailed chemical processes, as represented in the ADCHEM model, may yield results of sufficient relevance to climate impacts to justify the associated increase in computational expense when coupled with EC-Earth3.

It should also be noted that the scheme for organic–H₂SO₄ nucleation is simple, with no dependencies on temperature, scavenging sink or IPR (Eq. 1). Because of this, the modeled nucleation rate for this pathway does not fully correspond and/or respond to the ambient conditions and their changes. Additional uncertainties in the EC-Earth3 aerosol chemistry can also arise from the simplifications in modeling organic oxidation yields (from e.g. Jokinen et al. 2015) and growth processes. However, incorporating more complex chemical mechanisms is often constrained by the computational costs associated with most Earth system models.

Comparing the modelled mass concentrations in ADCHEM versus the EC-Earth3 versions, revealed that the surface organic aerosol mass is significantly higher for ADCHEM, especially during the month of July, shown in Fig. A4. This disparity in organic aerosol mass could be due to a combination of the factors mentioned above, e.g. the models' emission inventories, parameterization for SOA formation, and/or chemistry of the organic precursors. Noting the ADCHEM trajectories shows the prevailing air-masses are coming from continental areas with high organic emissions during this period (Wollesen de Jonge et al., 2024). Evaluating and refining the available condensable organic (and inorganic) vapor for secondary aerosol formation and growth in EC-Earth3 is crucial for future studies, as the current nucleation and growth schemes lead to significant underestimations of summer Aitken and accumulation mode concentrations at boreal stations. The overestimation by EC-Earth3 in winter might be due to the coarse resolution of EC-Earth3, which entails that anthropogenic emission which are quite far from the station are still emitted within the same grid box. There are also some differences in the emission inventories used by ADCHEM and EC-Earth3. However, these discrepancies are unlikely to be the primary source of emission-related differences observed at the two stations. A more significant factor is the spatial resolution at which each model reads the emissions. EC-Earth3 uses areal mean emissions from the CMIP6 inventory, interpolated over a coarse 2° × 3° (longitude × latitude) grid. In contrast, ADCHEM utilizes the CAMS inventory at a much finer resolution of 0.1° × 0.1° extracting emission values directly from individual grid cells.

A significant model uncertainty produced by TM5 is revealed in the hourly representations of near-surface aerosol concentrations. This is visible in Fig. 4 at noon (12:00), and occurs due to TM5's 6-hourly meteorological dynamics retrievals from IFS. The IFS dynamics controls the boundary layer height and consequential vertical mixing, and will maintain the 6:00 conditions up until new values are retrieved at 12:00 which causes an abrupt change in surface gas and aerosol concentrations following this time step. At the two stations, this effect is most prominent at noon in summer, when the boundary layer experiences its maximum fluctuation during the day. Similarly, at midnight (24:00) the surface aerosol concentration starts accumulating much faster as the boundary layer height is drastically decreased when TM5 retrieves the new IFS variables (from previous 18:00 conditions). The NH₃(g) concentrations (and likely many gas compounds) are also dramatically influenced by these shifts at noon and midnight, shown in Fig. 4. The unrealistic dynamics produced by TM5, resulting from this model assumption, introduce greater uncertainty in direct comparisons with hour-to-hour NPF events and aerosol populations in this study. ~~It would also add uncertainty to how aerosols near the surface potentially influences cloud-aerosol~~This artifact may also

introduce uncertainty regarding the influence of near-surface aerosols on cloud–aerosol interactions in EC-Earth3 within ~~this~~
the 6-hour coupling window.

4 Conclusions

The new CLUST implementation of $\text{H}_2\text{SO}_4\text{--NH}_3$ nucleation accounts for more spring and autumn NPF in the boreal regions,
380 where the default EC-Earth3 case is underestimating aerosol numbers. Comparing the CLUST cases versus the control run;
the total 2018 aerosol number concentration at Hyytiälä increased by 53 % for CLUST-High, 12 % for CLUST-Low, and
22 % for CLUST-Low+Riccobono. At Hyltemossa it increased by 91 % for CLUST-High, 28 % for CLUST-Low, and 34
% for CLUST-Low+Riccobono. From the resulting aerosol PNSD during spring, summer and autumn, we suggest the use
of CLUST-Low+Riccobono version with $\text{H}_2\text{SO}_4\text{--NH}_3$ and $\text{H}_2\text{SO}_4\text{--organic}$ pathways as a default option in future EC-
385 Earth3 studies to account (in principle) for both nucleation mechanisms in EC-Earth3. ADCHEM uses the same chemistry
data for the $\text{H}_2\text{SO}_4\text{--NH}_3$ pathway as implemented in the CLUST-Low scheme. The CLUST-Low scheme applies the most
recent quantum chemistry methods for the $\text{H}_2\text{SO}_4\text{--NH}_3$ nucleation rate and is considered to have more realistic sensitivity to
ammonia compared to the previous scheme (CLUST-High).

Furthermore, the ADCHEM model results and the global NPF modelled by Zhao et al. (2024), ~~suggests~~suggest the mean
390 atmospheric new particle formation rate is higher than $10^{-2} \text{ s}^{-1} \text{ cm}^{-3}$ between the second model layer in TM5 ($\sim 100 \text{ m}$) and
the upper troposphere. Our new CLUST nucleation mechanism accounts for ~~more NPF~~higher NPF rates in these regions for
the two boreal European mid-latitude stations, which has better agreement with detailed modelling in ADCHEM and NPF rates
presented in Zhao et al. (2024). The increase in Aitken mode particles may also be attributed to more nucleation occurring at
higher altitudes. This further suggests the inclusion of the $\text{H}_2\text{SO}_4\text{--NH}_3$ NPF pathway in Earth System Modelling to cover
395 nucleation and production of particles in low ELVOC environments and at higher altitudes, which have significant influence
on model clouds properties and radiative effects (Svenhag et al., 2024).

This study further ~~highlights the potential false representation from using solely annual medians when evaluating~~underscores
the potential for misrepresentation when relying solely on annual median values to evaluate the performance of EC-Earth3
~~'s-(and other ESMs)performance due to significant seasonal variations, due to pronounced seasonal variability~~. An annual
400 average of the two modeled stations in this study would merge the model's significant underestimations of summer aerosol
concentrations with its substantial overestimation in winter, resulting in a balanced median value. As the climate forcing from
aerosols have large seasonal variations (Carslaw et al., 2013; Forster et al., 2021), it would be crucial to have a representative
seasonal model depiction of aerosol formation and growth. The underestimated aerosol concentrations for EC-Earth3 in the
summer could have a significant impact on the radiative outcome with regard to the high sensitivity between CRE and sub-100
405 nm aerosols in EC-Earth3 (Svenhag et al., 2024). The observed mechanisms in these regional environments are not necessarily
transferable to different environments, which makes comparisons towards modelled NPF effects on a global scale challenging.
Another challenge is introduced by the uncertainty from the TM5's 6-hourly exchange of meteorological dynamics from the

IFS model, which alters the aerosol and precursor gas concentrations and conditions significantly near the surface at noon and midnight.

410 We have demonstrated that the detailed evaluation of seasonal boreal new-particle formation in the EC-Earth ESM can be accomplished by both representative field observations, and detailed aerosol simulations (ADCHEM). This combination aims to recognize the factors affecting the ability of the ESM to capture NPF trends. Such analysis can be applied (1) to assess seasonal trends and possible reasons for season-wise biases, and (2) to separate different error sources stemming from fundamental model schemes, e.g. inclusion/exclusion of given NPF mechanisms, and from ESM model approximations for
415 e.g. aerosol size distribution.

Appendix A

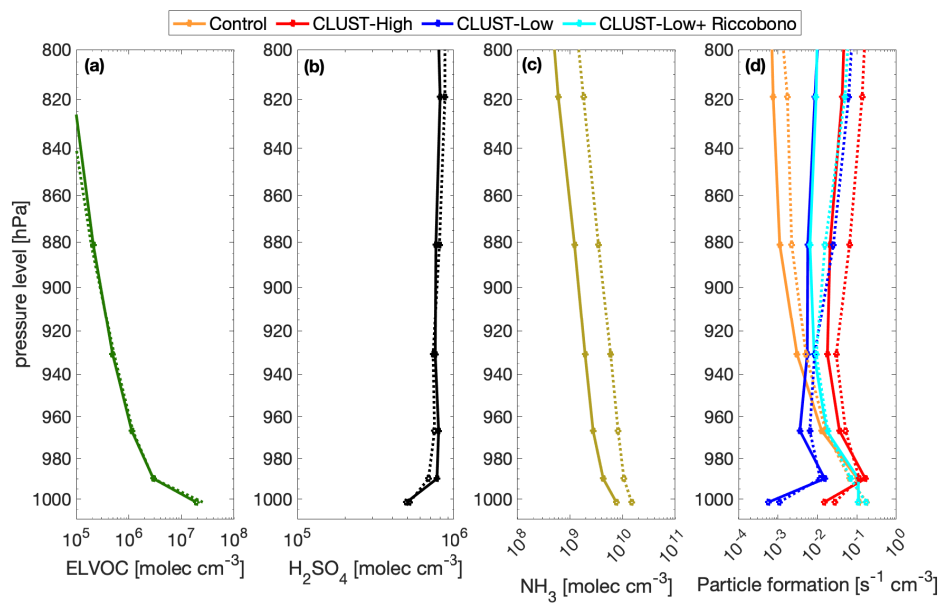


Figure A1. Same as Fig. 2 but at 800 - 1000 hPa.

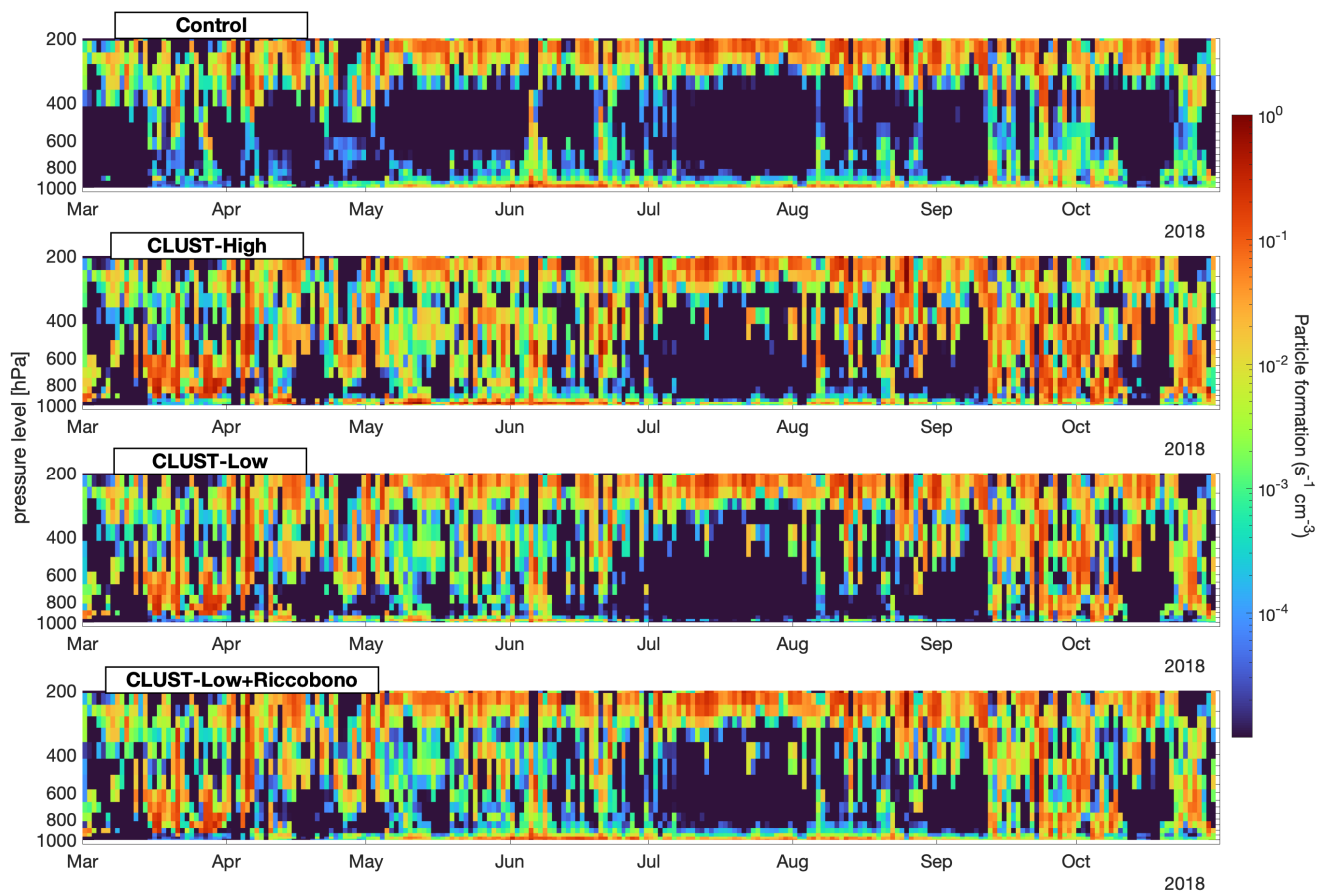


Figure A2. Vertical distribution of daily mean particle formation rates in the four EC-Earth3 cases at Hyytiälä for 2018, March to October.

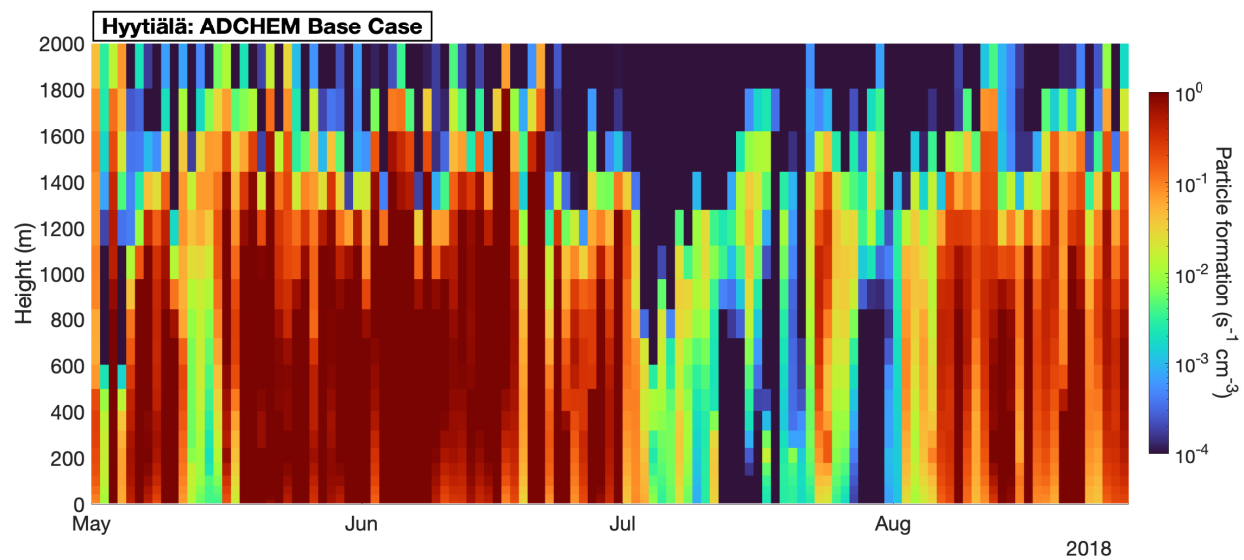


Figure A3. Vertical distribution of daily mean particle formation rates in ADCHEM Base Case at Hyytiälä 2018, May to August

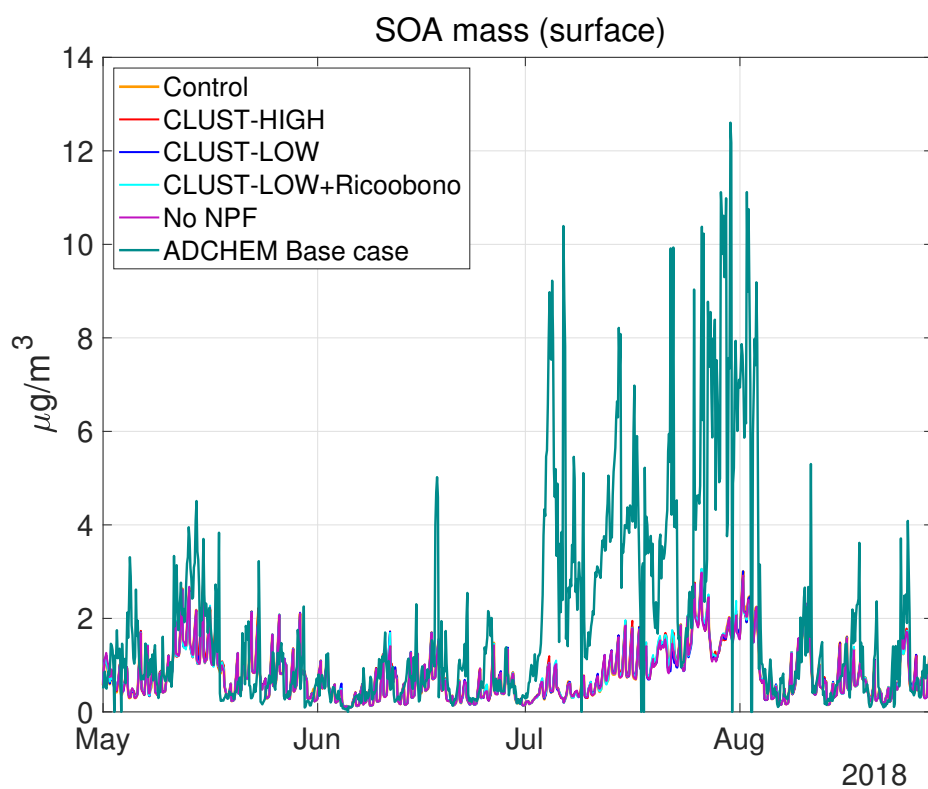


Figure A4. Modelled mass concentration of particulate organic matter (SOA) for EC-Earth3 cases and ADCHEM at Hyytiälä 2018, May to August

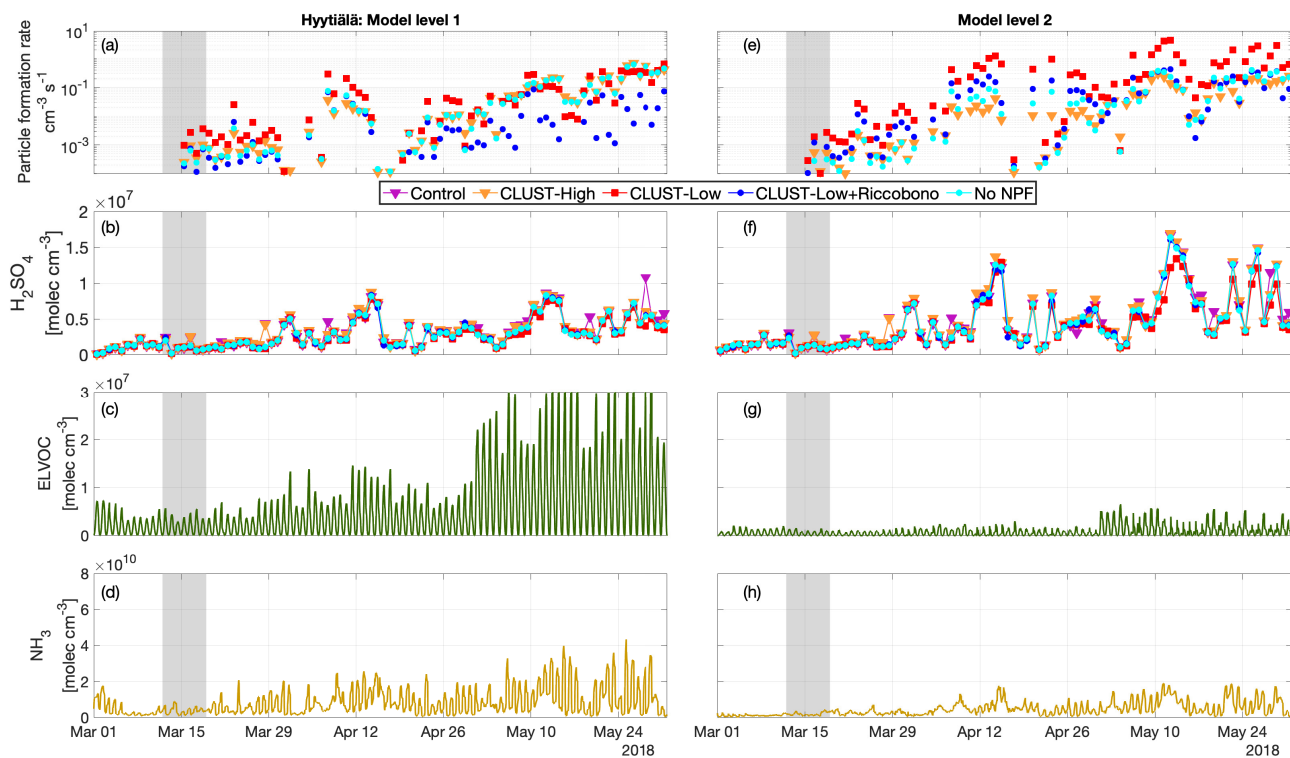


Figure A5. EC-Earth3 modelled layer 1 and layer 2 Hyytiälä springtime (a) daily maximum particle formation rate (b) H_2SO_4 (c) the No-NPF case ELVOC and (d) the No-NPF case NH_3 gas concentration. The shaded areas represent the studied weekly cases.

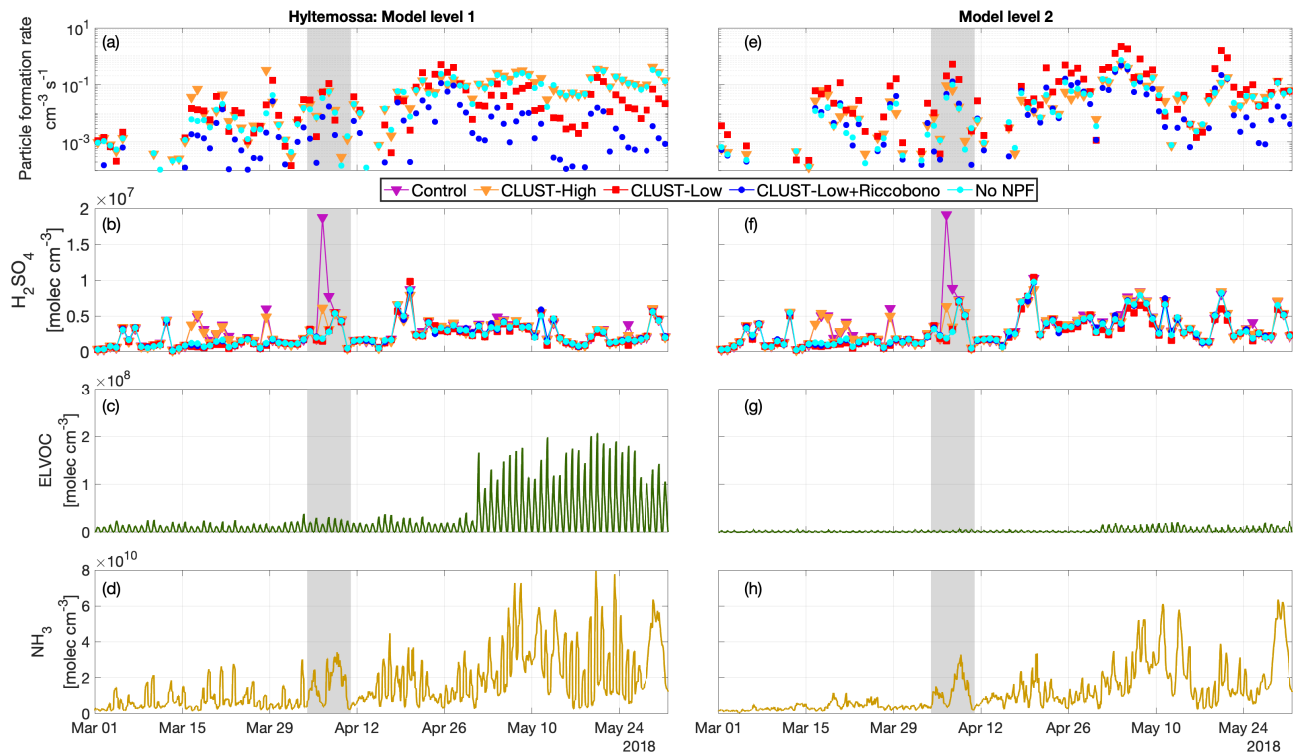


Figure A6. Same as Fig. A5 but for Hyltemossa. The ELVOC's y-axis scale magnitude in (c) is $\times 10$ greater here in reference to Fig. A5c.

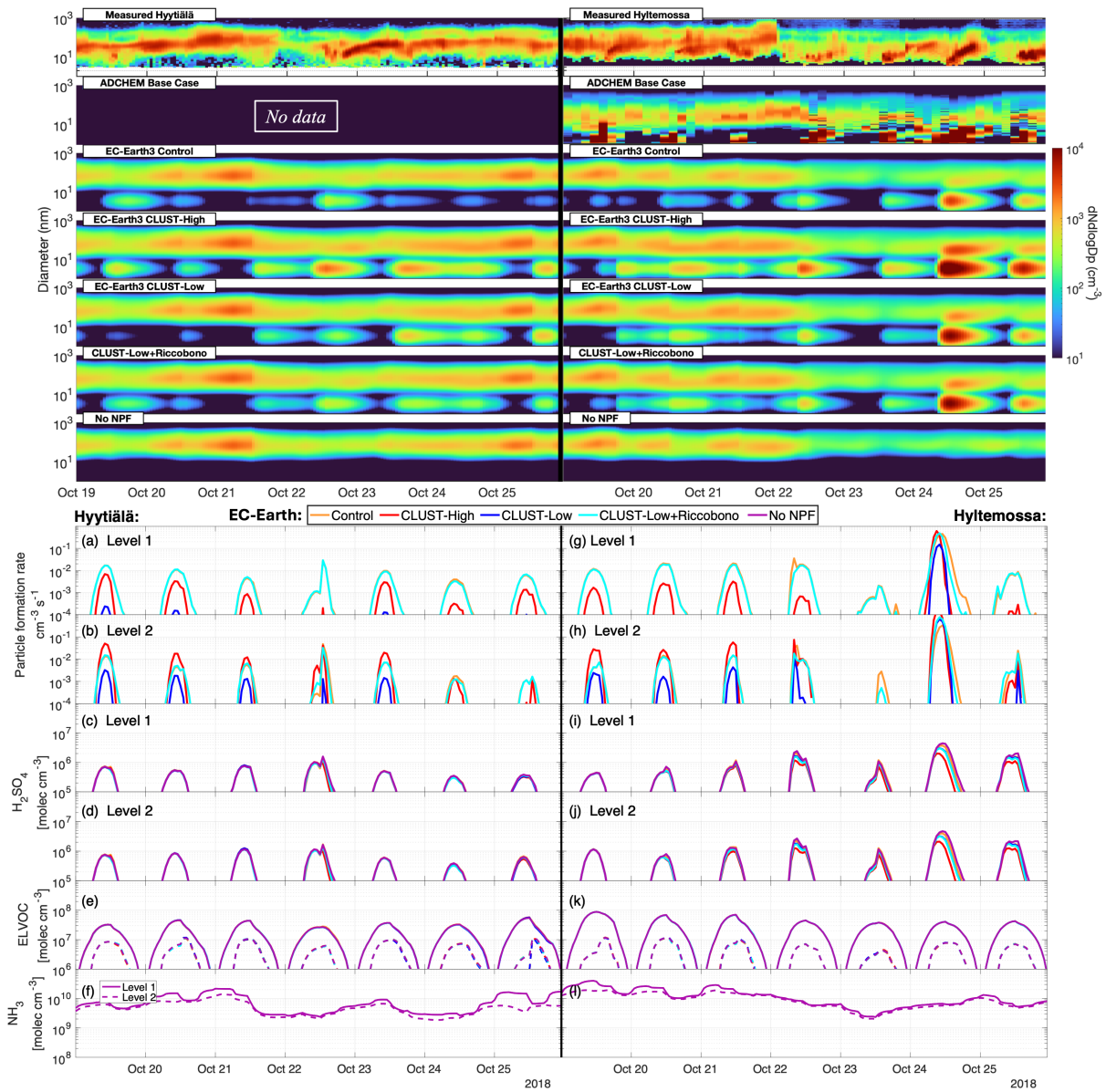


Figure A7. The top two figures shows the surface aerosol number size distribution over the autumn period for the 5 modelled EC-Earth3 cases, with the DMPS measured aerosols at Hyttiälä and Hyltemossa. ADCHEM simulations have no available hourly data outside the autumn months for Hyttiälä. The bottom section shows EC-Earth3 modelled layer 1 and layer 2 Hyttiälä and Hyltemossa autumn cases showing (a, b, g, h) daily maximum particle formation rate, (c, d, i, j) H₂SO₄, (e, k) ELVOC, (f, l) and NH₃ gas concentration. The concentrations for ELVOC and NH₃ level 2 are shown as dotted lines in e, f, k, l. The missing particle formation rates below 10⁻⁴ in panel a, b, g, and h are considered practically zero.

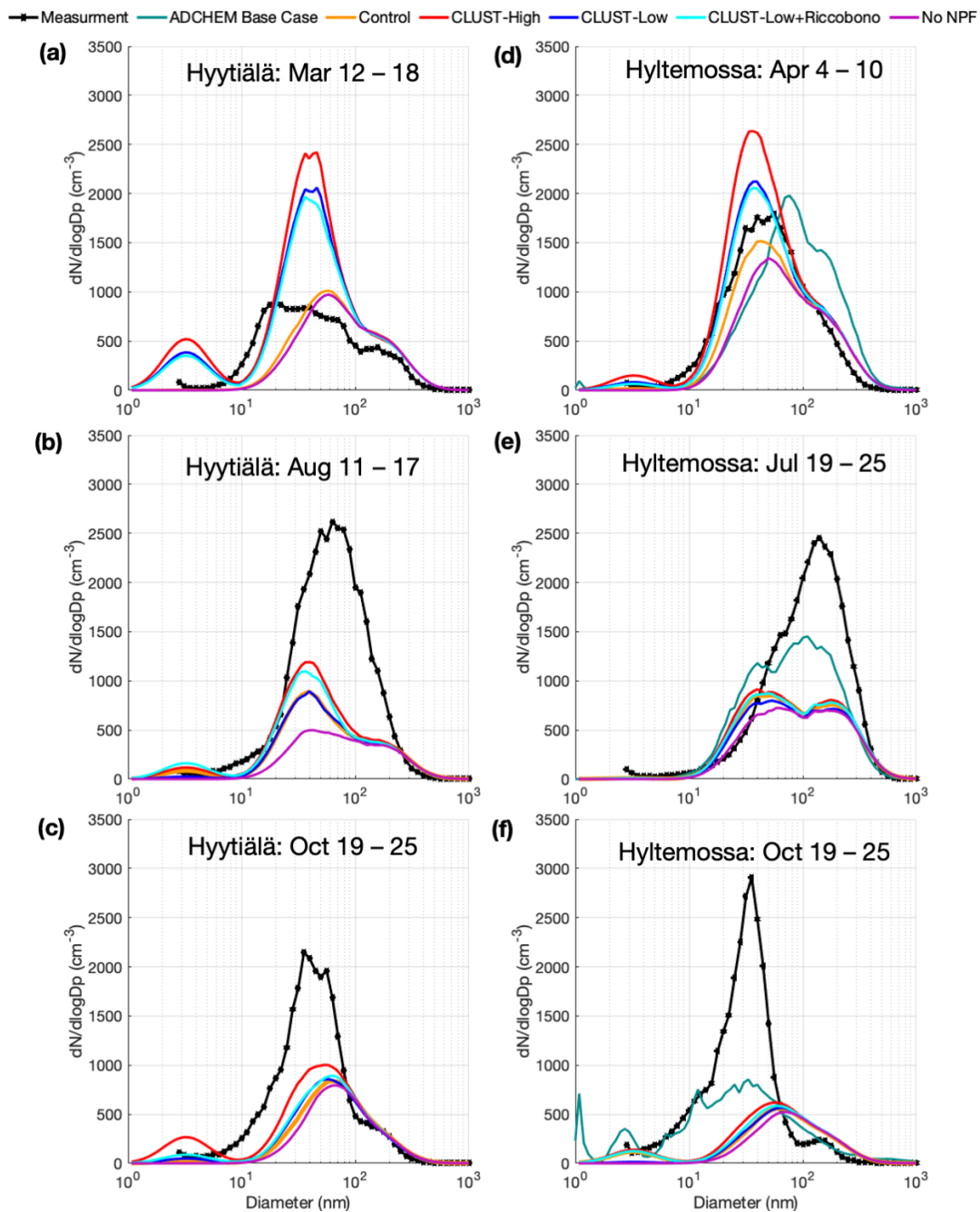


Figure A8. The surface median particle number size distribution of the studied seasonal weeks for spring, summer, and autumn. Showing The five EC-Earth3 cases, ADCHEM, and the station measurements

Code and data availability. Model code for TM5-MP version 1.2 with implemented CLUST look-up table is found at Svenhag (2024a). The model output datasets are found in Svenhag (2024b), post-process scripts are located in Svenhag (2024c). Codes for the J-GAIN v1.0 generator and the interpolator used for the CLUST look-up table in the experiments can be found at Yazgi and Olenius (2023a). Resources
420 for the IPR lookup table can be found in Yu (2019). The DMPS and SMPS measurement products at the stations are available for download at <https://ebas-data.nilu.no/Default.aspx>.

Author contributions. CS, MS, and PR designed the research idea, CS performed the EC-Earth3 simulations and post process handling. RWD and PR provided ADCHEM base case data and its model description. TO and DY developed the CLUST model, and CS performed the EC-Earth3 implementations of CLUST. SMB contributed with discussions for the analysis.

425 *Competing interests.* The authors declare that they have no conflict of interest.

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430 2022-06725 and 2018-05973.

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