



## Evaluation of European air quality modelled by CAMx including the volatility basis set scheme

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**Abstract.** Four periods of EMEP (European Monitoring and Evaluation Programme) intensive measurement campaigns (June 2006, January 2007, September–October 2008 and February–March 2009) were modelled using the regional air quality model CAMx with VBS (volatility basis set) approach for the first time in Europe within the framework of the EURODELTA-III model intercomparison exercise. More detailed analysis and sensitivity tests were performed for the period of February–March 2009 and June 2006 to investigate the uncertainties in emissions as well as to improve the modelling of organic aerosol (OA). Model performance for selected gas phase species and PM<sub>2.5</sub> was evaluated using the European air quality database AirBase. Sulfur dioxide (SO<sub>2</sub>) and ozone (O<sub>3</sub>) were found to be overestimated for all the four periods, with O<sub>3</sub> having the largest mean bias during June 2006 and January–February 2007 periods (8.9 ppb and 12.3 ppb mean biases respectively). In contrast, nitro-

gen dioxide (NO<sub>2</sub>) and carbon monoxide (CO) were found to be underestimated for all the four periods. CAMx reproduced both total concentrations and monthly variations of PM<sub>2.5</sub> for all the four periods with average biases ranging from  $-2.1$  to  $1.0 \mu\text{g m}^{-3}$ . Comparisons with AMS (aerosol mass spectrometer) measurements at different sites in Europe during February–March 2009 showed that in general the model overpredicts the inorganic aerosol fraction and underpredicts the organic one, such that the good agreement for PM<sub>2.5</sub> is partly due to compensation of errors. The effect of the choice of VBS scheme on OA was investigated as well. Two sensitivity tests with volatility distributions based on previous chamber and ambient measurements data were performed. For February–March 2009 the chamber case reduced the total OA concentrations by about 42 % on average. In contrast, a test based on ambient measurement data increased OA concentrations by about 42 % for the same

period bringing model and observations into better agreement. Comparison with the AMS data at the rural Swiss site Payerne in June 2006 shows no significant improvement in modelled OA concentration. Further sensitivity tests with increased biogenic and anthropogenic emissions suggest that OA in Payerne was affected by changes in emissions from residential heating during the February–March 2009 whereas it was more sensitive to biogenic precursors in June 2006.

## 1 Introduction

Air pollution is known to cause damage to human health, vegetation and ecosystems. It is one of the main environmental causes of premature death. Only in Europe, more than 400 000 premature deaths were estimated in 2012, with PM<sub>2.5</sub> (particles less than 2.5 µm in aerodynamic diameter) having the highest relative risk for health damage (WHO, 2014). Air quality models help to understand the processes taking place between emission sources and pollutant concentrations at receptor sites. They are very useful for defining control strategies for future legislation. In spite of large improvements in recent years, chemical transport models (CTMs) still have some uncertainties (Solazzo et al., 2012). Various air quality model intercomparison exercises were successfully carried out over the last decades to determine uncertainties in chemical and physical processes governing particulate matter and its precursors (Solazzo et al., 2012; Bessagnet et al., 2014). However, a large variability in particulate matter concentrations was found between different models, indicating process parameterization as one of the main reasons for such discrepancies. Moreover, recent studies based on AMS (aerosol mass spectrometer) measurements at different sites in Europe revealed that the organic fraction dominates the non-refractory PM<sub>1</sub> composition (Crippa et al., 2014). Organic aerosol (OA) can be found in the atmosphere from direct emission by various sources, such as fossil fuel combustion by road vehicle engines or residential wood combustion. Direct emissions of OA are typically referred to as primary organic aerosol (POA) whereas gas-to-particle conversion is referred to as secondary organic aerosol (SOA). Formation mechanisms of SOAs are not very well known yet and their representation in CTMs is still challenging (Hallquist et al., 2009; Fountoukis et al., 2011; Bergström et al., 2012; Li et al., 2013; Langmann et al., 2014; Tsigaridis et al., 2014). In one of our recent aerosol modelling studies we compared model PM<sub>2.5</sub> prediction with PM<sub>1</sub> AMS measurements for different sites (Payerne and Zürich) and periods (summer and winter) in Switzerland. We found that particulate matter was generally well reproduced by the model with the SOA fraction being underpredicted and POA overpredicted (Aksoyoglu et al., 2011). Traditional CTMs treat POA as nonvolatile. Some studies, however, have revealed the semivolatile nature of POA, through its dynamic

equilibrium of organic aerosol with its gas phase, and the importance of semivolatile (SVOC) and intermediate volatility (IVOC) organic compounds as SOA precursors (Donahue et al., 2006; Robinson et al., 2007; Cappa and Jimenez, 2010). To describe the absorptive partitioning and ongoing oxidation of the atmospheric material, a volatility basis set (VBS) where organic species are organized into surrogates according to their volatility was developed (Donahue et al., 2011, 2012a, b). Air quality models updated with the VBS scheme started being used (Lane et al., 2008; Murphy and Pandis, 2009; Hodzic et al., 2010; Fountoukis et al., 2011, 2014; Bergström et al., 2012; Murphy et al., 2012; Jo et al., 2013; Zhang et al., 2013; Athanasopoulou et al., 2013). Bergström et al. (2012) reported an EMEP model study over Europe for the 2002–2007 period using different assumptions regarding partitioning and aging processes. They could not reproduce the measured OA levels in winter, suggesting that residential wood combustion inventories might be underestimated in different parts of Europe. Fountoukis et al. (2014) applied the PMCAMx model to simulate EUCAARI (Kulmala et al., 2009, 2011) and EMEP (Tørseth et al., 2012) campaigns in Europe. They could reproduce most of PM<sub>1</sub> daily average OA observations within a factor of 2, with the February–March 2009 period having the largest discrepancies. Zhang et al. (2013) deployed the CHIMERE model with the VBS framework during the MEGAPOLI summer campaign in the Greater Paris region for July 2009. They found a considerable improvement in predicted SOA concentrations which might be even overestimated depending on the emission inventory used. In our study, we applied the regional air quality model CAMx with the VBS scheme for the first time in Europe within the framework of EURODELTA-III (EDIII) model intercomparison exercise. In addition to the base case configuration used in the exercise, more sensitivity tests with the VBS scheme for winter and summer episodes were performed together with a general evaluation of the four EMEP field measurement campaigns.

## 2 Method

### 2.1 The EURODELTA-III exercise

The EDIII framework is a European model intercomparison exercise between several modelling teams sharing both efforts and technical knowledge in order to reduce model uncertainties and to improve understanding of the performances. It contributes to the scientific work of the United Nations Economic Commission for Europe (UNECE) Task Force on Measurement and Modelling (TFMM) within the Convention on Long-range Transboundary Air Pollution (CLRTAP). In the first phase of the EDIII exercise, four periods of the EMEP field measurement campaigns were chosen in order to evaluate the model results:

– 1 June–30 June 2006

- 8 January–4 February 2007
- 17 September–15 October 2008
- 25 February–26 March 2009.

Multiple models were applied on a common domain and driven with the same input data provided by the National Institute for Industrial Environment and Risks (INERIS). However, for some models, different meteorology, boundary conditions and emissions data such as biogenic emissions were used (Bessagnet et al., 2014).

## 2.2 Modelling method

### 2.2.1 CAMx

The Comprehensive Air quality Model with extensions, CAMx-VBS (CAMx5.41\_VBS, kindly provided by ENVIRON before its public release), was used in this study. The model domain consisted of one grid with a horizontal resolution of  $0.25^\circ \times 0.25^\circ$ . The latitude and longitude grids extended from  $25.125^\circ$  W to  $45.125^\circ$  E and  $29.875$  to  $70.125^\circ$  N, resulting in  $281 \times 161$  grid cells covering the whole of Europe. Hourly four-dimensional meteorological fields for wind speed and direction, pressure, temperature, specific humidity, cloud cover and rain required by CAMx simulations were calculated from ECMWF IFS (Integrated Forecast System) data at  $0.2^\circ$  resolution. Vertical diffusivity coefficients were estimated following the Kz approach of O'Brien (1970) using planetary boundary layer depth profiles as available in IFS data. CAMx simulations used 33 terrain-following  $\sigma$ -levels up to about 8000 m above ground level, as in the original IFS data. The lowest layer was about 20 m thick. MACC (Monitoring Atmospheric Composition and Climate) reanalysis data were used to initialize initial and the boundary condition fields (Benedetti et al., 2009; Inness et al., 2013). Elemental carbon, organic aerosol, dust and sulfate were used to model aerosol species at the boundaries of the domain. One half of the OA was assumed to be SOA and the other half POA, as recommended in the EDIII exercise. Photolysis rate inputs were calculated using the Tropospheric Ultraviolet and Visible (TUV) radiative transfer and photolysis model (<https://www2.acom.ucar.edu/modeling/tropospheric-ultraviolet-and-visible-tuv-radiation-model>). The required ozone column densities to determine the spatial and temporal variation of the photolysis rates were extracted from TOMS data (NASA/GSFC, 2005). Removal processes as dry and wet deposition were simulated using the Zhang resistance model (Zhang et al., 2003) and a scavenging model approach for both gases and aerosols (Environ, 2011), respectively. For the gas phase chemistry the carbon bond (CB05) mechanism (Yarwood et al., 2005) with 156 reactions and up to 89 species was used. Partitioning of inorganic aerosols (sulfate, nitrate, ammonium, sodium and chloride) was performed using the ISORROPIA thermodynamic

model (Nenes et al., 1998). Aqueous sulfate and nitrate formation in cloud water was simulated as well, using the RADM aqueous chemistry algorithm (Chang et al., 1987).

### 2.2.2 Emissions

#### Anthropogenic emissions

Annual total gridded anthropogenic emissions were prepared and provided by INERIS for the EDIII exercise, which is based on a merging process of databases from different sources, i.e. TNO-MACC (Kuenen et al., 2011), EMEP (Vestreng et al., 2007) and GAINS (The Greenhouse Gas and Air Pollution Interactions and Synergies). For specific countries where TNO-MACC emissions were missing (Iceland, Liechtenstein, Malta and Asian countries), the EMEP  $0.5^\circ \times 0.5^\circ$  emissions were used and re-gridded using adequate proxies such as “artificial land-use” and EPER (European Pollutant Emission Register) data (<http://www.eea.europa.eu/>) for industries. Total primary particle emissions were made available by EMEP in two different size ranges: below  $2.5 \mu\text{m}$  (fine) and between  $2.5 \mu\text{m}$  and  $10 \mu\text{m}$  (coarse). Total emissions were later split to estimate the amount of elemental carbon and organic matter for each of the 10 SNAP codes (Selected Nomenclature for Air Pollution) and country. The final emission inventory thus compiled consisted of six gas species, namely methane, carbon monoxide, ammonia, sulfur oxides, non-methane volatile organic compounds and nitrogen oxides, and six categories of particulate matter classes: fine elemental carbon (EC2.5), coarse elemental carbon (EC10), fine POA, coarse POA, fine other primary particulate material (non-carbonaceous) and coarse other primary particulate material (non-carbonaceous).  $\text{PM}_{2.5}$  and  $\text{PM}_{10}$  emissions were provided by EMEP and they were split to elemental carbon and organic matter using the fractions given by IIASA (International Institute for Applied Systems Analysis) for each source and country. Total non-methane volatile organic compounds were split for the CB05 mechanism using the recommendations of Passant (2002). Hourly, weekly and monthly time profiles were applied to total annual anthropogenic emissions.

#### Biogenic emissions

Biogenic volatile organic compound (BVOC) emissions were calculated using the Model of Emissions of Gases and Aerosols from Nature (MEGANv2.1; Guenther et al., 2012). This model is driven by meteorological variables such as hourly temperature, solar radiation, humidity, wind speed, soil moisture and land cover data, including leaf area index and plant function type, as available in the Community Land Model 4.0. Eight-day average satellite data at  $0.25^\circ \times 0.25^\circ$  resolution were pre-processed and made available from the TERRA/MODIS satellite system. Sixteen plant function types including needle-leaved evergreen, needle-

leaved deciduous, broad-leaved evergreen, broad-leaved deciduous, grass and crop for different climatic zones were prepared for this study at  $0.25^\circ \times 0.25^\circ$  resolution together with the global emission factors of  $\alpha$ -pinene,  $\beta$ -pinene, 3-carene, isoprene, limonene, 232-methylbutenol, myrcene,  $\text{NO}_x$ , *t*- $\beta$ -ocimene and sabinene. Common BVOC species such as isoprene, terpene, sesquiterpene, xylene and toluene were obtained for each hour and cell in the domain.

### 2.2.3 VBS scheme

A new VBS scheme is available in the CAMx model to describe changes in oxidation state and volatility. A total of four basis sets simulate the evolution of organic aerosol in the atmosphere (Koo et al., 2014). POA emissions were split in hydrocarbon-like organic aerosol (HOA) and biomass burning organic aerosol (BBOA) emissions and allocated in two different basis sets. HOA-like emissions include emissions from all SNAP sectors except SNAP2 (nonindustrial combustion plants) and SNAP10 (agriculture), which were assigned to BBOA-like emissions. Two other sets were used in the model to allocate secondary organic aerosol from anthropogenic (ASOA) (i.e. xylene and toluene) and biogenic (BSOA) (i.e. isoprene, monoterpene and sesquiterpene) gaseous precursors. These two sets also allocate oxidation products of POA vapours from each of the two primary sets (HOA-like and BBOA-like). The 2-D volatility space retrieved by Donahue et al. (2011, 2012a, b) was used to distribute the organic molecular structures for each of the volatility bins and different sets (Table S1 in the Supplement). Five volatility bins represent the range of SVOCs ranging from  $10^{-1} \mu\text{g m}^{-3}$  to  $10^3 \mu\text{g m}^{-3}$  in saturation concentrations ( $C^*$ ). Oxidation processes are modelled by shifting  $C^*$  by a factor of 10 in the next lower volatility bin, increasing the oxidation state and reducing the carbon number to account for fragmentation. OH reaction rates are assumed to be  $4 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$  for the reaction of semivolatile primary vapours with OH and  $2 \times 10^{-11}$  for further aging of ASOA and POA vapours from HOA-like emissions. More details about the VBS parameterization in CAMx can be found in Koo et al. (2014). Further aging of BSOA is not considered in this study, based on previous modelling results showing overprediction of OA when such process is taken into account (Lane et al., 2008; Murphy and Pandis, 2009). This implies that also further aging of POA vapours from BBOA-like emissions was not considered since it is performed in the same basis set. In this work we focus on the effects of a VBS framework on the total OA fraction. Aging processes and alternative VBS implementations will be discussed together with SOA and POA components in a following paper (Ciarelli et al., 2016). Three sensitivity tests were performed with different assumptions on the volatility distributions (Table 1).

- NOVBS: primary organic aerosol was assumed to be nonvolatile. Biogenic (isoprene, monoterpenes and

sesquiterpenes) and anthropogenic (xylene, toluene and other aromatics) volatile organic compounds were used as precursors for secondary organic aerosol. Partitioning of condensable gases to secondary organic aerosol was calculated using a semivolatile equilibrium approach (Strader, 1999).

- VBS\_ROB: primary organic aerosol was assumed to be volatile and undergo chemical oxidation. The volatility distribution estimated by Robinson et al. (2007) was applied to HOA-like and BBOA-like emissions. Emissions of IVOCs were assumed to be 1.5 times those of POA as suggested by Robinson et al. (2007).
- VBS\_BC: primary organic aerosol was assumed to be volatile and undergo chemical oxidation using the approach of Shrivastava et al. (2011) and Tsimpidi et al. (2010). The total primary emissions are roughly 3 times higher than in VBS\_ROB. Different volatility distributions were applied for HOA- and BBOA-like emissions. IVOCs were assumed to be 1.5 times the amount of POA. This implies that for this scenario the SVOC + IVOC mass added is equal to 7.5 times the initial amount of POA. This represents the base case scenario used to evaluate gas phase and  $\text{PM}_{2.5}$  model performance.

Based on the VBS\_BC base case scenario, two other sensitivity tests were performed with respect to emissions:

- VBS\_BC\_2xBVOC increased BVOCs emissions by a factor of 2
- VBS\_BC\_2xBBOA increased BBOA-like emissions by a factor of 2.

### 2.3 Statistical methods

Statistical procedures as available in the Atmospheric Model Evaluation Tool (AMET; Appel et al., 2010) were used in this study to evaluate model performance. Daily ambient measurements of main gas phase species, i.e.  $\text{O}_3$ ,  $\text{NO}_2$ , CO,  $\text{SO}_2$  and fine particulate matter ( $\text{PM}_{2.5}$ ), were extracted from the AirBase database in Europe and statistics reported in terms of mean bias (MB), mean error (ME), mean fractional bias (MFB) mean fractional error (MFE) and correlation coefficient ( $r$ ).

Due to the coarse grid resolution, only rural-background stations, defined as stations far from city sources of air pollution with pollution levels determined by the integrated contribution from all sources upwind of the station (ETC/ACC, 2004/7), with at least 80 % daily average observations available were considered for the statistical analysis. For  $\text{PM}_{2.5}$  this resulted in 48 stations available for June 2006, 56 for January–February 2007, 90 for September–October 2008 and 110 stations for February–March 2009.  $\text{PM}_{2.5}$  components were further evaluated for the February–March 2009

**Table 1.** Volatility distributions used for different scenarios.

Scenarios	POA emission sources	Emission fraction for volatility bin with $C^*$ of				
		0 <sup>a</sup>	1	10	100	1000
NOVBS (nonvolatile CAMxv5.40)	HOA-like	1.00	–	–	–	–
	BBOA-like	1.00				
VBS_ROB (Robinson et al., 2007)	HOA-like	0.09	0.09	0.14	0.18	0.5
	BBOA-like	0.09	0.09	0.14	0.18	0.5
VBS_BC (Tsimpidi et al., 2010; Shrivastava et al., 2011)	HOA-like	0.40	0.26	0.40	0.51	1.43
	BBOA-like	0.27	0.27	0.42	0.54	1.50

<sup>a</sup> Properties of the lowest volatility bins refer to all OA with  $C^* \leq 0.1 \mu\text{g m}^{-3}$  (non-volatile OA).

period where comprehensive high-resolution AMS measurements at 11 European sites were available, i.e. at Barcelona, Cabauw, Chilbolton, Helsinki, Hyytiälä, Mace Head, Melnitz, Montseny, Payerne, Puy de Dôme and Vavihill (Crippa et al., 2014).

### 3 Results and discussions

#### 3.1 Model evaluation

Model performance metrics for gas phase species CO, NO<sub>2</sub>, O<sub>3</sub> and SO<sub>2</sub> as well as for PM<sub>2.5</sub> are reported in Table 2 and they refer to the base case VBS\_BC.

##### 3.1.1 NO<sub>2</sub> and O<sub>3</sub>

NO<sub>2</sub> was found to be underpredicted for all the four periods with mean fractional bias between  $-54$  and  $-28\%$  and NO<sub>2</sub> concentrations being particularly underpredicted during June 2006. Evaluation of the EDIII model intercomparison exercise showed that all models performed similarly for NO<sub>2</sub> in terms of correlation with  $r$  values in the range 0.6–0.7 and the spatial correlation was much higher in the range 0.7–0.9 for all models (Bessagnet et al., 2016) with a general underestimation in the afternoon. The NO<sub>2</sub> performance could be influenced by several factors.

- Uncertainties in the emission inventories. Although NO<sub>x</sub> emission estimates in Europe are thought to have an uncertainty of about  $\pm 20\%$ , the complete dataset used in the inventories has much higher uncertainty (Kuenen et al., 2014). A recent study identified a significant discrepancy between emission estimates and actual flux measurements, with the highest underestimation being a factor of two in central London mainly due to underrepresentation of real world road traffic emissions (Vaughan et al., 2016)

- The relatively coarse resolution of the domain may result in too low NO<sub>x</sub> emissions or isolated local events that the model cannot resolve. We report daily average time series of NO<sub>2</sub> for the period of February–March 2009 for stations in Table 2 as well as daily average time series of NO<sub>2</sub> for stations not exceeding 5 ppb (which represents 92% of the stations in Table 2) (Fig. S1 in the Supplement). The model performance for NO<sub>2</sub> significantly improved when the 5 ppb threshold was applied to the dataset. An emission map of NO for 1 March 2009 at 06:00 is reported in Fig. S2. High emissions of NO are predicted in the Benelux area, Po Valley, Germany and in some of the eastern European countries. High NO emissions due to ship traffic are also visible, especially in the Mediterranean Sea.
- Possible positive artefacts in the chemiluminescence methods for measuring NO<sub>2</sub> may also occur when NO<sub>2</sub> is catalytically converted to NO on the molybdenum surface, leading to an overprediction of measured NO<sub>2</sub> concentrations (Steinbacher et al., 2007; Villena et al., 2012)
- Moreover, an evaluation of planetary boundary layer height (PBLH) within the EDIII shows that although the PBLH was quite well represented in general in the ECMWF IFS meteorological fields, CAMx tends to underestimate the night-time minima and to overestimate some daytime peaks, overpredicting the dilution of daytime NO<sub>2</sub> concentrations, whereas the wind speed was relatively well reproduced (Bessagnet et al., 2016).

O<sub>3</sub> concentrations were found to be overpredicted for all the four periods with a mean fractional bias ranging from 2 to 48%. Especially in June 2006, when the photochemical activity is higher, the general underprediction of NO<sub>x</sub> in the whole domain reduces the O<sub>3</sub> titration potential during night-time.

Model performance for O<sub>3</sub> is also strongly influenced by long-range transport, especially during the winter periods

**Table 2.** Model gas phase and PM<sub>2.5</sub> performance for the EDIII field campaigns (base case VBS\_BC).

Species	Number of sites	Observed mean (ppb) ( $\mu\text{g m}^{-3}$ for PM <sub>2.5</sub> )	Modelled mean (ppb) ( $\mu\text{g m}^{-3}$ for PM <sub>2.5</sub> )	MB (ppb) ( $\mu\text{g m}^{-3}$ for PM <sub>2.5</sub> )	ME (ppb) ( $\mu\text{g m}^{-3}$ for PM <sub>2.5</sub> )	MFB [-]	MFE [-]	<i>r</i>	
June 2006									
CO	36	192.0	158.0	-34.2	80.7	-0.12	0.36	0.20	
NO <sub>2</sub>	320	4.1	2.3	-1.9	2.2	-0.54	0.68	0.55	
O <sub>3</sub>	460	42.3	51.2	8.9	10.8	0.21	0.24	0.57	
PM <sub>2.5</sub>	48	12.0	11.7	-0.3	4.5	-0.07	0.39	0.55	
SO <sub>2</sub>	263	1.0	1.2	0.2	0.7	0.14	0.67	0.52	
January–February 2007									
CO	45	248.0	191.0	-57.8	107.0	-0.11	0.37	0.21	
NO <sub>2</sub>	337	6.5	4.4	-2.2	3.2	-0.28	0.57	0.68	
O <sub>3</sub>	455	23.5	35.8	12.3	12.6	0.48	0.49	0.61	
PM <sub>2.5</sub>	56	11.7	12.8	1.0	6.1	-0.04	0.56	0.69	
SO <sub>2</sub>	271	1.3	1.7	0.4	1.1	0.36	0.75	0.46	
September–October 2008									
CO	53	208.0	136.0	-72.0	91.4	-0.31	0.48	0.27	
NO <sub>2</sub>	370	5.3	3.7	-1.7	2.5	-0.28	0.56	0.62	
O <sub>3</sub>	465	24.3	32.5	8.2	9.6	0.32	0.37	0.50	
PM <sub>2.5</sub>	90	13.0	14.1	1.0	5.7	< 0.01	0.46	0.76	
SO <sub>2</sub>	256	0.9	1.1	0.2	0.8	0.25	0.74	0.37	
February–March 2009									
CO	57	262.0	170.0	-91.6	119.0	-0.26	0.48	0.37	
NO <sub>2</sub>	380	6.0	3.9	-2.0	2.8	-0.33	0.56	0.61	
O <sub>3</sub>	488	32.7	33.0	0.2	7.1	0.02	0.23	0.55	
PM <sub>2.5</sub>	110	15.1	13.0	-2.1	6.4	-0.13	0.50	0.71	
SO <sub>2</sub>	257	1.0	1.3	0.3	0.9	0.23	0.76	0.45	

when the local chemical production of O<sub>3</sub> is limited. Figure S3 shows the model performance at the Mace Head station located on the western coast of Ireland for all the four periods. Especially in January–February 2007 O<sub>3</sub> concentrations were found to be overpredicted by about 10 to 20 ppb, indicating that boundary conditions for O<sub>3</sub> were probably not well represented. In June 2006 and September–October 2008 O<sub>3</sub> was relatively well captured at Mace Head, suggesting that the observed positive bias in O<sub>3</sub> concentrations might arise from insufficient NO<sub>x</sub> emissions to undergo titration during night-time as well as not correctly represented planetary boundary layer dynamics. In February–March 2009 the model tends to underpredict the O<sub>3</sub> concentration at Mace Head and overall the O<sub>3</sub> model performance shows the lowest bias (2 %). Eventually, the underprediction of O<sub>3</sub> in the boundary condition may counteract the already mentioned deficiencies related to insufficient NO<sub>x</sub> emissions.

### 3.1.2 SO<sub>2</sub> and CO

SO<sub>2</sub> concentrations were found to be slightly overpredicted for all the four periods, with a mean fractional bias ranging from 14 to 36 % for SO<sub>2</sub>. The daily variations of modelled and measured SO<sub>2</sub> concentrations for February–March 2009 are reported as well in Fig. S1 (lower panel) for the stations

in Table 2. In general, the daily variations of modelled and measured SO<sub>2</sub> concentrations agree relatively well with each other throughout the period.

Most of the SO<sub>2</sub> emissions arise from high stack point sources which have injection heights of a few hundred metres. It might be that the vertical distribution of SO<sub>2</sub> might affect the model performance in particular near the harbours and coastal areas where ship emissions were allocated in the second layer of the model domain (extending from ~20 to 50 m above ground level), whereas they can reach up to 58 m in deep draft vessels (Starcrest Consulting Group, 2004) and also undergo plume rise. Insufficient conversion to sulfate or too low deposition processes might also positively bias the model performance for SO<sub>2</sub>.

CO was slightly underpredicted for all periods (mean fractional bias between -11 and -31 %), with highest values during the September–October 2008 period (-31 %). The late summer–autumn period is known to be influenced by agricultural open field burning activities which might be missing from standard emission inventories.

In general, for both SO<sub>2</sub> and CO, the model showed lower correlation coefficients with respect to other gas phase species (*r* values from 0.20 and 0.37 for CO and from 0.37 to 0.52 for SO<sub>2</sub>).

### 3.1.3 PM<sub>2.5</sub>

Of all investigated variables, CAMx shows the best statistical performance for PM<sub>2.5</sub>. For all four periods the acceptable model performance criteria recommended by Boylan and Russell (2006) for aerosols were met ( $MFE \leq +75\%$  and  $-60\% < MFB < +60\%$ ). The fractional bias ranges from less than 1% in September–October 2008 up to  $-13\%$  in February–March 2009. Also the recommended model performance goals ( $MFE \leq +50\%$  and  $-30\% < MFB < +30\%$ ) were met for all periods except for January 2007. Modelled average PM<sub>2.5</sub> concentrations are shown in Fig. 1. A different spatial distribution is seen for summer and winter. In June 2006 the model predicts higher concentrations in the southern part of the domain, especially over the Mediterranean Sea and North Africa (up to  $35 \mu\text{g m}^{-3}$ ). In contrast, the highest concentrations were predicted in the Po Valley area (above  $40 \mu\text{g m}^{-3}$ ) and in the southern part of Poland during January–February 2007. During the two colder periods (2007 and 2009) elevated concentrations of around  $15 \mu\text{g m}^{-3}$  are also visible close to urban areas such as Paris and Moscow. Figure 2 shows PM<sub>2.5</sub> variations at AirBase rural-background sites in terms of medians, 25th and 75th percentiles. In all the four periods CAMx is able to reproduce the observed monthly variation very well with some overprediction occurring mainly from the 14 to the 17 of January 2007 and towards the end of the 2008 period.

### 3.2 Detailed evaluation of PM<sub>2.5</sub> components in February–March 2009

The modelled concentrations of non-refractory PM<sub>2.5</sub> components were compared against aerosol mass spectrometer measurements at 11 European sites for the February–March 2009 period (Crippa et al., 2014). Even though the AMS measures particles with a diameter  $D < 1 \mu\text{m}$ , the difference between the non-refractory PM<sub>1</sub> and total PM<sub>2.5</sub> mass is in general rather small as shown in Aksoyoglu et al. (2011), at least for situations without exceedingly high air pollution and situations when sea salt makes a large relative contribution to PM<sub>2.5</sub>. The modelled average total non-refractory PM<sub>2.5</sub> (sum of nitrate, sulfate, ammonium and OA) concentrations match the measurements quite well with a few exceptions (Fig. 3 and Table 3). The model is able to reproduce both high concentrations observed at the urban site Barcelona and low ones at remote sites like Hyytiälä, Finland. Concentrations of inorganic aerosols are overpredicted and OA are underpredicted at most of the stations (with similar behaviour during the other investigated periods; Figs. S4 and S5). Very similar results were also presented by other recent studies (Knote et al., 2011). The effect of different schemes to treat OA is discussed in Sect. 3.3. At the Cabauw site nitrate was the most dominant species (Mensah et al., 2012). Especially at this site the model strongly overpredicts, in particular the nitrate ( $\text{NO}_3^-$ ) fraction (by a factor of 3). A

sensitivity test with 50% reduction in ammonia emissions significantly improved the modelled  $\text{NO}_3^-$  concentrations at almost all sites (Table S2), suggesting potential uncertainties in  $\text{NH}_3$  emissions and their seasonal variability. Other potential reasons for the overprediction of  $\text{NO}_3^-$  could be related to uncertainties in the removal process of  $\text{HNO}_3$  as well as the dry deposition velocity of  $\text{NH}_3$ . Substantial overpredictions were found at the higher-altitude site of Montseny and Puy de Dôme when compared with first model layer concentrations ( $\sim 200$  and  $800 \text{ m a.s.l.}$ , respectively, at these sites). These sites located at about 720 and 1465 m a.s.l. are sometimes not within the PBLH during winter periods. At the Montseny site, the relatively coarse resolution of the model could also influence model performance since the site is located in a complex area about 50 km north-east of Barcelona (Pandolfi et al., 2014). Sulfate concentrations ( $\text{SO}_4^{2-}$ ) were overpredicted at almost all sites and especially at Mace Head, suggesting that long-range transport of  $\text{SO}_4^{2-}$  might be positively biased.

Modelled and observed hourly concentrations of  $\text{NO}_3^-$ ,  $\text{SO}_4^{2-}$ , ammonium ( $\text{NH}_4^+$ ) and OA at Payerne are reported in Fig. 4 for March 2009 together with meteorological parameters in Fig. S6. The model was able to reproduce the meteorological parameters very well for most of the time. The temperature was slightly underpredicted at both night- and daytime (with a maximum of  $-2^\circ\text{C}$ ), whereas both the monthly variation and the absolute values of wind speed and specific humidity were reproduced well with a few underpredictions of high wind speed (6 and 11 March and towards the end of the simulation). The model was able to capture the three  $\text{NO}_3^-$  and  $\text{NH}_4^+$  peaks observed around the 7, 18 and 23 March with a general slight overprediction throughout the whole period. Indeed, the underprediction in temperature during day- and night-time could partially explain the overprediction of the  $\text{NO}_3^-$  fraction with more  $\text{NO}_3^-$  partitioning to the aerosol phase, which also applies to the other stations used in this study. An evaluation of modelled temperature at the European scale for the February–March 2009 period confirmed that the model systematically underpredicted the 2 m surface temperature (Bessagnet et al., 2014). All the inorganic components were overpredicted during the first 4 days of March 2009 with a peak around 3 March, indicating that the PBLH was probably not correctly reproduced by the model during this period. Although the temporal variation was captured, concentrations of OA were underpredicted throughout all the simulation ( $4.1$  and  $1.8 \mu\text{g m}^{-3}$  observed and modelled average concentrations). Analysis of the OA fraction is discussed in the next section.

### 3.3 Organic aerosols

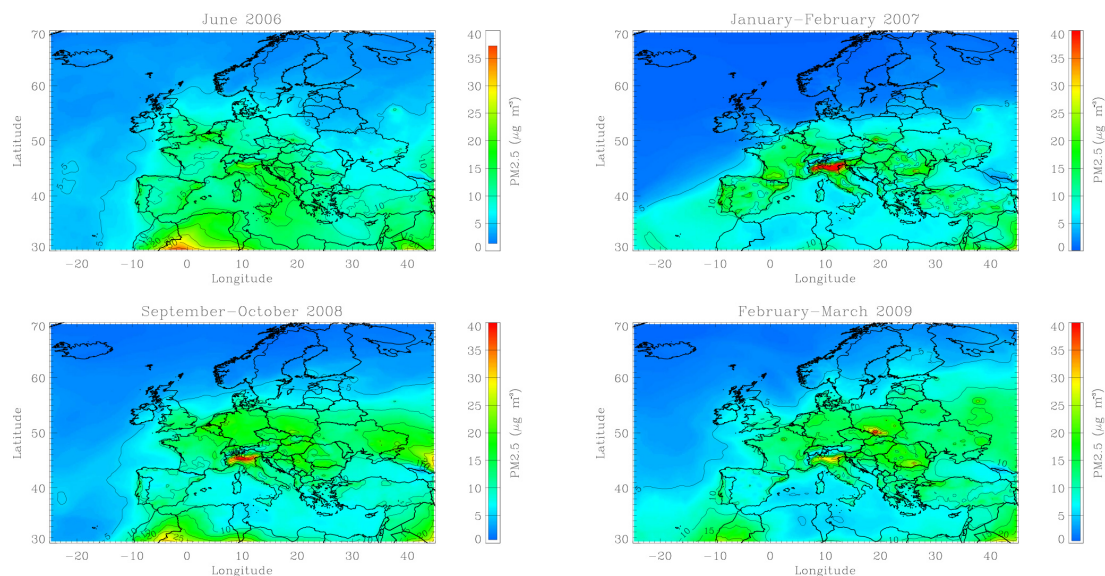
#### 3.3.1 Sensitivity of OA to the VBS scheme

In this section, effects of different parameterizations of the organic aerosol module on the modelled OA concentrations

**Table 3.** Statistical analysis of nitrate, ammonium, sulfate and organic aerosol for the base case (VBS\_BC) for February–March 2009 at different AMS sites.

Site	Mean observed ( $\mu\text{g m}^{-3}$ )	Mean modelled ( $\mu\text{g m}^{-3}$ )	MB $\mu\text{g m}^{-3}$	ME $\mu\text{g m}^{-3}$	MFB [-]	MFE [-]
$\text{NO}_3^-$						
Barcelona	3.6	5.8	2.2	4.0	0.35	0.98
Cabauw	2.2	6.7	4.5	4.6	0.85	1.01
Chilbolton	2.7	4.0	1.3	2.2	0.02	0.76
Helsinki	1.0	1.9	0.9	1.3	0.29	0.92
Hyytiälä	0.2	1.0	0.8	0.8	0.21	1.09
Mace Head	0.6	1.7	1.1	1.1	0.14	0.70
Melpitz	3.1	4.3	1.3	2.4	0.35	0.71
Montseny	3.1	5.9	2.8	4.3	0.38	1.00
Payerne	3.9	5.7	1.8	2.8	0.34	0.61
Puy de Dôme	0.9	2.7	1.8	2.2	1.13	1.30
Vavihill	2.8	3.7	0.9	2.2	0.14	0.78
$\text{NH}_4^+$						
Barcelona	1.6	2.5	0.9	1.4	0.42	0.71
Cabauw	1.0	2.7	1.7	1.8	0.95	0.97
Chilbolton	1.3	2.0	0.7	1.0	0.39	0.61
Helsinki	0.8	1.3	0.5	0.6	0.51	0.60
Hyytiälä	0.4	0.8	0.4	0.5	0.55	0.70
Melpitz	1.4	2.1	0.7	1.1	0.45	0.69
Montseny	1.7	2.6	0.9	1.6	0.39	0.74
Payerne	1.7	2.5	0.8	1.2	0.36	0.56
Puy de Dôme	0.7	1.2	0.5	0.9	0.83	1.07
Vavihill	1.6	1.9	0.4	0.9	0.17	0.56
$\text{SO}_4^{2-}$						
Barcelona	2.7	2.3	-0.4	1.3	-0.19	0.48
Cabauw	1.0	2.1	1.1	1.3	0.73	0.85
Chilbolton	1.3	2.2	0.9	1.3	0.45	0.70
Helsinki	2.4	2.2	-0.2	0.9	-0.04	0.43
Hyytiälä	1.4	1.7	0.3	0.7	0.09	0.58
Mace Head	0.4	1.2	0.8	0.9	1.04	1.12
Melpitz	1.1	2.2	1.1	1.4	0.54	0.76
Montseny	1.4	2.3	1.0	1.2	0.55	0.64
Payerne	1.1	2.1	1.1	1.2	0.62	0.70
Puy de Dôme	0.4	1.1	0.8	0.8	1.14	1.19
Vavihill	1.6	2.3	0.7	1.1	0.18	0.54
OA						
Barcelona	8.2	3.1	-5.1	5.2	-0.80	0.82
Cabauw	1.2	1.1	-0.1	0.5	-0.13	0.50
Chilbolton	2.4	0.7	-1.7	1.7	-1.09	1.10
Helsinki	2.7	2.9	0.3	1.6	0.08	0.62
Hyytiälä	1.3	1.0	-0.3	0.5	-0.48	0.60
Mace Head	0.8	0.4	-0.4	0.4	-0.29	0.70
Melpitz	1.5	0.5	-1.0	1.0	-0.94	0.97
Montseny	3.1	3.9	0.9	1.9	0.31	0.57
Payerne	4.1	1.8	-2.3	2.4	-0.85	0.90
Puy de Dôme	0.6	1.4	0.8	1.0	0.68	0.91
Vavihill	3.9	1.4	-2.5	2.5	-1.04	1.04





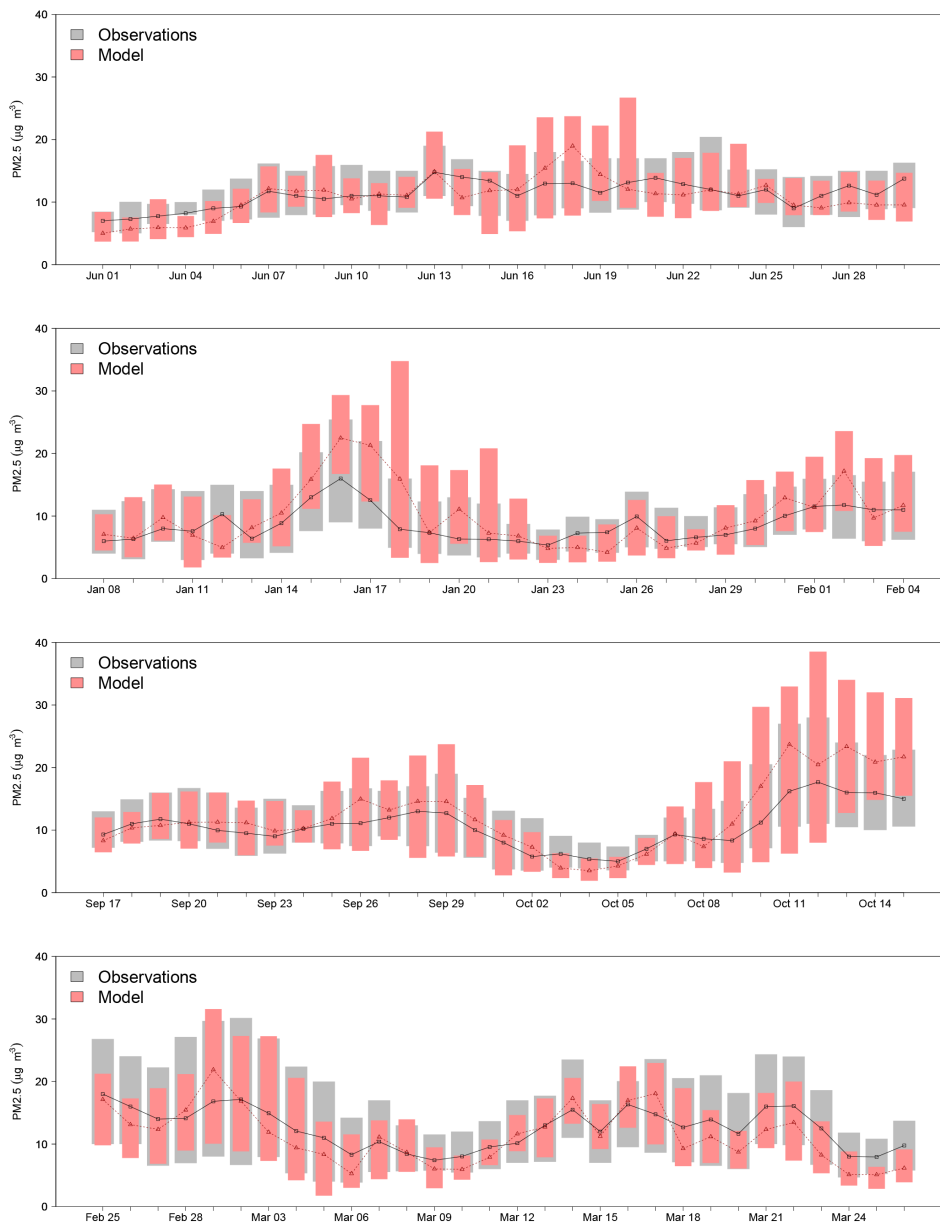
**Figure 1.** Modelled average  $\text{PM}_{2.5}$  concentrations for June 2006, January–February 2007, September–October 2008 and February–March 2009 (top to bottom and left to right) based on the base case (VBS\_BC). Note that the colour scale was limited to a maximum of  $40 \mu\text{g m}^{-3}$  to facilitate comparison of the panels.

**Table 4.** Statistical analysis of OA for NOVBS, VBS\_ROB and VBS\_BC scenarios for the 11 AMS sites for February–March 2009.

Scenario	Mean observed OA ( $\mu\text{g m}^{-3}$ )	Mean modelled OA ( $\mu\text{g m}^{-3}$ )	MB ( $\mu\text{g m}^{-3}$ )	ME ( $\mu\text{g m}^{-3}$ )	MFB [–]	MFE [–]
NOVBS	3.0	1.2	–1.8	2.0	–0.66	0.88
VBS_ROB	3.0	0.7	–2.3	2.4	–1.08	1.19
VBS_BC (base case)	3.0	1.7	–1.2	1.8	–0.47	0.79

are discussed. The scatter plots in Fig. 5 show a comparison of daily average OA concentrations against the same AMS measurements as in Table 3 during February–March 2009. Statistics for each scenario are reported in Table 4. When the semivolatile dynamics of primary organic aerosol is not taken into account (scenario NOVBS), the model underpredicts OA concentrations (MFB: –66%) with observed and modelled average concentrations of  $3.0$  and  $1.2 \mu\text{g m}^{-3}$ , respectively. In the VBS\_ROB scenario POA emissions are allowed to evaporate following the volatility distribution proposed by Robinson et al. (2007) and to undergo chemical oxidation. In this case modelled OA concentrations decrease by about 42% with respect to NOVBS, predicting an average OA concentration of  $0.7 \mu\text{g m}^{-3}$ . In contrast, the VBS\_BC scenario improves the OA model performance increasing the OA concentrations by about 42% with respect to NOVBS. Predicted OA concentrations are found to be  $1.7 \mu\text{g m}^{-3}$  on average (MFB: –47%). Similar behaviour during winter periods was also shown in recent studies where the same VBS scheme was applied in the US domain (Koo et al., 2014).

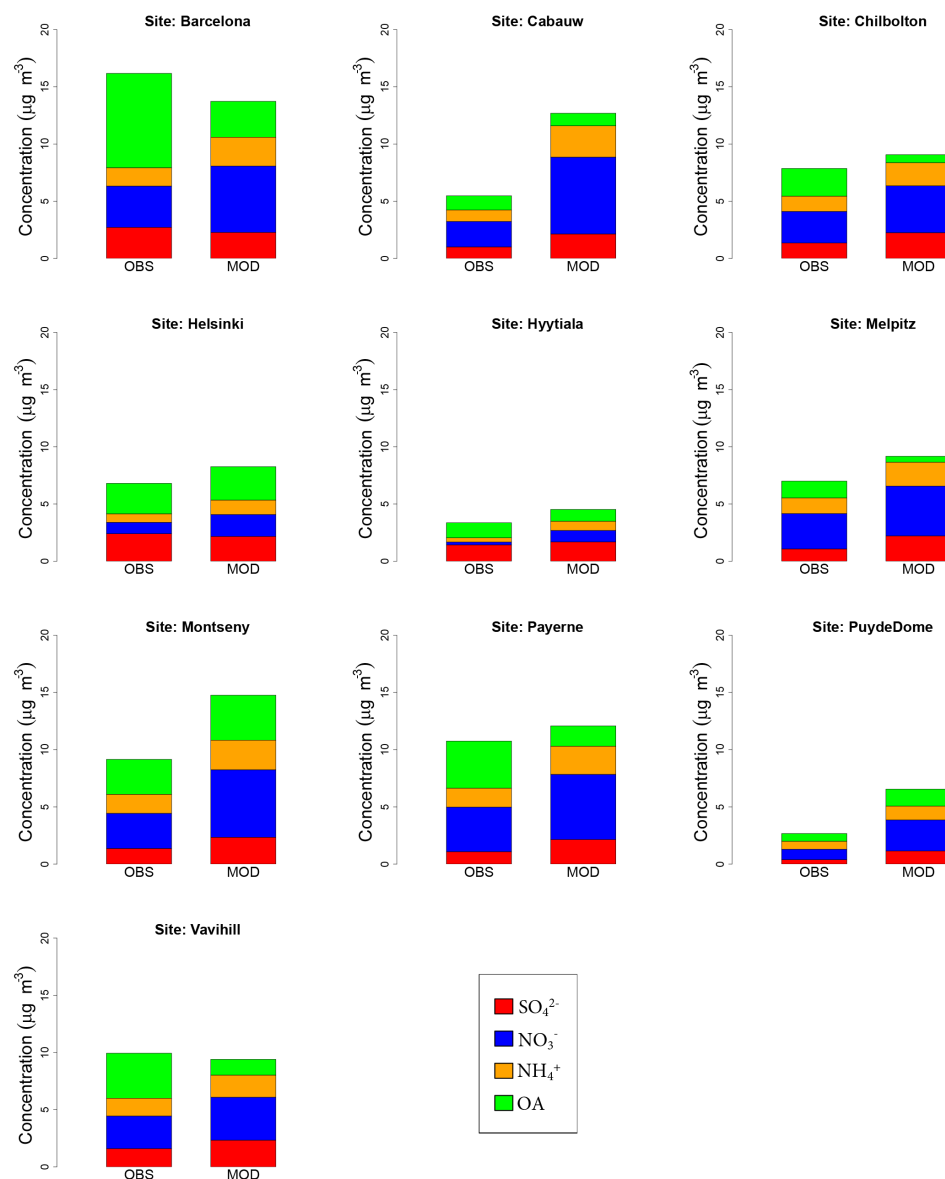
Figure 6 shows the modelled total OA concentration over Europe using NOVBS, VBS\_ROB and VBS\_BC scenarios. The model predicts high OA values in the eastern part of the domain as well as over Portugal, France and the Po Valley (VBS\_BC). Some hotspots around large urban areas are also visible, i.e. Paris and Moscow. Higher OA concentrations in the southern part of the domain are observed in the VBS\_BC case, likely because of higher temperature and more OH radicals available in that part of the domain leading to an increase in the total organic mass upon reaction with organic vapours. This is in line with the results of Fountoukis et al. (2014) for the February–March 2009 period even though their study predicts lower concentrations over the Po Valley. Even though model input data and parameterizations are not the same, the VBS\_BC case in particular uses a very similar volatility distribution to Fountoukis et al. (2014). Our study predicts relatively lower OA concentrations (MFB: –0.47; MFE: 0.79) compared to those reported by Fountoukis et al. (2014) (MFB: 0.02; MFE: 0.68) for February–March 2009. Unlike Fountoukis et al. (2014)



**Figure 2.** Comparison of modelled (red) and measured (grey)  $\text{PM}_{2.5}$  concentrations at AirBase rural-background sites. The extent of the bars indicates the 25th and 75th percentiles. The black and red lines are observed and modelled medians, respectively. The numbers of sites are 48, 56, 90 and 110 from top to bottom. Based on the base case (VBS\_BC).

our study does not include fire emissions and marine organic aerosol, which may partially explain the differences. Figure 7 shows hourly modelled and observed OA concentration at Payerne for March 2009 and June 2006. In March 2009 VBS\_ROB results are lower than those in NOVBS whereas OA concentrations in VBS\_BC case are higher (see Supplement Fig. S8 and Table S3 for average concentrations and statistics). In June 2006, the OA mass in VBS\_ROB is lower than those in NOVBS while VBS\_BC predicts similar concentrations as the NOVBS scenario ( $2.4$  and  $2.6 \mu\text{g m}^{-3}$ , respectively, Fig. S9 and Table S4). It has to be noted that the

NOVBS scenario predicts slightly lower OA concentration for June 2006 in Payerne with respect to our previous application (Aksoyoglu et al., 2011), mainly because of a different biogenic model being used which yields lower monoterpene and sesquiterpene emissions. Since both BVOCs and BBOA-like emissions are highly uncertain, sensitivity tests with increased biogenic and anthropogenic emissions were performed and results discussed in the next section (Sect. 3.3.2).



**Figure 3.** Comparison of observed (OBS) non-refractory  $\text{PM}_{10}$  and modelled (MOD) non-refractory  $\text{PM}_{2.5}$  at 10 AMS sites in Europe during February–March 2009. Mace Head is reported only in Table 3 since the ammonium component is not available.

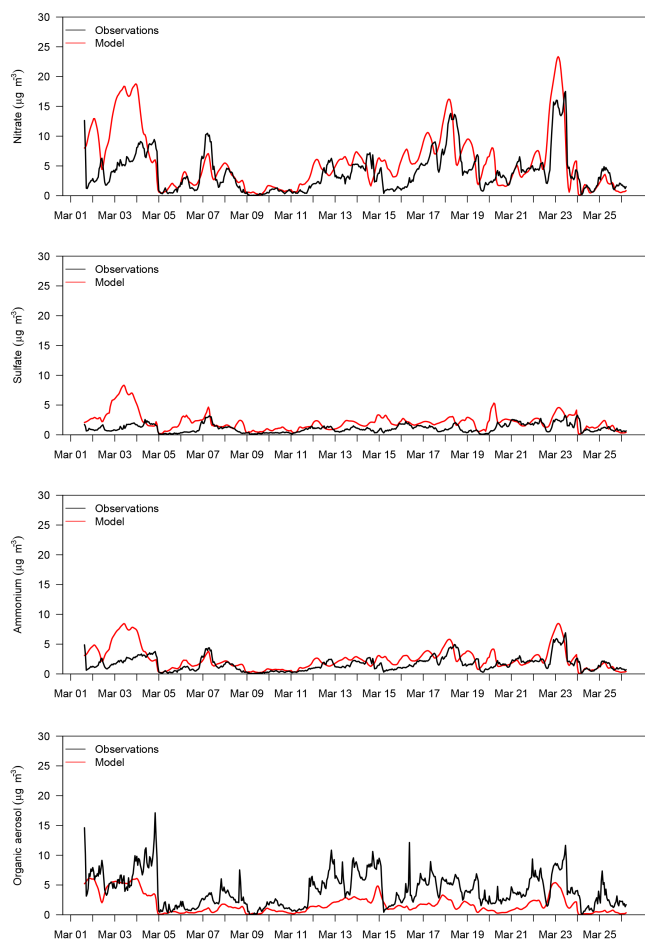
### 3.3.2 Sensitivity of OA to BBOA-like and BVOC emissions

Emissions of BVOCs compounds (i.e. monoterpenes, isoprene and sesquiterpenes) were doubled in scenario VBS\_BC\_2xBVOC, whilst primary organic aerosol emissions from SNAP2 and SNAP10 (BBOA-like) were doubled in scenario VBS\_BC\_2xBBOA, with other emissions and processes represented as in VBS\_BC. Figure 8 shows modelled and observed OA daily average concentrations for the VBS\_BC, VBS\_BC\_2xBVOC and VBS\_BC\_2xBBOA scenarios across the sites. Statistics for each scenario are reported in Table 5. Increasing biogenic emissions by a fac-

tor of 2 during February–March 2009 resulted in almost no change in the predicted total OA ( $1.7$  and  $1.8 \mu\text{g m}^{-3}$  for the VBS\_BC and VBS\_BC\_2xBVOC scenarios respectively). In contrast, doubling the BBOA-like emissions (VBS\_BC\_2xBBOA) during the same period strongly increased the predicted OA mass (up to  $2.8 \mu\text{g m}^{-3}$  on average). As a result the mean fractional bias decreased further, from  $-47$  to  $-12\%$  averaged across the sites. This could eventually confirm other studies in which substantial under-predictions in residential wood burning emissions were underlined (e.g. Bergström et al., 2012). A few points above the 2 : 1 lines in VBS\_BC\_2xBBOA mainly belong to the sites of Montseny, Puy de Dôme and Helsinki. During winter pe-

**Table 5.** Statistical analysis of OA for VBS\_BC, VBS\_BC\_2xBVOC and VBS\_BC\_2xBBOA scenarios for the 11 AMS sites for February–March 2009.

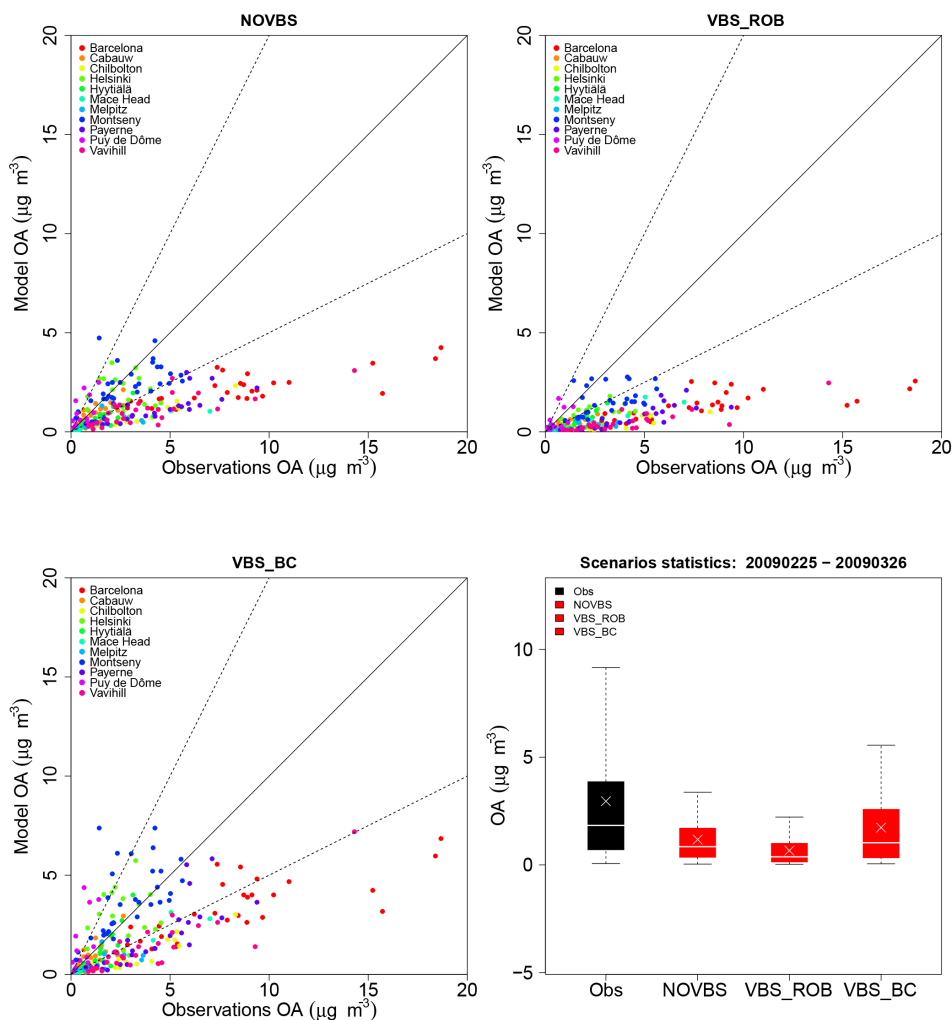
Scenario	Mean observed OA ( $\mu\text{g m}^{-3}$ )	Mean modelled OA ( $\mu\text{g m}^{-3}$ )	MB ( $\mu\text{g m}^{-3}$ )	ME ( $\mu\text{g m}^{-3}$ )	MFB [-]	MFE [-]
VBS_BC (base case)	3.0	1.7	-1.2	1.8	-0.47	0.79
VBS_BC_2xBVOC	3.0	1.8	-1.2	1.8	-0.46	0.78
VBS_BC_2xBBOA	3.0	2.8	-0.1	1.9	-0.12	0.69

**Figure 4.** Comparison of observed and modelled nitrate, ammonium, sulfate and organic aerosol at Payerne for March 2009.

riods, it is likely that elevated stations such Montseny and Puy de Dôme are most of the time above the PBLH, as suggested by previous studies for Puy de Dôme (Freny et al., 2011), whereas model concentrations are extracted from the first layer of the model. In Helsinki, BBOA emissions seem to be overestimated or the dispersion underestimated in the model.

Comparison with a warmer period in June 2006 is reported as well for Payerne, where AMS measurements were also available (Fig. 9). In February–March 2009,

increasing BBOA-like emissions (VBS\_BC\_2xBBOA) reduced the fractional bias from  $-85\%$  in VBS\_BC to  $-37\%$  (Table S3) with an overprediction occurring during 1–5 of March (Fig. 9, upper panel). As already discussed in Sect. 3.2, it is likely that the vertical mixing processes were not correctly represented by the model since also the inorganic components were overpredicted for the same period. Almost no change in the predicted OA mass was found when biogenic emissions were doubled (scenario VBS\_BC\_2xBVOC) (Fig. 9, upper panel) due to lower BVOCs emission during winter periods. Increasing BVOCs emissions in June 2006 increased the predicted OA mass at Payerne site, especially during the 12–16 June and towards the end of the simulation period, when higher concentrations and temperature (Fig. S7) were also observed (Fig. 9, lower panel). In contrast, similar OA concentrations were predicted in Payerne for VBS\_BC and VBS\_BC\_2xBBOA during June 2006 (with averages of  $2.4$  and  $2.8 \mu\text{g m}^{-3}$ , respectively). This is in line with a recent source apportionment study based on ACSM (Aerosol Chemical Speciation Monitor) measurements performed in Zürich for 13 months (February 2011–February 2012) which revealed substantial differences between the winter (February–March) and summer (June–August)  $f_{44}/f_{43}$  space (organic mass fraction measured at mass to charge ratio 44 and 43), indicating that summer OOA (oxygenated organic aerosol) is strongly influenced by biogenic emission and winter OOA by biomass burning emission (Canonaco et al., 2015). Increased OA concentrations at Payerne in June 2006 with increased biogenic emissions were also found in other modelling studies. Bergström et al. (2012) used the VBS framework with different assumptions regarding aging processes and compared the model results for June 2006 with the AMS results at Payerne. In their study the total OA was found to be underpredicted with lower bias observed when aging processes were taken into account and biogenic emissions were increased by a factor of 3. Even though their model differs from ours in various aspects (number of volatility bins, aging processes parameterization and input data) in two of their scenarios without aging of biogenic SOA, Bergström et al. (2012) predicted an average OA concentration ranging from  $2.6$  to  $3.4 \mu\text{g m}^{-3}$ , which is similar to our base case VBS\_BC and VBS\_BC\_2xBVOC scenario ( $2.4 \mu\text{g m}^{-3}$  and  $3.4 \mu\text{g m}^{-3}$ , respectively; Table S4).



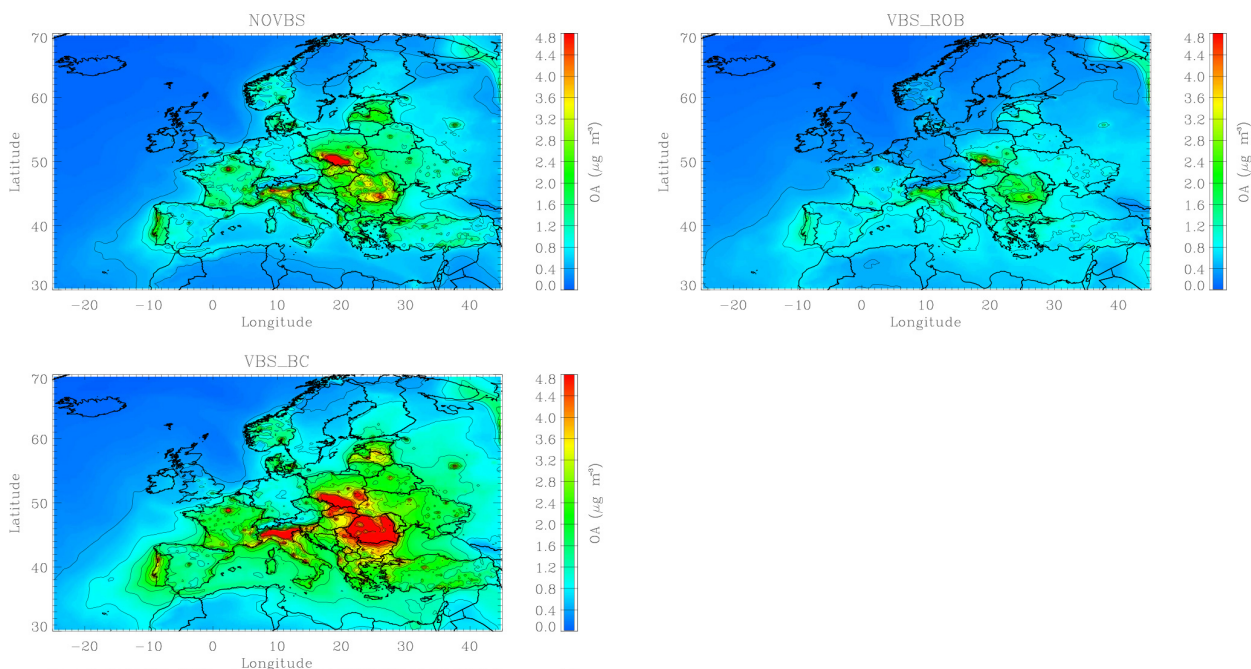
**Figure 5.** OA daily average scatter plots for NOVBS, VBS\_ROB and VBS\_BC scenarios for February–March 2009 for stations in Table 3. Solid lines indicate the 1 : 1 line. Dotted lines are the 1 : 2 and 2 : 1 lines. Box plots indicate medians, 5th, 25th, 75th and 95th quantiles for observations (black) and sensitivity tests (red). The crosses represent the arithmetic means.  $R^2$  is 0.55 for NOVBS, 0.64 for VBS\_ROB and 0.59 for VBS\_BC (excluding the elevated site of Puy de Dôme and Montseny).

### 3.3.3 OA components in summer and winter

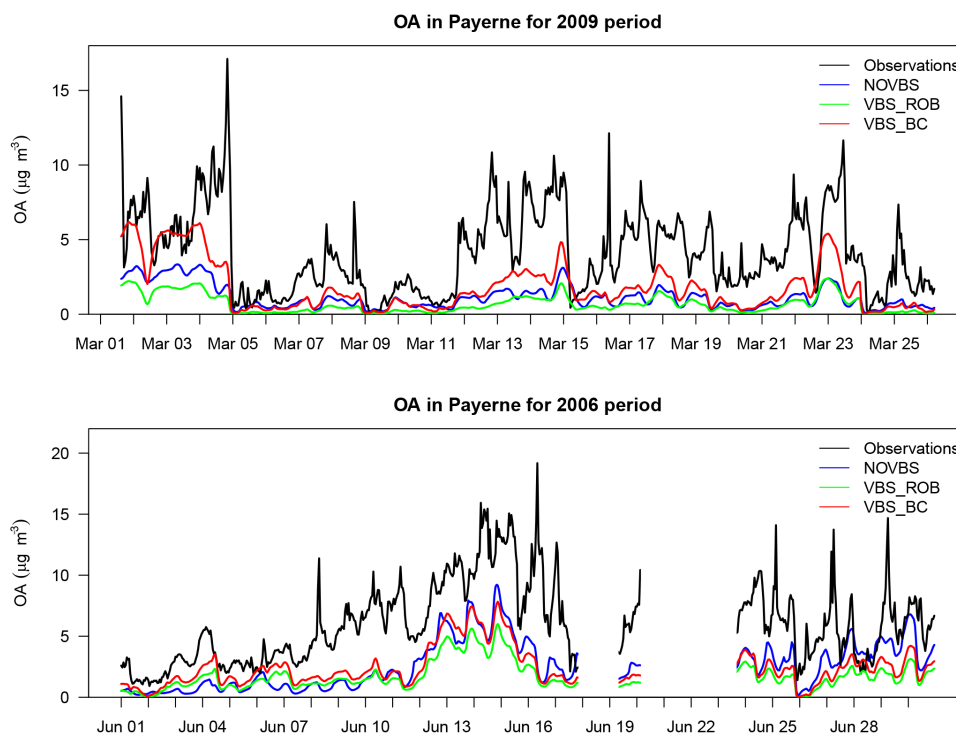
Comparisons of the primary and secondary organic fractions at the rural site of Payerne during summer (June 2006) and winter (February–March 2009) periods are reported in Fig. 10. During the winter period the VBS scheme better reproduced the primary and secondary organic aerosol components compared to the NOVBS case. In particular, for the VBS\_ROB base case, total OA concentrations were lower compared to the NOVBS case, consistent with the study of Woody et al. (2016) in which the same VBS scheme was applied to the US domain. The total OA concentrations in the base case (VBS\_BC) and in the scenario with increased biomass burning emissions (VBS\_BC\_2xBBOA) were higher compared to NOVBS case, even though SOA and POA fractions were not correctly reproduced. Higher

contribution from the primary fraction during winter periods was also predicted by the study of Koo et al. (2014) which deployed the same VBS scheme. Eventually, this might indicate that biomass burning precursors might be missing in this study or that the oxidation pathways of primary organic material need to be improved in the model (up to 86 % of the reacted primary organic material is still allocated in the primary set as oxidation proceeds, directly increasing the POA fraction).

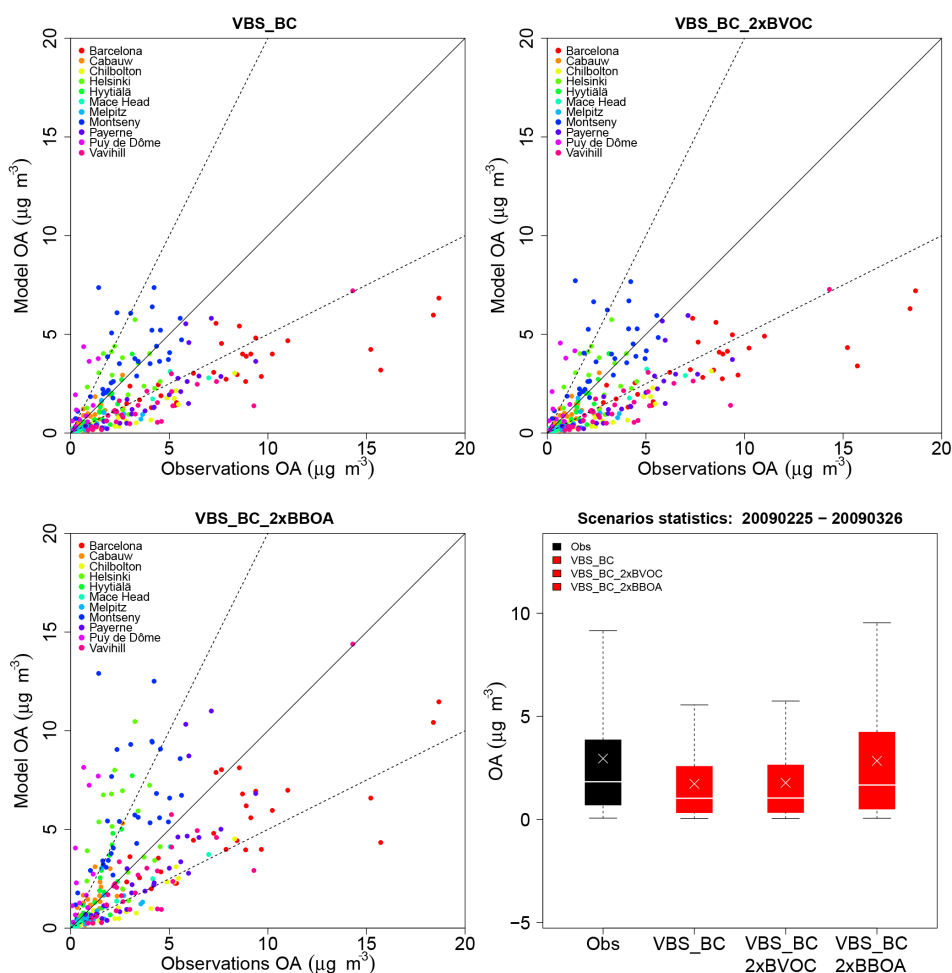
Different behaviour was observed for the summer period where the larger contribution of SOA to the total OA retrieved from measurements was also reproduced by the model, even though the total OA concentration was still underestimated. These results for summer are also in line with the study of Koo et al. (2014) for summer periods in the US domain using the same VBS scheme.



**Figure 6.** Predicted OA concentrations over Europe for the NOVBS, VBS\_ROB and VBS\_BC scenario in February–March 2009. Note that the colour scale was limited to a maximum of  $4.8 \mu\text{g m}^{-3}$  to facilitate comparison of the panels.



**Figure 7.** Predicted and observed total OA for scenarios NOVBS, VBS\_ROB and VBS\_BC in March 2009 (upper panel) and June 2006 (lower panel) at Payerne.



**Figure 8.** OA daily average scatter plots for VBS\_BC, VBS\_BC\_2xBVOC and VBS\_BC\_2xBBOA scenarios for February–March 2009 for stations in Table 3. Solid lines indicate the 1 : 1 line. Dotted lines are the 1 : 2 and 2 : 1 lines. Box plots indicate medians, 5th, 25th, 75th and 95th quantiles for observations (black) and sensitivity tests (red). The crosses represent the arithmetic means.

#### 4 Conclusions

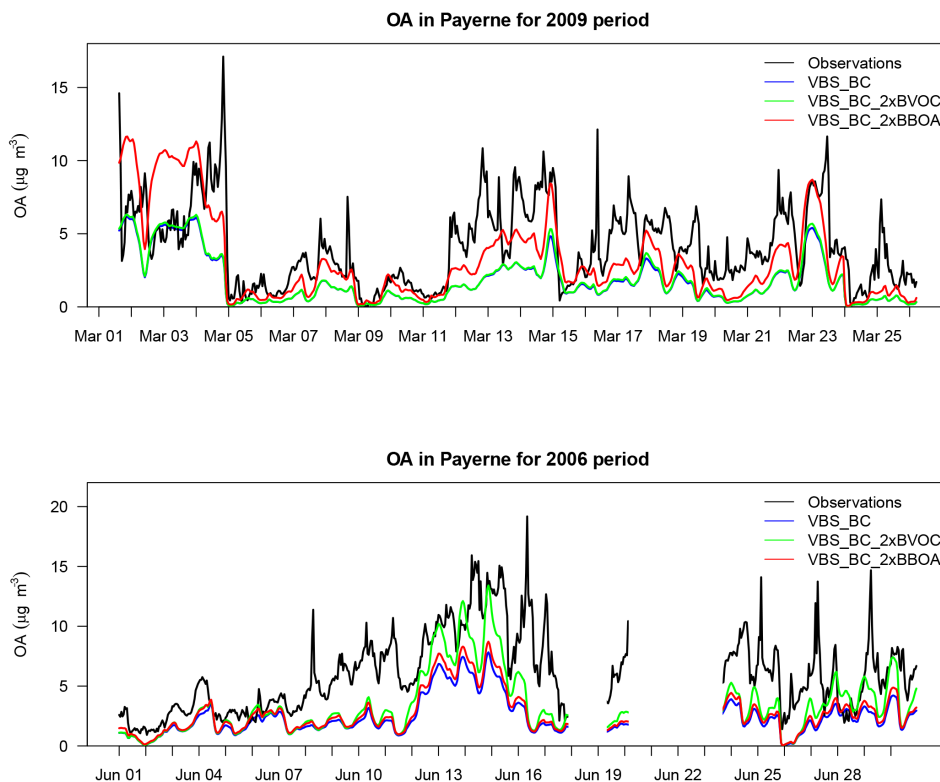
A modelling study using the regional air quality model CAMx with a VBS scheme was performed for the first time in Europe within the EDIII model intercomparison exercise. An evaluation for the main gas phase species and  $PM_{2.5}$  for four different periods was performed using the European air quality database AirBase as well as AMS measurements. The period in February–March 2009 was further analysed in more detail using different assumptions regarding the volatility of emitted organic aerosol and precursor emissions. The main findings of this study are summarized below.

Although total  $PM_{2.5}$  mass concentrations and its variations were well reproduced by the model in all four periods, comparisons with AMS measurements for the February–March 2009 period revealed that the good agreement between model and measurements was due, most of the time, to overestimation of the inorganic fraction, especially  $NO_3^-$ , and underestimation of OA. Sensitivity tests with reduced

$NH_3$  emissions generally reduced the positive bias in  $NO_3^-$ , suggesting potential uncertainties in  $NH_3$  emissions and their seasonal variability.

In general, for all the four periods, the model underpredicted  $NO_2$  and CO concentrations. In contrast,  $O_3$  was found to be overpredicted likely because of insufficient  $NO_x$  to undergo titration during night-time chemistry or not well-captured vertical mixing processes and concentrations at the boundaries.  $SO_2$  was slightly overpredicted, presumably mainly because of uncertainties in high stack point source representation in the model or too low deposition processes.

Including evaporation and oxidation processes of primary organic particles with the volatility distribution proposed by Robinson et al. (2007) lowered the modelled OA mass both in winter and summer periods. However, the adjustment of the scheme by Robinson et al. (2007) suggested by Shrivastava et al. (2011) and Tsimpidi et al. (2010) brought model and observations into better agreement by reducing the negative bias for OA by about 29 % (MFB) in winter.



**Figure 9.** Predicted and observed total OA for scenarios VBS\_BC, VBS\_BC\_2xBVOC and VBS\_BC\_2xBBOA in March 2009 (upper panel) and June 2006 (lower panel) at Payerne.

Sensitivity tests with increased BVOCs and BBOA-like emissions suggested that emissions from residential heating represent an important contributor to total OA during winter periods (February–March 2009). The model underpredicted the winter OA concentrations (MFB  $-47\%$  for base case VBS\_BC) more than gas phase pollutants, e.g.  $\text{NO}_2$  (Table 2). Eventually, increasing BBOA-like emissions by a factor of 2 brought model and observation to a reasonably good agreement even though the model still underpredicted the OA fraction ( $-12\%$  MFB). This underlines the necessity to better constrain emission inventories with a focus on residential heating. Also, the implementation of the VBS scheme for domestic wood burning, which substantially influences both the primary and secondary organic aerosol, should be evaluated.

A summer period was simulated as well and results were compared at Payerne. In June 2006, the current VBS implementation could not explain the discrepancy between modelled and observed OA. During this period the difference between the model and measurements was likely related to BVOCs emissions which are uncertain and difficult to constrain with measurements. In this case the model was sensitive to an increase in biogenic emissions especially during periods with higher temperature and OA concentrations. The latter could confirm the importance of BVOC precursors in

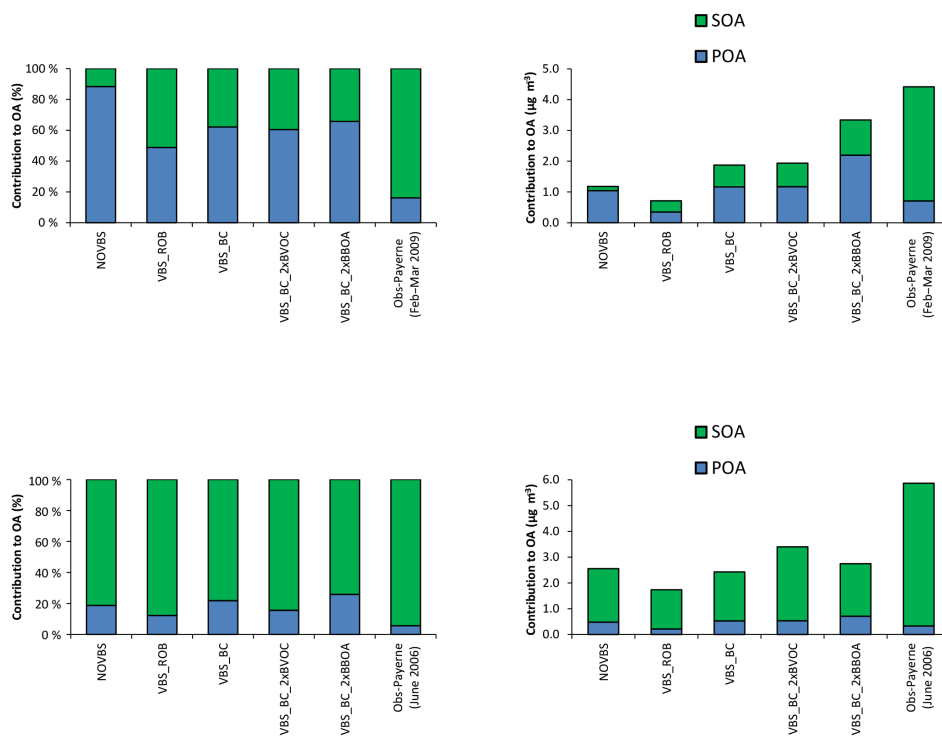
summer in Payerne and the way to correctly represent their evolution in the atmosphere.

## 5 Data availability

All data are available upon request from the corresponding authors. The CAMx model is available at <http://www.camx.com/>.

**The Supplement related to this article is available online at [doi:10.5194/acp-16-10313-2016-supplement](https://doi.org/10.5194/acp-16-10313-2016-supplement).**





**Figure 10.** Relative (left) and absolute (right) contributions of predicted and measured POA and SOA fractions to the total OA mass at Payerne for February–March 2009 winter period (upper panel) and June 2006 (lower panel) and different model scenarios. NOVBS (traditional nonvolatile POA), VBS\_ROB (Robinson et al., 2007), VBS\_BC (Tsimpidi et al., 2010; Shrivastava et al., 2011), VBS\_BC\_2xBVOC (increased biogenic emissions relative to VBS\_BC), VBS\_BC\_2xBBOA (increased biomass burning emissions relative to VBS\_BC), Obs-Payerne (AMS-PMF).

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