Geosci. Model Dev., 5, 1565–1587, 2012 www.geosci-model-dev.net/5/1565/2012/ doi:10.5194/gmd-5-1565-2012 © Author(s) 2012. CC Attribution 3.0 License.





How realistic are air quality hindcasts driven by forcings from climate model simulations?

G. Lacressonnière¹, V.-H. Peuch², J. Arteta¹, B. Josse¹, M. Joly¹, V. Marécal¹, D. Saint Martin¹, M. Déqué¹, and L. Watson¹

¹GAME/CNRM, Météo-France, URA1357, CNRS – Centre National de Recherches Météorologiques,
42 av. G.Coriolis, 31057 Toulouse, France
²ECMWF, Shinfield Park, Reading, UK

Correspondence to: G. Lacressonnière (gwendoline.lacressonniere@meteo.fr)

Received: 12 July 2012 – Published in Geosci. Model Dev. Discuss.: 31 July 2012 Revised: 13 November 2012 – Accepted: 19 November 2012 – Published: 12 December 2012

Abstract. Predicting how European air quality could evolve over the next decades in the context of changing climate requires the use of climate models to produce results that can be averaged in a climatologically and statistically sound manner. This is a very different approach from the one that is generally used for air quality hindcasts for the present period; analysed meteorological fields are used to represent specifically each date and hour. Differences arise both from the fact that a climate model run results in a pure model output, with no influence from observations (which are useful to correct for a range of errors), and that in a "climate" set-up, simulations on a given day, month or even season cannot be related to any specific period of time (but can just be interpreted in a climatological sense). Hence, although an air quality model can be thoroughly validated in a "realistic" set-up using analysed meteorological fields, the question remains of how far its outputs can be interpreted in a "climate" set-up. For this purpose, we focus on Europe and on the current decade using three 5-yr simulations performed with the multiscale chemistry-transport model MOCAGE and use meteorological forcings either from operational meteorological analyses or from climate simulations. We investigate how statistical skill indicators compare in the different simulations, discriminating also the effects of meteorology on atmospheric fields (winds, temperature, humidity, pressure, etc.) and on the dependent emissions and deposition processes (volatile organic compound emissions, deposition velocities, etc.). Our results show in particular how differing boundary layer heights and deposition velocities affect horizontal and vertical distributions of species. When the model is driven by operational analyses, the simulation accurately reproduces the observed values of O_3 , NO_x , SO_2 and, with some bias that can be explained by the set-up, PM_{10} . We study how the simulations driven by climate forcings differ, both due to the realism of the forcings (lack of data assimilated and lower resolution) and due to the lack of representation of the actual chronology of events. We conclude that the indicators such as mean bias, mean normalized bias, RMSE and deviation standards can be used to interpret the results with some confidence as well as the health-related indicators such as the number of days of exceedance of regulatory thresholds. These metrics are thus considered to be suitable for the interpretation of simulations of the future evolution of European air quality.

1 Introduction

The issues of climate change and air quality are intertwined; anthropogenic emissions contribute to climate change, and the evolution of the climate through changes in meteorological parameters (temperature, precipitation) impacts concentrations and distributions of pollutants in the atmosphere. In the lower troposphere, ozone (O₃) is a pollutant that affects human health (WHO, 2004; Schlink et al., 2006) and causes damages to crops (Fuhrer and Booker, 2003) and ecosystems. O₃ is a secondary pollutant; its principal precursors are carbon monoxide (CO), volatile organic compounds (VOCs) and nitrogen oxides (NO_x) emitted by both natural (biogenic) and anthropogenic (transport, industries) sources. Several studies have shown how O₃ photochemistry depends upon meteorological conditions such as temperature and precipitation. During summer, conditions of high temperature and low precipitation favor oxidant accumulation, and surface concentrations of O3 reach high values (Guicherit and van Dop, 1977; Sillman, 2000) and have the potential to exceed air quality standards. These conditions also favor the production of secondary pollutants such as sulphate and nitrate aerosols, and organic aerosols which can contribute to the high levels of particulate matter (PM) during summertime. Nevertheless, the frequency and intensity of pollution episodes vary considerably from year to year depending on weather; as an example the summer 2003 heat wave in Europe has been associated with exceptionally high O3 concentrations (Langner et al., 2005; Vautard et al., 2005, 2007; Guerova and Jones, 2007; Solberg et al., 2008). In fall and winter, stagnant conditions also enhance levels of primary pollutants (SO₂, NO_x) in the atmosphere and thus concentrations of PM₁₀ (particles with an aerodynamic diameter smaller than 10 µm), another pollutant of concern connected to air quality and health problems.

The interactions between climate change and air quality have been already extensively studied. At the global scale, studies (Prather et al., 2003; Dentener, 2006) have for instance evaluated the effects of changing emissions and climate on surface O₃ concentrations under an A2 scenario (IPCC AR4). Dentener (2006) showed that global mean surface O_3 may increase by about 4.3 ± 2.2 ppbv by the year 2030 and the area of global natural ecosystems exposed to critical nitrogen deposition may increase up to 25 % by this time. Regional models centered over the continental United States have been used to examine US air quality in the future due to climate change alone independently of evolution in emissions in North America and elsewhere (Hogrefe et al., 2004; Knowlton et al., 2004; Dawson et al., 2009). Hogrefe et al. (2004) concluded that the average daily summertime maximum 8-h O₃ concentrations will increase by 2.7 ppbv and 4.2 ppbv for summers in the 2020s and 2050s, respectively. In the literature, a set of regional models have similarly focused on the European region to isolate the impacts of climate change (Langner et al., 2005; Meleux et al., 2007; Giorgi and Meleux, 2007; Carvalho et al., 2010; Andersson et al., 2010; Katragkou et al., 2011; Huszar et al., 2011; Juda-Rezler et al., 2012). Precisely, Zlatev (2007) and Langner et al. (2005) presented the impacts of climate change on air quality over Europe with a constant emission rate and showed an increase in photochemical production in future climate scenarios. In Meleux et al. (2007), the authors isolated the impacts of European summer climate change on the increase in O_3 levels by using the same emissions and global chemical boundary conditions for the present day and future periods. Katragkou et al. (2011) investigated the sensitivity of surface ozone to the future climates of the 2040s and 2090s by studying changes in meteorological parameters under an A1B scenario. Andersson et al. (2010) suggested changes in surface ozone between -4 to 13 ppbv on average from 1961–1990 to 2071–2100, based on the A2 scenario, and highlighted the role of surface deposition processes. Carvalho et al. (2010) concluded that PM_{10} levels will be impacted by climate change depending on the month and region, with a maximum increase reaching $30 \,\mu g \,m^{-3}$ in September over Portugal. Szopa et al. (2006) investigated impacts of local anthropogenic emission changes and background O₃ changes. They estimated that the O₃ concentration in July may increase up to 5 ppbv across Europe by 2030. According to all these findings, the expectation of a warmer climate in the next decade may well affect air quality directly despite regulations to reduce the emission of pollutants (European Commission, 2008).

In most "climate" studies, air quality modeling systems are chemistry-transport models (CTM) that rely on global or regional climate models to provide the meteorological forcings for future periods. The purpose of this paper is to assess how realistic air quality hindcasts are when driven by forcings from climate models for the current period over Europe, in comparison to a reference obtained using instead analysed meteorological forcings (analyses), which include meteorological observations and are thus very realistic and specific for each single date. The results will be evaluated using a range of statistical tools and air quality indicators. In our study, we have used the chemistry and transport model MOCAGE of Météo-France (Peuch et al., 1999) for three multi-year simulations covering the present time (2004–2008) over a European domain. Comparisons between the simulations and the AirBase observations allow us to infer how a range of statistical indicators are affected when using different types of forcings. This work will provide guidance on how far to interpret air quality hindcasts relying on climate model outputs, which is essential for the study of future air quality.

In this paper, Sect. 2 describes the modeling approaches and the numerical experiment design. We also discuss the statistical indicators and the representativeness of the measurement stations used for this study. Section 3 compares simulations in order to evaluate separately how emissions and meteorological changes affect the distributions of pollutants over Europe. Finally, the experiments run with the analyses and the climate forcings are compared against observations in Sect. 4. The statistical indicators that produce similar results for the two experiments will be the most useful ones to consider when examining future trends.

2 Methodology

2.1 Model set-up and experimental design

The model used in this study is the three-dimensional multiscale chemistry-transport MOCAGE (Modèle de Chimie Atmosphérique à Grande Echelle), which simulates the interactions between the dynamical, physical and chemical processes in the troposphere and stratosphere (Peuch et al., 1999). The configuration allows for the representation of both long range transport of pollutants and regional impacts of pollutants on air quality. This model is used for operational air quality forecasting in France (http://www.prevair.org, Honoré et al., 2008) and in the context of the GMES atmospheric monitoring service (Hollingsworth et al., 2008), and has been evaluated during several campaigns, see for instance Dufour et al. (2004) and Bousserez et al. (2007).

MOCAGE uses a semi-Lagrangian advection scheme (Williamson and Rasch, 1989) to transport chemical species. On the vertical, the configuration has 47 hybrid levels from the surface up to 5 hPa with a resolution of about 150 m in the lower troposphere increasing to 800 m in the higher troposphere. Turbulent diffusion is parameterized with the scheme of Louis (1979) and convective processes with the scheme of Bechtold et al. (2001). The chemical scheme used in this study is RACMOBUS; it is a combination of the stratospheric scheme REPROBUS (Lefèvre et al., 1994) and the tropospheric scheme RACM (Stockwell et al., 1997). Overall, this chemical scheme includes 119 individual species with 94 prognostic variables and 377 chemical reactions. In our study, a sulfur cycle has been implemented; the oxidation reactions in the gaseous and aqueous phases lead to the formation of sulphate aerosols as in Ménégoz et al. (2012). These reaction mechanisms are provided in the Supplement. MOCAGE simulates the evolution of five types of aerosols: black carbon, sea salts, desert dusts, anthropogenic primary particulate matter and sulphates. They are compartmented in size bins (Martet et al., 2009) and divided into 6 bins for each aerosol compound: between 0.1 µm and 100 µm for dust aerosols, 0.001 µm and 10 µm for black carbon, 0.03 µm and $20\,\mu\text{m}$ for sea salt, $0.005\,\mu\text{m}$ and $10\,\mu\text{m}$ for anthropogenic particulate matter and 0.01 µm and 20 µm for sulphates. Nitrate and organic aerosols are not taken into account in this study. A negative bias is thus expected by design on total PM.

The model uses two-way nested domains on a $2^{\circ} \times 2^{\circ}$ horizontal grid over the globe and a $0.2^{\circ} \times 0.2^{\circ}$ horizontal grid over Europe (30° N-70° N; 15° W-35° E). Threehourly forcings are used for meteorology in this study, from either operational analyses from Météo-France (ARPEGE, Courtier et al., 1991) or from climate simulations obtained with ARPEGE-Climate, (version 5.1, Déqué et al., 1994), for the present decade. The resolution of ARPEGE is in a T798 spectral stretched grid (resolution of around 15 km over Europe and 60 km in the Pacific) while ARPEGE-Climate operates on a T63 triangular truncation, equivalent to a resolution of about 2.8°. Anthropogenic forcings of ARPEGE-Climate (GHG, aerosols) refer to the climatology of the present time. For the present simulation, ARPEGE-Climate is driven by prescribed observed SSTs (sea surface temperatures), and for the future simulations, the SSTs are thus from RCP8.5 scenario ocean-atmosphere coupled simulations. Meteorological forcings are interpolated horizontally on the two MOCAGE domains.

For the anthropogenic emissions, the inventory is the one (Visschedijk and Denier van der Gon, 2005; Pouliot et al., 2012) developed for the Global and regional Earth-system Monitoring using Satellite and in-situ data (GMES) project (Hollingsworth et al., 2008). This inventory has a high spatial resolution compatible with our model and a temporal resolution of 1 h. It is representative of the year 2003. We chose not to modify emissions depending on each specific year as this would be meaningless for runs driven by climate forcings. Biogenic emissions of isoprene and monoterpene are calculated offline with MEGANv2.04 model (Guenther et al., 2006). Two types of input files were required: the land cover variables (leaf area index, plant functional type and emissions factors) and the weather data. The land cover variables are available at a spatial resolution of $\sim 5 \text{ km}$ (150 s longitude \times 150 s latitude). The meteorological fields (temperature, solar radiation) are provided either by ARPEGE analyses or ARPEGE-Climate simulations.

The numerical experiment design is as follows (Table 1). Three five-year periods of the current climate were simulated using different meteorological forcings and surface processes. ANALY simulation acts as the reference. INT relies upon meteorological forcings from a climate model, and surface exchanges (weather-dependent emissions and deposition velocities) are the same as for ANALY. CLIM is driven by climate forcings, and surface processes are computed with meteorological conditions of ARPEGE-Climate. The summer heat wave of 2003 was uncharacteristic of the current climate, and studies have shown a similar pattern of heatwaves with future climate conditions (Meleux et al., 2007; Solberg et al., 2008). The climatological forcings from ARPEGE-Climate are representative of the current decade and do not reproduce the extremely hot and cold events. For this reason, year 2003 is not considered in the statistical comparisons. We chose the 2004–2008 period for our simulations. 5 yr is a short time on the one hand to represent the meteorological variability over Europe. On the other hand, we require that emissions do not evolve too much in time during the period, and 5 yr is certainly at the limit for such an assumption. The choice of 5 yr is thus clearly a compromise.

2.2 Statistical indicators

In Europe, air quality thresholds of acceptable levels of O_3 (European Commission, 2002), NO₂ and SO₂ (European Commission, 1999, 2001) and PM_{2.5} and PM₁₀ (European Commission, 1999, 2001, 2008) have been established in order to protect and inform populations, as described in Table 2. The impact of these air quality policies, in Europe and in the world as well, can be evaluated using numerical air quality modeling. As an example, in Europe, the CAFE (clean air for europe) program has been set up to assess the impacts of these policies on the pollutants' levels (Cuvelier et al., 2007).

Simulations	Periods	Meteorological		Deposition		
		forcings	anthropogenic	biogenic	desert dust, sea salt	velocities
ANALY	2004–2008	ARPEGE	GEMS	MEGANv2.4 /ARPEGE	ARPEGE	ARPEGE
INT	5 yr of 2000–2010 climate	ARPEGE-Climate	GEMS	MEGANv2.4 /ARPEGE	ARPEGE	ARPEGE
CLIM	5 yr of 2000–2010 climate	ARPEGE-Climate	GEMS	MEGANv2.4 /ARPEGE-Climate	ARPEGE-Climate	ARPEGE-Climate

Table 1. Simulations considered for the present study.

Table 2. Some regulatory European air quality thresholds of O_3 , NO_2 , SO_2 and PM_{10} levels.

Pollutant	Parameter	Threshold values
O ₃	hourly average	$180 \mu g m^{-3}$
	daily maxima of 8 h running	$120 \mu g m^{-3}$
NO ₂	hourly average	$200 \mu g m^{-3}$
	annual average	$40 \mu g m^{-3}$
SO ₂	hourly average	$350 \mu g m^{-3}$
	daily average	$125 \mu g m^{-3}$
PM ₁₀	daily average	$50 \mu g m^{-3}$

In order to forecast air quality, to understand the dynamics of air pollution and to develop regulations to reduce emissions, air quality modeling systems are needed. A variety of metrics has been used over the years to evaluate the performance of air quality models (US-EPA, 1984, 1991; Chang and Hanna, 2004; Boylan and Russell, 2006). Mean bias (MB), mean normalized bias (MNB), root mean square error (RMSE) and correlation coefficient (CORR) are common statistical parameters used by the modeling community. Furthermore, the mean normalized bias error (MNBE) and the mean normalized gross error (MNGE), normalizing the bias and error for each model-observed pair by the observations, are also useful parameters. For the evaluation of particulate matter concentration, Boylan and Russell (2006) suggested the consideration of the mean fractional bias (MFB) and the mean fractional error (MFE) parameters instead of MNBE and MNGE. They proposed that the model performance criteria would be met when both MFE \leq 75 % and MFB $\leq \pm 60$ %, respectively. The model performance goal would be met when both MFE \leq 50 % and MFB $\leq \pm$ 30 %. The US-EPA suggested several performance criteria for simulated O₃, such as MNBE $\leq \pm 15$ % and MNGE $\leq \pm 35$ % (US-EPA, 1991), while the EC proposes a modeling quality objective given as a relative uncertainty (%): 50% and 30% for PM₁₀/PM_{2.5}/O₃ and NO₂/SO₂ annual average, respectively (European Commission, 2008).

The model to data statistics MB, MNB, RMSE, correlation coefficient and sigma ratio are selected for the present **Table 3.** Definition of the metrics used in the evaluation of the MOCAGE model performance. *M* refers to the model, *O* refers to the observations. $\overline{M} = \frac{1}{N} \sum (M_i)$ and $\overline{O} = \frac{1}{N} \sum (O_i)$.

Mean bias	
	$MB = \frac{1}{N} \sum_{i=1}^{n} (M_i - O_i) = \bar{M} - \bar{O}$
Mean normalized bias	$1 \sum_{i=1}^{n} M_i = O_i$
Marchart	$MNB = \frac{1}{N} \sum_{i=1}^{n} \frac{m_i - \sigma_i}{O_i}$
Mean fractional error	MEE $- 2 \sum n M_i - O_i $
Mean fractional bias	$\mathbf{MFL} = \overline{N} \ \angle_{i=1} \ \overline{(M_i + O_i)}$
Weat fractional blas	$MFB = \frac{2}{2} \sum_{i=1}^{n} \frac{(M_i - O_i)}{(M_i - O_i)}$
Correlation coefficient	$N = M = 1$ $(M_i + O_i)$
	$r = \frac{\sum_{i=1}^{n} (M_i - \bar{M})(O_i - \bar{O})}{\sum_{i=1}^{n} (M_i - \bar{M})(O_i - \bar{O})}$
D	$ = \left\{ \sum_{i=1}^{n} (M_i - \bar{M})^2 \sum_{i=1}^{n} (O_i - \bar{O})^2 \right\}^{1/2} $
Root mean square error	$\sqrt{1 - 2}$
<i>.</i>	$\text{RMSE} = \sqrt{\frac{1}{N} \sum_{i=1}^{n} (M_i - O_i)^2}$
Sigma ratio	$\sqrt{1 - n}$
	$\sigma = \frac{\sqrt{\frac{1}{N}\sum_{i=1}^{n}(M_i - M)^2}}{\sqrt{\frac{1}{N}\sum_{i=1}^{n}(M_i - M)^2}}$
	$\sqrt{\frac{1}{N}\sum_{i=1}^{n}(O_i-\bar{O})^2}$

study. The definitions of these metrics are indicated in Table 3. We also considered the mean diurnal cycle and the temporal series. Mean diurnal cycles are averaged over all available days of concentrations for each 24 h period, while time series are based on the daily mean. Seasonal mean statistics are computed, with seasons corresponding to summer (June, July, August and September) and winter (December, January, February and March). We chose to study these two seasons of interest in air pollution while autumn and spring are rather transitional seasons. As summarized in Table 4, metrics are calculated for hourly values and daily averages for NO_x , SO_2 and PM_{10} , while the hourly value and daily maximum 8-h average concentration (M×8h) statistics are computed for O_3 , as the M×8h is one of the most important parameters to be considered for this species.

G. Lacressonnière et al.: Air quality hindcasts driven by forcings from climate model

Lable 4. Michiles considered in the evaluation of O_3 , $100x$, 50° and 10100 concentrations.
--

Indicators	Parameters		Codes		
		O ₃	NO _X	SO ₂	PM ₁₀
Mean bias	hourly value	MBO ₃ H	MBNO _x H	MBSO ₂ H	_
	daily mean	-	MBNO _x DM	MBSO ₂ DM	MBPM ₁₀
	$M \times 8h$	MBO3MAX	-	_	-
Mean normalized bias	hourly value	MNBO ₃ H	MNBNO _x H	MNBSO ₂ H	-
	daily mean	-	MNBNO _x DM	MNBSO ₂ DM	-
	$M \times 8h$	MNBO3MAX	-	-	-
Mean fractional bias	daily mean	-	-	_	MFBPM ₁₀
Mean fractional error	daily mean	-	-	_	MFEPM ₁₀
Correlation coefficient	hourly value	CORRO ₃ H	CORRNO _x H	CORRSO ₂ H	-
	daily mean	-	CORRNO _x DM	CORRSO ₂ DM	CORRPM ₁₀
	$M \times 8h$	CORRO3MAX	-	_	-
Roor mean square error	hourly value	RMSEO ₃ H	RMSENO _x H	RMSESO ₂ H	_
	daily mean	-	RMSENO _x DM	RMSESO ₂ DM	-
	$M \times 8h$	RMSEO3MAX	-	_	-
Sigma ratio	hourly value	$\sigma O_3 H$	$\sigma NO_{X}H$	σSO_2H	-
	daily mean	-	$\sigma NO_{X}DM$	σSO_2DM	σPM_{10}
	$M \times 8h$	$\sigma O_3 MAX$	_	_	_
SOMO35*	$M \times 8h$	SOMO35	_	_	-

* SOMO35: annual sum of excess of daily maximum 8-h mean ozone over the cut-off of 35 ppb.

2.3 Observations and representativeness

In order to evaluate the performance of MOCAGE and to be in position to investigate and put into context the differences between the simulations, we used AirBase (Version 5) measurement data. The AirBase metadata describes both the site area (urban, suburban or rural) and the site type (traffic, industrial or background). Giving the spatial resolution of our model $(0.2^{\circ} \times 0.2^{\circ})$, not all the reporting sites are representative enough. In Joly and Peuch (2012), an objective classification of the AirBase sites based on past measurements has been proposed in order to overcome issues of lack of homogeneity and erroneous information in the metadata. This classification allows for selection of the monitoring sites that are representative of the spatial resolution of our model. Through 10 classes, the less polluted stations (class 1) are distinguished from the very polluted sites (class 10). The robustness of this approach is obtained with a pollutant-specific classification, taking into account that transport, chemistry and lifetime are specific to each pollutant.

In order to highlight the effect of site representativeness, we have compared the summertime (JJAS) average diurnal cycles for classes 1–2 (1 and 2), 1–5, 1–10 and 6–10 over France for O_3 and NO_x (Fig. 1) for the simulation ANALY. The median diurnal cycles observed (red) and simulated with MOCAGE (black) are presented for the year 2007. As shown in Fig. 1, the modeled variability of ozone and NO_x is not as pronounced as in observations, whatever the type of sites. During nighttime hours, ozone levels decrease when polluted stations are added to the sample (statistics for classes 1–2



Fig. 1. Mean diurnal cycles of O_3 (**a**) and NO_x (**b**), as a function of hour, simulated with MOCAGE (dashed lines) and observed by Air-Base (solid lines) averaged for the summer (JJAS) of 2007. Diurnal cycles are represented for the classes 1–2, 1–5, 1–10 and 6–10.

versus for classes 1–10) due to titration by NO. Nevertheless, simulated O₃ values do not decrease as much as the observations for classes 6–10 and 1–10. As expected, a large variation in observed NO_x concentrations is seen when the stations considered are from different class types. Except when considering sites of classes 1–2, the amplitude of NO_x concentrations is generally underestimated by the model. The amplitude of the median diurnal cycles changed significantly between categories 1–2 and 1–5 both for NO_x and O₃. For classes 6–10 and 1–10, i.e. when most polluted sites are considered, the model does not reproduce the high observed values because, at least in part, of the model's horizontal resolution.

To conclude, the choice of the sites used for verification according to an objective classification allows for an



Fig. 2. From top to bottom: average summertime surface temperature (°C), precipitation (mm d^{-1}), humidity (g kg⁻¹) and planetary boundary layer height (m) for the summer period (JJAS) of ANALY and INT. Differences between ANALY and INT are shown on the right.

improvement in accounting for representativeness. For NO_x , which are short lived species, it is necessary to reduce the sample of sites to classes 1–2 only to focus on sites that are representative enough for the model grid size. Due to transport effects, 5 classes (1–5) can be used to evaluate simulations for longer-lived species such as O₃ in order to have a larger geographical basis. The spatial distribution of PM₁₀ (not shown) has the same behavior as O₃, and the same conclusion can be applied. In conclusion, the performances of MOCAGE will be assessed by comparing simulations against observations at sites of classes 1–5 for O₃ and PM₁₀, and of classes 1–2 for NO_x and SO₂ (not shown, but same behavior as NO_x). The number of sites finally taken

into account for each pollutant and country are summarized in Table 5.

3 Results

In the following section, we discuss MOCAGE's capability to simulate realistic air quality hindcasts when driven by forcings from climate modeling for the current period. We will evaluate statistical tools and air quality indices and compare how they evolve with different sets of forcings. Two main parts can be distinguished. First, in Sect. 3.1, comparisons between ANALY and INT, as well as between INT and

Table 5. Number of representative sites available by countries and species. Countries are, from left to right, France (FR), Spain (ES), England (GB), Germany (DE), Italy (IT) and Poland (PL).

Species	Classes	Countries									
		Europe	FR	ES	GB	DE	IT	PL			
03	1–5	970	201	191	33	202	54	47			
NO _x	1-2	354	68	115	_	97	9	_			
SO_2	1-2	342	31	101	7	101	17	7			
PM_{10}	1–5	824	168	151	47	259	-	28			

CLIM, allow us to detect the effects due to the meteorological forcings and to changes in surface exchange fluxes, respectively. As described previously, ARPEGE analyses and ARPEGE-Climate fields differ regarding horizontal resolution. Surface exchanges (emissions that depend upon meteorology, as well as surface deposition) have been computed with two sets of meteorological conditions (ARPEGE and ARPEGE-Climate). The differences are for the biogenic volatile organic compounds, desert dust and sea salt emissions, as well as for deposition velocities, which depend on meteorology. Section 3.2 presents statistical skill scores of ANALY and CLIM against AirBase data (for "representative sites" only).

3.1 Comparisons between ANALY, INT and CLIM

3.1.1 Impact of changes in meteorological fields on European air pollution levels

Figure 2 shows the mean differences in surface temperature, precipitation, humidity and planetary boundary layer height (PBL) for the JJAS period between ANALY and INT run. The purpose is to evaluate briefly how climate meteorological forcings differ from the analyses. For the comparisons, we have thus averaged spatially the analyses fields to a horizontal resolution similar to the climate run. Focusing on the temperature first, similar structures are found over Europe and an increasing gradient from the northern to the Mediterranean areas. However, over the northeastern part of the domain, the temperatures simulated by ANALY are locally significantly higher up to 4-5 °C. In contrast, over the Italian and Greek areas, INT displayed higher temperatures. The lower resolution of INT described previously explains partly the differences observed because ANALY displays more structured fields. Orography is smoothed in INT and the fields are representative of the current decade. Higher precipitations are simulated by INT over Europe and North Africa, but the spatial patterns are similar between the two simulations. Concerning the humidity, the spatial pattern is correctly reproduced by ARPEGE-Climate, with higher humidity over the Mediterranean Sea. The PBL is another important field to be considered as it affects the dilution of pollutants in the atmosphere. As depicted in Fig. 2, higher average PBLs in ANALY than in INT are seen over Spain and Greece.

In Fig. 3, we represent the average surface concentrations of O_3 (a), isoprene (b), NO_x (c), SO_2 (d) and sulphate aerosols (e) for the summertime 2004-2008 in ANALY and INT. Here, the observed changes in pollutant distributions are only due to differences in the meteorological conditions, as emissions and deposition velocities are identical. The spatial pattern of mean O₃ concentrations is similar for the two simulations over the European domain. The highest concentrations are found in Southern Europe, over the Mediterranean Sea (50–60 ppbv), caused by intense photochemical production of O₃ (EEA, 2005; Vautard et al., 2005). The meteorological fields such as temperature influence the production of O₃ (Meleux et al., 2007; Hedegaard et al., 2008). Fields of change in O3 present some similarities to the changes in temperature (Fig. 2). As noticed previously, over Spain, Africa and Northern Europe ANALY outputs higher temperatures up to 4-5 °C and higher O₃ concentrations (+6–8 ppbv). The highest positive temperature differences seen over Europe relate to the highest positive O₃ differences. Other studies have shown that among all the meteorological parameters, the one that causes the greatest impact on ozone is temperature (Dawson et al., 2007). Nevertheless, as explained in Katragkou et al. (2010), other variables such as differences in solar radiation, zonal and meridional winds and changes in atmospheric stability also impact ozone concentrations. Similar temperatures and O₃ spatial patterns with the opposite sign are observed over the Mediterranean basin (Italy, Greece).

High concentrations of isoprene, in the range of 2.5-3 ppbv, are simulated over North Africa and Greece with the simulation INT. The biogenic emissions calculated to drive ANALY and INT are the same, but the differences in simulated isoprene cannot be explained by the isoprene emissions or the temperature fields. A longitudinal cross-section (not shown) at a latitude of 36° N (across Africa) displays higher isoprene concentrations at the surface from 0 to 10° with INT. The accumulation of isoprene near the surface can mainly be explained by the boundary layer, which is less well-mixed in INT than in ANALY.

The simulated distributions of summertime average NO_x concentrations (NO+NO₂) show levels around 8–12 ppbv in the Netherlands, Belgium, central and eastern England and the industrial Po Valley (Fig. 3). In ANALY and INT, higher concentrations of NO_x are also found over major shipping routes (North Sea, Gibraltar) and near emissions sources. Tropospheric columns of NO_x (not shown) are identical in ANALY and INT; the differences seen over Europe at the surface are mainly explained by differing boundary layer mixing in the two experiments.

Simulated SO_2 concentrations over Europe display their highest levels over Spain, Eastern Europe (Poland, Romania, Greece) as well as Belgium and the UK. Over northern Spain (Fig. 3), the concentrations reach up to more than 8 ppbv



Fig. 3. From top to bottom: simulated average JJAS surface O_3 , isoprene, NO_x , SO_2 and sulphate fields for ANALY and INT. Differences between ANALY and INT are shown on the right. Units are in $\mu g m^{-3}$.



Fig. 4. On the left: latitudinal cross-sections at 30° E of sulphate concentrations (μ g m⁻³) averaged for the JJAS period. Levels are in hPa. On the right: sulphate column (DU) averaged for the summertime period of ANALY and INT.



Fig. 5. (a) Emissions of isoprene for the summertime period, averaged for 2004–2008 in the INT (left) and CLIM (middle) simulations. (b) Deposition flux (μ g m⁻²s⁻¹) of isoprene, averaged for the summertime of INT and CLIM simulations. Differences between INT and CLIM are shown on the right.



Fig. 6. Differences in simulated average surface O₃ (a), isoprene (b), NO_x (c), SO₂ (d), and sulphate (e) fields between INT and CLIM for the summertime (JJAS). Species are in units of μ g m⁻³.

(due to large plant sources in coastal Spain in the emissions dataset). Similar geographical distributions and ground levels of SO₂ are observed in ANALY and INT. Once emitted in the atmosphere, SO_2 leads to the formation of sulphate aerosols. Over the northeastern part of the domain, the levels of sulphate are higher in INT than in ANALY. Figure 2 shows higher precipitation and humidity in ANALY than in INT over this area. These differences imply enhanced transformation of SO2 into sulphate aerosols but also increased wet deposition; those two contrasting effects can explain the differences seen in sulphate concentrations for the two simulations. Figure 4 represents a latitudinal cross-section of sulphate at the longitude of 30° E, averaged for the summertime period of 2004–2008 (JJAS). From 60° N to 70° N, the vertical extent of the sulphate distribution is lower in INT than in ANALY. The difference in sulphate near the surface is due to differing PBL mixing properties in the two simulations. Tropospheric columns of sulphate (Fig. 4) indeed indicate very similar quantities in the two simulations.

To sum up, meteorological forcings (temperature, humidity, horizontal and vertical winds) differ in ANALY and INT. These differences lead to changes in the vertical and horizontal simulated distributions of pollutants. For all pollutants, primary as well as secondary, differences are primarily due to differing PBL mixing heights in the two simulations.

3.1.2 Impact of changes in surface exchanges on European air pollution levels

Comparisons between INT and CLIM indicate the contribution of surface processes on the pollutant level changes. The differences between these simulations are related to the biogenic emissions of isoprene and terpene, the desert dust and sea salt emissions, as well as the deposition velocities. In Fig. 5a, the spatial distributions of summer isoprene



Fig. 7. O₃ deposition velocity averaged for the summertime period simulated by INT (left) and CLIM (middle). Differences between INT and CLIM are shown on the right.

emissions in INT and CLIM are shown. Temperature and solar radiation are key driving variables that regulate emissions of isoprene and other biogenic volatil organic compounds (BVOCs) (Guenther et al., 1993, 1995, 2006). In accordance with the temperature field differences (Fig. 2), higher levels of isoprene are emitted in INT over central Spain, Central Europe, Scandinavia and the northeastern part of Africa than in CLIM. The changes in isoprene emissions induce corresponding changes in the geographical pattern of isoprene concentrations. Elevated concentrations of isoprene (> 1 ppbv) are observed over Central Europe, Greece and North Africa in INT compared to CLIM (Fig. 6b). The mean deposition fluxes ($\mu g m^{-2} s^{-1}$) of isoprene (Fig. 5b) show smaller differences between INT and CLIM.

Over Central Europe, O₃ deposition velocities are higher by up to 0.2-0.3 cm s⁻¹ in INT than in CLIM (Fig. 7). Average nighttime and daytime velocities have been calculated for both INT and CLIM; daytime is considered to be from 08:00 to 16:00 UTC and nighttime is considered to be from 20:00 to 04:00 UTC. For INT, daytime and nighttime mean deposition velocities reach 0.57 cm s^{-1} and 0.24 cm s^{-1} , respectively, over land $(0.06 \text{ cm s}^{-1} \text{ and } 0.05 \text{ cm s}^{-1} \text{ over sea})$. For CLIM, daytime and nighttime mean deposition velocities reach 0.54 cm s⁻¹ and 0.24 cm s⁻¹ over land (0.05 cm s⁻¹ and 0.04 cm s^{-1} over sea). Over land, similar deposition velocities are thus calculated in INT and CLIM. Higher velocities are found during the day as it is known that O₃ deposition velocity has a strong diurnal cycle due to increase in surface resistance at night. The mean deposition fluxes ($\mu g m^{-2} s^{-1}$) of O_3 , NO_x , SO_2 and sulphate have been computed for the summertime period (Fig. 8). In Fig. 6, the changes in concentrations between INT and CLIM follow the changes in deposition fluxes and velocities. Where higher deposition fluxes are seen in INT than in CLIM (parts of Spain, England, Italy and the north and west of France), higher concentrations of ozone are simulated in CLIM. On the contrary, over other areas (mainly in Central Europe) the mean deposition flux is higher in CLIM compared to INT, leading to higher concentrations in INT than in CLIM.

In contrast with O_3 , smaller differences in flux deposition are observed for NO_x , SO_2 and sulphate; nevertheless, these differences lead to differing concentrations between INT and CLIM. In the case of NO_x , higher concentrations observed in CLIM over the northern area (as in England, Belgium) are related to lower flux deposition at the surface. In addition, SO₂ concentrations rise by 1–1.5 ppbv in CLIM over northern Spain and Belgium (Fig. 6c); the simulated changes are related to the SO₂ deposition fluxes (Fig. 8c). Very similar distribution and levels of sulphate aerosol are observed in INT and CLIM (Fig. 6e) across Europe.

In summary, the comparisons between ANALY and CLIM represented in Fig. 9 have revealed the contribution of both meteorological and flux changes to simulated air pollutants. The differences linked to the meteorological parameters or surface processes are pollutant dependent. Depending on the species that are considered, the differences can be driven mainly by the meteorological fields or the emission inventories. The meteorological and surface process effects can also compensate each other. Over the whole domain, the changes in sulphate concentrations between ANALY and CLIM are mostly determined by chemical, physical and dynamical processes due to the meteorological fields (humidity, precipitation). The major changes in isoprene concentrations (Spain, North Africa, Greece) are attributed to both changes in atmospheric circulation and stability (ANALY vs INT), as well as to differences in surface emissions and deposition (INT vs CLIM). For the short lived species NO_x and SO₂, we see that the larger changes are localized near the high emission spots. In case of SO₂, the differences between ANALY and CLIM over Europe are explained by both the changes in deposition fluxes and by the meteorological fields. The O₃ concentration differences between the two simulations are partly related to the changes in meteorological fields (such as temperature) but are principally due to the changes in deposition velocities.

3.2 Statistical results: ANALY and CLIM against AirBase

First, in Sect. 3.2.1, we discuss the interannual variability simulated by the model. The statistical records are then described in detail in Sect. 3.2.2. To finish, Sect. 3.2.3 evaluates



Fig. 8. From top to bottom: (a) deposition flux of O_3 , (b) deposition flux of NO_x , (c) deposition flux of SO_2 and (d) deposition flux of sulphate. Deposition flux are in $\mu g m^{-2} s^{-1}$ and averaged for the summertime period of INT and CLIM simulations.

the impacts of chronology of pollution events on the skill scores.

3.2.1 Model interannual variability

Figure 10 represents the temporal series of the model (ANALY in black line; CLIM in gray line) and monthly measured AirBase data (red line) as an average of the daily mean O_3 (a), NO_x (b), SO_2 (c) and particulate matter PM_{10} (d) from 2004 to 2008 across the European domain. If we subtract the observed and simulated annual cycle averaged for the period 2004–2008 from these time series, positive and negative anomalies remain. The meteorology of ANALY is expected to follow the day-to-day variability in a more

realistic way than CLIM, which reproduces the climate of the decade 2000–2010. The interannual variability of O₃ simulated by ANALY follows the measured variability of meteorological events in terms of correlation and amplitude. The positive and negative anomalies estimated for ANALY are indeed mostly correlated with the observations (CORRO₃H = 0.61). This is obvious for the case of the summer 2006 heat wave. A positive anomaly, slightly underestimated, is calculated with ANALY, while this summer was particularly extreme with an intense production of O₃ (Struzewska and Kaminski, 2008) compared to the mean level of the period 2004–2008. Table 6 summarizes the statistics of the hourly mean and daily M×8h O₃ levels, averaged for the annual and seasonal period of 2004–2008 over all the European



Fig. 9. Same as Fig. 6 but comparing ANALY and CLIM.

stations considered (Table 5). If the annual trend and the interseasonal variability are reproduced, a systematic negative bias is detected with CLIM throughout the summers (MBO₃H = $-4.6 \,\mu g \, m^{-3}$, MBO₃MAX = $-5.2 \,\mu g \, m^{-3}$).

The model–observation comparisons of NO_x presented in Fig. 10b highlight a slightly overestimated annual amplitude of the concentrations for ANALY (with too high winter values and too low summer values), while the winter NO_x values simulated by CLIM are much more overestimated (MBNO_xDM = $8 \mu g m^{-3}$). The dynamics and the intensities of the anomalies are often well-captured by the simulation ANALY; the positive anomalies estimated at the beginning of 2005 and 2006 are for instance well correlated with the observations. Winter pollution characterized by high levels of NO_x (30–40 $\mu g m^{-3}$) is depicted on the time series. For ANALY, the statistics compiled in Table 7

show better performance in term of correlation in winter (CORRNO_xH = 0.42; CORRNO_xDM = 0.55) than in summer (CORRNO_xH = 0.29; CORRNO_xDM = 0.43). During winter, chemical processes that lead to O₃ production are less dominant compared to transport and could explain such differences (Bessagnet et al., 2004).

Time series of monthly mean concentrations of SO_2 from the AirBase stations and the model simulations are represented in Fig. 10c. Results show that the SO_2 concentrations are overestimated for both ANALY and CLIM. Overall, a good agreement is observed for the anomalies between ANALY and the observations in term of amplitude. From the year 2006, we notice a decrease in the observed SO_2 concentrations due to the regulations reducing anthropogenic emissions. Indeed, emissions have been reduced in the sector of power and heat generation with the emission abatement



Fig. 10. (1) Simulated (ANALY: black lines; CLIM: gray lines) and measured (at the AirBase stations; red lines) time series of monthly mean concentrations of O_3 (**a**), NO_x (**b**), SO_2 (**c**) and PM_{10} (**d**). The time series are plotted from 1 January 2004 to 31 December 2008 and averaged over the European domain. Concentrations are in $\mu g m^{-3}$. (2) Anomalies calculated when subtracting the average annual series from the time series in (1).

Table 6. Seasonal and annual statistics obtained with MOCAGE over Europe at the AirBase stations. Statistics are averaged for the 5yr period. Summer: June, July, August and September; winter: December, January, February and March. The calculated statistics are mean bias (MBO₃H, MBO₃MAX; μ g m⁻³), mean normalized bias (MNBO₃H, MNBO₃MAX, %), correlation coefficient (CORRO₃H, CORRO₃MAX), root mean square error (RMSEO₃H, RMSEO₃MAX; μ g m⁻³) and sigma ratio (σ H, σ MAX). Statistics are computed for the O₃ hourly value and O₃ M×8h. SOMO35 values are in μ g m⁻³ d.

O ₃ metrics	ANALY				CLIM			
	Year	JJAS	DJFM	Yea	ar JJAS	DJFM		
MBO ₃ H	2.7	-2.9	6.4	-5.	3 -4.6	-8.2		
MNBO ₃ H	5.1	-4.5	13.8	-9.	8 -7.7	-17.7		
CORRO ₃ H	0.61	0.59	0.58	0.	3 0.32	0.17		
RMSEO ₃ H	24.7	24.9	24.3	34.	3 35.4	34.1		
σH	0.87	0.73	1.04	0.9	8 0.97	1.08		
MBO ₃ MAX	0.01	-8.1	5.7	-6.	3 -5.2	-9.1		
MNBO ₃ MAX	0.04	-9.44	9.8	-8.	8 -5.9	-15.5		
CORRO3MAX	0.7	0.63	0.65	0.	4 0.04	0.25		
RMSEO ₃ MAX	21.3	21.4	21.5	32.	7 35.2	32.2		
σ MAX	0.85	0.66	1.09	0.9	9 0.90	1.22		
SOMO35	3925	1552	915	362	8 2043	531		

NO _x metrics	ANALY				CLIM			
	Year	JJAS	DJFM	Ŋ	lear	JJAS	DJFM	
MBNO _x H	-0.19	-2.5	2.2		2.1	-2.1	8.4	
MNBNO _x H	-1.2	-32.5	13.9	1	7.0	-28.8	55.1	
CORRNO _x H	0.46	0.29	0.42	C).14	0.08	0.05	
RMSENO _x H	13.2	8.8	15.9	2	20.5	10.6	29.2	
$\sigma NO_x H$	1.20	0.91	1.19	1	.64	1.00	1.78	
MBNO _x DM	-0.12	-2.4	2.2		2.1	-2.1	8.0	
MNBNO _x DM	-0.67	-32.4	14.6	1	7.5	-27.9	54.3	
CORRNO _x DM	0.61	0.43	0.55	0	0.20	0.09	0.07	
RMSENO _x DM	9.8	6.1	11.9	1	7.6	7.6	25.7	
$\sigma NO_{x}DM$	1.34	0.98	1.30	1	.92	1.07	2.08	

Table 7. Same as Table 6 for hourly value and daily mean NO_x concentrations.

Table 8. Same as Table 6 for hourly value and daily mean SO₂ concentrations.

SO ₂ metrics	ANALY				CLIM			
	Year	JJAS	DJFM	Year	JJAS	DJFM		
MBSO ₂ H	0.39	-0.06	0.79	0.46	-0.06	1.3		
MNBSO ₂ H	18.6	-2.9	36.7	24.2	-3.1	50.8		
CORRSO ₂ H	0.23	0.15	0.27	0.02	-0.01	-0.01		
RMSESO ₂ H	4.3	3.3	4.9	5.5	3.8	7.4		
σSO_2H	1.37	1.16	1.53	1.65	1.21	2.05		
MBSO ₂ DM	0.37	-0.07	0.79	0.45	-0.07	1.25		
MBNSO ₂ DM	17.8	-3.5	35.4	23.2	-4.0	49.7		
CORRSO ₂ DM	0.36	0.3	0.4	0.03	-0.01	-0.01		
RMSESO ₂ DM	3.2	2.3	3.7	4.5	2.8	6.29		
$\sigma SO_2 DM$	1.53	1.35	1.70	1.92	1.47	2.42		

strategies in some European countries during the period of 2000–2010 (EEA, 2007). In our simulations, we kept the same emissions inventory representative of the year 2003. According to the time series of anomalies, and as described in Table 8, the biases calculated in ANALY and CLIM are of the same order of value (MBSO₂DM = $0.37 \,\mu g \,m^{-3}$ for ANALY and MBSO₂DM = $0.45 \,\mu g \,m^{-3}$ for CLIM).

Although the simulation ANALY presents a persistent negative bias (Table 9), it has the capability to reproduce the dynamics of PM_{10} for each year. The underestimation of PM_{10} can be explained principally by the lack of secondary particulate and nitrate aerosols in our representation of PM_{10} . During summer, when photochemistry favors the formation of these particulates, the biases between simulated and observed concentrations become greater (MBPM₁₀DM = $-11.9 \,\mu g \,m^{-3}$, Fig. 10 and Table 9). In the case of CLIM, the time series of anomalies displays the capability of the model to reproduce the particulate matter events (CORRPM₁₀DM = 0.39), although the model hardly reproduces their amplitude.

3.2.2 Statistical results

The statistics of the model are spatially displayed in Fig. 11; we illustrate the mean biases for O_3 daily M×8h values, as well as for NO_x and SO₂ daily mean concentrations across the representative European stations. The scores are averaged for the summer season (JJAS). Regarding the results of simulation ANALY in the case of O_3 , two distinct spatial regimes can be distinguished from the figures; positive and low biases are shown over Germany while negative biases are noticed in Spain and Italy. The correlations (not shown) are also more elevated in the northern part of Europe, notably in Germany (0.6–0.8) while in Southern Europe (Italy and Spain), the performance of the model is rather low (Pay et al., 2010). Comparisons between the statistical metrics of ANALY and CLIM indicate comparable biases over Europe for the daily M×8h O₃. The spatial distribution and the amplitude of the negative and positive biases are mostly similar, except for the stations in Germany and France, which display higher positive $(30-40 \,\mu g \, m^{-3})$ and negative biases in CLIM than in ANALY, respectively. As shown in

Table 9. Same as Table 6 for the PM_{10} daily mean. The calculated statistics are MBPM₁₀DM (μ g m⁻³), MFBPM₁₀DM (%), MFEPM₁₀DM (

PM ₁₀ metrics	ANALY			CLIM			
	Year	JJAS	DJFM		Year	JJAS	DJFM
MBPM ₁₀ DM	-8.2	-11.9	-4.6		-5.3	-11.8	4.6
MFBPM ₁₀ DM	-59.5	-94.6	-33.6		-48.5	97.9	3.5
MFEPM ₁₀ DM	75.1	96.9	58.0		85.4	101.7	77.4
CORRPM ₁₀ DM	0.39	0.2	0.48		0.04	-0.04	0.03
RMSEPM ₁₀ DM	15.1	14.4	15.1		21.6	15.3	28.3
$\sigma PM_{10}DM$	0.96	0.49	1.02		1.39	0.60	1.59



Fig. 11. Spatial distribution of mean bias (μ g m⁻³) for daily M×8h O₃ (**a**), daily mean NO_x (**b**) and daily mean SO₂ (**c**) concentrations for the average summertime period of 2004–2008. The two rows represent ANALY (top) and CLIM (bottom).

Table 6, for ANALY, annual correlations of hourly values and daily M×8h O₃ concentrations reach 0.61 and 0.7, respectively. Also, annual and seasonal MNB values for hourly and daily M×8h O₃ show good performance in accordance with the recommendation of US-EPA (MNBE $\leq \pm 15\%$). In summer, the model tends to slightly underestimate the hourly levels of O₃ (MBO₃H = $-2.9 \mu \text{g m}^{-3}$) and daily M×8h concentrations (MBO₃MAX = $-8.1 \mu g m^{-3}$), while it overestimates concentrations in winter months ($MBO_3H =$ $6.4 \mu g m^{-3}$; MBO₃MAX = $5.7 \mu g m^{-3}$). Correlation values are lowest for both hourly values and daily M×8h over winter, 0.58 and 0.65, respectively. During winter, O₃ is controlled by processes other than photochemistry (such as boundary conditions, deposition, titration); O₃ is thus sensitive to physical and dynamic characteristics (higher resolution of forcings, emissions, deposition velocities). Higher negative biases are thus exhibited by CLIM in wintertime $(MBO_3H = -8.2 \mu g m^{-3}; MBO_3MAX = -9.1 \mu g m^{-3})$, as seen in Fig. 10, due to the differences in deposition velocities and in the concentrations in the lower atmosphere. Daily M×8h are also best reproduced by the model (correlations between 0.63 and 0.7); the variability of daily $M \times 8h$ is mainly driven by photochemistry as well as the boundary layer height. To examine if the model is able to simulate the variability of O₃ concentrations, we used the sigma ratio, which is the standard deviation of the modeled time series divided by the standard deviation of the observed time series (Table 3). In Table 6 the sigma ratio values are summarized, averaged for the period of 2004-2008. For both simulations, the model underestimates the observed variability of hourly and daily M×8h values, except for the winter season. In addition, two health related parameters are considered: SOMO35 and the number of exceedance days. SOMO35 corresponds to the yearly sum of the differences between daily maximum 8 h running average concentrations that are greater than 35 ppb (Amann et al., 2005). It is used as an indicator for O₃ health impact and is recommended by the World Health Organization (WHO). The values of SOMO35 are summarized in Table 6. When averaged over all of the European stations considered, the observed seasonal levels reach 2221 μ g m⁻³ d and 496 μ g m⁻³ d in summer and winter, respectively (4118 μ g m⁻³ d for the all year). Both simulations catch the levels of SOMO35 and the seasonal variation (van Loon et al., 2007). According to ANALY, about 40 % of the SOMO35 is produced during summer and 20 % during winter. Over the summer period, the number of days with O_3 exceeding the 120 µg m⁻³ threshold for the daily maximum 8-h average concentration is underestimated in ANALY (n = 5.7 days) and fairly well estimated in CLIM (n = 12 days), in comparison to the observations (n = 15.4)days) from the European stations. The mean ozone concentrations above the threshold of $120 \,\mu g \, m^{-3}$ for the M×8h simulated by ANALY are mostly in line with the observations or within the interannual variability. More elevated values are reached in CLIM than ANALY, as shown for the French and Italian stations (Fig. 12a). Figure 12b shows the percentile of daily O₃ maximum simulated by ANALY and CLIM simulations. The interval between the 20th and 70th percentiles display similar values for both simulations. The occurrence of extreme values (maxima) is underestimated by the model for both simulations. As seen previously, above the threshold of $180 \,\mu g \, m^{-3}$, CLIM simulates a higher number of occurrences than the observations. These figures depict, however, overall that MOCAGE driven by climate model outputs as forcings is able to simulate realistic ozone concentrations over Europe.

As exposed in Table 5, 354 stations were used to provide NO_x measurements throughout Europe. Considering the spatial distribution of mean biases for daily mean NO_x, statistical results show satisfactory seasonal mean bias for ANALY $(MBNO_xDM = -2.4 \mu g m^{-3} during summer)$ without spatial pattern between north and south. Similar geographical distributions and values of mean bias are displayed for CLIM (Fig. 11), while the summer mean bias reaches $-2.1 \,\mu g \, m^{-3}$ (Table 7). The spatial distribution of the correlation coefficients shows a large variability per station; while northern stations display high correlations (0.6 < r < 0.8), low correlations are observed in Southern Europe (r < 0.4). The performances of the model are reduced with CLIM for all seasons, notably during winter (Fig. 7) when the concentrations of NO_x are overestimated, as shown in Fig. 10. Thus, in winter, MNBNO_xH and MNBNO_xDM reach 55.1 % and 54.3 %, respectively; also, RMSENO_xH and RMSENO_xDM reach $29.2 \,\mu\text{g m}^{-3}$ and $25.7 \,\mu\text{g m}^{-3}$, respectively. In summer, the MNB values for hourly and daily mean are near the uncertainty proposed by EC and US-EPA for the ANALY simulation. Globally, the annual and seasonal daily mean statistics **Fig. 12. (a)** Summertime average ozone concentrations (over the stations available) above the threshold of $120 \,\mu g \, m^{-3}$ over 6 European countries. FR is France, ES is Spain, DE is germany, GB is England, IT is Italy, and PL is Poland. The standard deviation measuring the interannual variability is represented by the vertical bars. (b) Distribution of daily O₃ maximum percentiles for AirBase measurements (red) and MOCAGE simulations (ANALY in black, CLIM in gray).

present better performances in comparison with the hourly values.

For SO₂, low correlations are mainly concentrated in Southern Europe (coefficients under 0.2), as in Spain, while some northern stations display high correlations (r > 0.7) in regard to statistical results of ANALY. Averaged over all European stations, the summertime mean correlation of daily mean SO₂ reaches 0.3. Considering the mean bias for summer (Fig. 11), low biases are depicted across all the stations (MBSO₂DM = $-0.07 \,\mu g \,m^{-3}$ for ANALY and CLIM). Nevertheless, the stations located in Poland display high positive biases ($> 2 \,\mu g \,m^{-3}$). The uncertainties of the emissions inventory in Eastern Europe may contribute to the higher bias observed. In some stations in Spain, higher bias is also observed, due to the emission inventory in part. The regulation of SO₂ emissions in Spain have lead to an emission reduction of 50 % (Spain Environment Ministry, 2011). As shown



in Table 8, the highest correlations are obtained in winter (CORRSO₂H = 0.27; CORRSO₂DM = 0.40), but the concentrations are overestimated leading to a high value of MNB close to the EC criteria.

As seen in Fig. 10, the model presents a systematic negative bias for the simulated concentrations of PM₁₀. For the ANALY simulation, the correlation coefficient for the annual daily mean is 0.39, while it reaches 0.2 and 0.48 for the summer and winter season, respectively (Table 9). The spatial distribution of mean bias and correlations (not shown here) present a homogeneous pattern over Europe. The annual MFE (MFEPM₁₀DM = 75.1%) and MFB $(MFBPM_{10}DM = -59.5\%)$ calculated for ANALY does not meet the performance criteria or the performance goal proposed by Boylan and Russell (2006). The performance of the model is better during winter when the MBPM₁₀DM is about $-4.6 \,\mu\text{g m}^{-3}$ (against $-11.9 \,\mu\text{g m}^{-3}$ in summer) and the mean correlation reaches 0.48 (against 0.2 for the summer). The underestimation of PM_{10} can be explained by the lack of secondary particulate and nitrate aerosols in our representation of PM_{10} . Differences between the seasons are linked to chemical processes, dominant in summer, which favor the formation of these particulates and increase the bias between simulated concentrations (lacking these chemical processes) and observations.

Several air quality models operated in Europe have been evaluated either individually or in comparison to other models in the literature. In the following discussion, a quick comparison with other regional air quality models (Hass et al., 2003; van Loon et al., 2004, 2007) and MOCAGE will be carried out in order to situate our model among the community. For this reason, we used the studies similar with our simulation ANALY, which had a long time scale of 1 yr over the European domain on a regional scale with horizontal resolutions similar to MOCAGE. Also, these models were evaluated against ground observations at rural sites from AirBase and EMEP. Concerning O₃ daily M×8h, satisfactory performances are displayed with MOCAGE, in terms of annual MNB values: 0.04% versus -1 to 10%; correlations: 0.7 versus 0.69-0.84 (van Loon et al., 2007); and RMSE: $21.3 \,\mu g \, m^{-3}$ versus $18.1 - 25.5 \,\mu g \, m^{-3}$ (van Loon et al., 2004). Values for summer and winter daily M×8h O₃ are also in the range of other models, as for the correlations (0.63 versus 0.61–0.77 for summer; 0.65 versus 0.45–0.62 for winter) and the MNB (-9.44% versus -5 to 8% forsummer; 9.8 % versus -20 to 15 % for winter) according to the study of van Loon et al. (2007). The MOCAGE performances for NO_x can be compared with the performance of NO₂ in other models. The annual correlation of daily mean NO_x obtained in this study reaches 0.61, compared to 0.03-0.52 (Hass et al., 2003; van Loon et al., 2004) and the RMSE value is around $9.8 \,\mu g \, m^{-3}$ versus $8.5 - 13.9 \,\mu g \, m^{-3}$ (Hass et al., 2003; van Loon et al., 2004). As for O₃, the MOCAGE results for SO₂ show good performances in comparison with the other studies. The annual daily mean correlation is among the higher values (0.36 versus 0.24–0.49). The calculated RMSE reaches $3.2 \,\mu g \, m^{-3}$ against 2.7–10.9 $\mu g \, m^{-3}$ for the other models (Hass et al., 2003; van Loon et al., 2004). For PM₁₀, statistical results are in the same range as for other studies; nevertheless, the annual daily mean correlation is rather low and reaches 0.39 compared to 0.38–0.55; the annual RMSE is 15.1 $\mu g \, m^{-3}$ (versus 12.4–16.6 $\mu g \, m^{-3}$).

To summarize, MOCAGE performs well according to the comparisons between ANALY and AirBase observations, as discussed previously. The statistical scores of O₃, NO_x and SO₂ display satisfactory performances compared to other studies, while the accuracy in our representation of PM_{10} exhibits poorer results, which are expected by design. Comparisons between the simulations ANALY and CLIM have shown that the geographical distribution of mean biases are quite similar for each pollutant considered. In this section, the model to observation comparisons were based on a common approach, which consists in comparing each year of the simulation with the matching measured values from the Air-Base database. In CLIM, the meteorological forcings are representative of the current decade; there is no particular match in the sequence of years, and the representativeness of the skill scores can be assessed by permutations of all years. By doing the same for ANALY, the comparisons will allow us to determine which statistical tools are useful to consider for future studies.

3.2.3 Impacts of chronology of pollution events

We evaluated each year of the simulations with 5 yr of measurements, giving 25 model-to-date pairs of statistics for both ANALY and CLIM. On the time basis of the 2004–2008 period, we calculated every possible permutation, but on this period of 5 yr, we did not consider the same year of measurements more than once. We also filtered out the cases when one or more simulated years correspond with the same years of data. Finally, these conditions let us consider 44 realizations, which provide a large statistical basis.

In order to give a concise statistical summary, we used the Taylor diagrams, which indicate how well observed and simulated patterns match each other in terms of correlation and normalized standard deviation (NSD) (Taylor, 2001). The correlation coefficient R gives a measure of the co-variance of simulated and observed values. The NSD gives a measure of the amplitude of the variance in modeled values versus observed values. When NSD reaches a value lower than 1, it means that the temporal standard deviation in simulated values is lower than observed. Figure 13 shows the normalized Taylor plots that summarize the ANALY and CLIM permutations. The statistics are computed for the summertime daily $M \times 8h O_3$ concentrations (a) and daily averages of NO_x (b) and SO_2 (c). For each plot, "ANALY" refers to the reference model-observation analyses (black cross) whereas "ANALY-p" (blue symbols) and "CLIM-p" (red symbols) refer to the permutation cases. Considering



Fig. 13. Taylor plots of the comparison between modeled and observed $M \times 8h O_3$ concentrations, daily mean NO_x and daily mean SO_2 . The radial distance from the origin corresponds to NSD, and *R* corresponds to the azimuthal position.

Table 10. Seasonal JJAS statistics obtained with the permutations ANALY-p and CLIM-p. Statistics are the median values of the permutations. The calculated statistics are mean bias ($\mu g m^{-3}$), mean normalized bias (%), correlation coefficient, root mean square error ($\mu g m^{-3}$) and sigma ratio. Statistics are computed for the O₃ hourly value and daily M×8h concentrations.

O ₃ metrics			
	ANALY-p	CLIM-p	ANALY
MBO ₃ MAX	-7.9	-5.6	-8.1
MNBO3MAX	-9.1	-6.4	-9.4
CORRO3MAX	0.04	0.07	0.63
RMSEO3MAX	31	34.6	21.4
$\sigma O_3 MAX$	0.67	0.9	0.66
MBO ₃ H	-2.8	-5.0	-2.9
MNBO3H	-4.4	-7.9	-4.5
CORRO ₃ H	0.33	0.34	0.59
RMSEO ₃ H	30.9	35.1	24.9
σO ₃ H	0.75	0.98	0.73

the results of ozone first, comparisons between ANALY and ANALY-p confirm there is no day-to-day variability with the permutations of ANALY-p as shown by the correlations. As summarized in Table 10, for ANALY-p, the median value of the correlations reach 0.04 (against 0.63 for ANALY). The median RMSE of ANALY-p are not as good as the median RMSE for ANALY for both the hourly (RMSEO₃H = $31 \mu g m^{-3}$ for ANALY-p and $21.4 \mu g m^{-3}$ for ANALY) and daily M×8h (RMSEO₃MAX = $30.9 \mu g m^{-3}$ for ANALY-p and 24.9 μ g m⁻³ for ANALY) of O₃. Nevertheless, very similar values of MNBO3MAX and MNBO3H in the ranges of -9% and -4.5%, respectively, are obtained. The standard deviation values (σ) are similar in ANALY and ANALY-p and range around 0.66, meaning that the model underestimates the daily M×8h ozone variability. The same conclusions can be extended to the daily averages of NO_x, SO₂ (Table 11) and PM₁₀ (Table 12) concentrations. ANALY and ANALY-p only differ by the correlations while the MB, RMSE and variances are quite similar. For the daily mean NO_x (Fig. 13), the NSD are close to the reference, meaning that the amplitude of the simulated NO_x agrees with the observations. To sum up, these results point out that, for all the species, the correlations calculated for ANALY-p are weaker

NO _x metrics				SO ₂ Metrics			
	ANALY-p	CLIM-p	ANALY		ANALY-p	CLIM-p	ANALY
MBNO _x DM	-2.5	-2.0	-2.4	MBSO ₂ DM	-0.09	-0.04	-0.07
MNBNO _x DM	-32.6	-27.8	-32.4	MNBSO ₂ DM	-2.8	-1.0	-3.5
CORRNO _x DM	0.09	0.07	0.43	CORRSO ₂ DM	-0.004	0.005	0.3
RMSENO _x DM	7.1	7.6	6.1	RMSESO ₂ DM	2.6	2.7	2.3
$\sigma NO_{X}DM$	0.98	1.07	0.98	$\sigma SO_2 DM$	1.39	1.47	1.35
MBNO _x H	-2.5	-2.1	-2.5	MBSO ₂ H	-0.08	-0.04	-0.06
MNBNO _x H	-32.7	-28.6	-32.5	MNBSO ₂ H	-2.6	-0.68	-2.9
CORRNO _x H	0.09	0.07	0.29	CORRSO ₂ H	-0.008	-0.01	0.15
RMSENO _x H	9.9	10.5	8.8	RMSESO ₂ H	3.6	3.8	3.3
$\sigma NO_X H$	0.90	0.98	0.91	σSO_2H	1.18	1.21	1.16

Table 11. Same as Table 10 for hourly values and daily average of NO_x and SO₂.

Table 12. Seasonal JJAS statistics obtained with the permutations ANALY-p and CLIM-p. Statistics represent the median values of the permutations. The calculated statistics are mean bias (μ g m⁻³), mean fractional bias (%), mean fractional error (%), correlation coefficient, root mean square error (μ g m⁻³) and sigma ratio. Statistics are computed for the PM₁₀ daily mean concentrations.

PM ₁₀ metrics			
	ANALY-p	CLIM-p	ANALY
MBPM ₁₀ DM	-11.9	-11.8	-11.9
MFBPM ₁₀ DM	-93.8	-98.6	-94.6
MFEPM ₁₀ DM	97	102.2	96.9
CORRPM ₁₀ DM	0.01	0.01	0.2
RMSEPM ₁₀ DM	14.9	15.1	14.4
$\sigma PM_{10}DM$	0.49	0.59	0.49

than ANALY due to the permutations. The RMSE values increase by 19%, 11% and 8% for the hourly values of O₃, NO_x and SO₂, respectively, from ANALY to ANALYp. However, for the metrics MB, MNB and σ , similar values are computed.

For O₃ (Fig. 13), low and similar correlations of daily $M \times 8h$ levels are calculated for both ANALY-p and CLIM-p. As shown in Table 10, the correlations between the observed and simulated hourly values (0.33 for ANALY-p and 0.34 for CLIM-p) are higher than daily $M \times 8h$. The daily variability of ozone, characterized by higher levels during afternoon and lower values during nighttime hours is still captured with the permutations. The RMSE values show lower performances for CLIM-p compared to ANALY-p for both daily $M \times 8h$ (+10%) and hourly O₃ (+13%) levels. The σ values indicate the tendency of ANALY-p to underestimate ozone variance in summer. The CLIM-p simulations show better σ values (0.9 and 0.98 versus 0.67 and 0.75 for ANALY-p), which is unexpected and cannot be interpreted as greater performance. The MNB of M×8h ozone

show a tendency of model underestimation as the median reaches -9.1% in ANALY-p and -6.4% in CLIM-p. For the hourly and daily average NO_x and SO₂ concentrations, the simulations ANALY-p and CLIM-p are now well correlated (Table 11). During summer, ANALY-p and CLIM-p underestimate the daily mean and hourly NO_x concentrations. From ANALY-p to CLIM-p, the median MNBNO_xDM and MNBNO_xH change by about 14 % while the RMSENO_xDM and RMSENO_xH change by about 7 %. Concerning the amplitude of the NO_x variances (Fig. 13), a satisfactory agreement is observed; σ reaches 0.98 in ANALY-p and 1.07 in CLIM-p for daily mean NO_x (Table 11). For the SO₂ results, the variance is overestimated by the model for both ANALYp and CLIM-p. For the daily mean of PM₁₀ concentrations, the amplitude of the variances are in line for ANALY-p and CLIM-p simulations. The simulations underestimate the variability of the PM_{10} ($\sigma PM_{10}DM = 0.49$ for ANALY-p and $\sigma PM_{10}DM = 0.59$ for CLIM-p). MBPM_{10}DM reaches similar values for ANALY-p and CLIM-p. MFB and MFE metrics do not reach the performance criterion.

To summarize, the use of permutations have made the comparisons between the simulations suitable for discussion of the statistical results. Comparisons between the reference case "ANALY" with "ANALY-p" corroborate the incorrect phasing between the measurements and simulations when the day-to-day variability is not reproduced. Finally, the results allow us to conclude that statistical metrics such as variances, MB, MNB and RMSE give robust and sound information when climate forcings are used to drive the model.

4 Conclusions

This paper has investigated how different the hindcasts of an air quality modeling system are when using two different types of meteorological forcings: meteorological analyses or fields from a climate model. This is ground work needed to qualify and properly interpret statistical conclusions that can be drawn from simulations of air quality in a future climate.

The comparisons between three 5-yr experiments allow us to quantify the relative importance of changes in surface fields and upper air meteorology. We find that both elements contribute to changes in O_3 concentrations. Differences in sulphate aerosols and in isoprene (as a proxy for biogenic volatile organic compounds) are mainly related to changes in meteorology and mixing, while it is the contrary for SO_2 and NO_x , which essentially depend upon changes in surface fluxes.

The skill of the reference simulation (analysed forcings) to reproduce European surface observations is in the range of previous reference evaluation studies (Hass et al., 2003; van Loon et al., 2004, 2007). As expected, the simulation based upon forcings from a climate model is not as skillful at reproducing observations, since it cannot follow day-to-day variations by design. Nevertheless, the geographical distributions of the mean biases are similar in the two simulations. The comparisons of SOMO35 and the distribution of O₃ maximum percentiles support the capability of MOCAGE, driven by meteorological fields from a climate model, to simulate realistic European ozone levels.

The objective of this work was to determine useful statistical metrics that can be used for models driven by climate model meteorological parameters. For O₃, we show that simulations using either analyses or climate model forcings follow the same tendency: hourly and M×8h concentrations are slightly underestimated and the biases and RMSE are in the same range of values. Similar conclusions are observed for the daily averaged and hourly values of NO_x and SO₂, as well as daily PM₁₀. The amplitude of variance is accurately reproduced when the model is driven by climate fields. Finally, as for the standard deviation, statistical results of MB, MNB and RMSE can be interpreted with some degree of confidence.

Supplementary material related to this article is available online at: http://www.geosci-model-dev.net/5/ 1565/2012/gmd-5-1565-2012-supplement.pdf.

Acknowledgements. This work was partly founded by ADEME (Agence De l'Environnement et de la Maitrise de l'Energie, http://www.ademe.fr). We would like to thank N. Poisson (ADEME), L. Rouil (INERIS) and M. Beekman (CNRS/LISA) for useful discussions.

Edited by: P. Jöckel



The publication of this article is financed by CNRS-INSU.

References

- Amann, M., Bertok, I., Cofala, J., Gyarfas, F., Heyes, C., Klimont, Z., Schopp, W., and Winiwater, W.: Clean air for Europe (CAFE) programme final report, Tech. rep., 2005.
- Andersson, C. and Engardt, M.: European ozone in a future climate: Importance of changes in dry deposition and isoprene emissions, J. Geophys. Res., 115, D02303, doi:10.1029/2008JD011690, 2010.
- Bechtold, P., Bazile, E., Guichard, F., Mascart, P., and Richard, E.: A mass-flux convection scheme for regional and global models, Q. J. Roy. Meteor. Soc., 127, 869–886, 2001.
- Bessagnet, B., Hodzic, A., Vautard, R., Beekmann, M., Cheinet, S., Honoré, C., Liousse, C., and Rouïl, L.: Aerosol modeling with CHIMERE-preliminary evaluation at the continental scale, Atmos. Environ., 38, 2803–2817, 2004.
- Bousserez, N., Attié, J.-L., Peuch, V.-H., Michou, M., Pfister, G., Edwards, D., Emmons, L., Mari, C., Barret, B., Arnold, S. R., Heckel, A., Richter, A., Schlager, H., Lewis, A., Avery, M., Sachse, G., Browell, E. V., and Hair, J. W.: Evaluation of the MOCAGE chemistry transport model during the ICARTT/ITOP experiment, J. Geophys. Res., 112, D10S42, doi:10.1029/2006JD007595, 2007.
- Boylan, J. and Russell, A.: PM and light extinction model performance metrics, goals, and criteria for three-dimensional air quality models, Atmos. Environ., 40, 4946–4959, 2006.
- Carvalho, A., Monteiro, A., Solman, S., Miranda, A., and Borrego, C.: Climate-driven changes in air quality over Europe by the end of the 21st century, with special reference to Portugal, in: Environmental Science and Policy, Elsevier, Oxford, England, 445–558, 2010.
- Chang, J. and Hanna, S.: Air quality model performance evaluation, Meteor. Atmos. Phys., 87, 167–196, 2004.
- Courtier, P., Freydier, C., Geleyn, J. F., Rabier, F., and Rochas, M.: The ARPEGE project at Météo France, in: Atmospheric Models, vol. 2, Workshop on Numerical Methods, Reading, UK, 193– 231, 1991.
- Cuvelier, C., Thunis, P., Vautard, R., Amann, M., Bessagnet, B., Bedogni, M., Berkowicz, R., Brandt, J., Brocheton, F., Builtjes, P., Coppalle, A., Denby, B., Douros, G., Graf, A., Hellmuth, O., Honoré, C., Hodzic, A., Jonson, J., Kerschbaumer, A., de Leeuw, F., Minguzzi, E., Moussiopoulos, N., Pertot, C., Pirovano, G., Rouïl, L., Schaap, M., Stern, R., Tarrason, L., Vignati, E., Volta, M., White, L., Wind, P., and Zuber, A.: CityDelta: a model intercomparison study to explore the impact of emission reductions in European cities in 2010, Atmos. Environ., 41, 189–207, 2007.
- Dawson, J. P., Adams, P. J., and Pandis, S. N.: Sensitivity of ozone to summertime climate in the Eastern USA: a modeling case study, Atmos. Environ., 41, 1494–1511, 2007.

- Dawson, J. P., Racherla, P. N., Lynn, B. H., Adams, P. J., and Pandis, S. N.: Impacts of climate change on regional and urban air quality in the Eastern United States: role of meteorology, J. Geophys. Res., 114, D05308, doi:10.1029/2008JD009849, 2009.
- Dentener, F. E. A.: The global atmospheric environment for the next generation, Environ. Sci. Technol., 40, 3586–3594, doi:10.1021/es0523845, 2006.
- Déqué, M., Dreveton, C., Braun, A., and Cariolle, D.: The ARPEGE/IFS atmosphere model: a contribution to the French community climate modelling, Clim. Dynam., 10, 249–266, 1994.
- Dufour, A., Amodei, M., Ancellet, G., and Peuch, V.-H.: Observed and modelled "chemical weather" during ESCOMPTE, Atmos. Res., 74, 161–189, doi:10.1016/atmosres.2004.04.013, 2004.
- EEA: Air pollution by ozone in Europe in summer 2004, Technical Report, 3/2005, Copenhagen, Denmark, available at: http://www.eea.europa.eu/publications/technical_report_2005_3, 2005.
- EEA: Air pollution in Europe 1999–2004, Technical Report, 2/2007, Office for Official Publications of the European Communities, Luxembourg, 79 pp., 2007.
- European Commission: Council directive 1999/30/EC of 22 April 1999 relating to limit values for sulphur dioxide, nitrogen dioxide and oxides of nitrogen, particulate matter and lead in ambient air, Technical Report 1999/30/EC, L163, Off. J. Eur. Comm., 1999.
- European Commission: Commission decision of 17 October 2001 amending annex v to Council directive 1999/30/EC relating to limit values for sulphur dioxide, nitrogen dioxide and oxides of nitrogen, particulate matter and lead in ambient air (text with EEA relevance) notified under document number C (2001) 3091, Technical Report 2001/744/EC, 278, Off. J. Eur. Comm., 2001.
- European Commission: Directive 2002/3/EC of the European Parliament and of the Council of 12 February 2002 relating to ozone in ambient air, Technical Report 2002/3/EC, 167, Off. J. Eur. Comm., 2002.
- European Commission: Directive 2008/50/EC of the European Parliament and of the Council of 21 May 2008 on ambient air quality and cleaner air for Europe, Technical Report 2008/50/EC, L152, Off. J. Eur. Comm., 2008.
- Fuhrer, J. and Booker, F.: Ecological issues related to ozone: agricultural issues, Environ. Int., 29, 141–154, 2003.
- Giorgi, F. and Meleux, F.: Modelling the regional effects of climate change on air quality, C. R. Geosci., 339, 721–733, 2007.
- Guenther, A., Zimmerman, P., Harley, P., Monson, R., and Fall, R.: Isoprene and monoterpene emission rate variability: model evaluation and sensitivity analysis, J. Geophys. Res., 98, 12609– 12617, doi:10.1029/93JD00527, 1993.
- Guenther, A., Hewitt, C. N., Erickson, D., Fall, R., Geron, C., Graedel, T., Harley, P., Klinger, L., Lerdau, M., Mckay, W., Pierce, T., Scholes, B., Steinbrecher, R., Tallamraju, R., Taylor, J., and Zimmerman, P. A.: A global model of natural volatile organic compound emissions, J. Geophys. Res., 100, 8873–8892, doi:10.1029/94JD02950, 1995.
- Guenther, A., Karl, T., Harley, P., Wiedinmyer, C., Palmer, P. I., and Geron, C.: Estimates of global terrestrial isoprene emissions using MEGAN (Model of Emissions of Gases and Aerosols from Nature), Atmos. Chem. Phys., 6, 3181–3210, doi:10.5194/acp-6-3181-2006, 2006.
- Guerova, G. and Jones, N.: A global model study of ozone distributions during the August 2003 heat wave in Europe, Environ.

Chem., 4, 85–292, 2007.

- Guicherit, R. and van Dop, H.: Photochemical production of ozone in Western Europe (1971–1975) and its relation to meteorology, Atmos. Environ., 11, 145–155, doi:10.1071/EN07027, 1977.
- Hass, H., van Loon, M., Kessler, C., Stern, R., Matthijsen, J., Sauter, F., Zlatev, Z., Langner, J., Foltescu, V., and Schaap, M.: Aerosol modeling: results and intercomparison from European regional scale modeling systems, Technical Report, EUROTRAC 2 Report, EUREKA Environmental Project, GLOREAM, 2003.
- Hedegaard, G. B., Brandt, J., Christensen, J. H., Frohn, L. M., Geels, C., Hansen, K. M., and Stendel, M.: Impacts of climate change on air pollution levels in the Northern Hemisphere with special focus on Europe and the Arctic, Atmos. Chem. Phys., 8, 3337–3367, doi:10.5194/acp-8-3337-2008, 2008.
- Hogrefe, C., Lynn, B., Civerolo, K., Ku, J.-Y., Rosenzweig, J. R. C., Goldberg, R., Gaffin, S., Knowlton, K., and Kinney, P. L.: Simulating changes in regional air pollution over the Eastern United States due to changes in global and regional climate and emissions, J. Geophys. Res., 109, D22301, doi:10.1029/2004JD004690, 2004.
- Hollingsworth, A., Engelen, R., Textor, C., Boucher, O., Chevallier, F., Dethof, A., Elbern, H., Eskes, H., Flemming, J., Granier, C., Kaiser, J. W., Morcrette, J. J., Rayner, P., Peuch, V.-H., Rouïl, L., and the GEMS consortium: Toward a monitoring and forecasting system for atmospheric composition: the GEMS project, Bull. Amer. Meteor. Soc., 89, 1147–1164, doi:1S230.1175/2008BAM55.1, 2008.
- Honoré, C., Rouïl, L., Vautard, R., Beekmann, M., Bessagne, B., Dufour, A., Elichegaray, C., Flaud, J.-M., Malherbe, L., Meleux, F., Menut, L., Martin, D., Peuch, A., Peuch, V.-H., and Poisson, N.: Predictability of European air quality: assessment of 3 years of operational forecasts and analyses by the PREV'AIR system, J. Geophys. Res., 113, D04301, doi:10.1029/2007JD008761, 2008.
- Huszar, P., Juda-Rezler, K., Halenka, T., Chervenkov, H., Syrakov, D., Krüger, B. C., Zanis, P., Melas, D., Katragkou, E., Reizer, M., Trapp, W., and Belda, M.: Effects of climate change on ozone and particulate matter over Central and Eastern Europe, Climate Research, 50, 51–68, doi:10.3354/cr01036, 2011.
- Juda-Rezler, K., Reizer, M., Huszar, P., Krüger, B. C., Zanis, P., Syrakov, D., Katragkou, E., Trapp, W., Melas, D., Chervenkov, H., Tegoulias, I., and Halenka, T.: Modelling the effects of climate change on air quality over Central and Eastern Europe: concept, evaluation and projections, Clim. Res., 53, 179– 203, doi:10.3354/cr01072, 2012.
- Joly, M. and Peuch, V.-H.: Objective classification of air quality monitoring sites over Europe, Atmos. Environ., 47, 111–123, 2012.
- Katragkou, E., Zanis, P., Tegoulias, I., Melas, D., Kioutsioukis, I., Krüger, B. C., Huszar, P., Halenka, T., and Rauscher, S.: Decadal regional air quality simulations over Europe in present climate: near surface ozone sensitivity to external meteorological forcing, Atmos. Chem. Phys., 10, 11805–11821, doi:10.5194/acp-10-11805-2010, 2010.
- Katragkou, E., Zanis, P., Kioutsioukis, I., Tegoulias, I., Melas, D., Krüger, B. C., and Coppola, E: Future climate change impacts on summer surface ozone from regional climate-air quality simulations over Europe, J. Geophys. Res., 116, D22307, doi:10.1029/2011JD015899, 2011.

- Knowlton, K., Rosenthal, J. E., Hogrefe, C., Lynn, B., Gaffin, S., Goldberg, R., Rosenzweig, C., Civerolo, K., Ku, J. Y. and Kinney, P. L.: Assessing ozone-related health impacts under changing climate, Environ. Health Persp., 112, 1557–1563, 2004.
- Langner, L., Bergstrom, R., and Foltescu, V.: Impact of climate change on surface ozone and deposition of sulphur and nitrogen in Europe, Atmos. Environ., 39, 1129–1141, doi:10.1016/j.atmosenv.2004.09.082, 2005.
- Lefèvre, F., Brasseur, G. P., Folkins, I., Smith, A. K., and Simon, P.: Chemistry of the 1991–1992 stratospheric winter: three dimensional model simulations, J. Geophys. Res., 99, 8183–8195, 1994.
- Louis, J.-F.: Parametric model of vertical eddy fluxes in the atmosphere, Bound.-Lay. Meteorol., 17, 187–202, 1979.
- Martet, M., Peuch, V.-H., Laurent, B., Marticorena, B., and Bergametti, G.: Evaluation of long-range transport and deposition of desert dust with the CTM MOCAGE, Tellus, 61B, 449–463, 2009.
- Meleux, F., Solmon, F., and Giorgi, F.: Increase in summer European ozone amounts due to climate change, Atmos. Environ., 41, 7577–7587, 2007.
- Ménégoz, M., Voldoire, A., Teyssèdre, H., Salas Y Mélia, D., Peuch, V.-H., and Gouttevin, I.: How does the atmospheric variability drive the aerosol residence time in the Arctic region?, Tellus, 64, 11596, doi:10.3402/tellusb.v64i0.11596, 2012.
- Pay, M., Piot, M., Jorba, O., Gassó, S., Gonçalves, M., Basart, S., Dabdub, D., Jiménez-Guerrero, P., and Baldasano, J. M.: A full year evaluation of the CALIOPE-EU air quality modeling system over Europe for 2004, Atmos. Environ., 44, 3322–3342, 2010.
- Peuch, V.-H., Amodei, M., Barthet, T., Cathala, M.-L., Michou, M., and Simon, P.: MOCAGE, MOdéle de Chimie Atmosphérique à Grande Echelle, in: Proceedings of Météo France: Workshop on atmospheric modelling, Toulouse, France, 33–36, 1999.
- Pouliot, G., Pierce, T., Denier van der Gon, H., Schapp, M., Moran, M., and Nopmongcol, U.: Comparing emission inventories and model-ready emission datasets between Europe and North America for the AQMEII project, Atmos. Environ., 53, 4– 14, 2012.
- Prather, M., Gauss, M., Berntsen, T., Isaksen, I., Sundet, J., Bey, I., Brasseur, G., Dentener, F., Derwent, R., Stevenson, D., Grenfell, L., Hauglustaine, D., Horowitz, L., Jacob, D., Mickley, L., Lawrence, M., von Kuhlmann, R., Muller, J.-F., Pitari, G., Rogers, H., Johnson, M., Pyle, J., Law, K., van Weele, M., and Wild, O.: Fresh air in the 21st century?, Geophys. Res. Lett., 30, 1100, doi:10.1029/2002GL016285, 2003.
- Schlink, U., Herbarth, O., Richter, M., Dorling, S., Nunnari, G., Cawley, G., and Pelikan, E.: Statistical models to assess the health effects and to forecast ground-level ozone, Environ. Modell Softw., 21, 547–558, doi:10.1016/j.envsoft.2004.12.002, 2006.
- Sillman, S.: Ozone production efficiency and loss of NO_x in power plant plumes: photochemical model and interpretation of measurements in Tennessee, J. Geophys. Res., 105, 9189–9202, doi:10.1029/1999JD901014, 2000.
- Solberg, S., Hov, O., Sovde, A., Isaksen, I., Coddeville, P., De Backer, H., Forster, C., Orsolini, Y., and Uhse, K.: European surface ozone in the extreme summer 2003, 113, D07307, doi:10.1029/2007JD009098, 2008.

- Stockwell, W. R., Kirchner, F., Khun, M., and Seefeld, S.: A new mechanism for regional atmospheric chemistry modelling, J. Geophys. Res., 102, 25847–25879, 1997.
- Struzewska, J. and Kaminski, J.: Formation and transport of photooxidants over Europe during the July 2006 heat wave – observations and GEM-AQ model simulations, Atmos. Chem. Phys., 8, 721–736, doi:10.5194/acp-8-721-2008, 2008.
- Szopa, S., Hauglustaine, D., Vautard, R., and Menut, L.: Future global tropospheric ozone changes and impact on European air quality, Geophys. Res. Lett., 115, L14805, doi:10.1029/2006GL025860, 2006.
- Taylor, K.: Summarizing multiple aspects of model performance in a single diagram, J. Geophys. Res., 106, 7183–7192, 2001.
- US-EPA: Interim procedures for evaluating air quality models, Technical Report. EPA-450/4-91-013, Research Triangle Park, Environmental Protection Agency, Office of Air Quality Planning and Standards, NC, US, 1984.
- US-EPA: Guideline for regulatory application of the urban airshed model, Technical Report, EPA-450/4-91-013, Research Triangle Park, Environmental Protection Agency, Office of Air Quality Planning and Standards, NC, US, 1991.
- van Loon, M., Roemer, M., Builtjes, P., Bessagnet, B., Rouïl, L., Christensen, J., Brandt, J., Fagerli, H., Aasón, L., and Rodgers, I.: Model Inter-comparison in the Framework of the Review of the Unified EMEP Model, TNO Report, Technical Report R2004/282, 53 pp., 2004.
- van Loon, M., Vautard, R., Schaap, M., Bergstrom, R., Bessagnet, B., Brandt, J., Builtjes, P., Christensen, J., Cuvelier, C., Graff, A., Jonson, J., Krol, M., Langner, J., Roberts, P., Rouïl, L., Stern, R., Tarrasòn, L., Thunis, P., Vignati, E., White, L., and Wind, P.: Evaluation of long-term ozone simulations from seven regional air quality models and their ensemble, Atmos. Environ., 41, 2083–2097, doi:10.1016/j.atmosenv.2006.10.073, 2007.
- Vautard, R., Honore, C., Beekmann, M., and Rouïl, L.: Simulation of ozone during the August 2003 heat wave and emission control scenarios, Atmos. Environ., 39, 2957–2967, doi:10.1016/j.atmosenv.2005.01.039, 2005.
- Vautard, R., Builtjes, P., Thunis, P., Cuvelier, K., Bedogni, M., Bessagnet, B., Honoré, C., Moussiopoulos, N., Schaap, M., Stern, R., Tarrason, L., and van Loon, M.: Evaluation and intercomparison of ozone and PM₁₀ simulations by several chemistry-transport models over 4 European cities within the city-delta project, Atmos. Environ., 41, 173–188, 2007.
- Visschedijk, A. H. J. and Denier van der Gon, H. A.: Gridded European anthropogenic emission data for NO_x , SO_2 , NMVOC, NH₃, CO, PM₁₀, PM_{2.5} and CH₄ for the year 2000, TNO B O-A Rapport 2005/106, 2005.
- WHO: Health aspects of air pollution results from the WHO project "Systematic review of health aspects of air pollution in Europe", Tech. rep., 2004.
- Williamson, D. L. and Rasch, P. J.: Two-dimensional semilagrangian transport with shape-preserving interpolation, Mon. Weather Rev., 117, 102–129, 1989.
- Zlatev, Z.: Comprehensive air pollution studies by the Danish Eulerian Model, in: Air, Water and Soil Quality Modelling for Risk and Impact Assessment, edited by: Ebel, A. and Davitashvili, T., Nato Science for Peace and Security, Environmental Security, 293–302, 2007.