# Evaluation of operational on-line-coupled regional air quality models over Europe and North America in the context of AQMEII phase 2. Part I: Ozone

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#### 1. Introduction

Tropospheric ozone (O<sub>3</sub>) is an important secondary air pollutant produced by photochemical oxidation of volatile organic compounds (VOC) and carbon monoxide (CO) in the presence of nitrogen oxides (NO<sub>x</sub>). It has implications on climate and health and therefore its levels are subject to regulatory monitoring in Europe (EU) and North America (NA). The regulatory O<sub>3</sub> levels are still exceeded in a number of cities and are especially a concern in growing urban areas (European Environmental Agency, 2013). Air quality models (AQMs) are valuable tools to investigate the complex and dynamic interactions between meteorology and chemistry leading to O<sub>3</sub> pollution episodes at multiple temporal and spatial scales. In the last decade, AQM development started shifting from off-line-coupled models where the meteorological forcing for chemistry was produced off-line by a separate meteorological model, to fully-coupled online models, which are able to simulate the feedbacks between chemistry and meteorology, taking the advantage of increased computational power (Zhang, 2008; Baklanov et al., 2014). The use of on-line models for O<sub>3</sub> predictions is beneficial, as O<sub>3</sub> not only depends on emissions and chemistry but also on regional transport, clouds, photolysis and vertical mixing in the boundary layer, all of which can be more realistically represented in an on-line model (Wong et al., 2012; Zhang et al., 2013). The impact of aerosols on the radiation, and therefore temperatures and photolysis rates, can significantly impact the gas-phase chemistry affecting O<sub>3</sub> and secondary aerosol formation (Kim et al., 2009, 2011). Thus, simulating these feedbacks can lead to more realistic O<sub>3</sub>, NO<sub>x</sub> and aerosol levels that are relevant to policy applications. The wide use of regional AQMs for supporting policy, abatement strategies and forecasting justifies the increased need for online models, which can simulate feedback mechanisms, and especially account for the effect of aerosols on radiative balance and photolysis (e.g. Hodzic et al., 2007).

The Air Quality Model Evaluation International Initiative (AQMEII) served to promote policy-relevant research on regional air quality model evaluation across the atmospheric modeling communities in Europe and North America through the exchange of information on current practices and the identification of research priorities (Galmarini and Rao, 2011). As part of this collaboration, standardized observations and model outputs were made available through the ENSEMBLE system (http://ensemble2. jrc.ec.europa.eu/public/) that is hosted at the Joint Research Centre (JRC). This web-interface allows temporal and spatial analyses of individual models as well as their ensemble operators (Bianconi et al., 2004; Galmarini et al., 2012). The first phase of AOMEII was focused on the evaluation of off-line coupled atmospheric modeling systems against large sets of monitoring observations over Europe and North America for the year 2006 (Solazzo et al., 2012; Vautard et al., 2012; Solazzo et al., 2013; Hogrefe et al., 2014). As summarized in Schere et al. (2012), the intercomparison model results for O<sub>3</sub> suggested a strong influence of chemical boundary conditions for ozone, whose bias extends far into the interior of the modeling domains, especially during winter months. The observed variance as well as the daily ozone cycle was underestimated by the majority of models. Night-time, overcast, and stable conditions led to poor model skill in reproducing ozone mixing ratios over both continents. Stable atmospheric boundary layers have been notoriously difficult to simulate in numerical weather prediction models (Holtslag et al., 2013), but they are highly relevant in the context of air quality modeling. Due to the high sensitivity of air pollutants to the representation of stable boundary layers, online coupled modeling could be of great use to tackle this problem in the future.

The second phase of AQMEII extends this model assessment to on-line-coupled air quality models. In this study, we analyze  $O_3$ concentrations provided by eight on-line-coupled models, which have been run by sixteen independent groups from Europe and North America (while a companion study is devoted to the analyses of particulate matter, Im et al., 2015). The models made use of the same input emissions and chemical boundary conditions, in an effort to reduce the impact of uncertainties originating from these inputs to model results among different groups. The goal of the study is to evaluate the performances of widely used operational on-line coupled models in Europe and North America in simulating  $O_3$  levels on a sub-regional and seasonal basis employing an experimental set-up with common anthropogenic emission and boundary conditions. The surface levels and vertical profiles simulated by the individual models as well as their ensemble mean and median are compared with the observational data provided by the ENSEMBLE system.

# 2. Materials and methods

## 2.1. Participating models

In the context of AQMEII2, twelve modeling groups from EU and four modeling groups from NA (Table 1) have applied their modeling systems to simulate hourly O<sub>3</sub> concentrations for the year 2010 over the EU and NA continental scale domains (Fig. 1). Among all participants, seven groups from EU and two groups from NA applied the same model system (WRF-CHEM), but with different settings such as different shortwave radiation schemes, gas-phase chemical mechanisms and aerosol modules. The WRF-CHEM community applied a common horizontal grid spacing of 23 km over Europe and 36 km over North America. Other modeling groups applied different grid spacings, ranging from  $12 \times 12 \text{ km}^2$  to ~50  $\times$  25 km<sup>2</sup> as seen in Table 1. The simulations were conducted for continental-scale domains of Europe and North America covering continental U.S., southern Canada and northern Mexico (Fig. 1). To facilitate the cross-comparison between models, the participating groups interpolated their model output to a common grid with 0.25° resolution for both continents. Model values at observation locations were extracted from the original model output files for comparison to observations (described below).

#### 2.2. Emissions and boundary conditions

For the EU domain, the recently updated anthropogenic emissions for the year 2009 (http://www.gmes-atmosphere.eu/; Kuenen et al., 2015; Pouliot et al., 2015) were applied by all modeling groups and are based on the TNO-MACC-II (Netherlands Organization for Applied Scientific Research, Monitoring Atmospheric Composition and Climate – Interim Implementation) (CO), ammonia (NH<sub>3</sub>), total non-methane volatile organic compounds (NMVOC), nitrogen oxides (NO<sub>x</sub>), particulate matter (PM<sub>10</sub>, PM<sub>2.5</sub>) and sulfur dioxide (SO<sub>2</sub>) from ten activity sectors are provided on a latitude/longitude grid of  $1/8^{\circ} \times 1/16^{\circ}$  resolution. Emission inventories for the NA domain were provided by US EPA and Environment Canada. The 2008 National Emission Inventory (http://www.epa.gov/ttn/chief/net/2008inventory.html) and the 2008 Emission Modeling Platform (http://www.epa.gov/ttn/chief/ emch/index.html#2008) with year specific updates for 2006 and 2010 were used for the US portion of the modeling domain. Canadian emissions were derived from the Canadian National Pollutant Release Inventory (http://www.ec.gc.ca/inrp-npri/) and Air Pollutant Emissions Inventory (http://www.ec.gc.ca/inrp-npri/ donnees-data/ap/index.cfm?lang=En) values for the year 2006. These included updated spatial allocations for Canadian mobile emissions (Zhang et al., 2012) for the emissions of NH<sub>3</sub> (Makar et al., 2009), as well as other updates (Sassi et al., 2010). Mexican emissions were 2008 projected forward from a 1999 inventory (Wolf et al., 2009). Seven pollutants (CO, NO<sub>x</sub>, NH<sub>3</sub>, SO<sub>2</sub>, PM<sub>10</sub>, PM<sub>2.5</sub>, and VOC) were used to develop the model ready emission inventory. Further details and analyses of the anthropogenic emissions used in both domains are provided in Pouliot et al. (2015). Annually-integrated anthropogenic emissions for both domains are presented in Table 2 while the spatial distribution of NO<sub>x</sub> emissions for the EU and NA domains are depicted in Fig. 1. Table 2 shows that anthropogenic emissions per  $\rm km^2$  in EU are larger than those in NA, except for PM<sub>10</sub>. Particularly NO<sub>x</sub> and NH<sub>3</sub> emissions in EU are more than a factor of two larger than those in NA. Consistent temporal profiles (diurnal, day-of-week, seasonal) and vertical distributions were also made available to maintain consistency among different groups. NMVOC speciation factors were applied by all groups individually with a recommendation to follow the NMVOC speciation profiles for EU by Visschedijk et al. (2007). The temporal profiles for the EU anthropogenic emissions were provided from Schaap et al. (2005). Chemical and temporal profiles for the EPA anthropogenic emissions were based on the 2007v5 modeling platform (http://www.epa.gov/ttn/chief/emch/index.html#2008).

framework. Annual emissions of methane (CH<sub>4</sub>), carbon monoxide

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Modelling systems	participated	to AQMEII2 and	their configurations
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	Groups	5 Domain	Model	Grid spacing	First layer height (m)	Biogenic model	Gas phase	Photolysis	Model reference
<b>M</b> 1	AT1	EU	WRF-CHEM	23 km	24	MEGAN	RADM2 (Stockwell et al., 1990)	Fast-J (Wild et al., 2000)	Grell et al., 2005
M2	CH1	EU	COSMO-ART	0.22°	20	Guenther et al., 1993	RADM2K (Vogel et al., 2009)	GRAALS + STAR (Vogel et al., 2009)	Vogel et al., 2009
M	DE3	EU	COSMO- MUSCAT	0.25°	20	Guenther et al., 1993	RACM-MIM2 (Karl et al., 2006)	Fast-J	Wolke et al., 2012
M4	DE4	EU	WRF-CHEM	23 km	24	MEGAN	RADM2 modified (Forkel et al., 2015)	Fast-J	Grell et al., 2005; Forkel et al., 2014
M5	ES1	EU	WRF-CHEM	23 km	24	MEGAN	RADM2	Fast-J	Grell et al., 2005
M	ES2a	EU	NMMB-BSC- CTM	0.20°	45	MEGAN	CB05 (Yarwood et al., 2005)	Fast-J	Jorba et al., 2012
M7	ES3	EU	WRF-CHEM	23 km	24	MEGAN	CBMZ (Zaveri and Peters, 1999)	Fast-J	Grell et al., 2005
M	IT1	EU	WRF-CHEM	23 km	24	MEGAN	CBMZ	Fast-J	Grell et al., 2005
MS	IT2	EU	WRF-CHEM	23 km	24	MEGAN	RACM (Stockwell et al., 1997)	Fast-J	Grell et al., 2005
<b>M</b> 1	0 NL2	EU	RACMO LOTOS-EUROS	$0.5^{\circ} \times 0.25^{\circ}$	25	Beltman et al., 2013	CB-IV modified (Sauter et al., 2012)	Poppe et al., 1996	Sauter et al., 2012
M1	1 SI1	EU	WRF-CHEM	23 km	25	MEGAN	RADM2	Fast-J	Grell et al., 2005
<b>M</b> 1	2 UK4	EU	MetUM-UKCA RAQ	0.22°	20	TNO	UKCA RAQ (Savage et al., 2013)	Fast-J	Savage et al., 2013
M1	3 CA2f	NA	GEM-MACH	15 km	20.66	BEIS	ADOM-II (Lurmann et al., 1986)	Dave, 1972	Makar et al., 2015a,b
<b>M</b> 1	4 US6	NA	WRF-CMAQ	12 km	19	BEIS3.14	CB05-TU (Whitten et al., 2010; Sarwar et al., 2011)	Binkowski et al., 2007	Wong et al., 2012
<b>M</b> 1	5 US7	NA	WRF-CHEM	36 km	55-60	MEGAN	MOZART (Emmons et al., 2010; Knote et al., 2013)	fTUV (Tie et al., 2003)	Grell et al., 2005
<b>M</b> 1	6 US8	NA	WRF-CHEM	36 km	38	MEGAN	CB05	fTUV	Grell et al., 2005; Wang et al., 2014



**Fig. 1.** Annual NO<sub>x</sub> emissions (tonnes/grid) overlaid with the rural monitoring stations used for model performance evaluation in EU (a) and in NA (b). The red circles show EU1/NA1, yellow diamonds show EU2/NA2, green squares show EU3/NA3 and black triangles show EU4/NA4. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Each modeling group used their own biogenic emission module as detailed in Table 1. The majority of the models used the online MEGAN2 model (Model of Emissions of Gases and Aerosols from Nature version 2; Guenther et al., 2006), two groups used the BEIS v3.14 model (Biogenic Emission Inventory System; Schwede et al., 2005) and one group (NL2) used the Beltman et al. (2013) biogenic model. It should be noted that UK4 group used the offline simulated biogenic emissions provided by the Beltman et al. (2013) model. In addition to the biogenic emissions algorithm used in the models, they may also differ in the databases used for vegetation. Feedbacks may have a significant influence on biogenic emissions; reductions in biogenic isoprene emissions of 20% were found with the introduction of the aerosol indirect effect (Makar et al., 2015a). The biogenic isoprene emissions calculated on-line by each group show a large variability as shown in Table 2 that may lead to large differences in the simulated O<sub>3</sub> levels. Curci et al. (2009) showed that different biogenic emission models may lead to a factor of 2 differences in domain-integrated isoprene emissions over Europe while difference can be up to a factor of 5–6 locally. They estimated that these differences on average may lead to an increase of 2.5 ppb in domain-mean surface O<sub>3</sub> levels and up to 10–15 ppb locally in the Mediterranean. Hourly biomass burning emissions were provided by the Finnish Meteorological Institute (FMI) fire assimilation system (http://is4fires.fmi.fi/; Sofiev et al., 2009). More details on the fire emissions and their uncertainties are discussed in Soares et al. (2015). The fire assimilation system provides only data for total PM emissions. Emissions of other species (CO, NO, NH<sub>3</sub>, SO<sub>2</sub>, NMVOC) were therefore deduced based on mass ratios relative to PM following Andreae and Merlet (2001). NMVOC speciation followed Wiedinmyer et al. (2011) combined with the mapping to different chemical mechanisms proposed by Emmons et al. (2010). Note that the ES2a model does not include biomass burning emissions and as it does not contain aerosols leading to a lack of effect of aerosols on photolysis rate calculations and therefore producing overestimated O3 within the fire plumes (Badia and Jorba, 2015). Lightning NO<sub>x</sub> is included in the UK4 model (O'Connor et al., 2014) as well as in the global MACC model used for the boundary conditions as described below.

3-D daily chemical boundary conditions were taken from the MACC re-analysis (Inness et al., 2013). The MACC re-analysis (referred to as MACC hereafter) has been produced by assimilating satellite observations of  $O_3$ , CO and  $NO_2$  in the coupled system IFS-MOZART (Flemming et al., 2009). As pointed out in Inness et al. (2013), the assimilation of satellite-corrected  $O_3$  greatly improved the ozone total columns and stratospheric profiles but did not change significantly the surface levels because of the limited signal from this region in the assimilated satellite observations. The chemical species available in the reanalysis included

#### Table 2

Annual anthropogenic emissions (ktons km<sup>-2</sup> yr<sup>-1</sup>) provided by TNO-MACC-II inventory and biogenic isoprene emissions (ktons km<sup>-2</sup> yr<sup>-1</sup>) integrated over the EU and NA domains.

Species	EU	NA
СО	614	478
NO <sub>x</sub> <sup>a</sup>	277	120
NMVOC	230	85
NH <sub>3</sub>	109	31
SO <sub>2</sub>	109	70
PM <sub>2.5</sub>	49	29
PM <sub>10</sub>	69	76
ISOP <sup>b</sup>	2.4–24.9	0.02-8.1

<sup>a</sup> Only anthropogenic NO<sub>x</sub> is reported.

<sup>b</sup> The groups that provided isoprene emissions are AT1, CH1, DE3, IT2, NL2 and UK4 for the EU domain and CA2f, US6 and US7 for the NA domain.

 $O_3$ ,  $NO_x$ , CO,  $CH_4$ ,  $SO_2$ , NMVOCs, sea-salt, dust, organic matter, black carbon and sulfate. NMVOC species had to be lumped or disaggregated according to the individual models' chemical speciation and particulate matter size discretization.

#### 2.3. Observations

Measurements of hourly surface O3 concentrations for the year 2010 in EU were taken from the European Monitoring and Evaluation Programme (EMEP; http://www.emep.int/) and the European Air Quality Database (AirBase; http://acm.eionet.europa.eu/ databases/airbase/) and in NA from the Canadian National Atmospheric Chemistry (NAtChem) Database and Analysis Facility operated by Environment Canada (http://www.ec.gc.ca/natchem/) that contains measurements from the Canadian National Air Pollution Surveillance Network (http://maps-cartes.ec.gc.ca/rnspanaps/data.aspx), the Canadian Air and Precipitation Monitoring Network (http://www.ec.gc.ca/natchem/), the U.S. Clean Air Status and Trends Network (http://java.epa.gov/castnet/clearsession.do), the U.S. Interagency Monitoring of Protected Visual Environments Network (http://views.cira.colostate.edu/web/DataWizard/), and the U.S. Environmental Protection Agency's Air Quality System database for U.S. air quality data (http://www.epa.gov/ttn/airs/ airsaqs/detaildata/downloadaqsdata.htm). In the AQMEII2, rural, urban and suburban background stations were extracted from the EMEP and AirBase networks. Given the coarse native grid resolutions used in different models (Table 1), data from only rural background stations was used in the comparisons. Stations that have more than 90% data availability have been selected for the comparisons. Regarding the whole simulation domains, hourly surface O<sub>3</sub> observations were provided by 510 and 200 stations in EU and NA, respectively. A geographical break-down into four subregions for each continent has also been defined based on the climatological and source characteristics. The geographical breakdown of these stations overlaid with the annually-averaged anthropogenic NO<sub>x</sub> emissions is shown in Fig. 1. Model evaluation statistics were computed for the four sub-regions separately. The European sub-region EU1 is characterized by north-western European sources with a transition climate between marine and continental and hosts 102 stations. Sub-region EU2 covers the north-eastern and central Europe sources as well as Germany with 277 monitoring stations. Sub-regions EU3 and EU4 are characterized by a Mediterranean type climate. Sub-region 3 covers southwestern sources including Italy (30 stations) while sub-region 4 covers the East Mediterranean with 101 stations. The North American sub-region 1 (NA1) covers the western U.S. and south western Canada with 80 stations. It includes large emission sources along the coast as well as polluted hot spots like Los Angeles that are characterized by poor air quality. NA2 consists of U.S. plains and covers 36 monitoring stations and is characterized by a continental and humid climate. NA3 consists of north eastern NA and south central Canada and is characterized by the largest emissions in North America and contains 60 monitoring stations. Finally NA4 covers the south eastern part of U.S., consisting of 24 monitoring stations.

To evaluate the capability of the modeling systems to simulate the tropospheric distribution of  $O_3$  concentrations, comparisons against  $O_3$  soundings provided by the World Ozone and Ultraviolet Radiation Data Centre (WOUDC: http://www.woudc.org/) have been carried out. Ozone concentration data from nine stations in EU and six stations in NA have been used for the comparisons. For an optimal comparison with observations, model profiles were computed by averaging only over the available observation hours. The participants were required to provide their data at fixed heights up to 18 km above the ground in order to be comparable. However, due to the coarse vertical resolution of some models in the upper troposphere and not simulating the stratospheric chemistry, the analyses are performed only for the first 9 km above ground.

## 2.4. Statistical analyses

To score the individual model performances as well as those of the ensemble mean and median, the following statistical parameters have been calculated: Pearson's correlation coefficient (PCC: Eq. (1)), root mean square error (RMSE: Eq. (2)); normalized mean standard error (NMSE: Eq. (3)) and normalized mean bias (NMB: Eq. (4)).

$$PCC = \left[\frac{\frac{1}{N}\sum_{i=1}^{N} \left(O_{i} - \overline{O}\right) \left(P_{i} - \overline{P}\right)}{\sigma_{0}\sigma_{P}}\right]$$
(1)

RMSE = 
$$\sqrt{\frac{1}{N} \sum_{i=1}^{N} (P_i - O_i)^2}$$
 (2)

$$\text{NMSE} = \frac{\sum_{i=1}^{N} (P_i - O_i)^2}{N \times \overline{P} \times \overline{O}} \times 100$$
(3)

NMB = 
$$\frac{\sum_{i=1}^{N} (P_i - O_i)}{\sum_{i=1}^{N} O_i} \times 100$$
 (4)



**Fig. 2.** Observed and simulated annual mean diurnal profiles (a,d), box plots (b,e) and soccer diagrams (c,f) for surface levels ozone mixing ratios in EU (upper panel) and NA (lower panel). Mn and Md represent the mean and median ensembles, respectively. EU0 and NA0 represent the two respective continents. Different colors represent the different sub-regions. Note the differences in scales. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Table 3

Statistical comparisons of observed and simulated annual domain-mean hourly surface O<sub>3</sub> and domain- and annually-integrated O<sub>3</sub> dry deposition over EU and NA in 2010.

Members	r	NMSE (%)	NMB (%)	RMSE <sup>a</sup>	Dry deposition (Tg km <sup>-2</sup> )
M1/AT1	0.86	2.66	-4.92	9.57	NP
M2/CH1	0.82	8.03	-18.30	15.42	0.28
M3/DE3	0.68	6.37	-2.12	15.02	0.13
M4/DE4	0.83	3.17	-1.64	10.62	2.24
M5/ES1	0.86	4.08	-11.41	11.44	2.18
M6/ES2a	0.83	6.37	-7.71	14.59	2.79
M7/ES3	0.86	4.29	-12.07	11.69	1.82
M8/IT1	0.85	4.57	-12.45	12.03	NP
M9/IT2	0.84	6.21	-15.80	13.76	1.77
M10/NL2	0.89	2.83	-4.34	9.90	0.14
M11/SI1	0.87	2.38	-3.78	9.10	1.91
M12/UK4	0.85	7.88	2.30	17.08	NP
EU Mean	0.86	3.22	-7.70	10.37	
EU Median	0.86	3.23	-8.69	10.33	
M13/CA2f	0.85	1.45	2.43	4.02	0.09
M14/US6	0.84	2.15	1.14	4.85	0.10
M15/US7	0.78	4.36	-4.56	6.72	0.15
M16/US8	0.88	8.11	-22.36	8.26	3.05
NA Mean	0.83	3.70	-11.98	5.94	
NA Median	0.87	2.62	-9.51	5.07	

 $^a\,$  RMSE is in units of  $\mu g\,\,m^{-3}$  for EU and ppb for NA.

where *P* and *O* denote model predictions and observations, respectively. The PCC is a measure of associativity and allows gauging whether trends are captured, and it is not sensitive to bias; RMSE is a measure of accuracy and, because it is squared, is sensitive to large departures. NMSE and NMB are normalized



operators, useful for comparing scores coming from time series of different lengths, as those produced over different areas and/or with different time span. The comparison is performed individually for the two domains and their sub-regions for the whole year of 2010 and on a seasonal basis, in order to identify which regions and/or seasons lead to systematic errors.

# 3. Results and discussion

# 3.1. Surface ozone analyses

Observed and simulated diurnal cycles of surface  $O_3$  concentrations averaged over the whole simulation period (2010) are shown in Fig. 2a, d for EU and NA, respectively. Models are labeled by the ID of the respective modeling group, with each ID corresponding to a member of the overall model ensemble. In the same figures, the MACC IFS-MOZART global model (MACC) and the ensemble mean and median are also shown. Note that the MACC model is not considered in the ensemble calculations.

# 3.1.1. Europe

Most models capture reasonably well the shape of the annual diurnal cycle over Europe as seen in Fig. 2. The temporal variations on all time scales were captured successfully as seen in Table 3 (PCC > 0.80), although the predicted  $O_3$  levels are generally underestimated by up to 18%. Only one group (UK4) slightly overestimates the yearly-averaged observed surface  $O_3$  levels by 2% while the other groups have underestimations up to 18%. The largest underestimations are calculated for IT2 (by 16%) and CH1



(by 18%) groups. Other groups have mean normalized biases within the  $\pm 5\%$ -15% range suggested by Russell and Dennis (2000). Fig. 2a shows that the underestimations generally occur both during day and night hours, which is expected to some extent given the coarse horizontal resolution (Qian et al., 2010). The exceptions are AT1, DE4. SI1 and UK4 that overestimate the nighttime levels. The MACC model underestimates the nighttime levels as also reported in Inness et al. (2013). Overestimation of nighttime O<sub>3</sub> levels can be due to the overestimation of NO<sub>2</sub> concentrations under low-NO<sub>x</sub> conditions leading to overestimated O<sub>3</sub> concentrations (e.g. DE4). Fig. 2a shows that the underestimations generally occur both during day and night hours, which is expected to some extent given the coarse horizontal resolution (Qian et al., 2010). The exceptions are AT1, DE3, DE4, SI1, and UK4 that overestimate the nighttime levels. The MACC model underestimates the nighttime levels as also reported in Inness et al. (2013). The small overestimation of nighttime O<sub>3</sub> levels for AT1 and SI1 can be attributed to the underestimation of nocturnal ozone titration in urban areas with high NO<sub>x</sub> emissions for the QSSA solver that was applied for these simulations. For DE4, where a modified version of this solver (Forkel et al., 2015) has been applied, the overestimation of nighttime ozone can be attributed to a general overestimation of NO<sub>2</sub> concentrations under low-NO<sub>x</sub> conditions. This is also the case for the DE3 model during the nighttime, where this overestimation is probably related to difficulties of the meteorological model to simulate nighttime vertical mixing accurately and, furthermore, to comparatively small dry deposition fluxes for O<sub>3</sub> simulated by the model (see Table 3). It should be noted that the ES2a model does not include

anthropogenic aerosols and secondary aerosol formation and neither aqueous chemistry, leading to a more oxidized atmosphere. Furthermore, the heterogeneous formation of HNO<sub>3</sub> through N<sub>2</sub>O<sub>5</sub> hydrolysis, which is an important sink of NO<sub>2</sub> during night, is not considered in ES2a (Badia and Jorba, 2015). As a consequence, the ES2a model overestimated the annual domain-mean NO<sub>2</sub> levels by 15% while the rest of the models underestimate NO<sub>2</sub> by 9%-45%. The overestimation of surface O<sub>3</sub> levels by the ES2a model can also partly be due to the coarser vertical resolution of its first layer (45 m) compared to other models (Table 1). The general underestimation may be partly attributed to biases in meteorological variables, including an overestimation of surface wind speeds by all models by up to 60% and a general slight underestimation of surface temperatures by less than 1 K (Brunner et al., 2015). Such a small temperature bias, however, will affect ozone levels by no more than a few ppb (Sillman and Samson, 1995). A common feature of all groups is that the daily maximum is simulated earlier than the observed maximum. Differences in O<sub>3</sub> predictions between the WRF-CHEM models suggest that the choice of the chemical mechanism plays an important role in the model performance. WRF-CHEM runs using RADM chemical mechanism (AT1, ES1 and SI1) produced higher concentrations than runs using RACM (IT2) and CBMZ (ES3 and IT1) mechanisms (Baró et al., 2015). These differences may partly be attributed to VOC emission preprocessing. WRF-CHEM is designed to ingest VOC emissions for RADM2 and then, in case of other mechanisms, the emissions are chemically specified to the final scheme, possibly leading to a degradation of the reactivity in the VOC mixture. There are also differences in



Fig. 4. Geographical distributions of observed and simulated annual surface level ozone mixing ratios in NA. Note the differences in scales.



**Fig. 5.** Soccer diagrams for the seasonal and geographical model performances in EU: a) winter, b) spring, c) summer and d) autumn. Mn and Md represent the mean and median ensembles, respectively. EU0 and NA0 represent the continental levels. Different colors represent the different sub-regions. Note the differences in scales. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



**Fig. 6.** Soccer diagrams for the seasonal and geographical model performances in NA: a) winter, b) spring, c) summer and d) autumn. Mn and Md represent the mean and median ensembles, respectively. EU0 and NA0 represent the continental levels. Different colors represent the different sub-regions. Note the differences in scales. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



Fig. 7. Observed and simulated seasonal diurnal O<sub>3</sub> profiles in a) winter, b) spring, c) summer and d) autumn over EU2.

the microphysics schemes among the different WRF-CHEM configurations used, leading to different cloudiness and therefore to different temperature and radiation acting on the O<sub>3</sub> production (Brunner et al., 2015; Baro et al., 2015). Makar et al. (2015a) and Wang et al. (2015) found that models including the simulation of indirect effects tended to have lower O<sub>3</sub> concentrations during the summer production period than those with the direct effect only, or those with no feedbacks. This is due to the reduction of NO<sub>2</sub> mixing ratios during daytime and near-surface temperatures, resulting from the reduction of solar radiation (Wang et al., 2015). Dry deposition of O<sub>3</sub> is also investigated for the models that provided deposition data (CH1, DE3, DE4, ES1, ES2a, ES3, IT2, NL2 and SI1) in order to explain the differences in simulated O<sub>3</sub> levels among the models (Table 3). The results show a negative relation between underestimation and dry deposition; i.e. the underestimation increases with decreasing deposition, suggesting that other terms aside from deposition were controlling the O3 concentrations (chemistry, vertical diffusion etc.).

The model performances are also assessed against the observed variability in box-and-whisker plots of Fig. 2b and e. The plot shows the frequency distribution of observed and simulated surface  $O_3$  mixing ratios. The spread of the data in the European case is largest in CH1, ES2a and UK4 (Fig. 2b). The majority of other models show a much lower spread, which also tends to be lower than the observed spread. Data from MACC are associated with a larger spread compared to the observations in both domains, suggesting a better representation of local processes by regional models as well as an indication of an exaggerated seasonal cycle simulated by the MACC

model. The larger spread in some models as compared to others is partially related to the amplitude of the diurnal ozone cycle, which tends to be larger in models simulating a more stable and shallow nocturnal PBL such as the global MACC model (Inness et al., 2013). A larger amplitude may also be expected for models with a higher vertical resolution. The NMB vs NMSE plot (also known as the soccer diagram) for EU (Fig. 2c) shows that the models have mean biases below 30% and mostly below 15%. The geographical analyses for the EU domain presented in Fig. 3 show that for the majority of models, the underestimation is mainly originating from sub-region EU2 (north Eastern Europe) while in sub-region EU4 (East Mediterranean), most models overestimate the observed mean. The underestimation, particularly in EU1 and EU2 could be partly due to the chemical boundary conditions (Fig. 3) as discussed in more detail in Section 3.3.

#### 3.1.2. North America

The hourly  $O_3$  temporal variability over the whole simulation period is also well captured (PCC > 0.78) by all groups for the NA domain (Table 3). The CA2f model overestimates the nighttime surface  $O_3$  concentrations and underestimates the daytime levels with a slight overall overestimation of 2% while other groups underestimate the nighttime levels (Fig. 2d). NMSE values are below 10% for all the groups while NMB values are within ±15% except for the US8 model, which underestimates the surface  $O_3$  levels by 22%. The box plots for the NA case (Fig. 2e) shows that the MACC model has the highest variability while CA2f is characterized with the smallest spread. Larger biases in US7 and US8 can also be partly



Fig. 8. Observed and simulated seasonal diurnal O<sub>3</sub> profiles in a) winter, b) spring, c) summer and d) autumn over NA3.

attributed to their coarser resolution (36 km) compared to other NA models (Table 1). In the NA case, according to the soccer diagrams (Fig. 2f), all groups and sub-regions are characterized with biases lower than 25% except for US8. The geographical break down presented in Fig. 4 shows that the US8 model underestimates in all sub-regions. The MACC model also shows a general underestimation in all sub-regions except for NA4. Regarding the dry deposition of O<sub>3</sub> (Table 3), the results suggest that the large underestimation by US8 can be partly due to the relatively large O<sub>3</sub> dry deposition simulated by the model, acting as a significant sink. As analyzed in Yahya et al. (2015a,b) and Wang et al. (2015), other factors that contribute to underpredictions of O<sub>3</sub> by the US8 model include large underpredictions of afternoon temperatures, low MACC boundary conditions of  $O_3$ , the overpredictions of the  $NO_x$  titration effects on O<sub>3</sub> during nighttime, possible underestimates in biogenic VOCs and wildfire emissions, and the inclusion of aerosol indirect effects. The lower spread in CA2f seems to be due to overpredicting the lower end of the O<sub>3</sub> range compared to the observations, in regions NA3 and NA4.

#### 3.2. Seasonal vs. geographical surface ozone variations

#### 3.2.1. Europe

Inter-seasonal variations of surface  $O_3$  concentrations are analyzed for each sub-region in order to understand how the model bias varies depending on the region and season. The results for the EU domain are depicted in Fig. 5. The temporal variability in Europe is better captured in all models in summer and autumn

(PCC = 0.8 - 0.9) than in winter and spring (PCC = 0.6 - 0.8). There is a systematic overestimation of the observed concentrations in autumn by up to 35%, particularly by the DE4 model. In winter (Fig. 5a), O<sub>3</sub> mixing ratios in EU2 are underestimated by more than 50% by three groups (CH1, ES2a and UK4), which also underestimate systematically in other sub-regions, probably due to the bias from the boundary conditions from the MACC model. The MACC model underestimates by largest during winter (by 8%-55%) and overestimates by largest in autumn (by 8%-25%). Regarding EU1, all groups are within the 30% bias range. Spring and summer O<sub>3</sub> mixing ratios (Fig. 5b,c) in all EU sub-regions are similarly reproduced by all groups, with error below 30%. In autumn, the majority of the models are biased high. In northern Europe (EU1 and EU2), the majority of the models underestimate O<sub>3</sub> levels in all seasons with the DE4, UK4, and ES2a models overestimating during summer. There is a general overestimation in autumn in the EU1 subregion by all models except for CH1 and IT2. The models NL2, DE4, UK4 and ES2a overestimate the summertime O<sub>3</sub> levels in southern Europe. The East Mediterranean region (EU4) is characterized by overestimated O<sub>3</sub> levels, in particular during autumn. The results show that the largest underestimations were calculated for the EU2 region, which is characterized with large anthropogenic emissions in the Eastern Europe that may lead to overestimated O<sub>3</sub>titration by NO<sub>x</sub>.

## 3.2.2. North America

Inter-seasonal and geographical variations of the models performances in NA are presented in Fig. 6. US8 underestimates the 414

observations in all seasons and in particular in winter and spring, and much larger compared to other models. In sub-region NA1, US6 overestimates by up to 9% while US8 underestimates by up to 22% in all seasons. CA2f slightly overestimates the winter and autumn O<sub>3</sub> levels by 3% and 5%, respectively. In the sub-regions NA2 and NA3, there is a general underestimation of all O<sub>3</sub> in winter and spring and a general overestimation in summer and autumn except for the US8 model. The winter and spring underestimates may be the result of underpredictions of afternoon temperatures and excessive O<sub>3</sub> titration by NO<sub>x</sub> as NA3 can be characterized by the largest emission sources in NA. In NA4, summertime O<sub>3</sub> levels are overestimated by all models including the US8 model. Slightly lower correlation coefficients (PCC = 0.7-0.9) are calculated for winter in NA while other seasons are simulated with PCC values of ~0.8-0.9, with slightly lower PCC values calculated for US7 (not shown).

#### 3.3. Influence of chemical boundary conditions

The influence of the chemical boundary conditions on the simulated surface O<sub>3</sub> levels has also been investigated on a seasonal basis. The analysis is carried out for the EU2 (north Eastern Europe) sub-region for Europe assuming that it is the least affected by the dominant westerly transport and having large anthropogenic emissions, suggesting that O<sub>3</sub> levels are more strongly controlled by local processes than regional transport, compared to the other sub-regions. Following the same rationale, sub-region NA3 was selected for the NA domain. The results presented in Fig. 7a show that in winter, all models underestimate O<sub>3</sub> levels along with the MACC model that provides the boundary conditions suggesting that large scale circulation and chemistry dominates over the local O<sub>3</sub> production. In spring and in summer (Fig. 7b,c), the regional production is more important than transport due to increased photochemical activity. In autumn (Fig. 7d), transport becomes more effective over local production. The MACC model slightly overestimates the summer levels (NMB = 1%), and slightly underestimates the autumn levels (NMB = -5%) while it underestimates the winter and spring levels 55% and 21%, possibly leading to the systematic overestimation of the regional models in autumn. The impact of large-scale transport over NA is less pronounced compared to Europe (Fig. 8). The impact is the smallest during summer when photochemical production is the largest (Fig. 8c). At the same time, it is interesting to note that the MACC results in the winter for NA1 are the lowest of the models shown in Fig. 8a, with a deficit of 8 ppb relative to the observations at 0 LST. The implication is that local chemistry, physics, model resolution and/or emissions relative to the global model all account for an increase in the winter  $O_3$  levels for region NA1 of 8 ppb (28.5%). and these local effects are captured by the suite of regional models. This may be compared to findings from the HTAP experiment, which suggest a 20% reduction in emissions in Europe, South Asia and East Asia would result in a 0.9 ppb reduction in O<sub>3</sub> in North America (Reidmiller et al., 2009). Here, simulated O<sub>3</sub> levels seem to be much more sensitive to the local  $O_3$  chemistry than to the boundary conditions associated with long-range transport (winter being the dominant season for long-range transport effects). Over both continents, the nighttime differences in all seasons are particularly large, with the MACC model largely underestimating the nighttime O<sub>3</sub>. Similar results were reported by Solazzo et al. (2012 and 2013a) for the first phase of the AQMEII project. A more detailed analysis of the influence of the MACC boundary conditions on a range of simulated species is presented in Giordano et al. (2015).

AB calculâ	ted for vertical O <sub>3</sub> pro	files for each model {	group and ense	emble mean	and media	in for the W	OUDC sta	tions in EU.									
Stations	Station name	Country	Lat/Lon	AT1	CH1	DE3	DE4	ES1	ES2a	ES3	Ш1	IT2	NL2	SI1	UK4	Mean	Median
5TN043	Lerwick	United Kingdom	60.1 / - 1.2	-8.40	-12.14	-27.80	-2.39	-9.82	-11.46	-7.86	-6.91	-9.05	-3.32	-8.16	-11.13	-10.40	-7.40
5TN053	NCCLE	Belgium	50.8/4.4	-4.11	-10.09	-14.08	3.80	-6.02	-7.46	-4.14	-4.53	-7.23	-1.96	-3.95	-4.86	-5.50	-3.56
660NTS	Hohenpeissenberg	Germany	47.8/11.0	-10.65	-21.94	-23.98	-2.04	-12.15	-11.98	-8.96	-9.55	-11.47	0.17	-10.39	-8.43	-11.62	-9.63
5TN156	Payerne	Switzerland	46.5/6.6	1.18	-10.06	-11.71	11.77	-0.63	1.84	2.51	2.43	0.51	2.70	1.44	3.94	0.64	2.52
5TN242	Praha	Czech Rep.	50.0/14.5	-8.55	-16.18	-26.48	-1.72	-11.38	-8.82	-6.98	-6.97	-8.68	-4.77	-7.86	-5.06	-9.50	-7.35
5TN308	Barajas	Spain	40.5/-3.7	-6.02	-14.29	-9.83	1.91	-7.72	-4.77	-6.72	-6.32	-7.83	0.21	-5.67	-1.61	-5.95	-5.01
5TN316	De Bilt	Netherlands	52.1/5.2	-4.57	-5.83	-9.82	3.62	-6.14	-4.29	-4.99	-5.08	-7.29	1.15	-4.37	-0.59	-4.23	-3.76
5TN318	Valentia	Ireland	51.9/-10.3	-6.51	-10.56	-15.49	-0.44	-8.01	-9.30	-6.00	-2.93	-6.35	-5.74	-6.43	-5.97	-7.01	-5.04
5TN348	Ankara	Turkey	40.0/32.9	-11.48	-16.13	-12.94	5.76	-13.38	-4.28	-10.95	-11.12	-15.24	0.55	-11.36	2.41	-8.66	-9.74

Table 5NMB calculated for vertical O3 profiles for each model group and ensemble mean and median for the WOUDC stations in NA.

Stations	Station name	Country	Lat/Lon	CA2f	US6	US7	US8	Mean	Median
STN021	Stony Plain	Canada	53.4/-114.1	-9.82	1.58	-2.29	-4.71	-3.81	-2.85
STN107	Wallops Island	USA	37.9/-75.5	-10.19	1.77	-1.17	-13.52	-5.78	-6.30
STN338	Bratts Lake	Canada	50.2/-104.8	-14.29	0.27	-3.26	-9.46	-6.68	-4.47
STN456	Egbert	Canada	44.2/-79.8	-16.78	-1.40	-3.95	-15.01	-9.28	-8.54
STN457	Kelowna	Canada	49.9/-119.4	-10.09	13.61	4.95	-0.62	1.96	2.05
STN458	Yarmouth	Canada	43.9/-66.1	-17.76	-1.17	-5.95	-15.27	-10.04	-10.20

## 3.4. Multi-model mean and median

The combination of concentrations simulated by several models can enhance the skill when compared to those from individual models (Galmarini et al., 2004a,b), which has also been demonstrated by Solazzo et al. (2012) in the first phase of the AOMEII project. In the present study, we provide simple multi-model mean and median analyses. Therefore, the calculated multi-model mean and median presented in Tables 3-5 and in Figs. 2-11 can only provide a basic distribution of all models with respect to the observations and should not be treated as multi-model ensemble analyses as they represent the bias originating from each individual model. As shown in Solazzo et al. (2012, 2013b) and Kioutsioukis and Galmarini (2014), introducing correlated biases into ensembles and analysis of the redundancy of the datasets is essential. As detailed multi-model ensemble analysis is not the scope of this paper, further analyses have been performed by Kioutsioukis et al. (2014) for the EU case using the multi-model data presented in the present paper.

#### 3.5. Regulatory analysis based on 8-h maximum surface O<sub>3</sub>

Observed and simulated daily maximum 8-h averaged surface O<sub>3</sub> levels during the O<sub>3</sub> season (May–September), which is a regulatory metric used in EU and NA, are compared in order to understand how the model biases vary with O<sub>3</sub> levels. The results are shown in Fig. 9 (note that in Fig. 9, observed concentrations are presented by/10). Over EU, all models overestimate O<sub>3</sub> concentrations below 50  $\mu$ g m<sup>-3</sup> by ~40%–~80% while they underestimate values above 140  $\mu$ g m<sup>-3</sup> except for the UK4 model that overestimates the levels above 160  $\mu$ g m<sup>-3</sup>. Most models follow the MACC model up to a concentration of 200  $\mu$ g m<sup>-3</sup> with increasing variability towards higher concentrations. NL2 and UK4 models overestimate the 230–240  $\mu$ g m<sup>-3</sup> concentration bin where the spread is also largest among other models. The UK4 model defines the upper boundary while IT2 defines the lower boundary of the

envelope until 100  $\mu$ g m<sup>-3</sup> while above that, the highest differences are calculated for IT1. The CH1 model, which together with the IT2 model showed the largest negative biases in annual mean values, is more consistent with other models when considering 8-h maximum values. Above a concentration of 70  $\mu$ g m<sup>-3</sup>, ES2a, NL2 and UK4 models are associated with positive deviations from the MACC model while other models are below the MACC-simulated levels. Results show that depending on the station, there are underestimations by up to >200  $\mu$ g m<sup>-3</sup>.

Over NA (Fig. 9b), the biases are lower compared to EU. Note that for NA the values are reported in volume mixing ratios (ppb) rather than concentrations ( $\mu g/m^3$ ). The surface O<sub>3</sub> levels below 30 ppb are overestimated by all models by ~15-25% and levels above 60 ppb are underestimated by all models by up to ~80%. The largest biases are calculated for US8 except for the 120-130 ppb bin where US7 has the largest bias. US8 has the smallest bias below 50 ppb. The results show that models have a tendency to severely underpredict high O<sub>3</sub> values which are of concern for air quality forecast and control policy applications. Further improvement of model treatments (e.g., gas-phase chemistry, O<sub>3</sub> dry deposition and processes affecting afternoon temperature predictions) and inputs (e.g., boundary conditions, biogenic VOCs and wildfire emissions) as well as a better understanding of interplays among on-linecoupled atmospheric processes (e.g., the impact of aerosol indirect effects on  $O_3$  formation) are urgently needed.

## 3.6. Vertical ozone profiles

The model results from each group as well as the ensemble mean and median are compared with  $O_3$  soundings obtained from WOUDC for the EU and NA domains up to 9 km height above the ground. Figs. 10 and 11 show the observed and simulated vertical  $O_3$  levels at fixed heights over the EU and NA domains, respectively while Tables 4 and 5 present the normalized mean bias (NMB) for all the models and ensemble mean and median. On average, most models underestimate the observed vertical profiles



Fig. 9. Observed surface O<sub>3</sub> concentration bins against mean bias for the EU and NA domains for the O<sub>3</sub> season (May–September).



Fig. 10. Observed and simulated (models, mean and median) vertical O<sub>3</sub> profiles averaged over 2010 in the EU domain. Note the differences in scales.

by up to 22% over EU. The DE4 model generally has smaller biases compared to other groups except for the station STN156 where it overestimates by ~12% (Fig. 10). The ensemble mean/median improves the results compared to the majority of the models depending on the station. The ensemble mean results in smaller biases compared to the median. Over NA (Fig. 11), the CA2f model underestimates the vertical O<sub>3</sub> levels at all stations by10–17% (Table 5). US6 and US7 have the smallest biases in most stations but with overestimations of 14% and 5%, respectively, at STN457. The US8 model underestimates at all stations by 4-15% but overestimates at STN457 by 2%. The ensemble mean and median lead to improved results compared to CA2f at all stations above ~1000-2000 m and to US8 at STN107 and STN456 below 2000-3000 m. Over Europe, among others, STN318 station (Valentia Observatory, Ireland) can be considered as a site that is largely impacted by long-range transport and is associated with the largest underestimation (NMB = -11%) by the MACC model (not shown), suggesting that boundary conditions can partly contribute to the underestimated vertical profiles by a majority of the models. Results also show that the tropospheric biases in the MACC model (Figs. 10 and 11) are less pronounced than the surface bias as also shown by Inness et al. (2013).

## 4. Summary and conclusions

An operational evaluation of simulated ozone  $(O_3)$  levels over Europe (EU) and North America (NA) in 2010 using eight different on-line-coupled air quality models from sixteen groups has been conducted in the context of the AQMEII project. Seven groups from EU and two groups from NA applied the WRF-CHEM model, but with different settings. Anthropogenic emissions and chemical boundary conditions were prescribed while biogenic emissions were calculated online by each individual group. All groups interpolated their model output to a common output grid and a common set of receptor locations and uploaded the data to the ENSEMBLE system. The results are evaluated against surface and sounding observations, which are provided by operational over EU and NA, at continental and sub-regional levels on annual and seasonal basis.

All models capture, reasonably well, the shape of the domainaveraged annual diurnal cycle of  $O_3$  over both domains, while the sub-regional temporal variability are simulated from moderate to good depending on the season and the sub-region that the particular model is configured for. There is a general underestimation of the annual surface  $O_3$  by up to 18% and 22% over EU and



Fig. 11. Observed and simulated (models, mean and median) vertical O<sub>3</sub> profiles averaged over 2010 in the NA domain. Note the differences in scales.

NA, respectively. Differences in performance among models can be attributed partly to the chemical mechanism used in the models, partly to VOC preprocessing and different biogenic emissions, and partly to the differences in the microphysics, leading to different cloudiness and therefore to different photolysis, temperature and radiation acting on the O<sub>3</sub> production. The sub-regional analyses highlight the influence of the anthropogenic emissions while the seasonal analyses show a strong tendency to overestimate the autumn surface levels. The temporal variation and magnitudes are much better captured during summer compared to other seasons. The winter and spring underestimations may be resulting from underprediction of afternoon temperatures, excessive O<sub>3</sub> titration by too much  $NO_x$  as well as biases from the chemical boundary conditions. Boundary condition analyses show that wintertime levels are mostly driven by transport rather than local production due to limited photochemistry. The global MACC model providing the boundary conditions to the regional models largely underestimate the surface ozone levels particularly in winter, leading to a negative bias in the regional model simulations, while in most subregions, it largely overestimates the autumn O<sub>3</sub> levels in winter. leading to the systematic overestimations of surface autumn O<sub>3</sub> levels by the regional models. The inclusion of aerosol indirect effects in some online-coupled models also contributes in part to the underpredictions of O<sub>3</sub> mixing ratios. On average, most models underestimate the observed vertical profiles by up to 22% over EU and up to 17% over NA.

Comparison of observed and simulated daily maximum 8h averaged surface O<sub>3</sub> levels during the O<sub>3</sub> season (May–September), which is a regulatory metric used in EU and NA, show that over Europe, O<sub>3</sub> concentrations below 50  $\mu$ g m<sup>-3</sup> are overestimated by up to 80% while levels above 140  $\mu$ g m<sup>-3</sup> are underestimated. Over NA the surface O<sub>3</sub> levels below 30 ppb are overestimated by all models by up to 25% and levels above 60 ppb are underestimated by all models by up to 80%. This has implications for air quality forecast and policy applications.

Overall, the results show a slight improvement in the surface ozone level predictions over EU by the models that participated in the second phase of AQMEII compared to those that participated in the first phase. The NMB calculated for the whole domain and simulation period in the first phase ranged from -24% to 9% while in this second phase, the NMB range was calculated to be -18% to 2%. On the other hand over NA, there is a significant change between the two phases of the project: the overestimation of 3%-22% in the first phase shifted to a NMB range of -22% to 3%. These results, however, should not be considered as solely the difference between on-line and off-line models as different simulation years, different emissions, different sets of models, particularly for the NA case, and different boundary condition data should be taken into account. Additionally, as the results presented in this paper are temporally and spatially averaged, cases where feedback mechanisms are of importance must be further studied and evaluated.

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