A new chemistry option in WRF/Chem v. 3.4 for the simulation of direct and indirect aerosol effects using VBS: evaluation against IMPACT-EUCAARI data

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1 Abstract

A parameterization for secondary organic aerosol (SOA) production based on the volatility basis set 2 (VBS) approach has been coupled with microphysics and radiative schemes in the WRF/Chem 3 4 model. The new chemistry option called "RACM/MADE/VBS/AQCHEM" was evaluated on a cloud resolving scale against ground-based and aircraft measurements collected during the 5 6 IMPACT-EUCAARI campaign, and complemented with satellite data from MODIS. The day-today variability and the diurnal cycle of ozone (O₃) and nitrogen oxides (NO_x) at the surface are 7 8 captured by the model. Surface aerosol mass concentrations of sulphate (SO₄), nitrate (NO₃), 9 ammonium (NH₄), and organic matter (OM) are simulated with correlations larger than 0.55. 10 WRF/Chem captures the vertical profile of the aerosol mass concentration in both the planetary boundary layer (PBL) and free troposphere (FT) as a function of the synoptic condition, but the 11 12 model does not capture the full range of the measured concentrations. Predicted OM concentration is at the lower end of the observed mass concentrations. The bias may be attributable to the missing 13 aqueous chemistry processes of organic compounds and to uncertainties in meteorological fields. A 14 key role could be played by assumptions on the VBS approach such as the SOA formation 15 pathways, oxidation rate and dry deposition velocity of organic condensable vapours. Another 16 source of error in simulating SOA are the uncertainties in the anthropogenic emissions of 17 primary organic carbon. Aerosol particle number concentration (condensation nuclei, CN) is 18 overestimated by a factor 1.4 and 1.7 within PBL and FT, respectively. Model bias is most likely 19 attributable to the uncertainties of primary particle emissions (mostly in the PBL) and to the 20 nucleation rate. Simulated cloud condensation nuclei (CCN) are also overestimated, but the bias is 21 22 more contained with respect to that of CN. The CCN efficiency, which is a characterization of the ability of aerosol particles to nucleate cloud droplets, is underestimated by a factor of 1.5 and 3.8 in 23 the PBL and FT, respectively. The comparison with MODIS data shows that the model 24 overestimates the aerosol optical thickness (AOT). The domain averages (for one day) are 25 0.38±0.12 and 0.42±0.10 for MODIS and WRF/Chem data, respectively. Droplet effective radius 26 27 (R_e) in liquid phase clouds is underestimated by a factor of 1.5, cloud liquid water path (LWP) is overestimated by a factor 1.1-1.6. The consequence is the overestimation of average liquid 28 29 cloud optical thickness (COT) from few percent up to 42%. Predicted cloud water path (CWP) in all phase displays a bias in the range +41-80%, whereas the bias of COT is about 30 31 15%. In sensitivity tests were we excluded SOA, the skills of the model in reproducing the observed patterns and average values of the microphysical and optical properties of liquid 32 and all phase clouds decreases. Moreover, the run with SOA (NOSOA) shows convective clouds 33 with an enhanced content of liquid and frozen hydrometers, and stronger updrafts and downdrafts. 34

Considering that the previous version of WRF/Chem coupled with a modal aerosol module
 predicted very low SOA content (SORGAM mechanism) the new proposed option may lead to a
 better characterization of aerosol-cloud feedbacks.

1 **1** Introduction

2 It is well recognized that aerosol particles have a fundamental role in the climate system. They directly alter the budget of the radiation that reaches the Earth surface by scattering and absorbing 3 the incoming sunlight (Haywood and Boucher, 2000), and they indirectly affect cloud properties 4 and precipitation patterns, because they act as cloud condensation nuclei (CCN) (Rosenfeld et al., 5 6 2008; Lohmann and Feichter, 2005). Some aerosol species as black and brown carbon or mineral dust heat the atmosphere absorbing the solar radiation. The local warming may increase the 7 8 atmospheric stability, leading to a decrease in cloud cover through the so called semi-direct effect (Hansen et al., 1997). The change in global median radiative forcing associated to 9 10 anthropogenic aerosol particles since pre-industrial time is highly uncertain and it is estimated to be -0.9 W/m², with uncertainty within a range from -1.9 W/m² to -0.1 W/m² 11 (Boucher et al., 2013). 12

Experimental evidence of the influence of aerosols on cloud macrophysical and microphysical 13 properties have been shown in several works (Klarke and Kapustin, 2010; Christensen and 14 Stephen, 2011; Koren et al., 2011; Ten Hoeve et al., 2011; Li et al., 2012). Several modelling 15 16 studies show that aerosol particles have a strong impact not only on the climatic spatial-temporal scale, but also at short range on the regional scale (Baklanov et al., 2013). At regional scale, online 17 18 coupled mesoscale meteorology-chemistry models are useful tools to take into account aerosol feedback effects on both meteorology and atmospheric composition (Zhang et al., 2008; Baklanov 19 20 et al., 2013). WRF/Chem, which is the model used in this study, is one of such models (Grell et al., 2005; Fast et al., 2006; Chapman et al., 2009). In this work we present and evaluate some 21 22 developments of WRF/Chem for a better simulation of direct and indirect aerosol feedback.

The introduction of the aerosol-cloud-radiation feedback leads to non-linear chains and loops of 23 24 interactions between meteorological and chemical processes that are inhomogeneous in space and time (Baklanov et al., 2013). Furthermore, the prediction of meteorological variables significantly 25 improves when the direct and indirect aerosol effects are taken into account in numerical 26 simulation. For example, Yang et al. (2011) found that the inclusion of aerosol feedback produces 27 28 significant benefits in the simulated optical and microphysical properties of marine stratocumulus, and these improvements positively affect the simulation of the boundary layer structure and energy 29 30 budget. Yu et al. (2013) reported an improvement of the simulation of shortwave and longwave cloud forcing when the aerosol feedback is added to the model. 31

Recent studies conducted with global models, predict an important contribution of secondary organic aerosol (SOA) to direct and indirect aerosol feedback. O'Donnell et al. (2011) calculated an annual mean direct and indirect shortwave forcing of -0.31 W/m^2 and $+0.23 \text{ W/m}^2$, respectively. Biogenic SOA (BSOA) seems to play an important role on aerosol-cloud-radiation interaction. Scott et al. (2014) find that BSOA contribute for 4-21% to the global annual mean of CCN and 2-5% to global mean of cloud droplet concentration. They also attribute to BSOA a global mean indirect radiative forcing ranges from -0.77 W/m^2 to $+0.01 \text{ W/m}^2$.

Previous studies over USA and Europe demonstrated that the "traditional" configuration of 6 7 WRF/Chem (Grell et al., 2005) using the Secondary Organic Aerosol Model (SORGAM) (Shell et al., 2001), presents a negative bias of simulated PM_{2.5} mass, mostly attributable to a scarce 8 production of SOA (Grell at al., 2005; McKeen et al. 2007; Tuccella et al., 2012). Therefore, an 9 updated "chemistry option" with a more sophisticated treatment of SOA, fully coupled with 10 radiative and microphysics modules, is highly desirable. In section 2 of this work, we describe the 11 developments of WRF/Chem code carried out in order to simulate the direct and indirect effects 12 13 with the new SOA parameterization (based on the Volatility Basis Set, VBS, approach) recently implemented in the model by Ahmadov et al. (2012). In section 3, we describe the measurements 14 15 used to evaluate the model. In section 4, we evaluate the performance of the updated model through comparison with satellite data and with meteorological and chemical constituent measurements 16 17 performed in the frame of the European Integrated project on Aerosol Cloud Climate and Air quality interaction (EUCAARI) (Kulmala et al., 2011). The aim of the section 5 is to address the 18 two following questions: (1) Does the introduction of SOA particles interacting with radiation and 19 cloud processes improve the numerical prediction of cloud fields? (2) What is the potential impact 20 21 of SOA particles on cloud development? The final remarks are given in section 6.

1 2 WRF/Chem model

2 2.1 Description and upgrade

A pre-release of version 3.4 of Weather Research and Forecasting model with Chemistry model (WRF/Chem) (Grell et al., 2005) was used in this study. WRF/Chem is a community model that has many options for gas chemistry and aerosols. One of these has been updated in order to include a new chemistry option for simulation of direct and indirect effects with an updated parameterization for SOA production. The modifications introduced by Fast et al. (2006), Chapman et al. (2009), and Ahmadov et al. (2012) are the basis of our work. The technical details of the implementation are summarized in the Appendix A.

10 The gas-phase mechanism used is an updated version of the Regional Atmospheric Chemistry 11 Mechanism (RACM) (Stockwell et al., 1997). The inorganic aerosols are treated with the Modal Aerosol Dynamics Model for Europe (MADE) (Ackermann et al., 1998). The updated 12 13 parameterization for SOA production is based on the volatility basis set (VBS) approach (Ahmadov et al., 2012). MADE/VBS model has three log-normal modes: Aitken, accumulation and coarse. 14 The species treated are the sulphate $(SO_4^{=})$, nitrate (NO_3^{+}) , ammonium (NH_4^{+}) , elemental carbon 15 (EC), primary organic matter (POM), secondary organic aerosol (SOA, anthropogenic and 16 biogenic), chloride (Cl), sodium (Na), unspeciated PM_{2.5} (that includes the fine fraction of sea-salt 17 and soil dust), aerosol water, unspeciated coarse fraction of PM_{10} , soil dust and sea salt. 18

SOA parameterization implemented by Ahmadov et al. (2012) is based on a four bins 19 volatility basis set, with saturation concentration of 1, 10, 100, and 1000 µg/m³ at 300 K. 20 21 respectively. VOCs are oxidized by reactions with hydroxyl radical (OH), ozone (O₃), and nitrate radical (NO₃). Oxidized VOCs are anthropogenic (alkanes, alkenes, toluene, xylene, 22 23 and cresol) and biogenic (isoprene, monoterpenes, and sesquiterpenes). In each bin, organic mass is produced for two different regimes, high and low NO_x. In the former, organic peroxy 24 25 radicals react with nitrogen oxide (NO), conversely in the latter organic peroxy radicals react with other organic peroxy radicals. The organic matter produced is partitioned into aerosol 26 and gas phase assuming a pseudo-ideal partition. Organic condensation vapours (OCV) 27 produced by the oxidation of VOCs may be oxidized by reacting with OH, consequently 28 reducing the vapour pressure and shifting mass from high volatility bins to lower ones. The 29 default oxidation rate (or aging factor) used in the model is 1.0×10⁻¹¹ cm³/molec./s for both 30 anthropogenic and biogenic OCVs. The aging factor is one the key uncertainties in SOA 31 formation in the VBS approach. Other two factors important factors of uncertainty are those 32

related to the SOA formation pathways and to the dry deposition velocity of OCVs. The latter is assumed to be 25% of the dry deposition velocity of nitric acid (HNO₃).

3 The implementation of aerosol-cloud-radiation interaction within RACM/MADE/VBS follows the 4 methods described by Fast et al. (2006) and Chapman et al. (2009). We modified the WRF/Chem 5 code by preparing the inputs for the modules devoted to calculation of the aerosol optical properties 6 and the aerosol activation, starting from the mass of each aerosol type as predicted by new 7 chemistry package. In the approach of Fast et al. (2006), the three modes of the lognormal 8 distribution are divided into 8 bins, and at each chemical constituent of the aerosol mass is associated a complex refractive index. The refractive index is calculated for each bin with a volume 9 10 averaging. Mie theory is used to find the scattering and absorption efficiencies. Aerosol optical thickness (AOT), single scattering albedo and asymmetry parameter calculated with the optical 11 12 package developed by Barnard et al. (2010), are used as input to the radiative scheme (Goddard and RRTMG). Aerosol direct effect on longwave radiation is included following Zhao et al. (2010). 13

Aerosol-clouds interaction is a complex problem that involves the activation and resuspension of 14 the aerosol particles, aqueous chemistry and wet removal. Following Chapman et al. (2009), 15 aerosols within cloud drops are treated as "cloud borne". Aerosols that do not activate as cloud 16 droplets are treated as "interstitial". In WRF/Chem the activation process is based on the 17 18 parameterization developed by Abdul-Razzak et al. (2000, 2002). The number and mass concentration of the activated aerosols are calculated for each mode. The activation of aerosols is 19 20 based on a maximum supersaturation determined from a gaussian spectrum of updraft velocities and bulk hygroscopicity of each aerosol compound for all lognormal modes of 21 22 particles. Bulk hygroscopicity is based on the volume weighted average of the hygroscopicity of each aerosol component. In addition to the activated aerosols at environmental conditions, 23 24 the CCN spectrum is also determined, i.e. the aerosol particles acting as CCN at some given maximum supersaturations (0.02, 0.05, 0.1, 0.2, 0.5, and 1%) are calculated. 25

Within the dissipating clouds, the droplets evaporate and the cloud borne aerosols are resuspended to the interstitial state. Cloud borne aerosols and dissolved trace gases may be modified by aqueous chemistry. In this chemistry option, cloud chemistry is modelled using the scheme of Walcek and Taylor (1986). Wet deposition of trace gases and aerosols is treated in and below clouds. Within clouds the aerosols and trace gases dissolved in the water are collected by rain, graupel and snow. Below clouds the wet scavenging by precipitation is parameterized using the approach of Easter et al. (2004).

The simulation of the activation, resuspension and wet scavenging of the aerosol particles requires a 1 2 prognostic treatment of the cloud droplets. The prognostic treatment of the clouds droplet takes into account the losses due to collision, coalescence, collection and evaporation, and the source due to 3 nucleation. The Lin and Morrison microphysics schemes in WRF/Chem version 3.4 include the 4 prognostic treatment of the cloud droplet number concentration. The source due to nucleation is 5 parameterized following Ghan et al. (1997). Both microphysical schemes take into account the 6 7 autoconversion of cloud drops to rain dependent on the cloud droplet number. Therefore, aerosol activation affects both the rain rate and the liquid water content. The droplet number concentration 8 9 affects the calculation of the cloud droplet effective radius and cloud optical thickness (COT). The 10 interaction of clouds with the incoming shortwave radiation is done by linking the microphysics to 11 the radiation scheme. The reader should note that the contribution to SOA concentration by cloud chemistry is missing and the interaction of aerosol with ice nuclei is not taken into account in this 12 13 version of the model.

14 **2.2 Model configuration**

The simulations were conducted over three 1-way nested domains centred on The Netherlands, as shown in Figure 1. The coarse domain (D1) has 30 km horizontal resolution, domain 2 (D2) 10 km, and domain 3 (D3) is cloud resolving at 2 km resolution. In our runs we used 67 vertical levels extending up to 50 hPa.

The main physical and chemical parameterizations used are listed in Table 1. The model setup is the same for all three domains, except that no cumulus parameterization is used for D3. Wet scavenging and cloud chemistry from both parameterized updraft and resolved clouds are taken into account in D1 and D2. However in these domains the sub-grid cloud processes involve only the interstitial aerosol, i.e. the aerosol-cloud coupling is not considered in convective parameterization. Therefore, the indirect effects are well resolved for domains with resolution less than 10 km in the version of WRF/Chem used in this work.

As mentioned in Section 2.1, two key uncertainties in SOA production are deposition velocity and aging factor of OCVs. The first is assumed to be 25% (this value is called "deposition factor" in WRF/Chem) of dry deposition velocity of HNO₃, the second is set to 1.0×10^{-11} cm³/molec./s. Ahmadov et al. (2012) showed that reducing the aging factor of OCVs by 50%, daily average concentration of SOA decreases by 20%, and an increase of the dry deposition velocity of OCVs by a factor 4 the SOA concentration decreases by 50%. Deposition factor and aging are tunable parameters. After some sensitivity tests, we chose the default value for deposition factor and 5.0×10⁻¹¹ cm³/molec./s as aging of OCVs, because this combination
 minimizes the model bias with observations.

3 We simulated the period from 14 to 30 May 2008. We chose this period because aerosol and 4 cloud state-of-art measurements were available to evaluate the model (see Section 3). Moreover, during this period anticyclonic and cyclonic meteorological conditions were 5 observed which allows the evaluation of the model under varying conditions. The initial and 6 7 boundary meteorological conditions for D1 are provided by the European Centre for Medium range Weather Forecast (ECMWF) analyses with a horizontal resolution of 0.5° every 6 hours. The 8 chemical boundary conditions of D1 are taken from output of the global Model for Ozone and 9 10 Related Chemical Tracers (MOZART) (Emmons et al., 2010). MOZART output has been converted to RACM/MADE/SOA-VBS species by using the "mozbc" interface that may be downloaded from 11 12 the www.acd.ucar.edu/wrf-chem website.

A series of 30-h simulations were performed on each day starting at 00 UTC, with the first 6 h discarded as model spin up for meteorology. Meteorology of D1 is reinitialized from global analysis, while initial and boundary meteorology conditions for D2 and D3 are taken from D1 and D2, respectively. For all three domains, the chemical initial state is restarted from previous run, while the chemical boundary conditions of D2 and D3 are taken from D1 and D2, respectively. The first 13 days of May 2008 are also simulated to spin-up the chemistry.

19 2.3 Emissions

Anthropogenic emissions data are taken from TNO 2007 inventory (Kuenen et al., 2014). TNO is a gridded European inventory with resolution of 0.125°x0.0625°. It provides the anthropogenic emissions of NO_x, NMVOC, NH₃, SO₂, CO, primary PM_{2.5} and PM₁₀. EC and primary OC emissions are taken from a specific TNO database that is part of the EUCAARI project (Kulmala et al., 2011). These EC and OC emissions are size resolved, they are separated for particles less than 1 µm, particles in the range of 1–2.5 µm and 2.5–10 µm.

Horizontal and vertical interpolation, temporal disaggregation, NMVOC speciation, and aggregation of emissions into WRF/Chem species is done following Tuccella et al. (2012), with minor updates described in Curci et al. (2014a). In order to prevent spurious concentration of aerosol particles, the distribution of aerosol emissions into WRF/Chem modes is based on the low emission scenario of Elleman and Covert (2010). The 10% of the emitted aerosol mass is attributed to Aitken mode, and the 90% to the accumulation mode. 1 Biogenic emissions are calculated on line with Model of Emissions of Gases and Aerosols from

2 Nature (MEGAN) (Guenther et al., 2006). Dust and sea salt emissions from soil and seawater

3 are calculated on line in the simulations.

1 3 Measurements

We evaluated model performances in D3 against ground and aircraft based data collected in May
2008 during the Intensive Cloud Aerosol Measurement Campaign (IMPACT) in the frame of the
EUCAARI project (Kulmala et al., 2009). Model results were also evaluated against MODIS
satellite data.

An overview of the synoptic conditions of May 2008 over Central Europe is given by Hamburger et al. (2011). The first 15 days of May are characterized by an anticyclonic block, while the period from 16 to 31 is dominated by westerly wind and passage of several fronts. The days from 17 to 20 May are referred as "scavenged background situation" (Mensah et al., 2012), because they are dominated by northerly wind from North Sea associated to a low aerosol mass loading, due to wet scavenging. The period starting from 23 May is dominated by long range transport of dust from Sahara desert (Roelofs et al., 2010; Begue et al., 2014).

13 **3.1 Ground based measurements**

Meteorological and aerosol ground based measurements used in this study are collected in Cabauw (The Netherlands) at CESAR observatory Cabauw (Figure 1). CESAR observatory is a tower located at 51° 57'N, 4° 54' E, and -0.7 m a.s.l, at about 50 km south of Amsterdam. Measurements performed at CESAR observatory are typical of North-West Europe, and are representative of maritime and continental conditions depending on the wind direction (Mensah et al., 2012).

Standard meteorological variables are collected at 2 m, 10 m, 20 m, 40 m, 80 m, 140 m, and 200 m height. Besides, in this study we used the measurements of temperature and relative humidity profiles obtained with radiometer, and aerosol speciation from aerosol mass spectrometry (AMS) collected at 60 m (Mensah et al., 2012).

The model is also compared to ozone (O₃), nitrogen oxide (NO_x), nitric dioxide (NO₂), nitric oxide (NO), ammonia (NH₃), nitric acid (HNO₃), nitrous acid (HONO), and sulphur dioxide (SO₂) measurements issued by Cabauw Zijdeweg EMEP station (NL0011R). O₃ is measured with an ultraviolet absorbing ozone instrument, NO_x, NO and NO₂ with a chemiluminescence monitor, and NH₃, HNO₃, HONO and SO₂ with an online ion chromatograph.

Although Cabauw supersite provides very detailed measurements, it could not be enough to characterize the model performance near the surface. Therefore, WRF/Chem is also compared to daily PM_{10} data from 63 stations (10 rural, 25 suburban, and 28 urban) of AIRBASE network and to daily inorganic aerosol measurements issued at Bilthoven (NL0008R), Kollumerward (NL0009R), Vredepeel (NL00010R), and De Zilk (NL00091R)
 EMEP stations. SO₄ and NH₄ measurements are also perfomed at all these sites while NO₃
 measurements are available only at De Zilk.

4 **3.2 Aircraft measurements**

5 During May 2008, the French ATR-42 research aircraft performed 22 research flights (RF). In this 6 work we used 14 RF to evaluate the model. Their tracks are reported in Figure 1, while flight number, take-off and landing time are reported in Table S1. RF50, RF55, RF56, RF57, RF58, 7 8 RF61 and RF62 were conducted around Cabauw supersite, in order to study the origin and 9 characteristic of air masses collected at Cabauw. Other RFs were aimed at the study of aerosol 10 properties along a quasi-Lagrangian flight track, with west-east and north-south transects. ATR-42 was equipped with instrumentation suitable for aerosol-cloud interaction measurements. We used 11 12 the measurements from a condensation particle counter (CPC), the CPC3010 with a cutoff size of 15 nm, a Cloud Condensation Nuclei Counter (CCNC) for CCN number concentration 13 14 measurements, and an AMS. During the campaign a scanning mobility particle sizer (SMPS) was used to measure the number size distribution of aerosol particles with diameter in the 15 16 range of 0.02-0.5 µm, while the size distribution of aerosol particles larger than 100 nm was sampled with a passive cavity aerosol spectrometer probe (PCASP). SMPS and PCASP 17 measurements were combined in order to calculate the PM_{2.5} concentration using an average 18 aerosol density of 1.7 g/m³. A more exhaustive and detailed description of the whole campaign and 19 20 instrumentation is given by Crumeyrolle et al. (2013).

21 **3.3 Satellite measurements**

The model was also evaluated with MODIS-Aqua aerosol and cloud data. The Level 2 products used here are MYD04 and MYD06 collection 051 for aerosol and clouds, respectively. For ease of comparison with model output, both satellite and model data were regridded onto a common lat-lon regular grid. Model output is sampled at same time and location of each MODIS pixel, and then data are averaged in space and time over the same grid. **In this study the horizontal spacing of the common grid is 4 km.**

1 4 Model evaluation

Model results are compared to ground based and aircraft observations, as detailed in section 3. The statistical indices used are the Pearson's correlation coefficient (r), mean bias (MB), normalized mean bias (NMB) and normalized mean gross error (NMGE). The indices are defined in the Appendix and reported in Table 2.

6 4.1 Meteorology

7 Figure 2 shows the observed and modelled time series of hourly vertical profiles of temperature and 8 relative humidity at Cabauw supersite. WRF/Chem reproduces the day-to-day variation of temperature, before, after, and during the wet period. As shown by statistical indices (Table 2), 9 10 within the first 200 m, the model reproduces the temperature with a correlation of 0.93-0.95 and a mean bias of about -0.5°C. Looking at the free troposphere, we may realize that the model 11 underestimates the height of the 0°C isotherm (the black line on Figure 2a) in the first days of 12 simulation and during wet period by about 200-300 m (i.e., the model is colder than observed by 1-13 1.5 °C). Whereas immediately after the passage of the cold front, the temperature rise in the 14 simulation is slower than the observed one. The model performances in simulating surface 15 temperature are consistent with other European studies (e.g., Zhang et al., 2013a; Brunner et al., 16 2014). For example, Brunner et al. (2014) compared several meteorology-chemistry coupled models 17 with annual simulations at continental scale, and found that on Central Europe the predicted surface 18 temperature shows a correlation with observations in the range of 0.95-0.98, whereas the bias 19 20 ranges from -1 to 0.1°C.

The model reproduces the vertical structure of relative humidity (Figure 2b) over the whole period, 21 22 but it has the tendency to overestimate (underestimate) the higher (lower) observed values. This behaviour is more evident during scavenging days, when the relative humidity between 1000-2000 23 m is overestimated on average by 40%, but sometimes up to 60%. Errors of this magnitude in 24 simulating the vertical profile of RH were already found in previous works (Misenis and Zhang, 25 2011; Luo and Yu, 2011). Nevertheless, the model correlation and mean bias are 0.84 and +3.4%26 below 1000 m of altitude, 0.50 and +13% in the range of 1000-2000 m, 0.78 and +6.4% between 27 2000-3000 m. These values are comparable with those found by Fast et al. (2014) in the comparison 28 of WRF/Chem simulations to aircraft data. They have shown correlations in the interval of 0.49-29 30 0.70, while the bias from -7 to +0.1%. Near the surface, the relative humidity is simulated with a correlation of 0.87-0.92 and a positive MB of 3-4% (+6-8%). 31

The biases of the temperature and relative humidity could be due to a misrepresentation of soil (and 1 2 sea) temperature and soil moisture or by misrepresentation of the clouds and rain. These two problems are tightly coupled via land surface-atmosphere interaction. The errors in the simulation 3 of surface moisture and energy budget influences the fluxes of latent heat and moisture in the 4 atmosphere, affecting the local circulation, convective available potential energy (CAPE), cloud 5 formation and rain pattern (Pielke, 2001; Holt et al., 2006). Moreover, WRF/Chem tends to fail 6 7 simulating the thermodynamic variables near to coastline, because the uncertainties of land use data may play an important role (Misenis and Zhang, 2010). Initial and boundary meteorological 8 9 conditions may also play an important role. Bao et al. (2005) demonstrated that meteorological prediction is sensitive to used input data. They showed that varying the inputs used as initial and 10 11 boundary conditions, the mean daily model bias ranges from -2.71 to -0.65 K for the temperature and from -0.81 to 0.50 g/kg for vapour water content. 12

In Figure 3 we compare the time series of observed and predicted wind speed and direction at 13 several heights of Cabauw Tower. WRF/Chem captures the diurnal trend of wind speed, but it 14 overestimates the wind speed during night. Generally, we found the higher correlation at 10 and 200 15 m (0.78 and 0.76 respectively) and higher NMB between 20 and 40 m (+30-40%). The wind 16 direction is captured at all altitude levels of Cabauw tower, except of 18May when WRF/Chem 17 does not reproduce some rapid variations most likely due to local effects. The simulation of wind 18 direction tends to improve with height. Indeed, the correlation coefficient increases from 0.52 to 19 0.73 at 10 and 200 m, respectively, and the MB decreases from 27° below 40m to 17° at 200 m. The 20 performance in simulating the surface wind speed are consistent with those reported by Brunner et 21 22 al. (2014) in Central Europe. They have shown a correlation for 10 m wind speed in the range of 23 0.53-0.73 and a mean bias of 1-1.8 m/s. It is well recognized that WRF tends to overestimate the wind near the surface (e.g. Misenis and Zhang, 2010; Ngan et al., 2013; Brunner et al., 2014), but 24 25 the bias of the simulated wind speed could be also explained with uncertainties in the large-scale pattern of analysis used as input. Bao et al. (2005) showed that varying the meteorological inputs, 26 the mean daily model bias may range from -1.53 to -0.28 m/s and from -1.43 to 0.01 m/s for the u 27 28 and v component of the wind, respectively.

29 4.2 Surface gas phase and aerosol mass

Figure 4 displays the comparison between the observed and modelled hourly time series and
average diurnal cycles of O₃, NO_x, NO₂, NO, NH₃, HNO₃, HONO and SO₂ near the surface.

WRF/Chem reproduces the day to day variations of O_3 , capturing its decrease during scavenging period due to low photolysis rate caused by cloud presence, recovery in the following days, and a new decrease starting from 25 May. The average daily cycle is well reproduced with a morning minimum and an underestimated maximum in the afternoon. The model simulates the O_3 with a correlation of 0.72 and systematic negative bias on average about 3.4 µg/m³. This bias is observed in the afternoon and the evening, and is most likely due to the titration in these hours caused by higher NO_x concentration than observed.

8 WRF/Chem simulates the NO_x, NO and NO₂ time series with a correlation of 0.70, 0.65, and 0.66 9 respectively. The timing of NO_x daily cycle is reproduced. Indeed, the model captures the morning 10 and evening peaks as well as diurnal minimum of NO₂. The mean bias of modelled NO₂ is +1.25 11 μ g/m³ (+20%) and occurs in the afternoon and evening hours. Moreover WRF/Chem reproduces the 12 morning peak and diurnal decrease of NO, but the daily cycle is affected by an average positive bias 13 of 0.28 μ g/m³, with the average morning maximum overestimated of about 2 μ g/m³ (+33%).

Ammonia is reproduced with a correlation of 0.43. WRF/Chem underestimates the NH₃ during the 14 scavenging days and from 28 to 31 May. The model captures the daily cycle shape of NH₃ 15 concentration average, but the modelled NH₃ concentrations are constantly underestimated. The 16 negative mean bias over the whole period is on average about -4.75 $\mu\text{g/m}^3$ (-28%). WRF/Chem 17 reproduces the observed HNO₃ with a poor correlation. The measured mean diurnal cycle is flat, 18 conversely the model predicts a nocturnal minimum and diurnal maximum. The origin of model 19 bias in simulating NH₃ and HNO₃ is discussed below, together with a discussion on inorganic 20 aerosols. 21

22 The nitrous acid concentrations are not well captured by the model and are underestimated by 95%.

This bias could be partly explained by the inefficiency of NO oxidation, the only important reaction known to form HONO. Lin et al. (2014), indeed, demonstrated that the major sources of HONO are some unknown reactions that consume nitrogen oxides and hydrogen oxide radicals.

The model reproduces the measured SO₂ with a correlation of 0.48 and a positive bias of 0.68 μ g/m³ (+90%). The overprediction is most likely attributable to anthropogenic emissions. SO₂ is emitted mainly from isolated and elevated large point sources (Figure S1) that are immediately mixed in the model cell leading to overestimation outside of the local plume (Stern et al., 2008). This is demonstrated, for example, by the larger NMGE for SO₂ than NO_x (116% and 45%, respectively). NO_x is emitted near the surface by traffic and domestic heating. Therefore, NO_x emissions are subjected to a stronger temporal modulation than SO₂ point sources. The different uncertainties found for the involved species may depend on the choice of the chemical mechanism. Knote et al. (2014) compared several chemical mechanisms within a box model constrained by the same meteorological conditions and emissions, and found that the prediction of the O₃ diurnal cycle differs by less than 5% among the different mechanisms. Larger differences were found for other species. For example, the key radicals exhibit differences up to 40% for OH, 5% for H₂O₂ and 100% for NO₃ among the selected mechanisms.

Figure 5 shows the simulated and observed time series and diurnal cycle of aerosol sulphate, nitrate,
ammonium, and organic matter, at CESAR observatory. WRF/Chem simulates the measured SO₄,
NO₃, NH₄ with a correlation of 0.56, 0.68, and 0.66, respectively.

10 WRF/Chem captures the daily variations of SO₄ and its decrease during scavenging days. The shape of diurnal cycle is also reproduced, with the nighttime minimum and diurnal maximum. The mass 11 concentration of SO₄ is overpredicted for the whole period with a mean bias of 1.04 μ g/m³ (+90%). 12 The modelled SO₄ overestimation is directly attributable to SO₂ concentration overprediction. 13 Another potential source of the surplus of simulated SO₄ is related to an excessive production 14 within the clouds. Indeed, during scavenging days, the particulate sulphate is overestimated while 15 the predicted SO₂ does not show a bias in respect to the measurements. The overestimation of SO₄, 16 moreover, explains in part the negative bias of predicted NH₃. The excess of particle sulphate 17 18 consumes too much ammonia (Meng et al., 1997)

NO₃ is reproduced with a positive bias of 1 μ g/m³ (+72%). Looking at diurnal cycle, the modelled 19 nitrate is on average biased high in the daytime, with a peak in the afternoon. This maximum 20 21 appears to be correlated with HNO₃ maximum. Really, the HNO₃ peak is caused by evaporation of particulate nitrate formed in the upper PBL (where the conditions of lower temperature and higher 22 23 relative humidity are favourable to NO₃ formation), and mixed towards the surface by vertical mixing (Curci et al., 2014a; Aan de Brugh et al., 2012). Therefore, the unrealistic afternoon peak of 24 modelled nitric acid should result from a too rapid relaxation of aerosol-gas partitioning to 25 thermodynamic equilibrium (Aan de Brugh et al., 2012). 26

The behaviour of the simulated NH₄ is related to modelled trend of NO₃. It is biased high by 0.66 μ g/m³ (+66%). The NH₄ overestimation is related to NH₃ underprediction (Meng et al., 1997).

29 Similar performances are found in reproducing inorganic aerosols at other Dutch EMEP sites

30 (see Section 3.1). Daily SO₄ is simulated with an average correlation (3 stations) of 0.66 and a

positive bias of 35%. WRF/Chem simulates NH₄ with a correlation of 0.82 (4 stations) and a

bias of +43%. Predicted daily NO₃ (measured at only one station) is overestimated by 15%.

Organic matter is reproduced with a correlation coefficient of 0.75. WRF/Chem reproduces the right concentration during dry period, the decrease in the wet days, and following recovery. The mean bias is negative by about 0.4 μ g/m³ and it is attributable to days from 23 to 26 May. The discussion about the origins of OM bias is given in section 4.3, where we will discuss the model evaluation with aircraft data.

6 The reader should consider that aerosol composition measurements performed with the AMS 7 are representative of particles with diameter between roughly 100-700 nm, whereas the model 8 is evaluated with aerosol concentration representative of $PM_{2.5}$. Therefore, a bias could be 9 present in the comparison. This means that the bias found for inorganic aerosols could be 10 smaller than that reported above, conversely the OM bias could be larger of that found.

The model evaluation performed so far is representative of few points in the domain and does 11 not include other aerosol components like black carbon or primary PM. This could limit our 12 understanding of the model behaviour. In order to overcome this shortcoming, WRF/Chem is 13 also compared to daily PM₁₀ measurements of AIRBASE network (Figure S2). The model 14 captures the daily variations of PM₁₀, the PM₁₀ decrease during scavenging days, the 15 consequent recovery to reach back the background PM₁₀ concentration and the PM₁₀ 16 enhancement during long-range transport period. Indeed, the correlations are of 0.72, 0.73, 17 18 0.75 in rural, suburban, and urban zones, respectively. Model bias at rural stations is important in the last days of May 2008, indeed in these days (28-30 May) it is about -15 µg/m³ 19 20 (-30%). Conversely, at suburban and urban stations, the model presents a bias for the most part of the days of about 3-4 μ g/m³ (5-10%) that could partly explained by the missing source 21 22 of resuspension due to traffic

23 The results obtained here are statistically consistent with other modelling studies over Europe (e.g., Lecœur and Signeur, 2013; Zhang et al., 2013b; Balzarini et al., 2014). For example, the results of 24 European annual simulations of Balzarini et al. (2014) exhibited a correlation of 0.48, 0.60 and 0.56 25 for surface SO₄, NO₃ and NH₄, respectively. During EUCAARI campaign, Athanasopoulou et al. 26 (2013) reported a mean correlation of surface OM with observations of 0.56 and a mean bias of -0.5 27 $\mu g/m^3$, whereas Fountoukis et al. (2014) simulates the OM at Cabauw on May 2008 with a bias of 28 29 $0.3 \mu g/m^3$. Moreover, with regards to surface PM₁₀, our performances are comparable to those found for example by Im et al. (2014) over Europe with annual simulations in terms of 30 correlation, but are higher in terms of bias. Comparing PM₁₀ concentrations from several 31 models, Im et al. (2014) found correlations of 0.18-0.86 and 0.07-0.82, and bias of about -40% 32 and -50% for rural and urban sites, respectively. This improvement is most likely due to the 33

very high resolution used in this study with respect to the 23 km of Im et al. (2014), since the
 anthropogenic emissions inventory used is the same here and in Im et al. (2014) study.

3 4.3 Aloft aerosol mass concentration

The comparison of WRF/Chem to aircraft data is done interpolating the model output point by point along the flight track. Observed and modelled aircraft data are presented by using the box plots for planetary boundary layer (PBL) and free troposphere (FT) (see Figure 6). The height of the PBL was lower than 1600 m during the whole campaign (Crumeyrolle et al., 2013). Therefore, we considered for PBL and FT concentrations the data below and above 1600 m up to 3000-4000 m, respectively. This rough approximation of PBL height could affect the comparison of the model to data.

Figure 6 displays the observed and modelled box plots of the mass concentration of SO₄, NO₃, NH₄, and OM for PBL and FT. Their mean value, standard deviation, relative mass fraction, and correlation coefficients, averaged over the whole period of interest, are reported in Table 3.

14 The average concentrations of inorganic aerosols show little absolute error (2-8%) with respect to the observations in the PBL, while the NO₃ and NH₄ mean concentration presents a bias of +14% 15 and +20% (+0.3 and $+0.2 \mu g/m^3$), respectively, in the FT. The mean OM mass is biased low by a 16 factor 2 and 3 in the PBL and FT, respectively. The correlation coefficients of SO₄, NO₃, NH₄, and 17 OM are 0.39, 0.47, 0.43, 0.67 in the PBL and 0.23, 0.44, 0.42, 0.53 in the FT. These performances 18 are comparable with those found with WRF/Chem (but with a different chemistry package) in 19 California by Fast et al. (2014). They reported an absolute mean bias of about 0.01-0.2, 0.03-0.6, 20 $0.1-0.45, 0.2-0.57 \text{ }\mu\text{g/m}^3$ and a mean correlation of 0.42, 0.45, 0.44, and 0.72 for SO₄, NO₃, NH₄, 21 and OM, respectively. 22

Although the predicted aerosol mass of each species is within the range of the observed values for 23 most of the flights used in this study (see Figure 4), the model does not capture the full range of the 24 25 measured concentrations. This assertion is made quantitative by the standard deviations reported in 26 Table 3. The predicted standard deviations for each species are lower than observed. In the PBL, the observed and modelled standard deviations differ by 4-10% for inorganic ions and 55% for OM. In 27 28 the FT, the differences are higher than in the PBL. The model predicts standard deviations lower 29 than 10-40% for inorganic particles and lower than 65% for organic matter with respect to the measurements. 30

For the purpose of this analysis, it is also interesting to explore how the model reproduces the relative fraction of aerosol mass species with altitude (see Table 3). WRF/Chem overpredicts the relative fraction of the SO_4 and NO_3 by few percent in the PBL and about 10% in FT, while the relative mass of NH_4 is overestimated by 3% along the whole profile. The relative amount of OM is underpredicted by about 20% in both PBL and FT. The decrease of relative amount of NO_3 and increase of SO_4 with altitude is captured by the model. The modelled relative mass of NH_4 and OM is near constant with altitude as well as in the observations.

6 Looking at the individual flights, it is possible to note how the model captures the aerosol mass 7 trend as a function of the synoptic frame in both the PBL and FT, during the dry period, scavenging 8 days, and dust period characterized by southerly wind and passage of several fronts. The FT is a 9 layer mainly affected by long range transport and cloud contamination. Therefore, the relative small 10 bias in simulating aerosol inorganic mass in FT means that the model resolves quite correctly the 11 large scale transport and processes related to clouds.

Nevertheless, it should be noted that SO_4 is overestimated for 8 out of 14 RFs, while NO_3 and NH_4 are underpredicted for 7-8 out of 14 RF. This SO_4 overprediction is attributable to the SO_2 excess and to a potential overproduction within the cloud chemistry scheme. The negative bias of NO_3 and NH_4 could be explained by a low NH_3 regime, that limits the formation of the ammonium-nitrate.

The simulated OM concentration is always at lower end of the observed variability. Several factors 16 17 may explain this systematic bias. First of all, our simulations do not include the processing of organic compounds in aqueous chemistry. SOA may be produced in the clouds (Hallquist et al., 18 19 2009). Modelling studies suggest that the contribution of cloud chemistry to SOA budget is almost as much as the mass formed from the gas phase (Ervens et al., 2011). OM prediction is also affected 20 21 by meteorological errors. Bei et al. (2012) found that the uncertainties in meteorological initial conditions have significant impact on the simulations of the peaks, horizontal distribution and 22 23 temporal variation of SOA. The same authors demonstrated that the spread of the simulated peaks can reach up to 4.0 μ g/m³. Other uncertainties may be related to the VBS formulation. SOA 24 formation pathways is one these, indeed halving the SOA yields the concentration of SOA 25 decreases by 30% (Ahmadov et al., 2012). Moreover, the assumption on the deposition velocity 26 of the OCVs may play an important role in the uncertainties of SOA production. The OCV 27 deposition velocity in the version of the VBS implemented in WRF/Chem by Ahmadov et al. 28 29 (2012) is proportional to the deposition velocity of the HNO₃. The proportionality constant is a tunable parameter and in this work is set to the default value of 0.25. WRF/Chem prediction of 30 SOA mass is very sensitive to the choice of the proportionality constant (Ahmadov et al., 2012). 31 Previous studies have shown that SOA concentration is highly sensitive to the treatment of the 32 deposition velocity of OCV (Bessagnet et al., 2010; Knote et al., 2014b). Finally we note that OM 33

bias could be partially explained by the uncertainties in the anthropogenic emissions (e.g. bias or spatial distribution) of primary organic carbon and in the factor 1.6 used to convert them to primary OM (POM) (Turpin and Lim, 2001). The reader should also consider that the uncertainties in POM simulation affect the SOA formation. Indeed, the partition between OCV and SOA used in VBS approach depends on the total OM (Equation 1 of Ahmadov et al., 2012), thus if POM is underpredicted the resulting SOA could be underestimated.

7 In order to have a more complete overview of the model skill in reproducing the upper air 8 aerosol mass concentration, we compare also the observed and modelled PM_{2.5}. Figure 7 shows measured and predicted box plots of the PM_{2.5} concentration in PBL and FT. Modelled 9 10 PM_{2.5} is in the range of the observed values within the PBL expect during wet scavenging period when it is at the lower end of the observations. In FT conversely, predicted PM_{2.5} is at 11 12 the lower end of the observed profiles for the most part of the flights. The comparison between modelled and observed PM2.5 concentration within PBL and FT show good 13 correlations (0.75 and 0.80, respectively). 14

Model correlation with observations is high, 0.75 and 0.80 in PBL and FT, respectively. The absolute mean bias if of -7 μ g/m³ (30-35%) in both PBL and FT.

17 Although the box plot and statistical summary (Table 3) provide significant information on the model performances, the model skills in reproducing vertical profiles of aerosol properties 18 19 need to be evaluated. Therefore, we also look at model vertical profiles along the flight tracks. As an example, we have chosen the 14 May 2008 (RF50) for two reasons: first, there is a 20 21 relatively large contribution of OM, SOA (70-85% of OM), and CCN (see Figure 10), and 22 second, it is a day of high pressure, thus the interpretation of the results is not complicated by 23 cloud processes. Figure 8 displays the comparison of modelled vertical profiles of PM_{2.5}, SO₄, NO₃, NH₄ and OM along the flight track and measurements. WRF/Chem captures several 24 features present in the aircraft observations. Both observed and modelled PM_{2.5} exhibit a 25 maximum in a layer between 500 and about 2000 m. Model predicts the inhomogeneity of 26 chemical secondary species of PM2.5 displayed also in the observations: SO4 and OM 27 concentrations are relatively homogeneous within the PBL, whereas NO₃ and NH₄ show 28 enhanced concentrations in the upper levels of the PBL. For completeness, we note that 29 30 primary PM_{2.5} components (not shown) have the maximum close to the surface (first 500 m) and are diluted throughout the PBL. These results are qualitatively similar to those found by 31 Curci et al. (2015) above Milan (Italy). 32

1 4.4 Aloft aerosol particles

The comparison of WRF/Chem output with aircraft measurements of the number concentration of condensation nuclei (CN) and of cloud condensation nuclei (CCN) at 0.2% of supersaturation is done by using the boxplots as for aerosol mass. In this case the modelled and measured data are smoothed by using a 10 seconds running mean.

Figure 9 reports the comparison of observed and modelled CN within PBL and FT. The measured
and predicted average, standard deviation, and correlation of CN number over the whole period of
our analysis are reported in Table 3.

9 The model resolves the decrease of a factor 5-6 of CN concentration between the PBL and the FT. The differences in simulated concentrations between land and sea (RF51 and 52) are also captured 10 by the modelling system. Nevertheless, WRF/Chem overestimates, on average, the observed CN by 11 a factor 1.4 within PBL and 1.7 within the FT. The bias is less pronounced above the sea during the 12 (RF51 and RF52), where the anthropogenic sources are less important. Moreover, it should be noted 13 that in some cases, for example during the RF56, 57 and 58, the predicted CN are completely 14 15 outside the range of the observed values. In these cases the predicted CN are biased high by about a factor of 2-3. Predicted CN concentration shows a higher variability than measured, especially in 16 17 the free troposphere where the difference of modelled standard deviation is biased high by 155%. Anyway, the modelled CN concentration correlates well with the observed one (0.40 and 0.74 in 18 19 PBL and FT, respectively). These values are consistent with the 0.61 found by Luo and Yu (2011) studying the new particle formation and its contribution to CN with a version of WRF/Chem 20 21 including an advanced aerosol microphysical model.

Figure 10 shows the comparison of observed and modelled CCN at 0.2% of supersaturation. The measured and predicted average and standard deviation of CCN are showed in Table 3. The bias of simulated CCN_{0.2} appears more contained with respect to CN prediction, especially in the free troposphere. Figure S3 shows the comparison of the modelled vertical profile of CCN along the flight track of 14 May and observed CCN aboard the ATR42. WRF/Chem predicts relatively homogeneous profile of CCN in the PBL also shown by observations.

The aerosol particles that mostly contribute to CCN number are those of accumulation and coarse modes, and accumulation and coarse mode particles are also the major contributor to PM_{2.5} mass concentrations. Since PM_{2.5} is underestimated and CCN overestimated, CCN bias cannot depend on model errors in PM_{2.5}. The major uncertainties in predicted CCN arise from aerosol nucleation rate and primary emissions (Lee et al., 2013). Direct emission of aerosol particles is the key factor for CCN production in the PBL and near particle sources (Spracklen et al., 2006),

and account for 55% of the total global production (Merikanto et al., 2009). On the other hand, 1 nucleation and subsequent growth up to CCN size is an important mechanism of CCN formation in 2 many parts of the atmosphere (Sotiropoulou et al., 2006). Using several nucleation 3 parameterizations, Pierce and Adams (2009) showed that CCN on average varies by up to 4 12% within the PBL. At the same time, they also found that varying by a factor of 3 the primary 5 emissions, the CCN mean changes by 40% in the PBL. On the basis of these argumentations and 6 7 the correlation between predicted and observed CN being larger in the FT than in PBL (i.e. far from anthropogenic sources), we may speculate that the errors in the CCN prediction 8 9 arise mainly from the uncertainties in the primary emissions of the aerosol particles and in 10 their distribution in the lognormal modes.

The analysis of CCN efficiency reveals other interesting features in the model behaviour. The CCN 11 12 efficiency is given by CCN/CN ratio and represents the ability of aerosol particles to nucleate cloud droplets (Andreae and Rosenfeld, 2008). CCN efficiency observed during the studied RFs is in the 13 range of 0.02-0.33 for PBL and 0.18-0.41 in the FT, while the model predicts values of 0.03-0.17 14 and 0.04-0.18 for PBL and FT, respectively. In other words, WRF/Chem underestimates the CCN 15 efficiency by a factor of 1.5 and 3.8 within the PBL and FT, respectively. Moreover, the modelled 16 CCN efficiency, contrary to the observation, shows almost the same range of values within the 17 PBL and within the FT. 18

The calculated observed and modelled CCN efficiencies could be underestimated. In general, 19 20 the CCN efficiency should be computed with the aerosol population with size larger than the minimum activation diameter (Asmi et al., 2012). The latter depends on the aerosol type and 21 22 ranges from about 50 to 125 nm. We calculated the observed CCN/CN ratio with the measurements of CPC 3010 which gives the total number of particles larger than 15 nm, and 23 24 modelled CCN fraction is calculated with total particle number given by the sum of the three modes of the lognormal distribution (Aitken, accumulation and coarse). In order to better 25 26 characterize the relationship between CCN and corresponding aerosol population in the 27 model, predicted CCN efficiency was also calculated with particles of the accumulation and coarse modes (the most favored particles to act as CCN) and it was qualitatively compared to 28 observed efficiency during the IMPACT campaign computed with particles larger than 100 29 nm. Observed values of CCN efficiency are in the range of 0.28-0.4 and 0.38-0.6 in the PBL 30 and FT (Crumeyrolle et al., 2013), respectively. The simulated CCN fraction calculated with 31 the particles of the accumulation and coarse modes, is always underestimated with respect to 32 the observations, and it is in the range of 0.17-0.3 in PBL and 0.23-0.36 in FT. The model 33

deficiency in simulating the CCN/CN ratio could be attributable to the uncertainties in
 geometrical diameter and bulk hygroscopicity of the lognormal modes, and updraft velocity
 that lead to error in the prediction of minimum activation diameter of each mode.

4 4.5 Comparison with MODIS data

5 WRF/Chem output was also compared to aerosol optical thickness (AOT), cloud 6 microphysical and optical properties retrieved by MODIS-Aqua.

7 Figure 11 shows the comparison between the AOT at 550 nm measured by MODIS and the 8 corresponding AOT predicted by the model during the high pressure period on 14 May 2008. 9 WRF/Chem reproduces the spatial distribution of observed AOT, such as the lowest values in the southern part of the domain or the highest values around Cabauw, but underestimates the strong 10 11 gradient between eastern boundary and the centre of the domain. The model overestimates the regional mean of AOT, indeed the domain averages are 0.38±0.12 and 0.42±0.10 for MODIS and 12 WRF/Chem data, respectively. Unfortunately, good coverage of the D3 domain by MODIS was 13 achieved only on one day (14 May 2008), this does not allow us to have a general overview of 14 model skill in predicting AOT. In general, model intercomparisons revealed that a large part 15 of the uncertainties in simulating the AOT arises from the assumption on the mixing state. 16 For example, AOT computed with external mixing is larger by 30-35% of that calculated with 17 internal mixing assumption (Curci et al., 2014b). For typical atmospheric particle sizes and in 18 the visible wavelength range, the AOT is then expected to be lower under internal mixing 19 assumption (that is the assumption done in this work). Moreover, a 10% error in predicting 20 AOT may be attributable to the choice of species density, refractive index, and hygroscopic 21 growth factor (Curci et al., 2014b). 22

23 As the WRF microphysics scheme aerosol is only aware about liquid clouds, comparison among predicted cloud optical and microphysical properties with MODIS data was done 24 separately for liquid, excluding mixed clouds. Top liquid cloud effective radius (R_e) was 25 calculated from liquid water content (LWC) and cloud droplet number concentration 26 27 predicted by WRF/Chem. Liquid water path (LWP) was calculated by vertically integrating liquid cloud mixing ratios (water and rain water), while liquid cloud optical thickness (COT) 28 29 was estimated from LWC and Re. Since MODIS L2 data provide the total cloud water path (CWP), combined effective radius for all cloud types and total (water and ice) cloud optical 30 thickness (COT), the observed contribution to the liquid water cloud was separated by using 31 the retrieved top cloud phase, i.e. were discarded mixed clouds. 32

The comparison between the predicted and observed Re, LWP and liquid COT was done in 1 the scavenging background and long-range transport periods in the days when MODIS 2 cloudy pixel coverage was larger than 60%. Figure 12 shows the comparison of the averaged 3 values for 17-19 May 2008 period (1P). The same comparison has been done on averaged 4 values for 25-27 (2P) and 28-30 May (3P) and are reported in the Supplement (Figures S4 and 5 S5). WRF/Chem reproduces several features of the liquid cloud systems during 1P: the liquid 6 7 cloud distribution, the maximum values of Re close to the coast, maximum of LWP and liquid COT on the centre and at Nord-East of the domain. However, model results present a larger 8 9 spatial extension of liquid water cloud highest values off the Dutch and Belgian coast not observed in the MODIS data. During 2P the structure of the cloud system is not completely 10 reproduced by the model. Although R_e value magnitude is captured above the sea, 11 WRF/Chem underestimates the cloud droplet dimension on the land. Therefore LWP and 12 13 liquid COT structure on the sea is resolved by the model, whereas on the land they are too small compared to the observations. Finally, the model reproduces the average structure of 14 the cloud system in 3P, but LWP and liquid COT are overestimated on the Western part of 15 the domain. 16

As shown in Table 4, R_e values averaged over the entire domain is underestimated by the 17 model by a factor of about 1.5 during all three periods of interest. LWP, values averaged over 18 the entire domain, is also overestimated in all three cases by a factor of 1.1, 1.3 and 1.6 for 1P, 19 2P, and 3P, respectively. The R_e (LWP) underprediction (overestimation) may be due to the 20 overestimation of droplet number concentration that stems from overestimation of CN. 21 22 Another reason that could explain the positive bias of modelled LWP, is the inefficient 23 autoconversion of cloud water to rain, typical of the Morrison microphysical scheme (Saide et al., 2012). The consequence of the negative (positive) of R_e (LWP) is the overestimation of 24 25 average liquid COT by few percent for 1P and 3P, and 42% for 3P.

The biases found here are quite different from the WRF/Chem study by Yang et al. (2011) on the modelling of marine stratocumulus in Southeast Pacific. They have shown a bias of +30% in reproducing the COT, while LWP was underestimated by a factor 1.3. The reader should note that in Yang et al. (2011) the aerosol model adopted was different from the one used here and SOA formation was not included at all.

Figure 13 displays the distribution functions (DF) of $R_{e_{r}}$ LWP and liquid COT for 1P. The DF for 2P and 3P are reported in the Supplement (Figure S6 and S7). In all three periods analysed, the model captures the frequency of large droplets (R_{e} >18-20 µm), underestimates

the number of small droplets (8-10<R_e<18-20 μ m), and overestimates the frequency of very 1 small cloud droplets (R_e <8-10 µm) by more 30%. Whereas DF of the observed R_e presents the 2 maximum in the range of 8-13 µm, modelled DF shows the maximum values in 3 correspondence of the droplets with size less than 7-8 μ m. WRF/Chem captures the shape of the 4 distribution functions of LWP and liquid COT, but underestimates the maximum and 5 overestimates the higher and lower end of the distributions. Both variables show a variability higher 6 7 than the observations. The predicted standard deviations (Table 4) are about 2-3 and 1.5 times larger than those observed for LWP and liquid COT, respectively. This probably stems from the 8 9 large variability in simulated CCN.

Now it is interesting to analyse the model behaviour in reproducing the total CWP and COT given by contribution of all cloud phases. Modelled CWP was calculated by vertically integrating all cloud mixing ratio (water, rainwater, ice, snow and graupel). Predicted COT is given by the contribution of the liquid water and ice. The contribution of the liquid water was calculated as described above for liquid water cloud. The contribution of ice phase to COT was calculated following Ebert and Curry (1992).

Figure 14 displays the comparison between observed and predicted CWP and COT in P1, whereas the same figures for P2 and P3 are reported in the Supplement (Figures S8 and S9). Although for all three cases, the model reproduces with good approximation the shape and localizations of the cloud systems, CWP and COT are systematically overestimated (except COT in P2). As shown in Table 5, the predicted domain average of CWP presents, indeed, a bias of 62%, 41%, and 80% for P1, P2 and P3, respectively, whereas the bias of COT is about 15% in P1 and P3.

At this point of the analysis, although the aerosol-cloud interaction is a very complex nonlinear process, we are able to relate the model error in aerosol particles to the uncertainties in cloud prediction. The overestimation of CN leads to overprediction of the CCN. Higher number of CCN means clouds with higher number of cloud droplets, higher water content, smaller droplets and clouds optically deeper.

In addition to the uncertainties in aerosol particle simulation, one typical source of error in the prediction of cloud fields, are the choices related to the model setup. For example Otkin and Greenwald (2008) found a strong sensitivity of cloud properties while evaluating the response of WRF model to the permutation of several PBL and cloud microphysical schemes. Moreover, the same authors have shown that the low level clouds are sensitive to PBL parameterization, whereas the upper level clouds are sensitive to both PBL and microphysics schemes.

One element that may affect the model-satellite comparison are the uncertainties associated to the 1 retrieval. For example, in South-Pacific stratocumulus, MODIS overestimates the droplet effective 2 radius by 13-20% with respect to concomitant in situ measurements (Painemal and Zuidema, 2011; 3 King et al., 2013). The overestimation of COT by MODIS results in the overestimation of CWP 4 (King et al., 2013). Henrich et al. (2010) have shown systematic differences between MODIS data 5 6 and in situ observations. Indeed, analysing a system of thin cumulus clouds during EUCAARI 7 campaign, they also found that MODIS overestimates the droplet effective radius by a factor 2-3 and COT is 2-3 times lower than in situ measurements. 8 9

5 Impact of SOA particles on cloud prediction

The last part of this study focused on the evaluation of the impact of SOA on the simulation of 2 3 cloud fields. We performed sensitivity simulations during P1, P2 and P3 without the SOA (NOSOA), and compared them to the reference run (CTRL) discussed so far. NOSOA runs 4 5 are carried out only in the higher resolution domain (D3). The simulations of all three periods are initialized at 00 UTC with the same meteorological and chemical input data used for 6 7 CTRL, except chemical initial conditions that are restarted by previous NOSOA run. Each 8 period is preceded by 30 hours of simulation used as spin-up for D3 chemistry. The sensitivity simulation is performed zeroing the arrays pertaining to SOA. Thus, the SOA fields are not 9 affected by incoming SOA from domain boundaries or by local production. We did not 10 perform the sensitivity tests with SORGAM option because this model produces very little 11 SOA mass concentrations (Grell at al., 2005; McKeen et al. 2007; Tuccella et al., 2012). 12 Therefore, we may assume that simulations with SORGAM and without SOA (in VBS option) 13 are roughly equivalent. The advantage of this assumption is that the model is forced with 14 same initial meteorological conditions and boundary meteorological and chemical conditions 15 of the CTRL simulation. The use of SORGAM would require to run the model on all three 16 17 domains, leading to different results on D2 which is used to initialize D3. Finally, this would 18 introduce dependencies on the D3 input data making the comparison not directly comparable to the CTRL run. 19

The comparisons of R_e, LWP and liquid COT simulated of CTRL and NOSOA runs with 20 21 MODIS data are reported in Figures 12, S4 and S5. In general, the average spatial pattern of 22 these three variables is captured better by CTRL simulation with respect to NOSOA run, 23 especially in P1. Figures 13, S6 and S7 display the comparison between DFs of the cloud properties simulated by CTRL and NOSOA runs with those retrieved with MODIS 24 observations. The domain averages for each variable are reported in Table 4. NOSOA runs 25 show a domain averaged R_e larger than CTRL. DFs of the LWP are different between the 26 runs, but it is not clear if there are improvements in CTRL with respect to NOSOA run. Only 27 the domain averages allow to establish that LWP values predicted by CTRL run are closer to 28 the observed means than NOSOA. The presence of SOA in the numerical prediction improves 29 the DF of liquid COT with respect to NOSOA simulation in P1 and P3, whereas there are no 30 differences during P2. NOSOA has 10% and 3% more optically thin liquid clouds (liquid 31 COT<40) with respect to CTRL in P1 and P3, respectively. 32

Figures 14, S8 and S9 report the comparison of modelled CWP and COT of all cloud phases predict in CTRL and sensitivity runs with MODIS data. As well as for liquid phase, including SOA aerosol particles in the runs, the WRF/Chem skills to reproduce the observed pattern of observed CWP and COT increase. As shown in Table 5, domain averaged CWP and COT are larger up to about 50% in CTRL with respect to NOSOA.

6 Now it is interesting to explore the impact of SOA on the vertical structure of the cloud fields. As example we chose the 17 May because around 6 UTC, a frontal system associated to a trough 7 8 from North Sea crossed the Benelux area (Figure S10). In both runs, some isolated and shallow 9 clouds form during the night. When the cold front reaches Benelux around 5-6 UTC, a low pressure 10 centre forms (Figure S11). The winds rotate around the low pressure with speeds up to 14 m/s at 925 hPa height (Figure S12). A convective system develops around the vortex. Figure 15 shows the 11 12 maximum radar reflectivity (maximum dBZ) at 6 UTC for CTRL simulation, and the difference of maximum dBZ between CTRL and NOSOA runs. In general, the echo is larger for run with SOA, 13 i.e. the intensity of the storm is stronger in the CTRL run. Figures 16 and 17 show the vertical fields 14 of PM_{2.5} mass, vertical wind, liquid and frozen hydrometeor for both runs in the cross section A 15 displayed in Figure 15. These differences between both simulations (CTRL-NOSOA) along the 16 cross section A, are also displayed in Figures 16 and 17. The convection appears to be stronger in 17 the control simulation, with a larger amount of hydrometeors and stronger updrafts and downdrafts. 18 The larger differences in the simulated fields of vertical wind and hydrometeors occur in the same 19 location where occurs the enhancement of PM_{2.5} mass at cloud base (950-900 hPa), roughly at the 20 distance of 5-15 and 40-90 km away of the origin of the cross section A (Figures 16 and 17). The 21 22 results should be taken with caution because the aerosol-cloud interaction is treated only for liquid 23 clouds, the interaction of aerosol with ice phase is still missing in the model. Although the aerosolcloud interaction is a nonlinear process, it is possible to give an interpretation of the results with the 24 25 conceptual model for cloud invigoration proposed by Rosenfeld et al. (2008). The larger number of CCN in CTRL may curb the autoconversion rate of droplets to raindrops, therefore the beginning of 26 precipitation may be delayed with respect to NOSOA. This delay leads to a larger amount of 27 condensed water that crosses the freezing level and forms ice hydrometeors. The freezing process 28 warms the higher layers of the cloud through release of latent heat, whereas the melting process due 29 to falling of ice cools the lower levels. This thermodynamic disequilibrium enhances the upward 30 31 transport of heat. The enhanced conversion of CAPE to kinetic energy may yield the cloud invigoration found in the CTRL simulation. 32

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1 6 Summary and conclusions

Secondary organic aerosol particles play an important role in the aerosol-cloud-radiation interaction because they contribute to the global budget of radiation and cloud condensation nuclei (CCN). The introduction of SOA particles in numerical simulations has the potential to reduce the uncertainties on the prediction of meteorological fields and air quality. To this aim, a parameterization for SOA production based on the recent VBS approach was coupled with microphysics and radiative schemes in the WRF/Chem community model.

The performances of the updated model at cloud resolving scale (2 km horizontal resolution) were evaluated using ground- and aircraft-based measurements collected during the IMPACT-EUCAARI campaign and the data from the MODIS satellite instrument. The study focuses on the Benelux area, around the supersite of Cabauw, from 14 to 30 May 2008. The analysed period was characterized by few days of high pressure (14-15 May), followed by a scavenged background situation (17-20 May), and finally by long range transport of Saharan dust with the passage of southerly fronts (23-31 May).

The model reproduces the variations of meteorological variables as a function of the synoptic 15 16 frame. The model broadly captures the inter- and infra-diurnal variability of O₃ and NO_x at the surface. The concentration of NH₃ is underestimated. Concentrations of HNO₃ and HONO is 17 18 reproduced with poor correlation. Simulated SO_2 shows a positive bias of +90%, probably due to 19 overestimated point sources. Surface aerosol mass of SO₄, NO₃, NH₄, and OM is simulated with a correlation larger than 0.55. Their diurnal variations as a function of the synoptic frame are resolved 20 by the model. The bias of simulated inorganic aerosol mass is explainable together with error of 21 SO₂, NH₃, and HNO₃ in terms of anthropogenic emissions and the approximation to instantaneous 22 thermodynamic equilibrium. The performances in reproducing the surface aerosol mass found here 23 are comparable to other European studies where these variables are simulated with correlations 24 range from 0.5-0.7 (e.g., Lecœur and Signeur, 2013; Zhang et al., 2013b; Balzarini et al., 2014 for 25 inorganic species; Athanasopoulou et al., 2013; Fountoukis et al. 2014; Li et al., 2013; Knote et al., 26 27 2011; for organic aerosols).

The analysis of aircraft data reveals that WRF/Chem captures the aerosol mass trend both in the PBL and the free troposphere (FT). The predicted aloft aerosol mass of each species is within the observed values range, but the model does not capture the full range of the measured concentrations: the modelled standard deviations of aerosol mass are lower than those observed. Nevertheless, SO₄ (NO₃ and NH₄) mass is overpredicted (underpredicted) in more than half of the flights. SO₄ bias is attributable to the SO₂ excess and to a potential overproduction within the cloud

chemistry scheme. The negative bias of NO₃ and NH₄ could be explained by a low concentration of 1 NH₃ that limits the formation of the ammonium-nitrate. The simulated OM concentration is at 2 lower end of the observed mass. The bias is attributable to the missing aqueous chemistry processes 3 of organic compounds, uncertainties in meteorological fields, to assumptions on the VBS 4 approach such as the SOA formation pathways, oxidation rate and dry deposition velocity of 5 organic condensable vapours. Another source of error in simulating SOA are the 6 7 uncertainties in the anthropogenic emissions of primary organic carbon and in the factor (1.6) used to convert them to POM. In general, the statistical analysis reveals that the predicted average 8 9 concentrations of inorganic aerosols show absolute error of 2-8% in the PBL, while the NO₃ and NH₄ are simulated with a bias of +14% and +20% (+0.3 and +0.2 μ g/m³), respectively, in the FT. 10 The mean OM mass is underestimated by a factor 2 and 3 in the PBL and FT, respectively. These 11 biases are similar to those reported by Fast et al. (2014) comparing WRF/Chem (but with a different 12 chemistry package) to aircraft data performed over California. Indeed they found an absolute mean 13 bias of about 0.01-0.2, 0.03-0.6, 0.1-0.45, 0.2-0.57 $\mu g/m^3$ for SO_4, NO_3, NH_4, and OM, 14 15 respectively. However, we highlighted that the comparison of aerosol composition predicted by the model with AMS measurements could be affected by a bias because the model 16 17 concentrations are representative of PM_{2.5} particles and AMS collects aerosols with diameter only between 100-700 nm. 18

Condensation nuclei (CN) are overestimated by a factor of 1.4 and 1.7 in the PBL and FT, 19 respectively. However, in some cases, the predicted CN are overestimated by a factor of 3. 20 Predicted CN show higher variability than measurements. The model correlation with observed CN 21 22 is 0.40 and 0.74 in PBL and FT, respectively. These values are consistent with the 0.61 below 10 km of altitude found by Luo and Yu (2011) in the Eastern United States with WRF/Chem including 23 24 an advanced aerosol microphysical model. Model biases in predicting CN are attributable in large part to the uncertainties of primary particle emissions (mostly in the PBL) and to the nucleation 25 26 rate.

The bias of simulated CCN is more contained with respect to that of CN. The CCN efficiency (CCN/CN ratio) is underestimated by a factor of 1.5 and 3.8 in the PBL and FT, respectively. This could be due to a low number of particles in the accumulation and coarse mode or to uncertainties in the hygroscopicity of aerosol particles. CCN/CN ratio represents the ability of aerosol particles to nucleate in cloud droplets. Therefore, its misrepresentation may lead to issues in the simulation of cloud droplet number. In other words, the uncertainties in CCN efficiency is a general modelling problem that may prevent a correct representation of the amplitude of the aerosol-cloud interaction, i.e. the response of microphysical cloud properties to the variation of CCN load. This issue surely
 deserves and warrants further insight in the future.

3 The bias of simulated CN affects the prediction of droplet effective radius (Re), aerosol optical 4 thickness (AOT), cloud water path (CWP), and cloud optical thickness (COT). The comparison 5 with MODIS data shows that the model overestimates the AOT. The AOT averaged over the entire 6 domain on a single day are 0.38±0.12 and 0.42±0.10 for MODIS and WRF/Chem data, respectively. The domain averaged R_e of liquid cloud droplets is underestimated by a factor of 7 8 1.5 in all the periods examined in the main text. Modelled mean cloud liquid water path (LWP) is also overestimated by a factor 1.1-1.6. The consequence of the negative (positive) 9 10 bias of R_e (LWP) is the overestimation of average liquid COT by few percent up to 42%. CWP and COT of all cloud phases are systematically in 2 out of 3 periods analysed. Predicted 11 domain average of CWP presents a bias that ranges from 41-80%, whereas the bias of COT is 12 about 15% in P1 and P3. The overprediction of CWP could be due to the overestimation of 13 droplet number concentration that results from the overestimation of CN, and to inefficient 14 autoconversion of cloud water to rain. The reader should note that the model error found here are 15 different from the study conducted with WRF/Chem by Yang et al. (2011) on the modelling of 16 marine stratocumulus in Southeast Pacific where SOA formation was not included in the 17 simulations. Those authors reported a bias of +30% in reproducing the COT, while CWP was 18 underestimated by a factor 1.3. 19

In summary, the model behaviour of this new chemistry option in WRF/Chem in simulating the relationship between aerosol and cloud fields may be summarized by this way. The overestimation of CN results in the overprediction of the CCN. Higher number of CCN leads to clouds with higher number of cloud droplets, higher water content, smaller droplets and clouds optically deeper.

As test application of the new chemistry option, we performed a sensitivity simulation where SOA
 mass concentration is set to zero. The aim was to answer two questions:

1. Does the introduction of SOA particles improve the numerical prediction of cloud fields?

The introduction of SOA in the numerical simulations improves the predicted spatial pattern of microphysical and optical properties of cloud in liquid and all phases. NOSOA runs show an average R_e larger than CTRL. The analysis of LWP distribution function does not reveal a clear difference between CTRL and NOSOA simulations during the examined periods, only the domain averages allow to establish that LWP values predicted by CTRL run are closer to the observed means than NOSOA. 1 Conversely, including SOA in the numerical prediction improves the distribution 2 function of liquid COT with respect to NOSOA in 2 out of 3 cases. In these two cases, 3 NOSOA has up to 10% more optically thin liquid clouds (COT<40) with respect to 4 CTRL. Finally, with regards to CWP and COT (all phase), including SOA aerosol 5 particles in the runs, the WRF/Chem improves to reproduce the observed pattern of 6 observed CWP and COT.

7 2. What is the impact of SOA particles on cloud development?

8 The analysis was conducted on a convective system. The simulated radar reflectivity is 9 larger for run with SOA, i.e. the intensity of the storm is stronger in the CTRL run. The 10 CTRL simulation exhibits a larger amount of hydrometeors and stronger updrafts and 11 downdrafts. The larger differences in the simulated fields of vertical wind and hydrometeors 12 are associated to the larger differences of $PM_{2.5}$ mass located at the cloud base.

On the basis of the results discussed in this work, the option RACM/MADE/VBS coupled with cloud microphysics and radiation allows the WRF/Chem community to use a powerful tool for the study of the aerosol-cloud interactions, improved in terms of representation of the aerosol processes with respect to previous versions based on the RADM/MADE/SORGAM scheme.

For the future, there is still large space for improvements. For example, a more advanced treatment of deposition of organic condensable vapours is desirable. Moreover, the missing production of SOA in cloud is a gap that should also be filled. Finally, the extension of aerosol-cloud interaction to the ice-phase would lead to a complete representation of the aerosol indirect effects.

1 Code availability

The code updated, described, and evaluated here will be incorporated in the next available release of WRF/Chem. The users will be able to freely download the code from the WRF website (http://www2.mmm.ucar.edu/wrf/users/download/get_source.html). A general WRF/Chem user's guide is also available online (http://ruc.noaa.gov/wrf/WG11/).

6

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1 APPENDIX A: Technical details of coupling of VBS scheme with radiation and 2 microphysics schemes

The new chemistry option in namelist.input is *chem_opt=44*. It works with both *Lin* and *Morrison* microphysics scheme, *Goddard* and *RRTM* shortwave scheme, and *RRTM* longwave parameterization. Coupling of new scheme for SOA production with microphysics and radiative processes requires several modifications to code:

- The first step is to create a new chemistry option. The package racm_soa_vbs_aqchem_kpp
 (chemopt==44) has been added to /Registry/registry.chem together to some new model
 variables for the cloud-borne organic aerosols, called, for example, *asoa1cwi*, *asoa1cwj* etc.
- New chemistry package is a KPP option. Therefore, we created a new subdirectory in /*chem/KPP/mechanisms/racm_soa_vbs_aqchem* containing the files (*.spc, *.eqn, *.kpp, and *.def) where are defined the chemical model species and constants, chemical reactions in KPP format, model description, computer language, precision and integrator. The files are the same used in *racm_soa_vbs_kpp* package (chemopt==108).
- 3. The last step is to update the subroutines in *chem* subdirectory. In order to call necessary
 subroutines, the modules that we modified are:
- 17 chemics_init.F

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- module_input_chem_data.F
- mechanism_driver.F
- 20 cloudchem_driver.F
- 21 module_sorgam_aqchem.F
 - module_wetscav_driver.F
 - module_aerosols_soa_vbs.F
 - *aerosol_driver*.*F*
 - dry_dep_driver.F
 - module_mixactivate_wrappers.F
- emissions_driver.F
 - module_bioemi_megan2.F
- 29 optical_driver.F
 - module_optical_averaging.F
- 31 module_ctrans_grell.F

1 APPENDIX B: Statistical indices used in the model evaluation

2 Let Obs_i and Mod_i be the observed and modeled values at time *i*, and *N* the number of 3 observations.

• The Pearson's Correlation (r):

$$r = \frac{1}{N} \sum_{i=1}^{N} Z_i (Mod) \cdot Z_i (Obs)$$
$$Z(X) = \frac{X \cdot \langle X \rangle}{\sigma_X}$$

where X is a generic vector, Z(X) is its standard score, and σ_X is the standard deviation.

• Mean Bias:

8
$$MB = \frac{1}{N} \left(\sum_{i=1}^{N} Mod_i - Obs_i \right)$$

• Normalized Mean Bias (NMB):

10
$$NMB = \frac{1}{N} \sum_{i=1}^{N} \frac{Mod_i - Obs_i}{Obs_i} \times 100$$

• Normalized Mean Gross Error (NMGE):

12
$$NMGE = \frac{1}{N} \sum_{i=1}^{N} \frac{|Mod_i - Obs_i|}{Obs_i} \times 100$$

13

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2 TABLES

3 Table 1. Physical and chemical parameterizations used in the simulation

Physical Processes	WRF/Chem parameterizations
Cloud Microphysics	Morrison
Cumulus Cloud	New Grell (D1 and D2)
Longwave radiation	RRTM
Shortwave radiation	RRTM
PBL	MYNN
Surface Layer	Monin-Obukov
Surface	Noah LSM
Gas-phase Chemistry	Modified RACM-ESRL
Aerosol Chemistry	MADE/SOA-VBS
Biogenic Emissions	MEGAN

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2 T	Table 2.	Statistical	indices	of the	comparison	of WRF/Chem	to	observations	of tem	perature	(T).

3 relative humidity (RH), wind speed (WS), wind direction (WD), ozone (O_3) , nitrogen oxide (NO_x) ,

4 nitric dioxide (NO₂), nitric oxide (NO), ammonia (NH₃), nitric acid (HNO₃), nitrous acid (HONO),

sulphur dioxide (SO₂), particle sulphate (SO₄), particle nitrate (NO₃), particle (ammonium), and
particle OM, collected at Cabauw tower.

Variable	R	MB	NMB	NMGE
T (°C) at 2m	0.93	-0.66	-5.46	10.65
T (°C) at 10m	0.93	-0.67	-5.24	9.31
T (°C) at 20m	0.94	-0.56	-4.15	8.16
T (°C) at 40m	0.94	-0.46	-3.24	7.21
T (°C) at 80m	0.95	-0.32	-2.10	5.97
T (°C) at 140m	0.95	-0.26	-1.51	5.92
T (°C) at 200m	0.95	-0.45	-2.79	6.66
RH (%) at 2m	0.87	3.17	6.42	11.23
RH (%) at 10m	0.89	4.44	8.36	11.48
RH (%) at 20m	0.91	3.04	6.33	10.50
RH (%) at 40m	0.92	2.99	6.40	10.51
RH (%) at 80m	0.73	-1.04	2.13	13.34
RH (%) at 140m	0.92	2.81	6.40	11.19
RH (%) at 200m	0.91	2.99	7.44	12.50
WS (m/s) at 10m	0.78	0.67	27.90	35.56
WS (m/s) at 20m	0.66	1.27	40.89	49.32
WS (m/s) at 40m	0.67	1.26	32.64	42.04
WS (m/s) at 60m	0.72	1.23	24.42	38.99
WS (m/s) at 140m	0.74	1.21	28.66	41.55
WS (m/s) at 200m	0.76	1.13	27.79	41.48
WD (deg) at 10m	0.52	27.32	43.31	43.31
WD (deg) at 20m	0.53	24.80	40.48	40.48

WD (deg) at 40m	0.60	23.59	40.34	40.34
WD (deg) at 80m	0.67	20.22	30.34	30.34
WD (deg) at 140m	0.71	19.21	32.01	32.01
WS (deg) at 200m	0.73	17.18	28.46	28.46
O ₃ (μg/m ³)	0.72	-3.43	70.03	90.88
$NO_x (\mu g/m^3)$	0.70	0.43	19.76	44.77
NO (μg/m³)	0.65	0.28	116.08	167.59
NO ₂ (μg/m ³)	0.66	1.25	28.68	54.20
$NH_3(\mu g/m^3)$	0.43	-4.75	-27.72	42.94
HNO ₃ (μg/m ³)	0.21	-0.09	-1.22	108.65
HONO (μg/m³)	0.05	-0.56	-95.37	95.37
SO ₂ (µg/m³)	0.48	0.68	90.20	116.33
SO ₄ (μg/m ³)	0.56	1.04	92.2	95.4
NO ₃ (μg/m ³)	0.68	1.00	72.4	94.3
NH₄ (μg/m³)	0.74	0.66	63.0	67.3
OM (μg/m³)	0.75	-0.42	3.21	29.9

Table 3. Observed and modelled mean values, standard deviations, and relative amount (expressed
as percentage) of aerosol species, number of aerosol particles, cloud condensation nuclei, over the
whole period of the aircraft campaign in boundary layer and free troposphere.

	Bo	oundary Layer		F	ree Troposphere	
	Observation	WRF/Chem	r	Observation	WRF/Chem	r
SO ₄ (μg/m ³)	3.1±2.4 (19%)	3.2±2.1 (24%)	0.39	2.6±2.0 (29%)	2.5±1.2 (38%)	0.23
NO ₃ (μg/m ³)	4.5±5.4 (28%)	4.6±5.1 (34%)	0.47	1.3±3.0 (14%)	1.5±2.7 (23%)	0.44
$NH_4 (\mu g/m^3)$	2.6±2.2 (16%)	2.4±2.1 (19%)	0.43	1.4±1.5 (16%)	1.2±1.2 (19%)	0.42
OM (µg/m³)	6.1±5.8 (37%)	3.0±2.6 (23%)	0.67	3.7±4.5 (41%)	1.3±1.4 (20%)	0.52
PM _{2.5} (μg/m ³)	23±13	16±10	0.75	19±17	12±9	0.80
CN (10 ³ #/cm ³)	6.7±5.0	9.4±5.4	0.40	1.0±1.1	1.7±2.8	0.74
CCN (10 ³ #/cm ³)	0.6±0.5	0.9±0.8	0.45	0.3±0.3	0.3±0.3	0.73

1

2 Table 4. MODIS and modelled mean values and standard deviations of droplet effective radius at

3 cloud top, liquid cloud water path and liquid cloud optical thickness, on 17-19, 25-27, and 28-30

4 May 2008.

		R _e (μm)		LWP (g/m2)			COT		
	MODIS	CTRL	NOSOA	MODIS	CTRL	NOSOA	MODIS	CTRL	NOSOA
17-19 May	10.2±3.5	7.5±3.5	8.5±4.3	230±343	242±343	208±352	32±22	33±33	21±24
25-27 May	13.7±3.7	9.1±4.4	9.8±4.9	200±166	256±502	273±480	22±19	23±30	21±27
28-30 May	10.7±3.9	8.4±4.1	7.8±3.6	141±128	224±327	243±447	19±14	27±27	24±30

Table 5. MODIS and modelled mean values and standard deviations of cloud water path and cloud
optical thickness of clouds in all phases, on 17-19, 25-27, and 28-30 May 2008.

		CWP (g/m2)			СОТ	
	MODIS-Aqua	CTRL	NOSOA	MODIS-Aqua	CTRL	NOSOA
17-19 May	207±203	336±531	218±359	26±20	30±33	19±23
25-27 May	235±182	331±533	244±418	21±16	17±23	16±21
28-30 May	206±201	370±714	262±523	21±16	24±26	21±27

2 FIGURES

3

1



Figure 1. Panel (a) shows the three nested model domains used for simulations. D1 is 30 km
resolution, D2 10 km, and D3 2 km. The black star indicates the Cabauw supersite. Panel (b) is a
zoom of D3 and shows the track of every flights used in this study, yellow circle represents Cabauw

8 supersite.



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Figure 2. Observed and simulated time series of vertical profile of the temperature (a) and relative
humidity (b), observed at Cesar observatory. The black line on the temperature profile represents

- 4 the 0° C isotherm.
- 5





3 Figure 3. Observed (black lines) at Cesar Tower and simulated (red lines) time series of wind speed

4 (a) and wind direction (b) at different height.





3 Figure 4. Observed and simulated time series of gas-phase species (a) and their average diurnal

4 cycle (b), at Cabauw Zijdeweg EMEP station (NL0011R).





Figure 5. As Figure 4, but for aerosol mass speciation at Cesar Observatory observed at 60m height.





Figure 6. Box plot of SO4, NO3, NH4, and OM mass concentrations measured by AMS aboard the 4 ATR-42 (blue) and simulated by WRF/Chem (red) within boundary layer (panel a) and free troposphere (panel b). The x-axis reports the day of May 2008 (black) and the number of the 5 6 research flight (red). The whisker plots denote median, 25th and 75th percentiles, $1.5 \times$ (inter-7 quartile range), and outliers. The squares represent the mean values.



Figure 7. Same as Figure 6 but for observed and simulated $PM_{2.5}$ mass concentrations. The blue colour represents the observations will the model is displayed in red colour.



3 Figure 8. Vertical profiles (shadow) along the flight track of 14 May (RF50) of modelled PM_{2.5}(a),

4 SO₄ (b), NO₃ (c), NH₄ (d), and OM (e). The circles are the measurements collected aboard the 5 ATR42.



4 Figure 9. Same as Figure 6 but for observed and simulated condensation nuclei (CN) 5 concentrations. The blue colour represents the observations while the model is displayed in red 6 colour.



Figure 10. Same as Figure 6 but for observed and simulated cloud condensation nuclei (CCN)
concentrations at 0.2% of supersaturation. The blue colour represents the observations, the model is
displayed in red color.





Figure 11. Aerosol optical thickness at 500 nm from MODIS-Aqua (panel a) and WRF/Chem simulations (panel b) on 14 May 2008.



Figure 12. 17-19 May 2008 averages of droplet effective radius at cloud top (first row), liquid water
path (second row), and liquid cloud optical thickness (third row), retrieved using MODIS-aqua
observations (first column), predicted by model in the references run (CTRL, second columns) and
sensitivity test without SOA (NOSOA, third column).



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3 Figure 13. 17-19 May 2008 averages of observed and simulated distribution function of droplet

effective radius at cloud top (a), liquid water path (b) and liquid cloud optical thickness (c). The
black line represents the observations retrieved by MODIS, blue and red colours correspond to
model predictions from the reference run (CTRL) and sensitivity test without SOA (NOSOA),

7 respectively.



Figure 14. As Figure 12, but for clouds in mixed phase.





3 Figure 15. Maximum dBZ at 6 UTC of 17 May for CTRL run (a) and difference of maximum dBZ

4 between CTRL and NOSOA simulations (b). The solid black line represents the cross section A

5 used to plot vertical profiles reported in Figure 13.

6



3 Figure 16. Vertical profile of PM2.5 mass (color) and wind (vector) for CTRL run (a), NOSOA run

- 4 (b) and their differences (c). The fields are extracted along the cross section A (see Figure 12) at 6
 5 UTC of 17 May. The x-axis reports the west-east distance in km along the cross section.



3 Figure 17. As Figure 16, but for water (color), and frozen (black contours) hydrometeors.