- 1 Evaluation of the high resolution WRF-Chem (v3.4.1) air
- 2 quality forecast and its comparison with statistical ozone
- 3 predictions
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Abstract

An integrated modelling system based on the regional on-line coupled meteorology-atmospheric chemistry WRF-Chem model configured with two nested domains with horizontal resolution 11.1 km and 3.7 km has been applied for numerical weather prediction and for air quality forecast in Slovenia. In the study an evaluation of the air quality forecasting system has been performed for summer 2013. In the case of ozone (O₃) daily maxima, the first and second day model predictions have been also compared to the operational statistical O₃ forecast and to the persistence. Results of discrete and categorical evaluations show that the WRF-Chem based forecasting system is able to produce reliable forecasts, which depending on monitoring site and the evaluation measure applied can outperform the statistical model. For example, the correlation coefficient shows the highest skill for WRF-Chem model O₃ predictions, confirming the significance of the non-linear processes taken into account in an on-line coupled Eulerian model. For some stations and areas biases were relatively high due to highly complex terrain and unresolved local meteorological and emission dynamics, which contributed to somewhat lower WRF-Chem

- skill obtained in categorical model evaluations. Applying a bias-correction could further
- 2 improve WRF-Chem model forecasting skill in these cases.
- 3 **Key words**: Air quality, forecast, ozone, WRF-Chem, online-coupled model, statistical model

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1 Introduction

Real-time air quality forecasting (RT-AQF) is a relatively new discipline in atmospheric 6 sciences, which has evolved as a response to societal and economic needs, reflecting the 7 8 progress in scientific understanding of physical processes and numerical and computational 9 technologies (Zhang et al., 2012a). The first RT-AQF systems, developed for forecasting air 10 pollution in exposed urban regions, were either empirical methods based on persistence, 11 climatology, human expertise and meteorological forecast (e.g. Wolff and Lioy, 1978), or 12 statistical models taking advantage of links between pollutant concentrations, meteorological variables (wind speed and direction, temperature, cloudiness, moisture etc.) and physical 13 14 (emissions) parameters (e.g. McCollister and Wilson, 1975; Cobourn, 2007; Vlachogianni et al., 2011). The next step in evolution of RT-AQF systems was the use of sophisticated 15 chemical transport models that represent all major processes (meteorological and chemical) 16 that lead to the formation and accumulation of air pollutants. Many of these RT-AQF systems 17 18 consist of an offline coupled meteorological model and a chemical-transport model, where the meteorological model (e.g., ALADIN, ALADIN International Team, 1997; MM5, Grell et al., 19 20 1994; WRF, Skamarock et al., 2008) provides meteorological input for the chemical-transport model (e.g., EMEP, van Loon et al., 2004; CMAQ, Byun and Schere, 2006; CAMx, 21 22 ENVIRON, 2011; CHIMERE, Menut et al., 2013) with an output time interval typically around 1 hour. Examples are the EURAD (http://db.eurad.uni-koeln.de/index_e.html), 23 24 **SILAM** (http://silam.fmi.fi/), ForeChem (http://atmoforum.aquila.infn.it/forechem/), 25 CALIOPE (http://www.bsc.es/caliope/) forecast systems and others. The new generation of an online coupled models (e.g., MCCM, Grell et al., 2000; GATOR-GCMM, Jacobson 2001; 26 27 Meso-NH-C, Tulet et al. 2003; WRF-Chem, Grell et al., 2005; Enviro-HIRLAM, Baklanov et al., 2008; GEM-AQ, Kaminski et al. 2008; COSMO-ART, Vogel et al., 2009; WRF-Chem-28 29 MADRID, Zhang et al., 2010a) presents an alternative approach with one unified modelling system, in which meteorological and air quality variables are simulated together within the 30 31 same model. The online approach permits the simulation of two-way interactions between different atmospheric processes including emissions, chemistry, clouds and radiation, and a 32

better response of the simulated pollutant transport to changes of the wind field (Grell et al., 1 2 2004), and can thus provide a more realistic representation of the atmosphere. The use of 3 online coupled models can be particularly important in regions with high aerosol loadings and 4 cloud coverage (Otte et al., 2005; Eder et al., 2006), where physical processes in the 5 atmosphere may be modified by the aerosol direct effect on radiation or by aerosol cloud interactions. Several reviews summarized the strengths and limitations of offline and online 6 7 coupled models (e.g. Zhang 2008; Klein, 2012; Baklanov et al., 2014). There is an increasing 8 awareness that an integrated online approach is needed not only for assessment, forecasting 9 and communication of air quality, but also for weather forecasting (e.g. Baklanov, 2010; Grell 10 and Baklanov, 2011; Klein et al., 2012; Zhang et al., 2012b; Baklanov et al., 2014). 11 Nevertheless, there are several issues regarding the inclusion of chemistry into numerical weather prediction models. More evidence is required whether an integrated model can 12 13 produce a good climatology of the most important chemical species, and if such a model is, 14 considering many uncertainties, able to beat persistence forecasts of these species (Grell and 15 Baklanov, 2011). These questions are calling for further research and studies exploring the 16 performance of the models with an online coupled chemistry. In recent years extensive efforts have been devoted to develop air quality (AQ) forecasting 17 systems for Slovenia. In this study we explore the use of the state-of-the-science WRF-Chem 18 model (Grell et al., 2005) with coupled meteorological, microphysical, chemical, and 19 radiative processes for forecasting AQ in Slovenia during summertime conditions. In last 20 21 decade WRF-Chem has been increasingly applied to many areas worldwide (e.g., Misenis and 22 Zhang, 2010; Fast et al., 2009; Zhang et al., 2010a, 2010b; Li et al., 2011; Tie et al., 2009; Hu et al., 2012; Forkel et al., 2012, Žabkar et al., 2011a, 2013). In most of these studies WRF-23 Chem model has been successfully used to simulate historical poor AQ conditions in hindcast 24 25 approach. To our knowledge, only a few studies focused on using WRF-Chem for forecasting AQ, most of these have applied WRF-Chem forecast before and during field campaigns 26 27 (McKeen et al., 2005, 2007, 2009; Yang et al., 2011). Takigawa et al. (2007) evaluated O₃ 28 forecast for a 1 month time period from a one-way nested global-regional RT-AQF system 29 with full chemistry based on the global CHASER (Sudo et al. 2002) and regional WRF-Chem 30 models, while Saide et al. (2011) evaluated a forecast system based on WRF-Chem model for 31 simulating carbon monoxide (CO) as a PM10/PM2.5 surrogate over Santiago de Chile for 32 wintertime conditions. WRF-Chem-MADRID (Zhang et al., 2010a) with two additional gasphase mechanisms, sectional representation for particle size distribution and more advanced 33

1 model treatments compared to WRF-Chem, was applied by Chuang et al. (2011) and by

2 Yahya et al. (2014) for forecasting AQ over the Southeastern U.S.. In spite of a limited

3 number of evaluation studies published in the literature, an increasing number of real-time

4 weather and air quality forecasting systems based on WRF-Chem are implemented worldwide

5 (http://ruc.noaa.gov/wrf/WG11/Real_time_forecasts.htm).

In our study we explore the forecasting skill of WRF-Chem model over the topographically complex and geographically diverse area of Slovenia for three summer months (June - August 2013). Furthermore, in the case of O₃ we compare WRF-Chem predictions with a statistical model for predicting O₃ daily maxima, currently used at the Slovenian Environment Agency (SEA). Both first day (1-day) and second day (2-day) forecasts are considered, while a persistence model, which assumes that pollutant level today and tomorrow will be the same as yesterday, is used as a threshold for useful model prediction. Since the availability of accurate and reliable forecasting system could be useful to the local authorities and could help to advise the public the proper preventive actions, we want to answer the question whether WRF-Chem model outperforms the statistical model or persistence. Namely, considering many uncertainties related to one unified model, it may not be easy for models with online chemistry to be able to perform well enough to meet the required standards, and more research and studies are needed to investigate that (Grell and Baklanov, 2011). Due to the limited number of previous studies focused on online coupled forecasting systems, the aim of our study is also to provide a greater insight into potential that lies in the approach based on an unified model for forecasting weather and air pollution. Finally, identified strengths, limitations and deficiencies of analyzed RT-AQFs, are expected to present the basis for further research.

2 Methodology

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2.1 WRF-Chem forecast system

The RT-AQF system for Slovenia based on the WRF-Chem model version 3.4.1 is configured with two nested domains (Fig.1) with horizontal resolution 11.1 km and 3.7 km, and 151×100 and 181×145 grid points, respectively. A 1-way nesting is applied by two separate consecutive simulations, where outputs from the coarse grid integration are processed to provide boundary conditions for the nested run every 15 minutes. The vertical structure of the atmosphere is resolved with 42 vertical levels extending up to 50 hPa, with the highest

resolution of ~25 m near the ground. About 15 levels are located within the lowest 2 km to 1 2 assure high vertical resolution of the daytime planetary boundary layer (PBL). To produce the 48-hour forecast, the model is run every day, starting at 00 UTC, with meteorological initial 3 (ICs) and lateral boundary conditions (BCs) taken from the 0.5° data from the Global Forecast 4 5 System (GFS) operated by the US National Weather Service (NWS). For chemical BCs forecasts from global MOZART-4/ GEOS-5 (Emmons et al., 2010) RT-AQF system with 6 temporal availability of 6 h are used. The instantaneous outputs at the 24th hour of the 7 previous day forecast are used to initialize next day's forecasting simulation. An exception is 8 9 the very first day of the first 48-hour forecasting cycle, when global MOZART-4/ GEOS-5 10 fields were used also to initialize chemistry. A three day spin-up ahead of the first analyzed 11 forecast day is then taken into account to allow pollutants to accumulate in the air masses. 12 In the WRF-Chem model, several choices for parameterizations of physical and chemical 13 processes are available (Grell et al., 2005; Skamarock et al., 2008; Peckham et al., 2011), and 14 their choice can have a strong impact on the model predictions. In both domains we decided 15 to apply the same schemes as were used in simulation SI1 for Phase-2 of the Air Quality Model Evaluation International Initiative (AQMEII) (e.g., Balzarini et al., 2014, Baró et al., 16 17 Curci et al., 2014, Forkel et al., 2014, Im et al., 2014a and 2014b, Kong et al., 2014, 2014, 18 San Josè et al., 2014). These include Yonsei University (YSU) PBL scheme (Hong et al., 19 2006), NOAH land-surface model (Chen and Dudhia, 2001), Rapid Radiative Transfer 20 Method for Global (RRTMG) long-wave and short-wave radiation scheme (Iacono et al. 21 2008), Grell 3D ensemble cumulus parameterization scheme (Grell and Devenyi, 2002) with 22 radiative feedback, Morrison double-moment cloud microphysics (Morrison et al., 2008), 23 Fast-J photolysis scheme (Fast et al., 2006), RADM2 gas phase chemistry (Stockwell et al., 1990) and the MADE/SORGAM aerosol module (Ackermann et al., 1998, Schell et al., 24 25 2001). Current model implementation includes a modified RADM2 gas phase chemistry 26 solver as described in Forkel et al. (2014), which avoids under-representation of nocturnal O₃ 27 titration in areas with high NO emissions. According to Forkel et al. (2014) the modified 28 solver tends to over-estimate the low NO₂ concentration for pristine regions and in the free 29 troposphere, which results in an overestimation of O₃. Due to the focus on polluted regions 30 this deficiency was considered as less important than the advantage of better description of the titration. In addition, the comparatively small modelling domain (D1) ensures that the 31 32 boundary conditions constrain the high bias of the modified solver for O₃ and NO₂ in the free troposphere. Also according to our sensitivity tests (results not shown) the modified solver 33

- 1 showed better performance for O₃ daily maxima and O₃ nighttime minima than the QSSA
- 2 RADM2 solver supplied originally with WRF-Chem model.
- 3 Among feedbacks only the aerosol direct effects on radiation according to Fast et al. (2006)
- 4 and Chapman et al. (2009) are taken into account. As shown by Kong et al. (2014) for two air
- 5 pollution episodes, this degree of aerosol-meteorology interactions in 3.4.1 version of the
- 6 WRF-Chem improved model performance for high aerosol loads, while the representation of
- 7 the indirect effects needs to be further improved to be able to outperform simulations with
- 8 direct effects only.
- 9 Biogenic emissions are estimated using MEGAN (Model of Emissions of Gases and Aerosols
- 10 from Nature; Guenther et al., 2006) online model calculations, while dust emissions are
- modelled according to Shaw et al. (2008) with an adjustment to avoid high dust fluxes from
- some Dalmatian islands in Croatia. A detailed anthropogenic inventory for pollutants CO,
- NH₃, NOx, SO₂, and NMVOC, which has been for the purpose of AQ forecasting constructed
- 14 for year 2009 by SEA (SEA, 2014), is used to estimate anthropogenic emissions in Slovenia.
- 15 For areas outside Slovenia the recently updated anthropogenic emissions for the year 2009
- based on the TNO-MACC-II (Netherlands Organization for Applied Scientific Research,
- 17 Monitoring Atmospheric Composition and Climate Interim Implementation), the same as
- prepared for phase-2 of the AQMEII exercise (Pouliot et al., 2014), are being used. Daily
- 19 updates of the WRF-Chem based experimental AQ forecast are provided at
- 20 http://meteo.fmf.uni-lj.si/onesnazenje.

21 2.2 Statistical ozone daily maximum forecast

- 22 The statistical O₃ model (Žabkar, 2011b), currently used at SEA for forecasting O₃ daily
- 23 maxima at 8 measuring sites in Slovenia (Fig.3), is a multivariate regression tool combined
- 24 with clustering algorithms to take into account measured data, weather forecast data, as well
- as the predicted backward trajectories of each monitoring site. As regards measurements,
- yesterday (at 12, 15, 18 and 21 local time, daily maximum, daily minimum, daily average)
- and today early morning (7 local time) meteorological (pressure, relative humidity, direct and
- diffusive solar radiation, wind speed) and AQ data (O₃, NO_x, NO₂, CO, PM₁₀, SO₂) are used.
- 29 For meteorological predictions the 24-h ECMWF forecast variables at 12 UTC of the forecast
- day at different vertical levels (1000 hPa, 925 hPa, 850 hPa, 500 hPa, 300 hPa) above the
- 31 measuring sites are taken into account. Among all these variables by the use of stepwise

- technique, based on the F-statistic only significant variables were selected to be included in
- 2 multivariate regression equations for different monitoring sites (from 15 to 26 variables,
- 3 depending on monitoring site).
- 4 The important part of the statistical forecast is calculation of 24-h backward trajectories on
- 5 meteorological fields of ALADIN/SI forecast. The inclusion of 24-h predicted trajectories
- 6 into statistical model is based on the study (Žabkar et al., 2008) which showed, that the
- 7 highest O₃ daily maxima at monitoring sites in Slovenia are in general associated with short
- 8 (slow-moving) backward trajectories with a southwestern origin, while the lowest measured
- 9 daily maximum O₃ values for all the stations are associated with the clusters of long
- 10 northwestern trajectories. Clusters of similar trajectories were for the purpose of statistical
- forecast calculated by k-means clustering algorithms (Moody and Galloway, 1988; Žabkar et
- al., 2008) on 6 years (2004-2010) of data (ALADIN/SI trajectories). As an example, Fig. 2
- shows a mean O₃ daily maxima for clusters of similar trajectories for one of the monitoring
- sites. The same 6-year time period of training data was used in the stepwise multiple
- 15 regression procedure to determine the multiple regression prognostic equations associated
- with monitoring sites and trajectory clusters, from measurements, ECMWF forecast data,
- 17 average cluster O₃ daily maximum, and day-of-the-year variable.
- 18 The first step of the statistical O_3 prediction is the calculation of trajectories approaching the
- monitoring stations at 12 UTC of the forecast day. In the next step these backward trajectories
- 20 of each monitoring site are associated to the nearest pre-calculated cluster of similar
- 21 trajectories. Finally, the multiple regression equation of the associated group of trajectories is
- used to calculate the O₃ daily maximum prediction. It must also be noted, that the decision on
- 23 declaring O₃ episodes is only partially based on the results from this statistical model; it also
- involves a decision made by AQ forecasters.

2.3 Evaluation methodology

- We evaluate the 1-day and 2-day WRF-Chem meteorological and AQ forecasts on the high
- 27 resolution domain during a 3-month period (June August 2013). The main focus is on O₃
- predictions. In the case of air pollutants, the instantaneous lowest model level mixing ratios
- 29 (with grid point center about 12 m above model orography an exception is KRV station as
- and explained below) are compared to the hourly averaged concentrations measured at monitoring
- 31 stations (which have a typical inlet height of 3 m) from the national network and some other

environmental information systems in Slovenia. Figure 3 shows locations of these AQ 1 2 monitoring stations, and Tab. 1 lists the basic characteristics, including comparison of the 3 station altitude, the height of model orography, model analysis height, and pollutants with 4 higher than 75% availability of valid data during the analyzed time period for each of the AQ 5 monitoring site. In the case of the elevated alpine KRV station, AQ variables are evaluated for the 5th model layer instead of the first model layer. We made this exception for KRV, since 6 7 the height of the model topography was significantly underestimated there (Tab. 1), as well as 8 the station is known to be strongly influenced by the conditions of the free troposphere. The selection of the 5th model layer for KRV station is based on analyses performed for different 9 model layers (results not shown) and was found to reduce the negative bias for O₃ due to too 10 11 low WRF-Chem topography at this location. Although even for this model layer the location 12 of the grid point representing KRV station (1414 m) is still well below the true station altitude 13 (1740 m), the O₃ bias for KRV station is significantly smaller than for the first layer, while 14 the correlation coefficient between the measured and simulated O₃ levels remains similar in both cases (the 5th or the lowest model layer). Taking results from higher model layers would 15 further decrease the negative model bias, but would also worsen the correlation coefficient for 16 O₃ at this station due to decreased impact of surface processes. 17 All AQ stations are background, 7 of them are measuring urban background, 1 suburban and 18 19 9 rural conditions. Valid O₃ measurements are for the analyzed time period available for 13 AQ stations. When studying the general model performance, data from additional 4 stations 20 21 for two other pollutants (NO₂, PM10) are also analyzed to get a better picture of model 22 behavior over the domain, known for its large topographical and climate diversity. The 23 coverage of three climate zones in Slovenia (Mediterranean, sub-alpine and mountainous) with monitoring stations is the following: NG, KOP and OTL are Mediterranean sites, KRV is 24 25 a mountainous station, and the remaining stations are sub-alpine. As well as the elevated station KRV, the ISK, OTL and VNA stations are also influenced by regional transport of 26 27 pollutants. For evaluation of predicted meteorological variables, data from SEA meteorological stations 28 (MET, Fig. 3) for 2m temperature (T2m), 10 m wind speed (W10m), relative humidity (RH), 29 incoming shortwave radiation (SR) and precipitation (RR) are used. It must be noted, that 30 MET stations with lower spatial representativeness (e.g. alpine stations) were not a priori 31 excluded from the analyses, which needs to be taken into account when looking at evaluation 32

- 1 results. The reason for not excluding these stations was that some information about the AQ
- 2 forecast can also be gained by the evaluation of meteorological forecast for these stations.
- 3 Basic statistical measures (correlation coefficient (CORR), mean error (ME), mean absolute
- 4 error (MAE) and root mean square error (RMSE)) are used for evaluating model's forecasting
- 5 skills of meteorological and AQ variables. In the case of O₃, correlation coefficients are
- 6 presented also by Taylor diagrams (Taylor, 2001), which graphically summarize the similarity
- 7 between model forecasts and observations not only in terms of their correlation, but also with
- 8 their centered root-mean-square difference and the amplitude of their variations, represented
- 9 by their standard deviations. Furthermore, some additional discrete statistical measures,
- 10 including index of agreement (IOA), the mean normalized bias error (MNBE), and the mean
- 11 normalized gross error (MNGE) are calculated for O₃ daily maximum concentrations
- predicted by the different models. Finally, to evaluate the model's ability to predict
- exceedances and non-exceedances also several categorical indices including Equitable Threat
- 14 Score (ETS), Critical Success Index (CSI), Bias (B), False Alarm Ratio (FAR) and
- 15 Probability Of Detection (POD) are calculated for different thresholds. Definitions of
- statistical measures are shown in Appendix A.

2.4 Meteorology and air quality of June-August 2013

- 18 The analyzed period was marked by three heat wave events, which contributed to the summer
- 19 characterized by high temperatures, sunny weather and lack of precipitation in Slovenia. The
- 20 first heat wave event with measured temperature daily maxima up to 35 °C occurred after a
- 21 rather cold beginning of the month and lasted from June 15 21. The event was terminated by
- a cold front passage and followed by the pronounced cold episode during the end of June and
- 23 the beginning of July. Another heat wave event with temperatures above 35 °C observed in
- 24 the lowland, started on July 26 and was briefly interrupted on July 29, when thunderstorms
- 25 related to frontal passage were accompanied by exceptionally strong wind gusts. The most
- 26 remarkable of three extraordinary hot episodes was recorded from August 1 8. On the last
- 27 day of this episode, August 8, temperatures reached 40 °C at some measuring sites in
- 28 Slovenia, and many of them observed their highest temperature ever recorded.
- 29 As expected for summertime conditions, measured concentrations of most air pollutants,
- 30 including PM10, were in general low during the analyzed time period. The only exception
- 31 was O₃ with exceedances of 8-hour target value (120 μgm⁻³) measured at all AQ monitoring

stations during the three heat wave events, which is the reason why the main focus of the 1 2 present study is on this pollutant. During the second two events (in July and August) also threshold exceedances of 1-hour daily maxima were recorded for O₃. In spite of the hot and 3 sunny conditions during the first heat wave event in June 2013, measured daily O3 maxima at 4 5 the Slovenian stations did not exceed the 1-hour information threshold value (1h ITV; 180 μgm⁻³), but reached 171 μgm⁻³ at the Mediterranean OTL and the elevated alpine KRV 6 stations. During the second heat wave event 1-hour daily maxima exceeded 180 µgm⁻³ at 7 8 KRV, OTL, NG and KP (July 23 - 28), while the highest number of 1-hour exceedances (20) 9 has been in July measured at OTL station. Similarly, during the August heat wave event O₃ concentrations exceeded the 1h ITV at LJ, MB, OTL, NG and KP from August 2 - 7. To 10 11 summarize, the Mediterranean stations (NG, OTL, KP) due to very high O₃ concentrations 12 measured during the heat wave events (especially the second two events) exhibited the 13 poorest AO in Slovenia during the analyzed time period, while the legislation limit values 14 have been exceeded only occasionally for the sub-alpine stations.

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3 Results and discussion

3.1 Evaluation of meteorological variables

- 18 Table 2 shows conventional statistical scores evaluating the 1-day WRF-Chem forecast for
- 19 the basic meteorological variables, 2m temperature (T2m; for hourly values and daily
- 20 maxima), 10 m wind speed (W10m), relative humidity (RH) and incoming solar radiation
- 21 (SR). Results for three selected measuring sites (LJ, NG, MS) and overall result for all 24
- MET monitoring sites (shown in Fig. 3) are presented separately.
- 23 Incoming solar radiation is the main energy source that drives all atmospheric processes,
- 24 including PBL processes, and has a critical role also in atmospheric chemistry. For almost all
- sites the mean SR was overestimated by the model, with an overall ME of 16 W/m² and 11
- 26 W/m² for 1-day and 2-day forecast, respectively. CORR was higher for 1-day (0.77) than for
- 27 2-day (0.71) forecast, with a range of 0.64 to 0.90 for 1-day forecasts at different stations. The
- 28 larger positive bias during the first day than for the second day can be attributed to less cloudy
- 29 conditions during the first day of simulation.

In the case of T2m 1-day (2-day) WRF-Chem meteorological forecast showed an overall 1 2 correlation with measurements of 0.93 (0.94) for all 1-hour values and 0.97 (0.96) for 1-hour daily maxima. With an exception of three alpine stations with higher simulated positive bias, 3 4 daily T2m maxima were simulated with ME between -3.9 °C and -0.6 °C, depending on 5 station spatial representativeness. All meteorological variables, including soil temperature and soil moisture, are always initialized with GFS data. This explains higher negative bias for 6 7 T2m during the first day of simulation in spite of the overestimated of solar radiation. An 8 average systematic underestimation of T2m daily maxima was -2.1 °C both for 1-day and 2-9 day forecast. Nighttime T2m minima showed lower systematic bias for 2-day forecast, which resulted in overall bias for all hourly T2m values of -1.3 °C for 1-day and -0.8 °C for 2-day 10 11 forecast. Predominant weak wind conditions with variable direction at stations located in 12 complex topography were challenging to simulate. The general model tendency was to 13 overestimate W10m with overall ME of 0.8 m/s for 1-day and 2-day forecast, where for some 14 stations bias can be very low (e.g. LJ; Tab. 2) and much higher for some other stations due to their local positioning in complex topography (e.g. HRA located in valley with ME of 1.9 15 16 m/s). For hourly values the correlation is lower (Tab. 2), but for mean daily W10m values 17 Pearson correlation coefficient between 0.4 and 0.9 has been simulated, depending on 18 monitoring site. Relative humidity shows slightly better results for 1-day than for 2-day 19 forecast with CORR of 0.77 and low overall ME of 2 % for 1-day forecast, which for particular stations can be positive (e.g. KRV) or negative (e.g. LJ, NG; Tab. 2). 20 21 Precipitation (RR) has an important role in cleansing of the atmosphere by wet deposition and 22 scavenging. On average, the predicted precipitation underestimated the measured 3-month 23 accumulations by -55 mm (1-day) or -8 mm (2-day forecast), where the station averaged predicted 3-month precipitation was 145 mm for 1-day, and 194 mm for 2-day forecast 24 25 (results not shown). It must also be taken into account that the 3.4.1 model version does not 26 allow to include the information about hydrometeors at the boundaries of the nested domain 27 (in the applied 1-way nesting procedure), which contributes to the negative simulated bias of 28 precipitation. A large decrease in the precipitation bias from day 1 to day 2 suggests that 29 different initialization methodology (e.g. using 1 day spin-up for meteorology) could improve 30 the prediction of precipitation events.

3.2 Evaluation of air quality variables

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2 In this section we evaluate WRF-Chem predictions for O₃, NO₂ and PM10, as three of the most problematic pollutants in terms of harm to human health and compliance with EU limit 3 4 values (EEA, 2012). Table 3 shows the domain wide performance statistics for 1-day and 2-5 day forecasts of these pollutants, where in the case of O₃ 1-hour and 8-hour averages and 6 daily maxima are analyzed separately. The comparison of 1-day and 2-day forecasts shows 7 that concentrations of air pollutants were somewhat better forecasted 1-day than 2-days ahead 8 by means of almost all of statistics shown in Tab. 3, with higher impact on O₃ predictions. 9 Although the 2-day prediction was generally not worse for the majority of meteorological variables, the reason for better 1-day prediction in the case of O₃ could be somewhat stronger 10 11 simulated winds on the second day of simulation. Stronger winds impact the transport and 12 dispersion of pollutants, and have the greatest consequence for secondary pollutants (like O₃) which need time to be formed. 13 14 As shown in Tab. 3 the WRF-Chem simulations tend to overestimate the 1-hour and 8-hour O₃ values with ME of 14.5 μgm⁻³ and 14.6 μgm⁻³, respectively. Looking at MAE, RMSE and 15 CORR statistics, agreement with measurements is better for 8-hour (22.6 µgm⁻³, 28.1 µgm⁻³ 16 and 0.69) than for 1-hour O₃ values (25.1 µgm⁻³, 32.1 µgm⁻³ and 0.65), which is in line with 17 results of previous studies (e.g. Tong and Mauzerall, 2013) and suggests that the current 18 19 modeling system has problems simulating the small-scale fluctuations of O₃. On the other hand evaluations of predicted 8-hour and daily O₃ maxima, which are of most concern, show 20 a nice model performance (ME, MAE RMSE and CORR of -2.7 µgm⁻³, 13.3 µgm⁻³, 16.7 21 μgm⁻³ and 0.81 for daily maxima, respectively), in line or even better than obtained in some 22 previous studies (e.g. Tong and Mauzerall, 2006; Chuang et al., 2011; Yahya et al., 2014), 23 24 which could be to some extent related to higher model resolution. 25 To understand results of the domain wide statistics (in Tab. 3) we further analyze spatial and temporal characteristics of model O₃ predictions. Figure 4 shows a spatial pattern of average 26 27 simulated 1-day predictions for O₃, NO₂ and PM10 overlaid with measured averages, where 28 in the case of O₃ results for all hourly values and for daily maxima are shown separately. 29 Examples of forecasted and measured time series for O₃ at different stations are shown in Fig. 30 5. In Fig. 4a the elevated alpine KRV station is the only one with high negative bias (-12 μgm⁻ ³) in forecasted 1-hour O₃ concentrations at the lowest model layer, which can be explained 31 32 by the too low altitude of the KRV station in model topography. The high negative bias for

hourly O₃ concentrations at KRV station is reduced to a value of only -2 µgm⁻³ by using the 1 2 5th model layer concentrations as explained in chapter 2.3. The 5th model level predictions will be used for KRV in all analyses that follow. Besides KRV also the Mediterranean KOP 3 4 and OTL stations, as well as the rural ZAV site, are stations with comparatively high measured nighttime O₃ levels, which results in low overall bias for all hourly O₃ values for 5 these stations (from -2 to -7 µgm⁻³). Namely, WRF-Chem model cannot capture well the 6 7 profound nighttime O₃ reductions (shown also by Žabkar et al., 2013; Im et al., 2014a), which 8 contributes to the overall over-prediction of hourly O₃ concentrations (from 10 to 36 µgm⁻³) 9 for stations with very low measured nighttime O₃ concentrations. For sites with highest positive bias in 1-hour O₃ concentrations (TRB, ZAG, HRA and ISK, with bias of 36 µgm⁻³, 10 31 µgm⁻³, 26 µgm⁻³ and 32 µgm⁻³, respectively), this can also be partly explained by too high 11 altitude of the stations in model orography (Tab. 1), since the mean O₃ concentration 12 13 increases with height. 14 Looking at O₃ daily maxima (Fig. 4b), the under-predictions occur at alpine KRV (-16 µgm⁻³ for the lowest model level shown in Fig.4) and at three Mediterranean stations (OTL, NG, 15 KOP; from -14 to -11 µgm⁻³). For Mediterranean stations the underestimations of daily 16 maxima are most probably due to inaccurate representation of costal processes in model, 17 18 which are crucial for PBL height evolution and accumulation of pollution in the near ground 19 air layers. For TRB station located in narrow valley of the very complex terrain that cannot be appropriately resolved in the current model topography, the model over-predicts O₃ daily 20 maxima for 14 μgm⁻³. For other sub-alpine stations the bias of O₃ daily maxima predictions is 21 22 lower. 23 To some extent the previously mentioned model over-predictions of nighttime O₃ minima 24 could be explained by model error in predicted NO₂ levels. When evaluating the primary 25 pollutants one must be aware that in the model the instantaneous emissions are spread over an 26 entire grid box, which results in underestimated emissions and concentrations close to the source regions and overestimated emissions and concentrations at rural locations adjacent to 27 the source regions, and can thus cause a combined effect of negative and positive biases at 28 urban and rural sites. Comparisons of WRF-Chem predicted NO₂ levels with measurements 29 30 show that in spite of the high spatial resolution the concentrations of the small urban areas are 31 insufficiently represented by the model (Fig. 4c). In Slovenia many towns are located in basins or very narrow valleys, usually poorly or even not resolved in model topography. 32

Smoothed local emissions for these towns show significant underestimations of NO₂ 1 2 concentrations (e.g. ZAG in Fig. 6). In combination with poorly reproduced meteorological 3 processes (calm and stable nighttime conditions in valleys and basins) this results in an 4 underestimation of the O₃ loss by titration. This can explain the positive nighttime bias of O₃ 5 found at these sites. The situation is better for bigger cities, located in wider basins, like LJ or CE (LJ; Fig. 6), while at rural sites NO₂ is either well simulated (e.g. MOH; Fig. 6), or 6 7 slightly over-predicted due to increased emissions from adjacent urban area (e.g. ZAD; Fig. 8 6). The overall agreement of hourly NO₂ predictions with measurements was good for rural 9 sites, while urban sites experienced under-predictions, which were highest for small cities, especially for NG (ME of -13 µgm⁻³) and ZAG (ME of -14 µgm⁻³). 10

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Also interesting to discuss are the results for predicted PM10 concentrations (Tab. 3 and Fig. 4d), showing slight over-prediction of daily PM10 levels at all stations which is somewhat surprising due to the fact that nearly all current off-line and on-line coupled chemical transport models show large systematic PM10 underestimations. For example, within AQMEII exercise, where seventeen modeling groups from Europe and North America were brought together, running eight operational online-coupled air quality models over Europe and North America, the rural PM10 concentrations over Europe were underestimated by all models (model configurations) by up to 66% while for the urban PM10 concentrations the underestimations were even much larger (up to 75%) (Im et al., 2014b). The reason for slight over-prediction of PM10 levels could be to some extent attributed to the high model spatial resolution used in our study. Further, CORR for daily PM10 concentrations is rather low (0.34 and 0.37 for 1-day and 2-day forecasts, respectively; Tab. 3), which is partly due to the low temporal dynamics of measured daily PM10 concentrations during the analyzed time period (no recorded PM10 exceeding), and partly due to the simulated PM10 overestimations during the heat wave events. These over-predictions contributed also to the overall positive bias of predicted PM10 levels. As shown in Fig. 7 for two monitoring sites, there was a significant PM10 over-prediction simulated on June 10 (day 8 in Fig. 7), related to the prefrontal advection of polluted air-masses coming from the north-western part of the domain D2 (coming from domain D1). The next significant PM10 over-prediction occurred during the first heat wave episode (June 17-22), when during the hot and low wind conditions (after June 17) the PM10 levels started to build up in the PBL over entire domain D2 (and over southwestern parts of domain D1), and reached the maximum concentrations in Slovenia again with prefrontal advection of polluted air masses. Both over-predictions contributed to

- an overall positive bias in forecasted PM10 concentrations. Detailed analyses showed that 1 2 high concentrations in domain D1 originated from boundary conditions, and appear to be a consequence of overestimated advection of Saharan dust in MOZART model predictions. The 3 increase in PM10 concentrations over Slovenia was also simulated during the prefrontal 4 5 advection related to the cold front which terminated the next two heat wave events in July and August (days 56-57 and days 67-68 in Fig. 7), but during these days predicted PM10 levels 6
- 7 were close to the measured PM10 concentrations.

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Evaluation and comparison of different methods for O3 daily maximum predictions

In this section we want to answer the question: "how accurate is the 1-hour O₃ daily maximum WRF-Chem forecast in comparison to the statistical model prediction or to persistence?". According to Zhang et al. (2012a) statistical models are known to be generally more suitable for complex site-specific relations between concentrations of air pollutants and predictors. With appropriate and accurate predictors they have a higher accuracy as compared to deterministic models, which is, along with their computational efficiency their main advantage (Zhang et al., 2012a). Among the strengths of the deterministic models are that they give prognostic time- and spatially-resolved concentrations under typical and atypical scenarios, and can give scientific insights into pollutant formation processes (Zhang et al., 2012a). Furthermore, they also allow forecasts for locations which are not monitored due to their complete spatial coverage. In spite of simplified descriptions of physical and chemical processes in the deterministic models and inaccuracies and uncertainties in model inputs (in particular the emissions), some previous studies already suggested that deterministic models can also have skills close to statistical forecasting tools (e.g. Manders et al., 2009). In addition to evaluation and comparison of O₃ daily maxima predictions with WRF-Chem and the statistical model, we decided to add a persistence model as a threshold for useful model prediction. Persistence works well under stationary conditions, but because it cannot handle changes in weather and emissions, fails at the beginning and at the end of the episodes (Zhang et al., 2010a). Regarding the extremes, models of all types are known to have problem to accurately predict them, while persistence predicts extremes with a 1-day (2-day) time lag. Figure 8 compares discrete statistics site by site for 1-day and 2-day model predictions of 1-

- hour O₃ daily maxima. Similarly, Tab. 4 shows these statistics for all data with different 31
- thresholds applied (only for WRF-Chem and persistence, because a statistical forecast is not 32

available for all stations), and separately for different types of stations (sub-alpine urban, 1 rural, Mediterranean urban) with an available statistical forecast, Looking at ME persistence 2 gives results close to zero as long as no threshold is applied, while with threshold of 140 µgm⁻ 3 ³ (Tab. 4) ME of 1-day persistence (-10.2 μgm⁻³) is very close to the WRF-Chem model for 1-4 day predictions (-11.2 μgm⁻³), and for 2-day predictions WRF-Chem (-13.8 μgm⁻³) already 5 beats persistence (-19.4 μgm⁻³). Site-by-site comparison (Fig. 8) shows that for most stations 6 7 the statistical forecast has a lower ME than WRF-Chem forecast, but there are also stations 8 (ISK, HRA, LJ, KRV) with lower or equal ME for WRF-Chem than for statistical model, 9 indicating the possible occurrence of atypical conditions not resolved by the statistical model. 10 Looking at MAE and RMSE, at all stations except those with highest ME (TRB, KOP) WRF-11 Chem outperforms the persistence already in the 1-day forecast. Among sites with available statistical forecast there are only two (OTL, KOP) with WRF-Chem performing worse than 12 13 the statistical forecast. CORR is one of the parameters that suggest how much the model is able to follow the true nature of processes regardless the possible bias. For almost all stations 14 WRF-Chem shows higher CORR than persistence for 1-day and 2-day forecasts. Only at the 15 16 KRV station the 1-day statistical forecast (CORR=0.80) somewhat outperforms WRF-Chem 17 (0.74), and at NG and KOP CORR for WRF-Chem and statistical model is very similar. 18 The Taylor diagrams in Fig. 9 show CORR together with the centered root-mean-square 19 difference (RMSD) between model forecasts and observations, and the amplitude of their 20 variations (standard deviation). The ideal model would have a correlation coefficient of 1 and 21 a standard deviation equal to the observations, which means that it would be co-located with 22 the black dot on the diagram. WRF-Chem gives higher CORR and lower RMSD for all types 23 of stations, while standard deviation of WRF-Chem O₃ daily maxima predictions is underestimated and lower than for other model forecasts. The latter shows that the variability 24 25 in WRF-Chem model predictions is not as large as that in observed values. MNBE in Fig. 8 26 has very similar results to ME. For all forecasts except WRF-Chem for the TRB site (with 27 MNBE of 16%) which is located in a narrow valley that is not resolved in the current model 28 resolution, MNBE is below the ± 10 -15%, which is the U.S. EPA (US EPA, 1991) 29 recommended threshold for the models used for regulatory applications. For MNGE the U.S. 30 EPA recommendation below 30-35% for O₃ applications is met by all forecasts, even in the 31 case of 2-day persistence model. With exception of the MS and KOP sites MNGE is lower for 32 WRF-Chem than for statistical forecast, while for KOP and KRV sites 1-day persistence gives best results, followed by the statistical forecast or WRF-Chem. Very similar are results for 33

IOA with the range of 0-1, and score 1 indicating perfect model agreement with the 1 2 observations. We can conclude that for most stations the WRF-Chem predictions are in line or 3 even outperform the statistical model. With the exception of the stations with high bias due to very complex local topography (TRB) or unresolved coastal processes (KOP), the WRF-4 5 Chem forecasts are more accurate than persistence. Here we recall that high negative bias in WRF-Chem forecast for alpine KRV site due to too low altitude of the station in model 6 topography was compensated by taking prediction from the 5th model level. 7 8 The key requirement for a forecast system is to be able to predict O₃ concentration levels 9 greater than a given threshold. Thus, in addition to the discrete evaluation just presented, also 10 the contingency-table-based statistics are an important metric of forecast performance. Table 11 5 summarizes the categorical evaluation results for three different thresholds (120, 140, 160 μgm⁻³) of elevated O₃ levels, which pose a greater risk to human health. Namely, it is 12 important to take into account that results of categorical statistics are very sensitive to the 13 14 threshold chosen, as well as to the overall pollution levels during the analyzed months. 15 Equitable Threat Score (ETS) measures the fraction of observed and/or correctly predicted events, adjusted for the frequency of hits that would be expected to occur by random chance. 16 17 Although this score takes into account the climatology it is not truly equitable. It ranges from -1/3 to 1, where the minimum value depends on climatology (it is near 0 for rare events). 18

Looking at Tab. 5 ETS shows equal skill for WRF-Chem and statistical forecast, higher than persistence for the 120 µgm⁻³ threshold (1-day and 2-day forecast). ETS decreases with increasing the threshold for both WRF-Chem and statistical forecast, indicating the challenge that both models have to accurately predict the extremes. In the case of 140 µgm⁻³ threshold, WRF-Chem has the same ETS as persistence, higher than the statistical model for 1-day forecast, while for 2-day forecast WRF-Chem outperforms the statistical model, followed by persistence. In the case of 160 µgm⁻³ threshold persistance has the highest ETS for a 1-day forecast, followed by statistical model and WRF-Chem, while in the case of 2-day predictions, statistical model shows the highest skill and WRF-Chem the lowest. Another measure, the critical success index (CSI), is similar to ETS, except that it does not take into account the climatology of the events and thus gives poorer scores for rarer events. It measures the percentage of cases that are correctly forecasted out of those either forecasted or observed, and ranges from 0 to 1 (1 indicating the perfect forecast). Similar as ETS, CSI gives higher scores for persistence in the case of 1-day forecast for the higher two thresholds, while on the second day WRF-Chem or the statistical model already performs better. Bias (B)

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- determines whether the same fraction of events are both forecasted and observed. A tendency
- 2 of the statistical model and of WRF-Chem to under-predict O₃ threshold exceedances shows
- 3 as a B below 1 for these two models. The false alarm ratio (FAR) that measures the
- 4 percentage of forecast high O₃ events that turn out to be false alarms, gives highest skill for
- 5 WRF-Chem, followed by statistical model and persistence. The probability of detection
- 6 (POD) is a measure of how often a high threshold occurrence is actually predicted to occur,
- 7 and is relatively low for WRF-Chem with respect to other models.
- 8 It must be noted, that in categorical evaluations systematic biases like those obtained with
- 9 WRF-Chem for some stations (e.g. KOP), significantly impact the model performance. For
- 10 example, if KOP station was excluded from categorical evaluations, WRF-Chem performance
- improved by means of all statistical measures (results not shown). If correction techniques,
- based on observations and the previous day's forecast (e.g., McKeen et al., 2005, 2007; Kang
- et al., 2008) were to be applied to correct the systematic biases, WRF-Chem forecasts might
- outperform the other two models even in categorical evaluations.

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4 Summary and conclusion

- 17 A high resolution modelling system based on an on-line coupled WRF-Chem has been
- applied for numerical weather prediction and for forecasting air quality in Slovenia. In the
- 19 study the evaluation of the forecasting system has been conducted for three summer months.
- 20 Since the selection of physical or chemical parameterization schemes influences and possibly
- 21 changes the outcomes, we decided to apply schemes which are well documented and have
- previously been used in other applications (e.g. AQMEII). Both 1-day and 2-day predictions
- of meteorological and air quality variables have been analyzed. The focus has been on O₃ as
- 24 the only pollutant with recorded exceedances of legislation limit values during the three heat
- wave events in June, July and August 2013. WRF-Chem daily O₃ maximum predictions have
- also been compared to the operational statistical model and persistence forecasts to answer the
- 27 question how skillful are the WRF-Chem model predictions compared to these two models.
- 28 1-day and 2-day WRF-Chem PM10 forecasts showed a very low bias. Exceptions were two
- events with significantly over-predicted PM10 levels due to prefrontal advection of polluted
- 30 air masses from neighboring regions. Knowing that majority of the current chemical transport
- 31 models show large negative biases in simulated PM10 concentrations, these results present a

- 1 good starting point for studying the importance of aerosol feedbacks with realistic model
- 2 aerosol concentrations, left for future research.

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- 3 The overall agreement of WRF-Chem NO₂ forecast with measurements was good for rural
- 4 sites, while urban sites experienced model under-predictions, which were highest for small
- 5 towns. One important reason is that many small towns are located in basins or very narrow
- 6 valleys, usually poorly presented in model topography. Smoothed local emissions result in
- 7 model underestimations of NO₂ concentrations for these towns. This in combination with
- 8 insufficiently reproduced calm meteorological conditions in basins and valleys during the
- 9 nighttime hours explains also WRF-Chem over-predictions of nighttime O₃ concentrations.
 - Evaluations of predicted 1-hour and 8-hour daily O₃ maxima, which are in the case of this pollutant of the highest interest, show good WRF-Chem model performance. Nevertheless, there are also stations which experience high over- or under-predictions of O₃ daily maximum levels. For Mediterranean sites the under-predictions of the daily maxima are most probably due to inaccurate representation of costal processes in model, which are crucial for the PBL height evolution and accumulation of pollution in the near ground air layers. For some subalpine stations the reason for the higher bias in O₃ daily maximum predictions is their location either at elevated mountainous or coastal regions, or in narrow valleys which cannot be appropriately resolved in the current model resolution - that impacts how accurately model simulates the local processes responsible for the level of local pollution. Comparisons of WRF-Chem O₃ daily maximum forecasts with persistence and with statistical model predictions show that with respect to some statistical parameters the deterministic WRF-Chem forecast can outperform the other two for both 1-day and 2-day predictions. For example, correlation coefficient shows highest skill for WRF-Chem model, confirming the importance of complex processes as taken into account in an on-line coupled Eulerian model. Further improvement of WRF-Chem forecasting skill could be obtained by applying one of the bias-correction methods in order to account for unresolved topographical and coastal effects, as well as emission patterns. Chemical data assimilation, although currently still in its infancy for online coupled meteorology-chemistry models (Bocquet et al., 2014), could in future also be used as an efficient method for improving prediction of chemical concentration fields. For WRF-Chem model a technical note on the implementation of the aerosol assimilation and a guidance for prospective users has been recently published by Pagowski et al. (2014).

1 Appendix A: Statistical measures

- 2 For i-th observed (O_i) and the corresponding modelled (M_i) value of variable, discrete
- 3 statistical measures are calculated as follows:
- 4 Mean error:

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$$ME = \frac{1}{N} \sum_{i=1}^{N} (M_i - O_i)$$

7 Mean absolute error:

$$8 \qquad MAE = \frac{1}{N} \sum_{i=1}^{N} \left| M_i - O_i \right|$$

10 Root mean square error:

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

13 Correlation coefficient:

16 Index of agreement:

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$$IOA = 1 - \frac{\sum_{i=1}^{N} (M_i - O_i)^2}{\sum_{i=1}^{N} (M_i - \overline{O}) + |O_i - \overline{O}|^2}$$

20 Mean normalized bias error:

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$$MNBE = \frac{1}{N} \sum_{i=1}^{N} \frac{M_i - O_i}{O_i} \times 100$$

24 Mean normalized gross error:

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$$MNGE = \frac{1}{N} \sum_{i=1}^{N} \frac{|M_i - O_i|}{O_i} \times 100$$

For categorical evaluation all model predictions are first classified into four groups (*a*, *b*, *c* and *d*):

a prediction is above, but observation is below the threshold

b prediction and observation are above the threshold

c prediction and observation are below the threshold

33 d prediction is below, but observation is above the threshold

Categorical statistics are calculated as follows:

- Equitable threat score: $ETS = \frac{b a_r}{a + b + d a_r}$, where $a_r = \frac{(a + b)(b + d)}{a + b + c + d}$
- 3 Critical success index: $CSI = \frac{b}{a+b+d}$
- 4 Bias: $B = \frac{a+b}{b+d}$

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- 5 False alarm ratio: $FAR = \frac{a}{a+b}$
- 6 Probability of detection: $POD = \frac{b}{b+d}$

Acknowledgements

- 9 Centre of Excellence for Space Sciences and Technologies SPACE-SI (OP13.1.1.2.02.0004)
- 10 is part financed by the European Union, European Regional Development Fund and Republic
- of Slovenia, Ministry of Education, Science and Sport. The authors thankfully acknowledge
- 12 TNO for providing the TNO-MACC-II anthropogenic emissions. Statistical model predictions
- and measurement data used in the study were kindly provided by Slovenian Environmental
- 14 Agency and Electroinstitute Milan Vidmar. The support through COST Action ES1004
- 15 EuMetChem is gratefully acknowledged.

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Table 1: AQ monitoring sites.

Monitoring site	Abbreviation	Type of zone	Altitude (m)	Model orography (m)	Model analysis height (m)	Pollutants
Celje	CE	Urban	240	300	313	O ₃ , PM10, NO ₂
Hrastnik	HRA	Urban	290	540	552	O_3 , SO_2
Iskrba	ISK	Rural	540	579	591	O ₃ , NO ₂
Koper	KOP	Urban	56	72	85	O ₃ , PM10
Kovk	KOV	Rural	608	516	528	NO_2
Krvavec	KRV	Rural	1740	1272	1414	O_3
Ljubljana	LJ	Urban	299	287	300	O ₃ , PM10, NO ₂ ,
Murska Sobota	MS	Rural	188	189	202	O ₃ , PM10, NO ₂
Nova Gorica	NG	Urban	113	150	163	O ₃ , PM10, NO ₂
Otlica	OTL	Rural	918	874	886	O_3
Sv. Mohor	МОН	Rural	394	254	266	NO_2
Trbovlje	TRB	Suburban	250	459	471	O ₃ , PM10, NO ₂
Velenje	VEL	Urban	389	461	474	O_3 , SO_2
Vnajnarje	VNA	Rural	630	468	480	NO_2
Zadobrova	ZAD	Rural	280	275	287	PM10, NO ₂
Zagorje	ZAG	Urban	241	431	443	O ₃ , PM10, NO ₂
Zavodnje	ZAV	Rural	765	678	690	O ₃ , NO ₂

Table 2: Statistical scores for 1-hour values of 2m temperature (T2m), 10 m wind speed (W10m) and relative humidity (RH), and for daily average incoming solar radiation (SR). Shown are results for 1-day forecast, calculated separately for three measuring sites (LJ, NG, MS) and for 24 MET monitoring stations (ALL) during the 3-month period. In the case of temperature results for daily maxima are also shown.

Variable	Station	NoCases	Mean	ME	MAE	RMSE	CORR
T2m 1h (°C)	LJ	2129	20.3	-1.6	2.3	2.9	0.91
	NG	2184	21.8	-1.1	2.1	2.5	0.94
	MS	2184	19.2	-2	2.3	2.8	0.95
	ALL	47836	18.7	-1.3	2.3	2.9	0.93
T2m max (°C)	LJ	89	26.5	-1.6	1.8	2.1	0.98
	NG	90	26.8	-3	3	3.3	0.96
	MS	90	26.2	-1.7	1.8	2	0.98
	ALL	1976	24.2	-2.1	2.7	3.2	0.97
W10m (m/s)	LJ	2129	1.5	0	0.7	1	0.58
	NG	2183	2.7	1	1.4	1.9	0.35
	MS	2184	2.3	0.4	1.1	1.4	0.53
	ALL	43378	2.4	0.8	1.4	1.9	0.36
RH (%)	LJ	2066	62	-2	8	10	0.85
	NG	2121	62	-1	12	15	0.75
	MS	2121	69	3	8	11	0.88
	ALL	48556	68	2	11	14	0.77
SR (W/m2)	LJ	90	276	19	31	43	0.84
	NG	90	278	4	32	43	0.77
	MS	90	273	15	26	37	0.9
	ALL	1710	273	16	35	49	0.77

Table 3: Domain wide performance statistics for 1-day and 2-day forecast in μgm^{-3} . For different pollutants statistics for all hourly (hour), 8-hour averages (8h), 8-hour daily maximum (8h max), daily maximum (max) or daily average (day) concentrations are shown.

		NoCases	Mean	ME	MAE	RMSE	CORR
O ₃ (hour)	1 day	28391	94.8	14.5	25.1	32.1	0.65
	2 day	28391	95.0	14.5	25.5	32.5	0.64
O ₃ (8h)	1 day	28072	94.8	14.6	22.6	28.1	0.69
	2 day	28072	95.0	14.6	23.0	28.5	0.68
O ₃ (8h max)	1 day	1157	111.5	-0.1	13.2	16.5	0.77
	2 day	1157	111.6	-0.2	13.7	17.0	0.75
O ₃ (max)	1 day	1170	116.5	-2.7	13.3	16.7	0.81
	2 day	1170	116.6	-3.1	14.0	17.5	0.78
NO ₂ (hour)	1 day	26178	7.3	-5.1	7.5	10.8	0.3
	2 day	26178	7.5	-4.9	7.6	10.8	0.3
PM10 (day)	1 day	718	29.0	7.1	12.0	18.8	0.34
	2 day	718	29.1	7.2	12.0	19.1	0.37

Table 4: Discrete evaluation of 1-hour daily maximum ozone predictions.

Stations	Threshold,	Forecast	Mean	ME	MAE	RMSE	CORR	MNBE	MNGE	IOA
	NoCases		(μgm ⁻³)	(µgm ⁻³)	(μgm ⁻³)	(μgm ⁻³)		(%)	(%)	
All	>0	F 1day	116.5	-2.6	13.3	16.7	0.81	-0.05	11.7	0.86
	1170	F 2day	116.6	-3.1	14.0	17.5	0.78	-0.1	12.3	0.84
		PER 1day	119.5	-0.4	15.8	21.1	0.65	1.6	14.5	0.81
		PER 2day	119.8	-0.4	21.7	27.7	0.39	2.8	19.6	0.65
	> 140	F 1day	144.1	-11.2	15.2	17.9	0.52	-6.8	9.5	0.57
	1102	F 2day	141.4	-13.8	16.5	19.4	0.42	-8.6	10.5	0.48
		PER 1day	145.0	-10.2	15.6	19.6	0.41	-6.5	10.0	0.52
		PER 2day	135.8	-19.4	24.76	29.2	0.31	-12.4	15.9	0.38
Sub-alpine	>0	F 1day	115.3	1.1	10.7	14.0	0.84	3.4	11.1	0.91
urban with SF	180	F 2day	115.4	0.8	12.0	15.2	0.80	3.5	12.2	0.88
(LJ, HRA)		PER 1day	114.3	-0.3	16.7	21.7	0.64	2.2	16.5	0.80
		PER 2day	114.6	-0.3	21.9	27.8	0.41	3.9	21.6	0.65
		SF 1day	114.0	-0.5	11.9	15.7	0.81	1.6	11.2	0.88
		SF 2day	116.2	0.6	13.4	17.1	0.75	3.2	12.7	0.84
Rural with SF	>0	F 1day	117.6	-5.6	13.3	16.3	0.80	-3.0	10.8	0.86
(MS, ISK,	360	F 2day	117.4	-6.4	14.2	17.4	0.76	-3.4	11.4	0.84
KRV, OTL)		PER 1day	123.6	-0.3	15.0	20.7	0.65	1.4	13.1	0.81
		PER 2day	124.1	-0.4	21.6	27.8	0.37	2.4	18.5	0.64
		SF 1day	121.5	-2.9	15.0	19.4	0.74	-0.7	12.2	0.83
		SF 2day	122.9	-1.8	15.8	20.5	0.67	0.5	13.2	0.79
Mediterranean	>0	F 1day	123.5	-11.8	17.4	22.5	0.76	-6.9	12.5	0.80
urban with SF	179	F 2day	124.5	-11.2	17.2	21.8	0.77	-6.5	12.4	0.82
(KOP, NG)		PER 1day	135.9	-0.5	17.4	23.0	0.68	1.2	13.8	0.83
		PER 2day	136.0	-0.2	25.2	31.5	0.41	2.8	19.7	0.66
		SF 1day	129.3	-7.0	15.9	20.7	0.75	-3.6	11.6	0.83
		SF 2day	131.6	-4.5	15.6	20.4	0.74	-1.6	11.6	0.84
	<u> </u>	<u> </u>					<u> </u>			

1 Table 5: Categorical evaluation of 1-hour daily maximum ozone predictions for different

2 thresholds, calculated for 8 monitoring sites with available statistical forecast.

Threshold	Forecast	ETS	CSI	В	FAR	POD	a	b	c	d
> 120	F 1day	0.42	0.63	0.81	0.13	0.70	39	253	313	107
	F 2day	0.39	0.61	0.79	0.14	0.68	41	245	303	115
	PER	0.31	0.59	0.99	0.25	0.74	91	267	249	93
	1day									
	PER 2day	0.17	0.49	1.00	0.34	0.65	123	235	209	124
		0.42	0.67	1.02	0.21	0.01	<i>c</i> 7	257	242	<i>c</i> 1
	SF 1day	0.42	0.67	1.02	0.21	0.81	67	257	243	61
	SF 2day	0.38	0.65	1.03	0.23	0.80	77	264	225	66
> 140	F 1day	0.40	0.50	0.64	0.15	0.551	19	111	490	92
	F 2day	0.37	0.47	0.66	0.19	0.53	25	108	476	95
	PER	0.40	0.53	1.00	0.31	0.69	62	141	435	62
	1day									
	PER	0.19	0.35	1.00	0.48	0.52	97	106	391	97
	2day									
	SF 1day	0.30	0.43	0.73	0.29	0.52	40	99	398	91
	SF 2day	0.30	0.43	0.70	0.27	0.51	37	98	403	94
> 160	F 1day	0.19	0.22	0.38	0.34	0.25	10	19	626	57
	F 2day	0.17	0.20	0.34	0.35	0.22	9	17	619	59
	PER	0.40	0.45	1.00	0.38	0.62	29	47	595	29
	1day									
	PER	0.22	0.28	1.00	0.56	0.43	43	33	572	43
	2day									
	SF 1day	0.23	0.27	0.49	0.35	0.32	13	24	539	52
	SF 2day	0.25	0.29	0.63	0.41	0.37	19	27	540	46

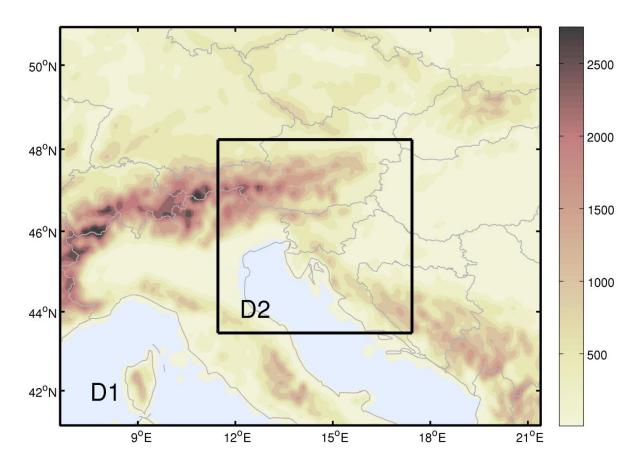


Figure 1: Modelling domains (D1, D2) used in WRF-Chem RT-AQF system. Orography (in meters) is shown in resolution of D1 domain (11.1 km).

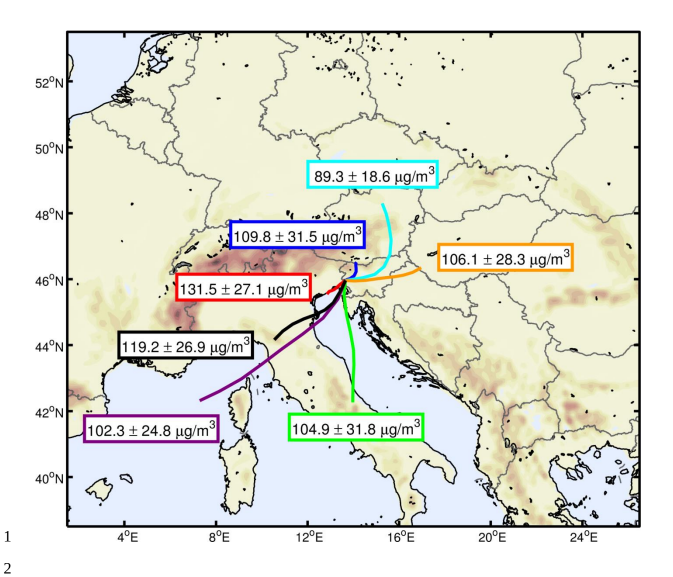


Figure 2: Example of ozone analysis for the Nova Gorica (NG) monitoring site (average daily maximum ± standard deviation) for 7 clusters of similar trajectories, as used in the statistical ozone daily maximum forecast for the NG station.

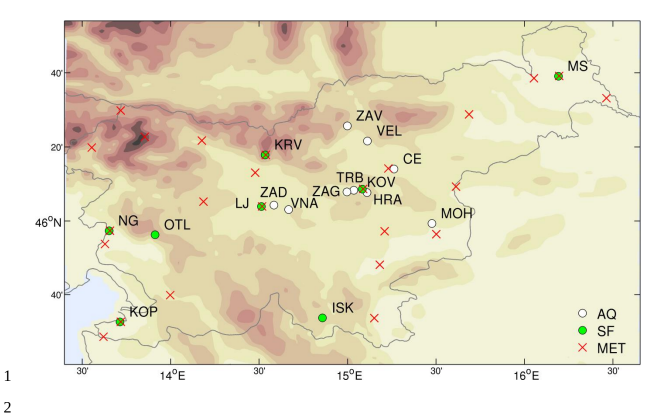


Figure 3: Locations of monitoring stations used in evaluation of air quality variables (AQ stations; shown are also station abbreviations) and meteorological variables (MET stations). Green dots indicate measuring sites with available ozone daily maximum statistical forecast (SF). For the meaning of abbreviations of AQ sites see Tab. 1.

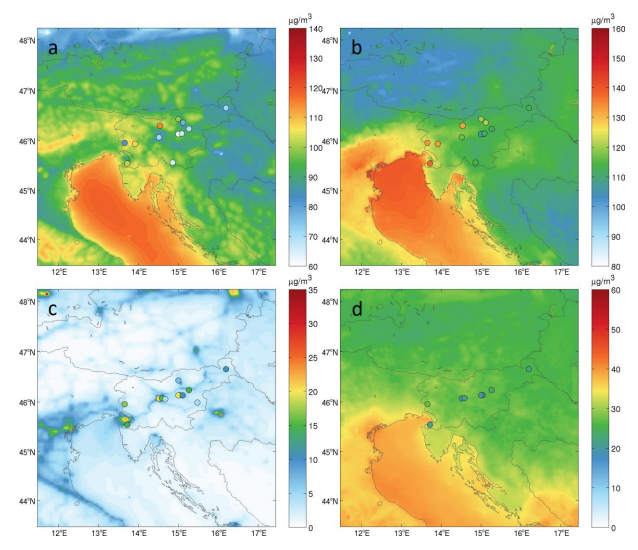


Figure 4: 3-month average 1-day predictions of a) hourly O₃, b) O₃ daily maximum, c) hourly NO₂, and d) daily PM10concentrations for the first model layer, overlaid with measurements.

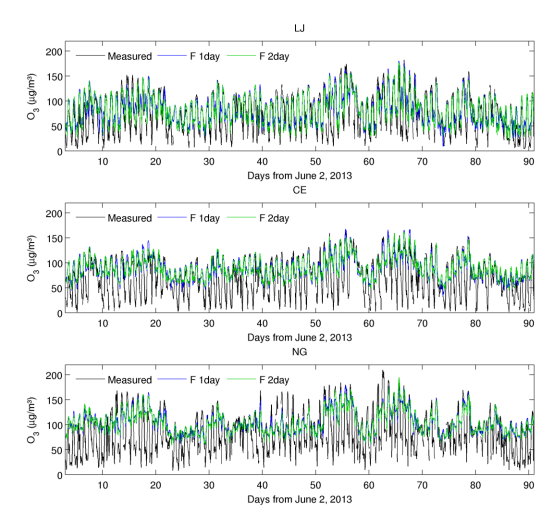
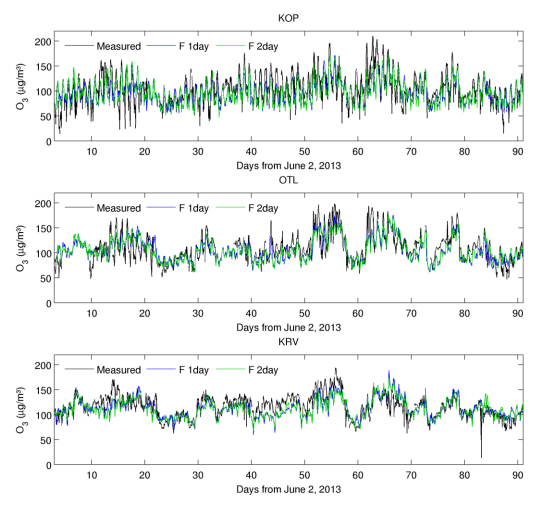
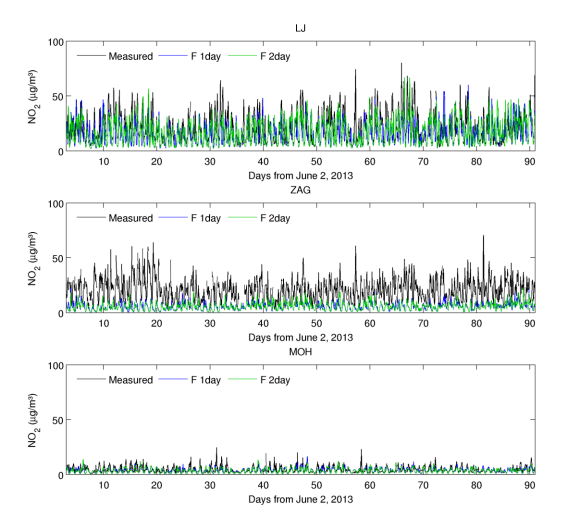


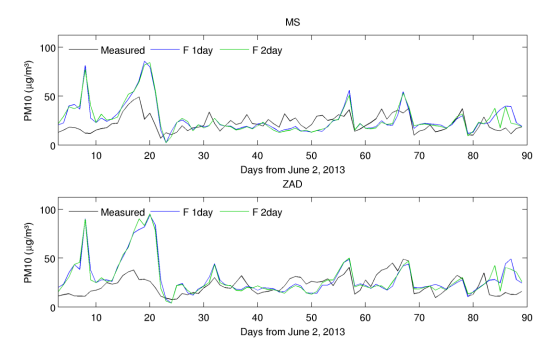
Figure 5: Time evolution of hourly ozone concentrations for 1-day (F 1day) and 2-day (F 2day) WRF-Chem predictions and measurements for some stations during the 3-month period. (continued)



1 2 Figure 5: (continued)



2 Figure 6: The same as Fig. 5 but for NO₂ at LJ, ZAG and MOH stations.



2 Figure 7: The same as Fig. 5, but for daily PM10 concentrations at MS and ZAD stations.

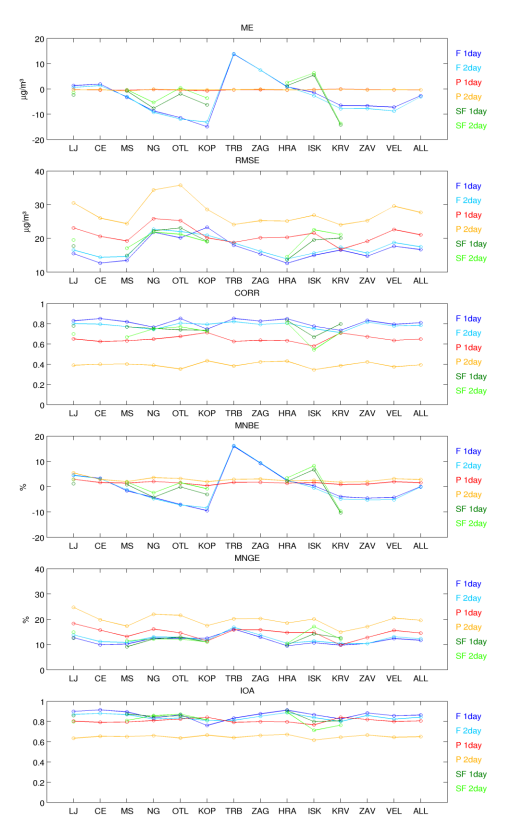


Figure 8: Site-by-site comparison of discrete statistics for 1-day and 2-day WRF-Chem (F 1day, F 2day), statistical (SF 1day, SF 2 day) and persistence model (P 1day, P 2day) predictions of ozone daily maxima during the 3 analyzed summer months.

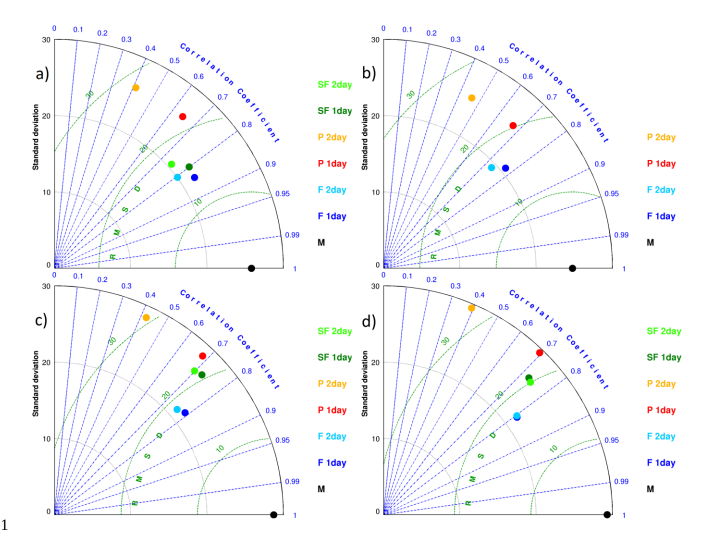


Figure 9: Taylor diagrams comparing 1-day and 2-day ozone daily maximum statistical forecast (SF), persistence (P) and WRF-Chem forecast (F) for a) sub-alpine urban stations with SF (LJ, HRA), b) sub-alpine urban stations without SF (CE, TRB, ZAG, VEL), c) rural stations with SF (MS, ISK, KRV, OTL) and d) Mediterranean urban stations (NG, KOP).