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Air quality across a European hotspot

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6	AIR QUALITY ACROSS A EUROPEAN
7	HOTSPOT: SPATIAL GRADIENTS,
8	SEASONALITY, DIURNAL CYCLES AND
9	TRENDS IN THE VENETO REGION, NE ITALY
10	
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27 ABSTRACT

28 The Veneto region (NE Italy) lies in the eastern part of the Po Valley, a European hotspot for air pollution. Data for key air pollutants (CO, NO, NO₂, O₃, SO₂, PM₁₀ and PM_{2.5}) measured over 7 29 years (2008/2014) across 43 sites in Veneto were processed to characterise their spatial and 30 temporal patterns and assess the air quality. Nitrogen oxides, PM and ozone are critical pollutants 31 frequently breaching the EC limit and target values. Intersite analysis demonstrates a widespread 32 pollution across the region and shows that primary pollutants (nitrogen oxides, CO, PM) are 33 significantly higher in cities and over the flat lands due to higher anthropogenic pressures. The 34 spatial variation of air pollutants at rural sites was then mapped to depict the gradient of background 35 pollution: nitrogen oxides are higher in the plain area due to the presence of strong diffuse 36 anthropogenic sources, while ozone increases toward the mountains probably due to the higher 37 levels of biogenic ozone-precursors and low NO emissions which are not sufficient to titrate out the 38 photochemical O₃. Data-depth classification analysis revealed a poor categorization among urban, 39 traffic and industrial sites: weather and urban planning factors may cause a general homogeneity of 40 41 air pollution within cities driving this poor classification. Seasonal and diurnal cycles were investigated: the effect of primary sources in populated areas is evident throughout the region and 42 drives similar patterns for most pollutants: road traffic appears the predominant potential source 43 shaping the daily cycles. Trend analysis of experimental data reveals a general decrease of air 44 pollution across the region, which agrees well with changes assessed by emission inventories. This 45 study provides key information on air quality across NE Italy and highlights future research needs 46 and possible developments of the regional monitoring network. 47

49	Keywords:	air pollution,	nitrogen	oxides,	particulate ma	utter, V	/eneto, trend	s.
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53 1. INTRODUCTION

Since the mid-90s, the European Community has adopted increasingly stringent standards for 54 abating emissions and for improving the air quality. The main steps in the legislative process were 55 56 the Framework Directive 96/62/EC, its subsequent daughter Directives and the more recent Directive 2008/50/EC. As a consequence, a general improvement of air quality has been recorded in 57 the last decade. However, current and future EU standards are still breached in some European 58 59 regions, the so-called hotspots, e.g., Northern Italy, Benelux and some Eastern Countries (Putaud et al., 2004;2010). Under this scenario, the development of additional successful strategies for 60 emission mitigation and the implementation of measures for air quality control are two major 61 62 questions addressed by policy makers and in scientific research, respectively.

63

Following the implementation of the EC Directives, local and national authorities are required to monitor air quality. Measurement data are primarily managed by local agencies to assess the extent of air pollution, check if standards are complied with, and, in case of the exceeding of Limit Values or even lower assessment thresholds, to inform the population about potential health impacts. Beyond their original regulatory purpose, such data also represent a valuable resource: if historical data are available, long-term trends can be investigated to obtain real feedback upon the successes and failures of past and current mitigation strategies.

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The Po Valley (N Italy) is neither a city nor an administrative unit, but it can be considered a
megacity (Zhu et al., 2012). Currently, it is one of the few remaining and most worrying European
hotspots: high levels of hazardous pollutants are commonly recorded over a wide area (~48·10³
km²) hosting ~16 million inhabitants. The anthropogenic pressure and some peculiar
geomorphological features (a wide floodplain is enclosed by the Alps and Apennine mountains)
lead to frequently breaches of limit and target values imposed by EU Directives (EEA, 2016) for
nitrogen oxides, ozone and particulate matter (PM).

The Veneto Region (Figure 1) is located in the eastern part of the Po Valley and extends over 79 $\sim 18.4 \cdot 10^3$ km² ranging from high mountain environments (29% of the territory), to intermediate hill 80 zones (15%), large and flat plain areas (56%) and ~95 km-long coastlines. From an administrative 81 82 point of view, it is subdivided in 7 Provinces: Belluno (BL), Treviso (TV), Vicenza (VI), Venice (VE), Padova (PD) and Rovigo (RO). Heavy anthropogenic pressures are present almost 83 continuously: a total of ~ $4.9 \cdot 10^6$ inhabitants are resident in some large cities (> $2 \cdot 10^5$ inhabitants: 84 Verona, Padova and Venice-Mestre) and in a number of minor towns and villages which form a 85 86 continuum "sprinkled" network of urban settlements. Consequently, emissions from road traffic and domestic heating are spread throughout the region. Some industrial areas are also present, mainly 87 88 close to the main cities and with distinctive features and different plant types. In addition, the Veneto lies in a strategic location linking Central, Eastern Europe and continental Italy: a dense 89 network of international E-roads and transport hubs attract large amounts of road and intermodal 90 traffic, mostly heavy duty diesel-powered trucks. A large percentage of the region includes 91 agricultural fields, mostly located in the plain (intensive farming) and hilly areas (vineyards, 92 93 orchards), while rural environments are present mainly in hilly and mountain regions. A land use map is provided in Figure 1: this composite landscape inevitably makes the emission scenario of the 94 region extremely complex and its spatial variations quite unpredictable. 95

96

This study aims to examine and describe the spatial variations, temporal trends and seasonality of 97 air quality across the Veneto Region over 7 years (2008-2014). Datasets used in this study include 98 99 the mass concentration of key air pollutants as required by European air quality standards: nitrogen oxides (NO+NO₂=NO_x), ozone (O₃), carbon monoxide (CO), sulphur dioxide (SO₂) and PM with 100 aerodynamic diameter less than 10 μ m (PM₁₀) and less than 2.5 μ m (PM_{2.5}). Air quality data were 101 measured by ARPAV (Veneto Environmental Protection Agency) through a well-established 102 103 network of 43 sampling stations (ARPAV, 2014) covering a large portion of the territory (Figure 1). A series of chemometric procedures are used to assess the extent of air pollution, to verify the 104

effectiveness of site categorization and to find patterns commonly recorded across the Veneto or identify sites with anomalous pollution levels. The gradients of concentrations are depicted and seasonal/diurnal/weekly patterns are investigated. The long-term trends are seasonally decomposed and then processed to find the general orientations and drifts in air pollution over sites with a different categorization. Furthermore, long-term trends are coupled with changes in the emission inventories to verify if estimated emissions match with experimentally recorded levels of key air pollutants across the region.

112

113 2. MATERIALS AND METHODS

114 2.1 Sampling sites

The map of sites is shown in Figure 1, while Table 1 summarises their general characteristics and 115 measured pollutants. Figure 1 also shows the political, relief and land use maps. Sites are identified 116 by the initials for the province and their category. Sites were selected to fulfil some specifications: 117 (i) data availability must cover at least four years in 2008/14; (ii) sites must be representative of 118 119 most important pollution climate scenarios, such as citywide pollution, regional background or traffic and industrial hotspots and (iii) sites must be representative of differing environments 120 (mountain, hilly, plain, coastal areas). In particular: 121 • At least one site was selected as rural background (RUR sites) for each Province (total 9 sites). 122

i.e. in areas not directly influenced by trafficked roads and/or urban and industrial settlements. In
particular, BL.RUR and VI.RUR are located in remote locations at high altitudes. RUR sites are
fundamental to assess the background gradients at the regional scale;

- Eighteen sites were categorised as suburban (SUB) and urban (URB), i.e. broadly representative
 of citywide levels of air pollutants;
- A total of 8 sites were selected as representative of traffic hotspots (TRA) and placed at kerbside
 locations in cities experiencing heavy traffic and/or frequent road congestion events;

130	• Eight sites were set as representative of main indust	trial (IND) areas. Each site has peculiar
131	1 characteristics. VE.IND is located downwind of Por	rto Marghera, one of the main industrial
132	2 zones in Italy extending over $\sim 12 \text{ km}^2$ and including	g a large number of different installations
133	3 (thermoelectric power plants burning coal, gas and	refuse derived fuels, a large shipbuilding
134	4 industry, an oil-refinery, municipal solid waste inci	nerators and many other chemical,
135	5 metallurgical and glass plants). VI.IND1 and VI.IN	D2 are representative of small and medium-
136	6 sized tannery industries, PD.IND1 and PD.IND2 ar	e set in an area potentially affected by a
137	7 municipal solid waste incinerator plant, PD.IND3 w	vas selected to monitor the fall-out from a
138	8 steel mill and VR.IND and PD.IND4 are representa	tive of emissions from cement plants.

139

140 2.2 Experimental

141 All selected sites are equipped with fully automatic analysers set to collect data on hourly (gaseous pollutants) or hourly/bihourly bases (PM₁₀ and PM_{2.5}). QA/QC of measurements is guaranteed by 142 ARPAV internal protocols, which fully comply with the standards required by EC Directives in-143 force: EN 14626:2012 for CO, EN 14211:2012 for NO, NO₂, and NO_x, EN 14212:2012 for SO₂, 144 EN 14625:2012 for O₃. CO, NO_x, O₃ and SO₂ instruments were calibrated every day. Hourly- or 145 146 bihourly- resolved PM₁₀ and PM_{2.5} were measured with beta gauge monitors: validation 147 experiments were routinely conducted between gravimetric (EN 14907:2005) and automatic methods; several tests were also performed routinely (at least 1 test every week) to keep a constant 148 149 check on the beta gauge samplers. Pairs of filters were measured with both methods and the results 150 were checked to ensure that they are within the variation margins imposed by the technical protocols adopted in UNI EN 12341:2001. Some sites were not equipped with hourly PM monitors 151 152 (Table 1), but may provide daily gravimetric-measured PM₁₀ levels. In this case, PM was collected by low-volume samplers on filters and the mass concentration was measured by gravimetric 153 determination (EN 14907:2005) at constant temperature (20±5°C) and relative humidity (RH, 154

50±5%). Consequently, no diurnal patterns are investigated in such sites, but only interannual and
seasonal trends.

157

158 2.3 QA/QC and data handling

Data have been validated by ARPAV through a well consolidated internal protocol (ARPAV, 2014) 159 and according to the European standards. The full dataset was therefore used for exploratory 160 161 statistics. However, the aim of this study is to detect the general behaviour of air pollution and some clearly identified high pollution episodes which occurred in 2008/14 across the region. Examples 162 are the burning of a thousand folk fires on the eve of Epiphany (Masiol et al., 2014a) or fireworks 163 164 for Christmas and other local celebrations. Consequently, preliminary data handling and clean-up are carried out to check the datasets for robustness, outliers and anomalous records. For this 165 purpose, data greater than the 99th percentile were included for exploratory analysis but not in the 166 trend estimation. Data were also adjusted to account for the shift in anthropogenic emissions due to 167 the changes between local time (UTC+1) and daylight savings time (DST). This latter correction 168 169 helps in investigating daily patterns of anthropogenic emission sources.

170

171 Data were analysed using R (R Core Team, 2016) and a series of supplementary packages,

including 'openair' (Carslaw and Ropkins, 2012; Carslaw, 2015), 'PMCMR' (Pohlert, 2015) and

173 'localdepth' (Agostinelli and Romanazzi, 2011;2013).

174

175 2.4 Data depth classification analysis

A classification analysis was used to check the accuracy of the site categorisation, i.e., to verify whether the sampling sites in a category (RUR, SUB+URB, TRA, IND) are characterised by a general homogeneity in air pollutant levels. This task is accomplished by applying a new classification technique based on statistical data depth (DD). This technique is well reviewed elsewhere (Mosler and Polyakova, 2012 and the reference therein) and recently was extended to 181 functional and multivariate data (Ramsay and Silverman, 2006; Lopez-Pintado and Romo, 2009;

182 Lopez-Pintado and Romo, 2011; Claeskens et al., 2014; Cuevas, 2014).

183

184 The depth of a point relative to a given dataset measures how deep the point lies in the data cloud, i.e. it measures the centrality of a point with respect to an empirical distribution (Mosler and 185 Polyakova, 2012). DD provides an order to the observations and the rank system provided by DD 186 can be used to perform unsupervised and supervised classification analysis. A statistical depth 187 function should be invariant to all (non-singular) affine transformations; it reaches the maximum 188 value at the centre of symmetry for symmetric distribution and it becomes negligible when the norm 189 190 of the point tends to infinity. Another important property is ray-monotonicity, i.e. the statistical depth does not increase along any ray from the centre (Liu, 1990; Zuo and Serfling, 2000). The 191 classical notion of data depth has been extended to functional data and the different available 192 implementations aim to describe the degree of centrality of curves with respect to an underlying 193 probability distribution or a sample. Some advisable properties of functional depths are the (semi-) 194 continuity, the consistency, and the invariance under some class of transformations, which tends to 195 vanish when the norm of a curve tends to infinity (Mosler and Polyakova, 2012). 196

197

A classification analysis was performed by using the DD-plot tool and a functional multivariate 198 simplicial depth. First, a (empirical) simplicial depth of a *point* y with respect to a *set of points* x =199 $(x_1, ..., x_n)$ in the Euclidean space of dimension p is introduced. $S_i = S(x_{i1}, x_{i2}, ..., x_{in}, x_{in+1})$ represents 200 the simplex obtained using p+1 points in the sample. A simplex of p+1 points in a space of 201 dimension p is the convex hull of those points. Then $d(y; x_n)$ is the fraction of simplicials S_n 202 contains y overall combinations of the p+1 indices $i_1, i_2, ..., i_p, i_{p+1}$ in the set 1,..., n. This is a 203 consistent estimator of the simplicial depth, which is the probability that a random simplex $S(X_1, X_2, X_3)$ 204 ..., X_p , X_{p+1}) will cover the *point* y when X_1 , X_2 , ..., X_p , X_{p+1} are identical and independent copies of 205 206 the random variable X from where the sample is drawn.

Thus, we can consider a set of (multivariate) curves $x_i(t)$, (j=1, ..., J) measured at time $t_1, t_2, ..., t_N$, 207 i.e. $x_i(t_i)$ is a point in the Euclidean space of dimension p. In our context, $x_i(t)$ is the multivariate 208 time series at location *j* (sampling site) for a set of *p* pollutants concentration measured at time *t*. 209 210 Following Claeskens et al. (2014), we defined the depth of a given curve y(t) according to the sample $x = (x_1(t), ..., x_n(t))$, denoted d(y(t), x) as the weighted average of the empirical simplicial 211 depth evaluated at any given time $t_1, t_2, ..., t_N$. The weights proportional to the fraction of available 212 213 observations at each time is then set. This is important to be considered for the presence of missing 214 values.

215

The classification is performed using the depth-versus-depth plot (DD-plot) (Li et al., 2012; Cuevas, 216 2014). Considering two groups of curves $x=(x_1(t), ..., x_J(t))$ and $z=(z_I(t), ..., z_K(t))$, the goal is to 217 classify a curve y(t) in one of these. Hence, we evaluate d(y(t),x) and d(y(t),z) that represents the 218 depth of y(t) according to x and to z. We assign y(t) to the first group if d(y(t),x) > d(y(t),z) and to 219 the second group otherwise. In particular, two groups of curves will be well separated if $d(x_i(t),x) >$ 220 $d(x_i(t),z)$ for each observation x_i and $d(z_k(t),x) < d(z_k(t),z)$ for each observation z_k . Finally, we can 221 plot on a Cartesian axis the points $(d(x_i(t),x), d(x_i(t),z)), j=1,..., J$ and the points $(d(z_k(t),x), d(z_k(t),z)), d(z_k(t),z))$ 222 k=1,...,K, i.e. the DD-plot. Curves (sampling sites) well classified in the first group will be plotted 223 at the right bottom corner of the graphics, whereas curves well classified in the second group will 224 appear at the left upper corner. Curves equally well classified in both groups will lie around the 225 226 bisector line of the plot.

227

In this study, four site categories are used to group the sites: rural (RUR), urban and sub-urban (URB), traffic (TRA) and industrial (IND). For each couple of categories, the DD-plots are evaluated on the basis of distributions of one or more pollutants.

231

233 **3. RESULTS**

A summary of data distributions during the whole study period (all available data) is provided as boxplots in Figure 2 and maps in Figures SI1. Concentrations are expressed as mass concentration; NO_x is expressed as NO₂, as required by EU standards. Moreover, Table SI2 reports annual average concentration and exceedances for NO₂, O₃, PM₁₀ and PM_{2.5}. CO and SO2 are not included because no exceedances were observed.

239

240 3.1 Carbon monoxide and sulphur dioxide

In Veneto, CO and SO₂ are not critical pollutants. CO values are well below the EC limit value and
WHO guidelines (WHO, 2000), i.e. 10 mg m⁻³ as daily maximum over eight hours. The highest
hourly average levels of CO over six years were recorded at VR.URB1, PD.IND2, VR.TRA3,
PD.URB, VE.TRA1 and PD.IND1 (all ~0.6 mg m⁻³); remaining sites showed concentrations
between 0.2 and 0.6 mg m⁻³.

246

 SO_2 does not show exceedances of European standards both for hourly averages (350 µg m⁻³) and 247 for 24-h averages (125 μ g m⁻³). The 2008/14 average concentrations across Veneto were very low, 248 ranging from ~0.6 μ g m⁻³ (often below instrumental detection limits) at VI.URB3 and RO.SUB to 249 \sim 4 µg m⁻³ at some sites near Venice (VE.IND, VE.URB1). The higher levels in Venice sites can be 250 likely associated to harbour and industrial emissions (i.e. thermal power plant, oil refinery and 251 municipal solid waste incinerator). While the contribution of the harbour to the levels of SO₂ in 252 Venice is still debated (Contini et al., 2015), the role of industrial emissions is well supported by the 253 emission inventories. In 2005, about 76% of overall SO₂ emissions in the Veneto were released in 254 the Province of Venice, of which 69% (19,742 Mg y^{-1}) came from combustion in energy and 255 transformation industries (ARPAV, 2011). 256

257

259 3.2 Nitrogen oxides

When considering the short-term metrics, NO₂ does not represent a special risk for human health in 260 Veneto: the 1-hour alert threshold of 400 μ g m⁻³ was never breached, while the 1-hour average of 261 $200 \ \mu g \ m^{-3}$ not to be exceeded more than 18 times over a calendar year was exceeded only at 262 VR.TRA3 in 2008. However, NO_x becomes pollutants of concern when looking at the long-term 263 metrics. The minimum average levels were found at two rural sites located in high mountain 264 environments, while maxima were measured at a traffic site in Verona located very close to a 265 logistic intermodal freight transport hub spanning over 4 km² and moving 26 million tons of goods 266 annually (VR.TRA3). The average concentrations of NO varied from 0.5-0.6 μ g m⁻³ (VI.RUR, 267 BL.RUR1) to 50 μ g m⁻³ (VR.TRA3), while NO₂ ranged from 4 μ g m⁻³ (BL.RUR1) to 53 μ g m⁻³ 268 (VR.TRA3) and NO_x from 5 μ g m⁻³ (BL.RUR1) to 130 μ g m⁻³ (VR.TRA3). Comparing results 269 averaged over 2008/14 with the annual EC limit value for NO₂ (40 μ g m⁻³ averaged over one year), 270 the limit was substantially exceeded at six traffic sites (VE.TRA2, VE.TRA1, VI.TRA, VR.TRA2, 271 PD.TRA, VR.TRA3). 272

273

274 3.3 Ozone

In rural sites, average O₃ levels ranged from 48 μ g m⁻³ (RO.RUR) to more than 90 μ g m⁻³ (BL.RUR1, VI.RUR), while urban and suburban sites varied between ~40 and ~60 μ g m⁻³. Despite the low number of sites measuring ozone in traffic and industrial environments, it is evident that polluted environments generally exhibit the lower average concentrations: the minimum levels were recorded at VR.TRA3 (36 μ g m⁻³), which, inversely, shows the higher NO concentrations (it leads close to a large logistic intermodal freight transport hub and, thus, it is affected by heavy traffic of diesel-powered trucks).

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Ozone is the subject of several regulations: the alert threshold (maximum 1-hour level of 240 μ g m⁻³) was never breached in 2008-2012, but was exceeded five times in 2013 at PD.RUR1. This

result recalls the anomaly of PD.RUR1 already reported for PM₁₀-bound polycyclic aromatic
hydrocarbons (Masiol et al., 2013). Despite PD.RUR1 being originally located far from direct
emission sources, it probably suffers from a local unknown source of air pollution. In addition, it is
located between and equidistant from three main urban settlements (Mestre, Padova and Treviso).
Further studies should be carried out to detect the potential sources; in the meanwhile, its
categorisation should be revised.

The information threshold (180 μ g m⁻³ over one hour) was frequently exceeded at most of the sites. VI.RUR deserves special attention because it is affected annually by 39–126 exceedances; it is located in a remote area (1366 m a.s.l.) characterised by grass- and wood-lands and is not affected by direct anthropogenic sources. Consequently, high O₃ levels may be linked to the transport of air masses containing ozone or ozone-precursors from the nearby highly populated plain areas or to the local biogenic emission of ozone-precursors from plants. In addition, since this area is not impacted by traffic, titration of photochemically produced ozone by primary NO is negligible.

299

The EC long-term target value of $120 \ \mu g \ m^{-3}$ and the WHO air quality guideline of $100 \ \mu g \ m^{-3}$ measured as maximum daily 8 hour running averages (not to be exceeded more than 25 days over 3 years for the EC target) were also frequently breached at almost all the sites. Similarly, the long term objective value for the protection of vegetation (AOT40) calculated over the warm period (May to July) is also amply breached at all the rural sites, posing a serious risk to high-quality agriculture promoted by regional policies.

306

307 It is therefore evident that ozone is a critical pollutant across the Veneto. The standards for ozone 308 are more difficult to achieve across the warmer regions (southern Europe) because of the larger 309 magnitude of summer photochemical O₃ episodes. In addition, O₃ levels are also influenced 310 substantially by the extent of reactions with local NO emissions, which are currently falling in

Europe (Colette et al., 2011) in order to comply with the increasingly stringent emission standards required for road vehicles. As a result, recent decreases in NO emissions across Europe may limit reaction with O_3 , which is the main sink of O_3 in the polluted atmosphere.

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- 315

3.4 NO_x partitioning and total oxidants (OX)

Recent changes in emission trends for some air pollutants and the complex photochemistry of the 316 317 NO-NO₂-O₃ system are further evaluated through two derived parameters. The partitioning of nitrogen oxides was investigated through the NO₂/NO_x ratio (Figure 2 and Figure SI1b) and total 318 oxidants (OX=NO₂+O₃, expressed as ppb) are often used to describe the oxidative potential (Kley et 319 320 al., 1999; Clapp and Jenkin, 2001). Results are highly variable, but higher ratios are generally recorded at rural sites because of the lower primary NO emissions. The average levels of OX show 321 little variation across the Region with distributions typically showing interquartile ranges between 322 25 and 50 ppby. The highest site averages are seen at rural sites due to photochemical ozone 323 creation, and at traffic sites, presumably caused by high emissions of primary NO₂. 324

325

326 3.5 Particulate matter

PM₁₀ and PM_{2.5} are critical air pollutants in Veneto. Average PM₁₀ levels over 2008/14 varied from 327 less than 20 μ g m⁻³ in BL.RUR2 and VR.RUR to more than 40 μ g m⁻³ in VI.URB1 and VI.TRA. 328 Some sites breached the European annual limit value of 40 μ g m⁻³ (PD.IND3, VI.URB1, PD.URB, 329 VR.TRA2, VE.TRA1, PD.TRA and PD.IND1), except in 2013 and 2014 when no exceedances 330 were recorded. In such sites, the annual average concentrations are usually close to the European 331 limit value: this way, small fluctuations in PM10 levels may have a large effect in marking them as 332 "fulfilling" or "not fulfilling" the EC standard. However, every year more than 20 sites exceeded 333 the daily mean of 50 μ g m⁻³ for more than 35 times in a calendar year. Among these sites, 334 VI.URB1, VE.TRA2, VE.IND and VE.URB3 showed the highest number of daily exceedances for 335 a minimum of 77, 66, 64 and 60 times in a calendar year, respectively. 336

The $PM_{2.5}$ monitoring network started its operation in 2009, when the Directive 2008/50/EC entered into force, and continued to grow over the following years. This way, only 8 sites provide sufficient data to be included in this study. The annual limit value (25 µg m⁻³) was frequently breached (five years out of six) at PD.URB, PD.IND1, PD.IND2, VE.IND and VI.URB1. Similar to PM_{10} , in 2013 only 6 stations breached the annual average value, while no exceedances were recorded in 2014.

343

344 4. DISCUSSION

345 4.1 Differences among sites

Maps showing the average levels of recorded pollutants over the 2008/14 period are reported in Figures SI1a,b for different site categories. Levels of CO are generally low across the region and do not show any evident trend among site categories. On the contrary, nitrogen oxides, SO₂, PM_{10} and $PM_{2.5}$ show levels increasing from rural to urban to traffic sites, while concentrations in industrial sites are highly variable and reflect the specific characteristics of each site (Figure 2). Such patterns are opposite to ozone (higher levels at rural sites in high mountain environments and lower at sites polluted with primary emissions).

353

Motor vehicles are major sources of NO (Keuken et al., 2012; Kurtenback et al., 2012) and a series 354 of volatile organic compounds (VOCs) (Gentner et al., 2013). Munir et al. (2012) have shown that 355 emissions from cars, buses and heavy vehicles have strong effects on urban decrements of ozone 356 levels. Due to this, the lower O3 levels in most anthropogenically affected sites in Veneto are related 357 to the primary emissions of NO. This fact is further well supported by the partitioning of nitrogen 358 359 oxides (Figure SI1b): lower NO₂/NO_x ratios are recorded at urban and hot-spot sites, indicating that high relative concentrations of NO lead to ozone depletion. Another reason is linked to the impact 360 of natural sources of VOCs, such as biogenic isoprene and terpenes, i.e. known effective ozone-361 362 precursors (Duane et al., 2002). The land cover map (Figure 1) shows that the hilly and mountain

areas of N Veneto are covered by forests. Thus, biogenic VOCs are expected to be elevated in rural and mountain environments and enhance the generation of ozone. In this context, modelling studies in Europe indicate that biogenically-driven O_3 accounts for ~5% in the Mediterranean region (Curci et al., 2009).

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The Kruskal-Wallis analysis of variance by ranks (KW_{test}) was applied as a global non-parametric 368 369 test for depicting statistically significant inter-site variations. The null hypothesis is rejected for 370 p < 0.05, meaning that the sites in a category are statistically different. In this case, the post-hoc test after Nemenyi (Demšar, 2006) was applied for multiple sample comparison to point out the pairs of 371 372 sites which differ significantly in the pollutant level. Generally, results indicated that air pollutants are uniformly distributed across the region: no pairs of sites differ in the levels of CO, while only 373 the two remote RUR sites (BL.RUR1 and VI.RUR) present statistically significant differences for 374 NO_x and ozone from a large number of other sites. The application of comparison tests for PM_{10} is 375 complicated by the lack of data at the two remote sites and by the availability of differently time 376 377 resolved data (hourly and daily). Despite such limitations, results indicate that only BL.RUR2 exhibits significantly lower concentrations than other sites. 378

379

380 4.2

4.2 Differences amongst site categories

KW tests indicated that the concentrations of many pollutants measured across the Veneto are 381 statistically similar even at sites which are categorised differently (except for the two sites located 382 in extremely remote mountain areas). Moreover, the results point to an important conclusion: air 383 pollutants are almost uniformly distributed across the region. However, these results also indicate 384 385 that there is an apparent homogeneity amongst sites, i.e. sites having different categorisation may experience statistically similar levels of air pollutants. This hypothesis was tested statistically using 386 data depth (DD) analysis. However, there are some limitations to its application: the important 387 388 fraction of missing data for some pollutants/sites and the lack of the full set of pollutants at most

sites limits the possibilities for its use. Therefore, data-depth classifications were separately performed for: NO, NO₂ and NO_x, (ii) PM_{10} , (iii) CO. Since few TRA and IND sites measured O₃ and SO₂, the DD analysis was not possible for these pollutants because not all the site categories would be adequately represented. Application to one or a few pollutants has the disadvantage of processing pollutants separately; however, it also has the advantage that it ensures the presence of a reasonable high number of sites, which make the results more robust and the spatial extent of the analysis more extensive.

396

The DD classification for nitrogen oxides (Figure 3) shows that most of RUR and URB sites are 397 398 generally well separated, i.e. they are plotted far from the 1:1 bisecting line. Rural sites are also well separated from traffic and industrial ones. However, some exceptions are found. BL.SUB, 399 VI.URB3, VI.URB4 and RO.SUB are more similar to RUR than to URB sites, i.e. NO_v levels are 400 generally lower than other urban sites, as also confirmed by boxplots in Figure 2. On the contrary, 401 PD.RUR1 and RO.RUR lie on the URB-side of the DD plot, i.e. they are more comparable to URB 402 than RUR sites. Although the anomalously high levels of air pollution at PD.RUR1 were already 403 recognised and discussed (Masiol et al., 2013), this result also shows that two rural sites probably 404 405 experience relatively high levels of NO_x with respect to remaining RUR sites.

406

Although levels of pollutants in sites categorized as rural generally differ from other categories,
there is not a clear separation among URB, TRA and IND sites. In fact, the DD classification
analysis has demonstrated that most of the TRA and IND sites can be grouped with URB sites
(points lay around the bisector of the plot). DD-plots of PM₁₀ also show this behaviour, while no
clear classification was possible using CO (Figure SI2 and SI3), i.e. CO levels are quite similar at
all site categories.

EU directives assume that URB sites are representative of the exposure of the general population, while TRA and IND sites should be representative of areas where the highest concentrations may occur. However, results show that this condition is rarely fulfilled in Veneto. In this context, a recent position paper (JRC-AQUILA, 2013) has pointed out the need to implement criteria for improving the classification and representativeness of air quality monitoring stations in Europe. There are several reasons that may explain failure of the classification:

• The widespread distribution of emissions. Since the flat areas of the Po Valley host a 'sprinkled' 420 continuum of urban settlements with different sizes, densities and uses (Romano and Zullo, 421 2015), the rather constant presence of anthropogenic emissions leads to a widespread distribution 422 of emission sources. As a consequence, air pollutants emitted by different sources mix in the 423 atmosphere and the limited atmospheric circulation in Po Valley further limits their dispersion. 424 The similarity of URB and TRA sites probably results from urban planning. Most large cities 425 have ancient origins and contain medieval or even Roman city centres with historic buildings 426 427 and narrow streets. A rapid and intense urbanization was experienced in Italy after World War II with fast build-up of large urban/suburban areas all around the ancient city centres, often without 428 a well-informed approach to urban planning. Consequently, some city configurations 429 inevitability limit the movement of road traffic and are characterised by busy streets which are 430 frequently congested during rush hour periods, i.e. when traffic is widespread over the city 431 432 centres and not properly channelled into main orbital or bypass roads. The high density of emissions leads to a high level of pollution from road traffic across cities and to the consequent 433 similar levels of pollutants between URB and TRA sites. 434

The poor differentiation between URB and IND sites may result from the lack of large industrial zones (except in VE). Most of Veneto cities host small and medium-sized industries in several sectors: glass, cement, food products, wood and furniture, leather and footwear, textiles and clothing, gold jewellery, chemistry, metal-mechanics and electronics. Large industrial zones are only present in VE (Porto Marghera). In addition, during the last 20 years, a large number of

companies have relocated their plants abroad, while the global financial crisis (2007/9) has 440 overwhelmed the industrial and economic sectors with a consequent decrease in the production 441 of goods and/or the collapse, closure or downsizing of many companies/industries. Today, 442 443 small/medium sized industrial areas are scattered across the region, with many of the more significant industries located close to major cities. As a result, some IND sites can be more 444 affected by urban sources than industrial emissions. For example, Porto Marghera, the main 445 industrial area of Veneto, lies SE of the city of Mestre. VE.IND (representative of Porto 446 447 Marghera) is located just east of the industrial area (Figure SI4). During winter frequent temperature inversions are responsible for the build-up of air pollutants (mostly NO_x, PM) and 448 449 VE.IND lies just downwind of the urban area of Mestre under prevailing wind regimes (wind rose in Figure SI4), while it is rarely downwind of the industrial area. Consequently, in winter 450 VE.IND is likely to be more affected by the plume from the urban area than by the industrial 451 emissions. This hypothesis is also confirmed by previous studies (Squizzato et al., 2014; Masiol 452 et al., 2012; 2014b), which reported similar concentrations of PM-bound species between 453 454 VE.IND and the city centre of Mestre. Further modelling studies are needed to inform enhancements to the monitoring network and to better sample the industrial emissions. 455

456

457 4.3 Spatial gradients

The characteristics of anthropogenic pressures and the peculiar topography strongly influence the 458 pollutant distribution: this make the quantification of pollutant gradients very challenging. RUR 459 sites can be used as good estimators for determining gradients across the region because they can be 460 considered as representative of the regional pollution as suggested by Lenschow et al. (2001) for 461 PM: they are located away from large sources and are also fairly uniformly distributed across the 462 region. Average concentrations at RUR sites have therefore been spatially interpolated to 463 investigate the gradients of background levels of air pollutants across the region. Semi-variograms 464 465 were investigated, showing that all pollutants do not all have the same direction, and ordinary

kriging was selected as the best model to interpolate the data. In this study kriging analysis merely 466 aims to shape the background gradients because it was applied over a limited number of points (9). 467 Maps for pollutants recorded at more than six sites (NO_x, O₃, OX) are provided in Figure 4, while 468 the standard errors of predictions are shown in Figure SI5. All variables show marked gradients. 469 Nitrogen oxides increase from the mountain environments in the north to the coastal plain areas; 470 however the direction of maximum slopes in the increases differ slightly: NO from NW to SE, 471 while NO₂ is from NNW to SSE. The gradient for ozone is opposite to NO, with maximum 472 concentrations at high mountain rural sites and minima in coastal areas. The clear anti-correlation 473 between the gradients of NO and O₃ clearly depicts their relationship: at rural sites, NO emissions 474 475 from anthropogenic sources are low and therefore, the reaction with ozone is its main sink, and the main sink for ozone. Due to the low concentrations of NO₂ at RUR sites when compared to ozone, 476 gradients for OX are similar to those of ozone. 477

478

479 **4.4** Seasonal patterns

The monthly-resolved distributions of air pollutants are reported in Figure 5. No evident seasonal patterns are found for the two high-mountain sites (BL.RUR1 ad VI.RUR). At the remaining sites, all the pollutants except SO₂ exhibit clear seasonal cycles. CO, NO, NO₂, NO_x, PM₁₀ and PM_{2.5} show significantly higher levels in colder months (KW_{test} at p<0.05). Rapid increases occur between mid-September and December; falls in concentration occur between March and mid-April. This pattern can be attributed to the interplay of some covariant causes:

• The lower mixing layer heights in winter, which limit the dispersion of pollutants emitted

locally. The typical planetary boundary layer height in the Po Valley is 450 m in winter, and

rises up to 1500-2000 m in the warmer months due to the thermal convective activity (Di

489 Giuseppe et al., 2012; Bigi et al., 2012).

- Ambient temperature controls the gas-particulate phase partitioning of semivolatile compounds
- 491 (ammonium nitrate and part of organic carbon). In Veneto nitrate accounts for $0.1-0.5 \ \mu g \ m^{-3}$

492	(0.7–2.5% of PM _{2.5} mass) in August and for 5–10 μ g m ⁻³ (16–25%) in February (Masiol et al.,
493	2015) and organic carbon in $PM_{2.5}$ varies from about 2.6 μ g m ⁻³ in June and 11.4 μ g m ⁻³ in
494	December (Khan et al., 2016).
495	• Increased emissions in the coldest months mainly driven by increasing energy demand for
496	domestic heating. Domestic heating is regulated at national level: generally, the switching on is
497	fixed to occur on 15 October, while the switching off is on15 April, i.e. when the fastest changes
498	occur. However, such dates can change according to the weather. Wood smoke from domestic
499	heating may contribute appreciably to higher winter concentrations of PM.
500	• the drop of actinic fluxes in winter and the consequent reduction of hydroxyl radical, ozone, and
501	the oxidative activity which is a sink for many pollutants.
502	
503	Ozone shows a strong seasonal pattern. Ozone shows the highest values during the warmest period
503 504	Ozone shows a strong seasonal pattern. Ozone shows the highest values during the warmest period (generally Apr–Sep), when the solar radiation is higher (Figure 5) and the atmospheric
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504 505 506	(generally Apr–Sep), when the solar radiation is higher (Figure 5) and the atmospheric photochemistry is more active. Generally, ozone levels reach approx. 20 μ g m ⁻³ in winter, but never drop below 50 μ g m ⁻³ at RUR sites located in high mountain environments (BL.RUR1, VI.RUR,
504 505 506 507	(generally Apr–Sep), when the solar radiation is higher (Figure 5) and the atmospheric photochemistry is more active. Generally, ozone levels reach approx. 20 μ g m ⁻³ in winter, but never drop below 50 μ g m ⁻³ at RUR sites located in high mountain environments (BL.RUR1, VI.RUR, VR.RUR >800 m above sea level). The patterns of OX are dominated by ozone both in rural and

510

Sulphur dioxide lacks a clear seasonal pattern. Industrial emissions are expected to be quite constant through the year, but highest emissions of SO_2 are may occur in summer due to an increase in energy production from coal power plants to meet air conditioning demand. However, the higher mixing layer height and the enhanced photochemistry driving S(IV) to S(VI) conversion lead to similar concentrations to the cold period.

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518

519 4.5 Daily and weekly patterns

Figure 6 reports the diurnal cycles, while Figure SI6 shows the weekly patterns. As for the seasonal 520 521 patterns, daily cycles are the result of interplays among the strength of sources, photochemical processes and weather factors. Although minor changes due to peculiar local conditions are found, 522 523 almost all the sites exhibit rather similar daily and weekly cycles, except at the three high mountain sites. CO and nitrogen oxides show typical daily cycles linked to road traffic at almost all the sites, 524 525 with two daily peaks corresponding to the morning and evening rush hours (7–9 am and 6–8 pm). The morning and evening peaks are split by a minimum, which is assumed to be the result of: (i) 526 527 lower emissions (less traffic); (ii) larger availability of ozone driven by the daylight photolysis of NO_x and the oxidation of VOC and CO (iii) higher convective activity leading to a deeper mixed 528 layer, which enhances the atmospheric mixing. Weekly patterns are also linked to road traffic: 529 generally, average levels increase from Mondays to Thursdays, while a fast drop is measured during 530 weekends, when road traffic reach minimum volumes and heavy duty vehicles over 7.5 tonnes are 531 532 subject to some limitations (over the whole weekends in summer and on Sundays in winter).

533

Ozone and total oxidants (OX) show evident daily peaks in the mid-afternoon, i.e. the hours 534 experiencing the higher solar irradiation, and lower levels are experienced between 6 and 9 am local 535 time (DST time-corrected in summer). These patterns are also enhanced in summer due to the 536 higher solar irradiation. It is evident that daily peaks of OX are delayed 2-3 hours with respect to 537 ozone, corresponding to the increased NO to NO₂ oxidation and primary NO₂ emissions in evening 538 rush hours. The weekly patterns are the mirror images of CO and NO_x with higher concentrations 539 540 during the weekends, following the drop of fresh anthropogenic emissions. However, such marked diurnal patterns are not found at the three high mountain sites, which show rather constant and high 541 levels throughout the day. This "flatter" pattern has also been observed at other high-mountain sites 542 543 in the Apennines (Cristofanelli et al., 2007) and Alps (Vecchi and Valli, 1998) and is likely related

to the lack of anthropogenic sources of freshly emitted ozone-precursors and the presence of higher
levels of biogenic ozone-precursors, which do not follow anthropogenic cycling. However, the
levels of ozone in high mountain environments are also known to be strongly affected by (i) the
transport of polluted air masses by local wind systems (valley and slope winds) from the Po Valley
and surrounding cities inside the Alps (Kaiser et al., 2009), Foehn wind events (Seibert et al., 2000),
large scale synoptic air pollutant transport (Wotawa et al., 2000) and stratospheric inputs
(Vingarzan, 2004). Nocturnal dry deposition is also less effective at mountain sites.

551

Daily cycles of SO₂ are quite different thorough the region and between site categories and show 552 553 two different patterns. Most sites present NO_x-like patterns, i.e. two weak peaks corresponding to the morning and evening rush hours. On the contrary, all RUR sites plus all sites in VE (with 554 different categorizations) and VI-URB3 (190 m a.s.l.) exhibited diurnal patterns very similar to 555 ozone, i.e. higher levels in the middle of the day. The two patterns remain different throughout the 556 year: different source emissions and weather factors may explain such patterns. The first cycle is 557 558 related to road traffic and is more evident at VR-TRA3 (close to a large logistic intermodal freight transport hub), where the highest levels of NO_x are recorded. In Europe, sulphur content in 559 automotive gasoline and diesel is now limited to <10 ppm (since 2009), however it is clear that 560 561 large volumes in traffic and congestion during rush hours may have a key effect in shaping the diurnal SO₂ levels. The second pattern was previously seen in Venice-Mestre (Masiol et al., 2014c) 562 and was related to local emissions from the nearby industrial zone hosting a large coal-fired power 563 plant and a large oil refinery, which are well known strong sources of SO₂, as indicated by local 564 emission inventories (ISPRA, 2015; ARPAV – Regione Veneto, 2015). The closeness of VE sites 565 566 to the sea drives the presence of sea/land breezes (mainly in warmest periods) and has a strong influence of the local circulation pattern and in bringing air masses from the industrial zone to the 567 site locations. The daytime increase in SO₂ was also reported for a background site in London by 568

Bigi and Harrison (2010) who attributed it to the downward mixing of plumes from elevated point
sources as the boundary layer deepens during daylight hours.

Generally, PM₁₀ exhibits higher concentrations overnight and clear minima in the early afternoon.
This pattern is consistent with the diurnal dynamics of the mixing layer. However, a secondary
cause may be related to the volatilisation of the more volatile aerosol compounds (e.g., nitrate)
during the early afternoon, i.e. when the air temperature is higher and relative humidity lower.
Minor peaks of PM₁₀ concentrations can be found just before noon at a few sites affecting by very
different emission scenarios (VR.URB2, VR.URB3, VE.URB3, PD.RUR2, PD.IND4). Their
interpretation is not clear and may be related to the local characteristics of the sites.

578

579 4.6 Long-term trends

The long-term trends were analysed using the monthly-averaged data: since missing data can 580 significantly affect the trend analysis, only months having at least 75% of available records were 581 processed. In addition, only trends over periods extending more than 4 consecutive years were 582 computed. The quantification and the significance of the trends were evaluated by applying the 583 Theil-Sen nonparametric estimator of slope (Sen, 1968; Theil, 1992). This technique assumes linear 584 trends and is therefore useful to estimate the interannual trends, but it is irrespective of the shape of 585 586 trends. The slopes were also deseasonalized by using the seasonal trend decomposition using loess (STL) to exclude the effect of the seasonal cycles. Details of STL are provided in Cleveland et al. 587 (1990). The statistics of the linear trends are listed in Table SI1 as percentage y^{-1} , while Figure SI7 588 shows all the single trends expressed in concentration y^{-1} along with the upper and lower 95th 589 confidence intervals in the trends and the *p*-values, which indicate the statistical significance of the 590 591 slope estimation.

592

593 The trends observed at each site reflect both regional and local changes and are affected by the 594 particular characteristics of the site locations. However, when trends observed at individual sites are aggregated at provincial or regional levels, their relationships with current and past emissions
inventories can be examined. The slopes calculated with the Theil-Sen method are then compared
with changes in primary emissions as reported in various emission inventories (EIs) available at
both regional and provincial level.

599

For each pollutant, statistically significant (p < 0.05) trends expressed in percentage y^{-1} are provided 600 in Figure 8 along with: (i) the differences in estimated emissions between 2010 and 2005 at 601 602 provincial level provided by the EI SINAnet top-down (ISPRA, 2015) and (ii) the differences between 2010 and 2007/8 at regional scale provided by INEMAR (ARPAV - Regione Veneto, 603 2015). Differences of EIs (Δ EIs) are also expressed as percentage v^{-1} for an easier comparison with 604 measured data (this choice considers a linear trend throughout the considered periods). Generally, 605 most of the trends are negative for all the species and the changes in measured concentrations of 606 CO, NO_x , PM_{10} , $PM_{2.5}$ agree well with EIs. However, some exceptions are also found (note that not 607 all sites revealed statistically significant slopes). 608

609

CO concentrations decreased significantly at 12 of 18 sites with more than four years of available 610 data, with slopes between -20.5% y^{-1} (TV.RUR) and -1% y^{-1} (PD.IND1). This result is in line with 611 the average drops by -6.5% reported by INEMAR for the whole Veneto and with the EIs provided 612 by SINAnet at province scale. However, some sites show the opposite behaviour, i.e. increased 613 concentrations: BL.URB (+3.5% v^{-1}), VR.URB2 (+9.6% v^{-1}) and VE.TRA2 (+18.2% v^{-1}). Since 614 falls of CO in EIs are mainly driven by road transport and non-industrial combustion (including 615 biomass burning), such sites have possibly experienced local increases of road traffic volumes or 616 emissions from the use of wood for domestic heating, which is widely used in mountain areas 617 (mainly BL). 618

The average decrease of NO_x levels across the whole region was estimated as -4.1% y^{-1} by 620 INEMAR and was mainly attributed to "road transport" and, secondarily, to "combustion in energy 621 and transformation industry" and "combustion in industry" sectors. The average slope from data 622 measured experimentally at all the sites was -3.5% y^{-1} , ranging between -12.3% y^{-1} (VR.IND) and -623 1.5% y^{-1} (VI.TRA). Statistically significant trends were negative at all sites except at VI.IND 624 $(+1.5\% \text{ y}^{-1})$. This result is a positive finding meaning that abatement strategies and improvement of 625 626 technologies had an effect in reducing the ambient levels of NO_x. However, in recent years there 627 has been increasing interest in NO_x emissions and NO/NO₂ partitioning in Europe. Evident discrepancies have been found between achieving NO_x emission reductions and NO₂ ambient 628 629 concentrations, which do not meet the targets in many locations (e.g., Grice et al., 2009; Cyrys et al., 2012). These concerns have been related to the recent increase in NO₂ levels in Europe due to 630 the increased proportion of diesel-powered vehicles, which are known to have higher primary 631 (direct) emissions of NO₂ (Carslaw et al., 2007). Since the recent boom of diesel vehicles was also 632 experienced in Italy (Cames and Helmers, 2013), the relationship between significant trends of NO_x 633 634 and NO₂ needs to be further investigated.

635

Analysis of trends in SO₂ requires particular care. Based on EIs estimations, a significant drop (-636 14% y^{-1}) was experienced across the whole region, while at province level SO₂ emissions decreased 637 between -18% y^{-1} (in RO) and -6% y^{-1} (in BL). These drops in EIs are principally attributed to the 638 sector of combustion in energy transformation industries. In Veneto, this sector is mostly 639 represented by the energy production in coal-fired power stations, which is present in VE province 640 (Porto Marghera). In addition, the sector of 'other mobile sources' has also experienced significant 641 642 drops, particularly of ship emissions (present only in coastal areas of VE province): from January 2010, ships in the harbours are requested to use fuels with sulphur content <0.1%. According to the 643 EIs, experimental data show that all sites in the VE province and PD.IND2 have negative trends, 644 while four sites have experienced statistically significant increases in SO₂ levels, i.e. VI.URB3 645

646 $(+24.5\% \text{ y}^{-1})$, VR.RUR $(+11.4\% \text{ y}^{-1})$, PD.TRA $(+9.9\% \text{ y}^{-1})$ and RO.TRA $(+7.4\% \text{ y}^{-1})$. While the 647 decline at VE sites may be attributed to the concurrent fall in industrial and maritime source 648 emissions at a local scale, the increases at other sites deserve further investigation and are possibly 649 due to long-range transport of polluted air masses. In this context, a recent study (Masiol et al., 650 2015) has reported that Eastern Europe is a potential source area for PM_{2.5}-bound sulphate, i.e. the 651 major sink for atmospheric SO₂.

652

Measured data fit well with changes in EIs for PM₁₀, except in VI province. All statistically 653 significant trends calculated from field data show decreases in PM₁₀ concentration ranging between 654 -5.6% v^{-1} in BL.RUR2 and -1.9% v^{-1} in RO.TRA. PM₁₀ emissions estimated by EIs dropped 655 between -4.4% y^{-1} in RO province and -2.4% y^{-1} in VR, with a slight increase in VI (+1.1% y^{-1}). At 656 a regional scale, the INEMAR inventory reports an overall decrease of -5.7% y^{-1} . Results for PM_{2.5} 657 are quite variable, mostly due to the short series of available experimental data: only 3 sites present 658 more than 4 years of data, of which only 2 have statistically significant trends. Hence, no further 659 information can be extracted for PM_{2.5}. 660

661

662 Conclusions

Air quality data from 43 monitoring sites have been used as input for a number of statistical tools to assess the extent of air pollution across the Veneto. This paper is the first one providing information from a large number of sites over a wide region of N Italy. The main findings can be summarised as follows:

Carbon monoxide and sulphur dioxide show low levels across the region and, therefore, are not
 considered as critical pollutants. While CO does not show any evident spatial trend, SO₂ levels
 are higher in VE Province, particularly around Venice-Mestre. This anomaly was linked to the
 industrial zone (coal power plant, oil refineries, other industrial installations) and harbour
 activities;

Nitrogen dioxide, ozone and particulate matter (both PM₁₀ and PM_{2.5}) are critical pollutants in view of protecting human health, i.e. the EC limit and target values are frequently breached at some sites. Those air pollutants deserve special attention because of their known adverse effects
 upon public health: future mitigation strategies should focus on reducing concentrations of such key pollutants;

Air pollutants are quite uniformly distributed across the region: no pairs of sites have statistically
 significant differences in the levels of CO, while only the two remote rural sites present
 statistically significant differences from a large number of other sites for NO_x and ozone;

The current site categorization was tested by applying a data depth classification analysis: results
 show that sites categorized as rural generally differ from other categories, while there is not a
 clear separation among urban background, traffic and industrial sites. Probable causes of the
 poor classification are discussed; some insights for improving the monitoring network are
 provided;

Spatial trends were investigated by interpolating average concentrations at the rural sites: despite
 NO and NO₂ having a slightly different direction of maximum slopes, nitrogen oxides generally
 increase from the mountain to the coastal plain environments, while ozone presents maxima
 concentrations at high mountain rural sites and minima in coastal areas;

Seasonal pattern analysis revealed that CO, NO, NO₂, NO_x, PM₁₀ and PM_{2.5} show significantly
 higher levels in colder months and minima in summer. This pattern is mainly attributed to the
 lower mixing layer heights, the limited oxidation potential and the emissions from domestic
 heating. The volatilization of semi-volatile aerosol compounds during the warmer seasons is
 another reason of this behavior for PM. On the contrary, ozone has an opposite seasonality with
 maxima in summer due to its increased photochemical generation. No seasonal patterns are
 found for SO₂;

Diurnal and weekly cycles were investigated. Generally, similar patterns are observed across the
 region for all the measured species. A strong potential effect of road traffic emissions was found

- for CO and nitrogen oxides: one/two daily peaks are commonly found at urban and hotspot sites 698
- and were related road traffic emissions during rush hours. Ozone cycles were shaped by the 699
- photochemistry and by the interplay with NO. PM cycles generally show higher levels overnight 700
- 701 and, therefore, are mostly shaped by the mixing layer height dynamics;
- An overall decrease of all measured species (except ozone) was observed throughout the region. 702
- 703 Generally, results of trend analysis well fit with changes in emission inventories. However,
- some sites with opposite trends have been identified; the reasons for increasing concentrations 704
- in such sites needs to be further investigated. 705
- 706
- Disclaimer 707
- The views and conclusions expressed in this paper are exclusively of the authors and may not reflect 708
- 709 those of ARPAV.
- 710

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852	Table 1. Characteristics of the sampling sites and analysed pollutants. Municipality refers to the territorial district where the sampling site is located: if
853	the site is not located in the main town of a municipality, the city/town is given in brackets.
854	

Province	No.	Code	Lat.	Long.	Height (m)	Municipality (City/Town)	Site full name	Area type	Monitored pollutants
BL	1	BL.RUR1	46.339	11.802	2020	Falcade	Passo Valles	Rural background	NOx, O3
	2	BL.RUR2	46.163	12.361	615	Pieve d'Alpago	Pieve d'Alpago	Rural background	NOx, O3, SO2, PM10,
	3	BL.SUB	46.031	11.906	263	Feltre	Area Feltrina	Suburban	CO, NOx, O3, SO2, PM10
	4	BL.URB	46.144	12.219	401	Belluno	BL-città	Urban background	CO, NOx, O3, SO2, PM10
TV	5	TV.RUR	45.837	12.51	14	Treviso (Mansuè)	Mansuè	Rural background	CO, NOx, O3 PM10
	6	TV.URB1	45.672	12.238	15	Treviso	TV-Via Lancieri	Urban background	CO, NOx, O3, SO2, PM10, PM2.5
	7	TV.URB2	45.89	12.307	72	Conegliano	Conegliano	Urban background	CO, NOx, O3, SO2, PM10
VI	8	VI.RUR	45.849	11.569	1366	Asiago	Asiago-Cima Ekar	Rural background	NOx, O3
	9	VI.URB4	45.759	11.736	114	Bassano	Bassano del Grappa	Urban background	NOx, O3 PM10
	10	VI.URB1	45.56	11.539	36	Vicenza	VI-Quartiere Italia	Urban background	CO, NOx, O3
	11	VI.URB2	45.532	11.522	33	Vicenza	VI-Ferrovieri	Urban background	CO, NOx, O3, SO2
	12	VI.URB3	45.714	11.368	190	Schio	Schio-via Vecellio	Urban background	NOx, O3 PM10, PM2.5
	13	VI.TRA	45.545	11.533	35	Vicenza	VI-San Felice	Urban-traffic	CO, NOx SO2, PM10
	14	VI.IND1	45.536	11.294	154	Chiampo	Chiampo	Urban-industrial	NOx
	15	VI.IND2	45.465	11.386	61	Montebello Vicentino	Montebello Vicentino	Suburban-Industrial	NOx
VR	16	VR.RUR	45.589	11.037	824	Boscochiesanuova	Boscochiesanuova	Rural background	CO, NOx, O3, SO2, PM10
	17	VR.SUB	45.462	10.911	91	Verona (Cason del Chievo)	VR-Cason	Suburban	CO, NOx, O3, SO2, PM10, PM2.5
	18	VR.URB1	45.443	11.007	64	Verona	VR-Piazza Bernardi	Urban background	CO, NOx
	19	VR.URB2	45.399	11.285	30	Verona (San Bonifacio)	San Bonifacio	Urban background	CO, NOx, O3, SO2, PM10, PM2.5
	20	VR.URB3	45.183	11.311	25	Legnago	Legnago	Urban background	NOx, O3 PM10
	21	VR.TRA1	45.444	10.963	62	Verona	VR-Borgo Milano	Urban-traffic	CO, NOx SO2, PM10
	22	VR.TRA2	45.41	10.989	60	Verona	VR-San Giacomo	Urban-traffic	CO, NOx SO2
	23	VR.TRA3	45.416	10.969	65	Verona	VR-ZAI	Urban-traffic	CO, NOx, O3, SO2
	24	VR.IND	45.543	10.886	183	Fumane	Fumane	Industrial	NOx SO2, PM10,
VE	25	VE.RUR	45.694	12.786	5	Concordia Sagittaria	Concordia Sagittaria	Rural background	NOx, O3
	26	VE.URB1	45.428	12.313	1	Venezia	VE-Sacca Fisola	Urban background	NOx, O3, SO2, PM10
	27	VE.URB2	45.5	12.261	1	Venezia (Mestre)	VE-Parco Bissuola	Urban background	CO, NOx, O3, SO2, PM10
	28	VE.URB3	45.629	12.59	3	San Donà di Piave	San Donà di Piave	Urban background	CO, NOx, O3 PM10, PM2.5

	29 VE.TR	A1 45.49	12.218	2 Ve	enezia (Mestre)	VE-Via Tagliamento	Urban-traffic	CO, NOx SO2, PM10
	30 VE.TR	A2 45.474	12.22	2 Ve	enezia (Marghera)	VE-Via Beccaria	Urban-traffic	CO, NOx , PM10
	31 VE.IN	D 45.438	12.205	2 Ve	enezia (Marghera)	VE-Malcontenta	Industrial	CO, NOx SO2
PD	32 PD.RU	R1 45.594	11.909	24 Sa	anta Giustina in Colle	S. Giustina in Colle	Rural background	CO, NOx, O3
	33 PD.RU	R2 45.289	11.642	18 Pa	adova (Cinto Euganeo)	Parco Colli Euganei	Rural background	NOx, O3, SO2, PM10
	34 PD.UR	B 45.371	11.841	13 Pa	adova	PD-Mandria	Urban background	CO, NOx, O3, SO2, PM10
	35 PD.TR	A 45.433	11.89	11 Pa	adova	PD-Arcella	Urban-traffic	CO, NOx, O3, SO2, PM10
	36 PD.IN	D1 45.395	11.909	10 Pa	adova	PD-APS-1-Ignoto	Urban-industrial	CO, NOx, O3, SO2, PM10, PM2.5
	37 PD.IN	D2 45.415	11.907	10 Pa	adova	PD-APS-2-Carli	Urban-industrial	CO, NOx, O3, SO2, PM10, PM2.5
	38 PD.IN	D3 45.378	11.94	8 Pa	adova	PD-Granze	Urban-industrial	PM10
	39 PD.IN	D4 45.227	11.666	12 Es	ste	Este	Suburban-Industrial	CO, NOx, O3, SO2, PM10
RO	40 RO.RU	J R 45.103	11.554	8 Ba	adia Polesine	Polesine-Villafora	Rural background	CO, NOx, O3, SO2
	41 RO.SU	B 44.95	12.333	1 Po	orto Tolle	Porto Tolle	Suburban	NOx SO2, PM10, PM2.5
	42 RO.UI	RB 45.039	11.79	3 Ro	ovigo	RO-Borsea	Urban background	CO, NOx, O3, SO2
	43 RO.TH	RA 45.074	11.782	7 Ro	ovigo	RO-Centro	Urban-traffic	CO, NOx, O3, SO2, PM10

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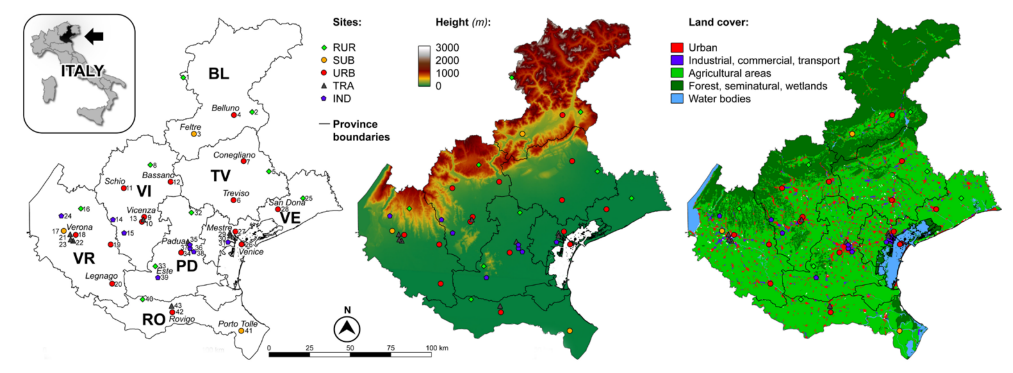


Figure 1. Map of the Veneto Region: administrative (left); terrain relief (centre); land use and cover from CORINE Land Cover 2006 data (right).
Sampling sites are also represented as diamonds (RUR sites), dots (SUB and URB sites), triangles (TRA sites) and pentagons (IND sites).

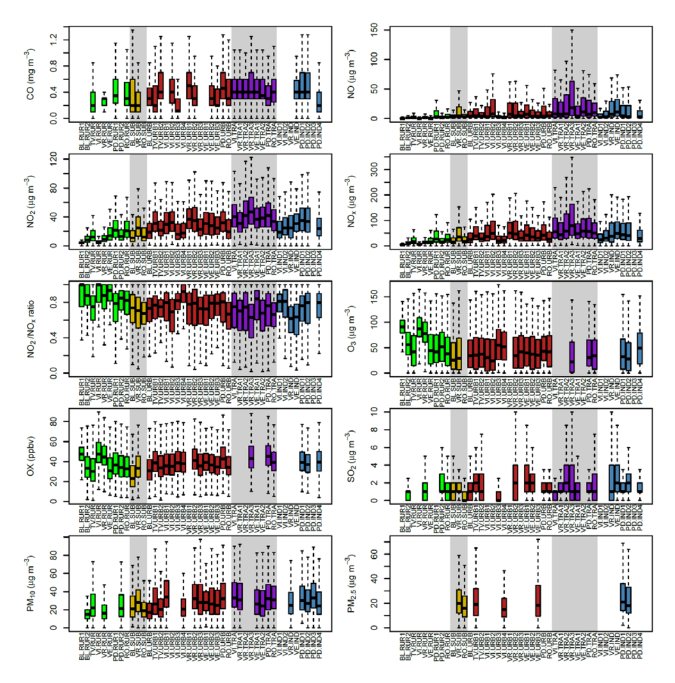


Figure 2. Average and ranges of concentrations of the analyzed pollutants as boxplots of raw data (line= median, box= inter-quartile range, whiskers= ± 1.5 *inter-quartile range). Sites are clustered following their categorisation.

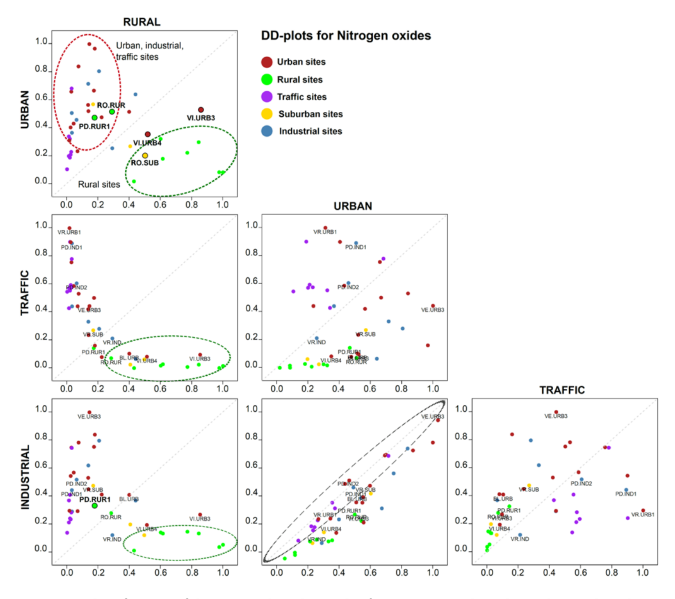


Figure 3. Classification of the stations based DD-plot for nitrogen oxides. The analysis is based on multivariate functional simplicial data depth using nitrogen oxides (NO, NO₂ and NO_x).

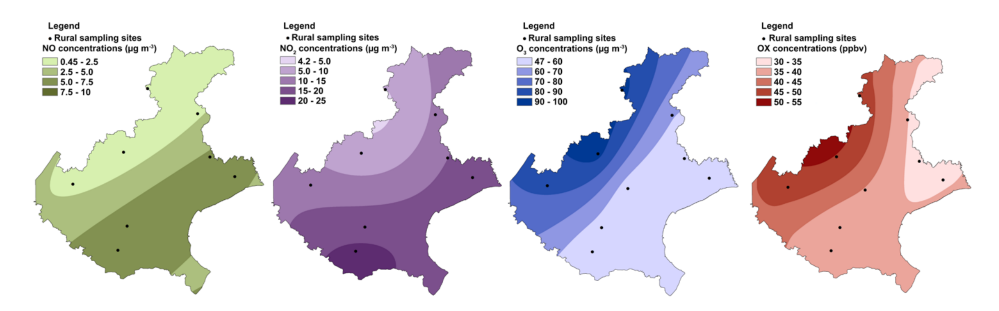


Figure 4. Maps of concentrations for NO, NO₂, O₃ and OX measured at the RUR sites. All RUR sites were used to interpolate data for O₃ and OX, while PD-RUR1 was not used for NO and NO₂ because of the anomalously high levels recorded. Ordinary kriging method was used for interpolation. Maps showing the standard errors associated with predictions are shown as Figure SI5.

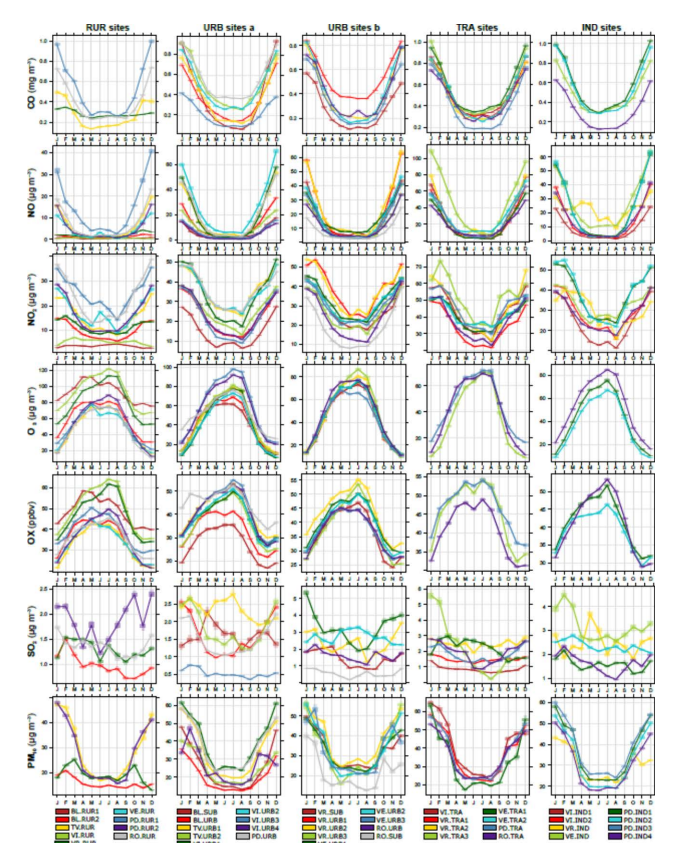


Figure 5. Seasonal variations of the monitored pollutants. Each plot reports the monthly average levels as a filled line and the associated 75th and 99th confidence intervals calculated by bootstrapping the data (n=200).

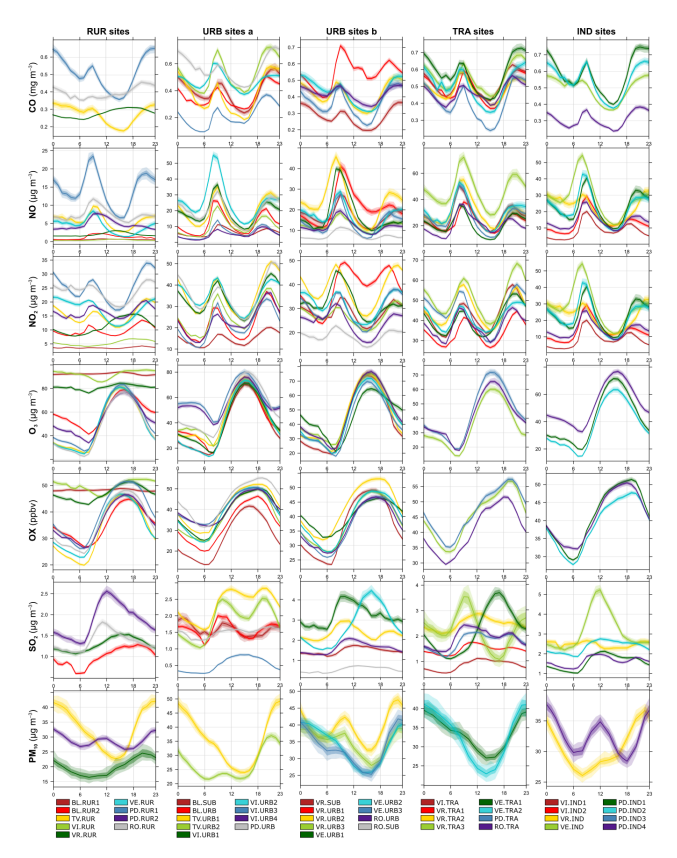


Figure 6. Diurnal variations of levels of measured pollutants computed over the hourly averaged data during the sampling period. Each plot reports the average level as a filled line and the associated 75th and 99th confidence intervals calculated by bootstrapping the data (n=200). Data are corrected for DST.

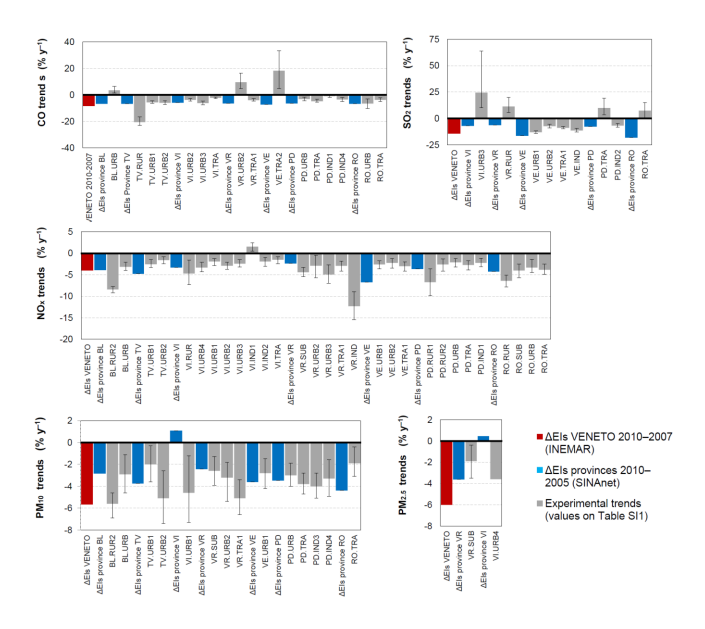


Figure 7. Results of the Theil-Sen analysis of trends (grey bars). The estimated trends are expressed as percentage; the confidence intervals of trends are given by error bars. Only sites showing significant trends (*p*-values < 0.05) are shown. The percentages of change in emissions provided by the most recent emission inventories are also provided and expressed as % y⁻¹: changes of EIs for the whole Veneto region (Δ _Veneto, red bars) refers to the difference of EIs between 2010 and 2007; changes of EIs for each province (Δ _BL, Δ _TV, Δ _VI, Δ _VR, Δ _VE, Δ _PD, Δ _RO, blue bars) refers to the difference of EIs between 2010 and 2005.