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6 **Spatial, seasonal trends and transboundary transport of PM_{2.5}**
7 **inorganic ions in the Veneto region (Northeastern Italy)**
8

9 Mauro Masiol^{a*}, Francesca Benetello^b
10 Roy M. Harrison^{a†}, Gianni Formenton^c
11 Francesco De Gaspari^c, Bruno Pavoni^b
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* To whom correspondence should be addressed. Email: m.masiol@bham.ac.uk

† Also at: Department of Environmental Sciences / Center of Excellence in Environmental Studies, King Abdulaziz University, PO Box 80203, Jeddah, 21589, Saudi Arabia

33 **SPATIAL, SEASONAL TRENDS AND**
34 **TRANSBOUNDARY TRANSPORT OF PM_{2.5}**
35 **INORGANIC IONS IN THE VENETO**
36 **REGION (NORTHEASTERN ITALY)**

37
38 **Mauro Masiol^{a*}, Francesca Benetello^b**
39 **Roy M. Harrison^{a†}, Gianni Formenton^c**
40 **Francesco De Gaspari^c, Bruno Pavoni^b**

41
42
43 **^aDivision of Environmental Health and Risk Management**
44 **School of Geography, Earth and Environmental Sciences**
45 **University of Birmingham**
46 **Edgbaston, Birmingham B15 2TT**
47 **United Kingdom**

48
49 **^bDipartimento di Scienze Ambientali**
50 **Informatica e Statistica, Università Ca' Foscari Venezia**
51 **Dorsoduro 2137, 30123 Venezia, Italy**

52
53 **^cDipartimento Provinciale di Padova**
54 **Agenzia Regionale per la Prevenzione e Protezione Ambientale**
55 **del Veneto (ARPAV), Via Ospedale 22, 35121 Padova, Italy**

* To whom correspondence should be addressed. Email: m.masiol@bham.ac.uk

† Also at: Department of Environmental Sciences / Center of Excellence in Environmental Studies, King Abdulaziz University, PO Box 80203, Jeddah, 21589, Saudi Arabia

60 **ABSTRACT**

61 The Veneto Region lies in the eastern part of the Po Valley (Italy). This is one of the hotspots
62 in Europe for air quality, where efforts to meet the European standard for PM_{2.5} according to
63 current and future legislation have been generally unsuccessful. Recent data indicating that
64 ammonium, nitrate and sulphate account for about one third of total PM_{2.5} mass show that
65 secondary inorganic aerosol (SIA) plays a key role in the exceedence of the standards. A
66 sampling campaign for PM_{2.5} was carried out simultaneously in six major cities (2012-2013).
67 The water soluble inorganic ions were quantified and data processed to: (1) investigate the
68 seasonal trends and the spatial variations of the ionic component of aerosol; (2) identify
69 chemical characteristics at the regional-scale and (3) assess the potential effects of long-range
70 transport using back-trajectory cluster analysis and concentration-weighted trajectory (CWT)
71 models. Results indicated that PM_{2.5} and SIA ions have an increasing gradient in
72 concentrations from North (mountain) to South (lowland) and from East (coastal) to West
73 (more continental), whereas K⁺ and Ca²⁺ levels are quite uniformly distributed. Similar
74 seasonal trends in PM_{2.5} and ions are seen across the region. Simultaneous daily changes
75 were observed and interpreted as a consequence of similar emission sources, secondary
76 pollutant generation and accumulation/removal processes. Sulphate and nitrate were not
77 directly related to the concentrations of their precursor gases and were generally largely, but
78 not completely, neutralised by ammonium. The clustering of back-trajectories and CWT
79 demonstrate that the long-range movement of the air masses has a major impact upon PM_{2.5}
80 and ion concentrations: an area spreading from Eastern to Central Europe was identified as a
81 main potential source for most ions. The valley sites are also heavily influenced by local
82 emissions in slow moving northerly air masses. Finally, two episodes of high nitrate levels
83 were investigated to explain why some sites are experiencing much higher concentrations
84 than others. This study identifies some key features in the generation of SIA in the Po Valley,

85 demonstrating that SIA generation is a regional pollution phenomenon and mitigation
86 policies are required at regional, national and even European scales.

87

88 **Keywords:** PM_{2.5}, Ionic composition, Secondary inorganic aerosol, Long-range transport, Po
89 Valley

90 1. INTRODUCTION

91 Although most elements of the periodic table and many thousands of different organic
92 compounds are found in airborne particulate matter (PM), a few major components usually
93 make up a large percentage of the total mass. Ammonium (NH_4^+), nitrate (NO_3^-) and sulphate
94 (SO_4^{2-}) are among the major components of aerosol in the lower troposphere and their
95 average mass percentages in fine PM (aerodynamic diameter less than $2.5 \mu\text{m}$, $\text{PM}_{2.5}$)
96 account for $\sim 7\%$, $\sim 9\%$ and $\sim 15\%$, respectively in southern Europe (Putaud et al., 2010).
97 These ions can be directly emitted from various sources, including sea salt, mineral dust,
98 traffic, biomass combustion, industries and other anthropogenic processes. However, the
99 dominant mechanisms for their presence in the particulate-phase are the oxidation of
100 precursor gases, i.e. nitrogen oxides ($\text{NO} + \text{NO}_2 = \text{NO}_x$) and sulfur dioxide (SO_2), to nitric
101 (HNO_3) and sulfuric (H_2SO_4) acids, respectively. The subsequent neutralisation with
102 ammonia (NH_3) forms salts such as ammonium nitrate (NH_4NO_3), ammonium sulphate
103 ($(\text{NH}_4)_2\text{SO}_4$) and ammonium bisulphate ($(\text{NH}_4)\text{HSO}_4$) (Seinfeld and Pandis, 2006; Holmes,
104 2007; Benson et al., 2011). These salts are commonly referred to as secondary inorganic
105 aerosol (SIA).

106

107 $\text{PM}_{2.5}$ has clearly demonstrated adverse effects upon human health (WHO, 2006), and
108 reducing human exposure to PM is, therefore, of primary importance. In particular, it is a key
109 objective in the few hot-spots left in Europe, such as the Po Valley, where the current
110 standards for PM are not met. Several large cities (e.g., Milan, Turin, Bologna, Verona and
111 Venice-Mestre) and a myriad of minor urban agglomerations, industrial areas, agricultural
112 and rural environments are spread over a $\sim 48 \cdot 10^3 \text{ km}^2$ -wide alluvial lowland. A total of ~ 16
113 million inhabitants and the related road traffic and energy production cause heavy
114 anthropogenic emissions across the entire valley. In addition, enclosure by the Alps and

115 Apennine mountains surrounding the valley from the North, West and South (only the eastern
116 side is opened to the Adriatic Sea) forms a barrier for the dispersion of pollutants and has a
117 negative impact on air quality, with a buildup of PM and nitrogen oxides mainly during the
118 cold season. Sampling at a rural site in the south-eastern Po Valley (San Pietro Capofiume),
119 Decesari et al. (2014) found that sulphate and nitrate contributed appreciably to particulate
120 matter mass. Their analysis of the association of particulate matter concentrations with
121 meteorological factors revealed a complex interplay of local and long-range transport
122 influences.

123

124 The European Directive 2008/50/EC imposed a PM_{2.5} annual average concentration of 25 µg
125 m⁻³ as a *target* value to be achieved by 2010. As the *target* value will become the European
126 *limit* value to be met by 2015, this standard has to be achieved with the current and future
127 legislation. However this concentration is not met in many locations of Veneto Region,
128 Eastern Po Valley (EEA, 2013): in 2012, eight of the 14 sites included in the main monitoring
129 plan for PM_{2.5} of the local environmental protection agency (ARPAV) breached the target
130 value (ARPAV, 2013). These sites are located in a number of major cities of the region and
131 generally the PM_{2.5} concentrations were 3–7 µg m⁻³ above the target value. In addition, PM_{2.5}
132 levels exceeding the target value were also recorded in rural environments demonstrating that
133 even the background pollution is high.

134

135 Almost all the literature available for the SIA pollution in the Veneto is based on studies
136 carried out in the Municipality of Venice (Squizzato et al., 2012;2013; Masiol et al., 2014a).
137 Results have shown that about 25–35% of the total PM_{2.5} mass in Venice-Mestre is made up
138 of SIA, which is therefore a key component when the target values in the eastern Po Valley
139 are exceeded. Consequently, successful policies should include not only the reduction of

140 direct (primary) sources, but also the reduction of precursor gases to prevent the formation of
141 secondary particles (de Leeuw, 2002; Andreani-Aksoyoglu et al., 2004; Wu et al., 2008).

142
143 However, data collected in a single coastal city, Venice, are not sufficient to depict the key
144 characteristics of SIA pollution across the Veneto, the territory of which extends from Alpine
145 environments to foothills to flat plain areas in the North-South axis and extends from
146 continental to coastal environments in the West-East axis.

147
148 In view of this, the present study investigates the levels, spatial distribution and sources of
149 SIA in six major cities of Veneto, which have been carefully selected to be representative of
150 different environments of the region. The investigated territory extends to ~125 km on the
151 North-South axis and ~60 km from West to East. The inorganic ionic composition of $PM_{2.5}$
152 was quantified at six sites located in major cities for one year (2012–2013). The seasonal and
153 spatial variations were examined using a series of statistical tests and chemometric
154 approaches. Starting from the experimental data, the SIA formation at a regional-scale in
155 Veneto is described and the potential local and external sources are investigated. This study
156 has identified some key features that can improve the understanding of the generation of
157 secondary inorganic particles in the entire Po Valley.

158

159 **2. MATERIALS AND METHODS**

160 **2.1 Site Selection**

161 A multiple-site $PM_{2.5}$ sampling campaign was carried out according to the EN 14907:2005
162 standard from April 2012 to March 2013 in 6 major cities: Belluno (BL), Conegliano (TV),
163 Vicenza (VI), Venice-Mestre (VE), Padua (PD) and Rovigo (RO) (Figure 1a). Stations
164 managed by ARPAV, were placed in high density residential areas and can be considered as

165 representative of city-wide background levels. In Table 1 some site characteristics are
166 summarised. Since the Veneto region includes a northern Alpine zone (29% of the territory),
167 an intermediate hilly one (15%), a wide southern flat lowland (56%) and an eastern coastline
168 (95 km long), the cities were also selected to represent most of the differing environments
169 and features of the territory. BL (36,600 inhabitants) is located in an Alpine valley
170 surrounded by mountains, with no large industries or heavy traffic, but biomass burning
171 emissions are intense in winter, as wood is largely used for domestic heating. TV (35,700
172 inhabitants) is in a foothill region and is therefore representative of the transition between the
173 mountain and lowland; many factories process stainless steel, produce appliances and
174 electrical equipment, but a large part of the land is used for agriculture, especially for
175 vineyards. VI (115,900 inhabitants) is an important city with intense traffic and small to
176 medium-sized mechanical, textile, tanning and jewelry manufactures. VE (271,000
177 inhabitants) is a conurbation extending from the coastal lagoon of Venice to the mainland
178 with a complex emission scenario. This includes heavy road, maritime and airport traffic, an
179 industrial zone hosting chemical and steel plants, an oil-refinery, incineration facilities,
180 thermoelectric power plants and others. PD (214,200 inhabitants) is the most densely
181 populated municipality of the region, with many medium-sized factories mainly in the
182 engineering, technological and building sectors, but it also suffers from intense traffic due to
183 the presence of a large intermodal and logistics hub. RO (52,800 inhabitants) is located in a
184 flat lowland midway between the Alps and the Apennines and is the biggest processing center
185 of Veneto for agricultural products. Demographic data refer to 2011 and to the whole
186 municipalities.

187

188

189

190 **2.2 Experimental**

191 PM was collected on quartz fiber filters (Whatman QMA), starting at midnight for 24 h
192 continuously using low-volume samplers installed in air conditioned cabins (temperature
193 $<20^{\circ}\text{C}$). $\text{PM}_{2.5}$ masses were gravimetrically determined (sensitivity $0.1\ \mu\text{g}$) after
194 preconditioning at constant temperature ($20\pm 1\ ^{\circ}\text{C}$) and relative humidity ($50\pm 5\%$). Sampled
195 filters were stored in clean Petri slides in the dark and at $-20\ ^{\circ}\text{C}$ until analyses to prevent
196 losses, photochemical reactions and biological processes. The entire set of collected samples
197 covers most of the year (total 2190). The quantification of the water soluble inorganic ions
198 was limited to a subset of 60 samples per site (total 360) collected in 6 periods of 10
199 consecutive days in the middle of April, June, August, October, December and February.
200 Periods were chosen to be representative of all the seasons and include the dates when home
201 heating was switched off (15 April) and on (15 October) as established by the national
202 legislation. A $\sim 2\ \text{cm}^2$ -wide subsample of each filter was extracted in vials with 10 mL MilliQ
203 water (resistivity= $18.2\ \text{M}\Omega\cdot\text{cm}$ at 25°C , Millipore) and sonicated for 50 min. Vials were
204 capped to avoid artifacts and sample evaporation. Extracts were pre-filtered on microporous
205 ($0.45\ \mu\text{m}$) PTFE membranes and injected in two Metrohm (Switzerland) ion chromatographic
206 systems with conductivity detectors to quantify the concentrations of five anions (F^{-} , Cl^{-} ,
207 NO_3^{-} , PO_4^{3-} , SO_4^{2-}) and five cations (Na^{+} , NH_4^{+} , K^{+} , Mg^{2+} , Ca^{2+}). Anions were separated on
208 a Metrosep A Supp 7–250/4.0 column applying a isocratic flow ($0.8\ \text{mL min}^{-1}$) of $360\ \text{mM}$
209 Na_2CO_3 (Sigma-Aldrich, ACS $\geq 99.8\%$) eluent. Cations were determined using a Metrosep C
210 3–150/4.0 column and a $1\ \text{mL min}^{-1}$ isocratic flow of $3\ \text{mM}$ ultrapure HNO_3 (Fluka,
211 TraceSELECT, $\geq 69\%$). Single-ionic standards were prepared from pure salts and used to test
212 the linearity and calibrate the instrumental responses. The analyses were routinely checked by
213 using certified liquid standards (Fluka, TraceCERT) diluted in MilliQ water. The relative
214 repeatability of each ion determination (standard deviation of 10 replications) was $<5\%$. Field

215 blanks were prepared and analysed together with the samples and the values obtained were
216 routinely subtracted. Limits of detections (LODs) were calculated as three times the standard
217 deviation of field blanks: data below the LODs were substituted by LOD/2.

218

219 Other chemical parameters were automatically determined on hourly or bihourly basis in each
220 site following European standards: NO, NO₂, NO_x (EN 14211:2012); SO₂ (EN 14212:2012);
221 O₃ (EN 14625:2012); PM₁₀ and PM_{2.5} with automatic beta-attenuation monitor systems. A
222 comprehensive list of measured parameters in each site is provided in Table 1.

223

224 **2.3 Sampling Artifacts**

225 A number of studies have reported that potential artifacts can occur during air sampling
226 because of ambient conditions and the interactions between collected particles and gaseous
227 compounds with each other or with the filter medium (e.g., Appel et al., 1984; Dasch et al.,
228 1989; Harrison et al., 1990; Harrison and Kitto, 1990; Koutrakis et al., 1992; Zhang and
229 McMurry, 1992; Cheng and Tsai, 1997; Pathak et al., 2004a; Schaap et al., 2004a; Pathak and
230 Chan, 2005). Generally, the most evident artifact is the evaporation of nitrate due to its gas-
231 particle partitioning (negative artifact), which is further enhanced by higher temperatures and
232 drier air. Also, pressure drop across the filter and mixing of acidic and alkaline particles on
233 the filter may perturb the gas-particle equilibrium. On the contrary, absorption of gas-phase
234 nitric acid may also occur (positive artifact) mainly driven by the presence of sea-salt
235 particles.

236

237 Studies conducted in the Po Valley (Putaud et al., 2002; Schaap et al., 2004a) have reported
238 that nitrate volatilization generally dominates over absorption. In particular, Schaap et al.,
239 (2004a) concluded that quartz filters have a full retention of nitrate at temperatures <20°C. In

240 this study, all the samplers were installed into air conditioned cabins with an internal
241 constant temperature below 20°C. Subsequent filter transport, handling and analysis were
242 carried out under the same controlled conditions, while filter storage was at -20 °C.
243 Moreover, the prevailing high relative humidity recorded at all of the sites (average >70%
244 RH) during the sampling periods further decreased the potential nitrate loss. For these
245 reasons, negative artifacts of nitrate can be considered negligible. Positive artifacts are also
246 expected to be small: concentrations of Na⁺ and Cl⁻ (as tracers of sea-salt) and Mg²⁺ and Ca²⁺
247 (as tracers of crustal particles) during the study were very low.

248

249 Another potential positive artefact can be caused by the absorption of SO₂ on collected
250 particles, which can be subsequently oxidized to sulphate (Pathak and Chan, 2005). Due to
251 the very low concentrations of SO₂ in Veneto (ARPAV, 2013) and according to data obtained
252 with and without the use of denuders by Vecchi et al. (2009), sulphate can be considered a
253 conserved specie in the Po Valley (i.e. not subject to adsorption or volatilisation).

254

255 In summary, sampling conditions and chemical results indicate that potential artifacts in this
256 study are small. For this reason, all the chemometric analyses have been performed on raw
257 data.

258

259 **2.4 Back-Trajectory and CWT Analysis**

260 Back-trajectories were computed to study the history of air masses during the sampling days.
261 Set-up: HYSPLIT model (Draxler and Rolph, 2013; Rolph, 2013); 96 h backward; starting
262 height at 20 m a.s.l.; 4 trajectories per day at 3, 9, 15 and 21 UTC calculated separately for all
263 the sites. A clustering algorithm using the Euclidean distance measure (Carslaw, 2014) was

264 applied to gain information on pollutant species with similar chemical histories by grouping
265 back-trajectories into clusters depending on their potential origin.

266

267 CWT is a back-trajectory-based hybrid receptor model used to assess potential source areas
268 affecting air pollution at a receptor site. Briefly, each grid cell ij in a grid domain was used to
269 compute the weighted concentration obtained by averaging sample concentrations that have
270 associated trajectories passing the grid cell according to:

$$C_{ij} = \frac{1}{\sum_{k=1}^N \tau_{ijk}} \sum_{k=1}^N (C_k) \tau_{ijk}$$

271 where i and j are the coordinates of grid, k the trajectory index, N the number of trajectories,
272 C_k the pollutant concentration measured at the receptor site upon arrival of the trajectory k ,
273 and τ_{ijk} represents the residence time of trajectory k in the ij cell. Further insights are provided
274 in Seibert et al. (1994) and Hsu et al. (2003). Cluster analysis and CWT were computed using
275 R and the ‘Openair’ package (Carslaw and Ropkins, 2012; Carslaw, 2013).

276

277 **3. RESULTS**

278 **3.1 Overview of Results**

279 Table 2 summarises the annual average concentrations of PM_{2.5} and ions and also gives
280 statistics for SIA (as sum of ammonium, nitrate and sulphate) and Σ WSII (sum of all the
281 analysed water soluble inorganic ions). Due to the high percentage of samples below the
282 LODs, F⁻, Mg²⁺ and PO₄³⁻ were excluded from the statistics. A comprehensive list of results
283 for each month is provided as Supplementary Information Table S11. The PM_{2.5} annual
284 average concentrations (365 days) ranged from a minimum of 16 $\mu\text{g m}^{-3}$ in BL and a
285 maximum of 28 $\mu\text{g m}^{-3}$ in PD. In the study period, the European annual average target value
286 of 25 $\mu\text{g m}^{-3}$ (2008/50/EC Directive) was breached in three sites (PD, RO, VI). On an annual

287 basis, Σ WSII accounted for a significant fraction of the total $PM_{2.5}$ mass, ranging from 30%
288 (BL) to 41% (RO) and generally showed a slightly increasing trend from north to south.
289 Annually, the most abundant ion in all the sites (Figure 1b) was nitrate, ranging from 36%
290 (BL) and 47% (VI) of the Σ WSII, followed by sulphate 22% (VI)–29% (BL, VE),
291 ammonium 17% (BL, TV)–21% (VI, RO) and potassium 3% (RO, VI)–5% (BL). Sodium
292 varied from 1% (VI) and 5% (TV), while the remaining single ions never exceeded 2%. The
293 annual levels of $PM_{2.5}$ and $PM_{2.5}$ -bound nitrate, sulphate and ammonium in this study are
294 very similar to those recorded in other urban sites in the Po Valley (Table SI2).

295
296 Gaseous pollutants were recorded for all the year on hourly basis and data were averaged to
297 give daily mean values (Table 2 and Table SI1). The annual average concentrations of NO
298 during the selected periods varied from $12 \mu\text{g m}^{-3}$ (TV) to $27 \mu\text{g m}^{-3}$ (PD); NO_2 from $23 \mu\text{g}$
299 m^{-3} (BL) to $37 \mu\text{g m}^{-3}$ (PD and RO); NO_x from $45 \mu\text{g m}^{-3}$ (BL and TV) to $79 \mu\text{g m}^{-3}$ (PD);
300 O_3 from $46 \mu\text{g m}^{-3}$ (RO) to $61 \mu\text{g m}^{-3}$ (PD); SO_2 from $1 \mu\text{g m}^{-3}$ (PD) to $2.8 \mu\text{g m}^{-3}$ (VE).
301 These mean concentrations are very close to the annual average levels and demonstrate that
302 the selected periods are representative of the annual concentrations.

303
304 The annual average NO_2 levels never exceeded the Limit Value fixed by the European
305 Directives ($40 \mu\text{g m}^{-3}$). In Veneto the emission inventory (EI) for 2007/8 (ARPAV and
306 Regione Veneto, 2013) reported that road transport was the main source of NO_x (52111 Mg
307 y^{-1}), followed by combustion in manufacturing industry (15119 Mg y^{-1}), other mobile
308 sources and machinery (13793 Mg y^{-1}), combustion in energy and transformation industries
309 (7322 Mg y^{-1}) and non-industrial combustion plants (7187 Mg y^{-1}), while remaining
310 EMEP/EEA sources (production processes, agriculture, waste treatment and disposal, solvent
311 and other product use, extraction and distribution of fossil fuels and geothermal energy and

312 other sources and sinks) accounted for 3216 Mg y⁻¹. The annual average levels of SO₂ were
313 very low at all the sites and well below the European limit value. The EI reported that in
314 2007/8 the main contributors in Veneto were (in Mg y⁻¹): combustion in energy and
315 transformation industries (5077)> combustion in manufacturing industry (4578)> other
316 mobile sources and machinery (2340)> production processes (1879)> non-industrial
317 combustion plants (1327)> sum of other EMEP/EEA sources (165).

318

319 It should be noted that most of the NO_x was emitted at ground level by mobile sources,
320 whereas most of SO₂ emissions originated from stationary sources via chimneys. SO₂ may
321 disperse widely from elevated sources, but the NO_x sources are themselves widely
322 distributed.

323

324 **3.2 Seasonal Variations**

325 The PM_{2.5} time series are reported as Supplementary Information Figure SI1 and exhibit
326 seasonal trends at all of the sites, i.e. higher levels during winter and lower in summer, as
327 commonly observed in most sites in the Po Valley (e.g., Marcazzan et al., 2003; Vecchi et al.,
328 2004; Perrone et al., 2012; Tositti et al., 2014). The seasonality is strongly linked to weather
329 conditions, such as prolonged atmospheric stability, shallower mixing layers, wind calm
330 periods and low temperatures, which favor the accumulation of atmospheric pollutants at the
331 ground level (Ferrero et al., 2010). The increased use of wood for domestic heating in winter
332 and the burning of biomass such as straw and crop residues in the harvest season (late
333 autumn) may also have a role in raising the PM_{2.5} levels. The semi-volatility of ammonium
334 nitrate may also be important. The time series also showed a number of single peaks of
335 concentration at various sites. In most cases these peaks occurred at individual sites and were
336 therefore linked to local and occasional phenomena. However, it is evident that the highest

337 concentrations were recorded on January 6th for all stations except BL, when thousands of
338 folk fires of wooden material were lit in most of the Veneto region for a local religious
339 celebration. This episode was extensively reported by Masiol et al. (2014b) and recorded
340 extremely high daily concentrations of PM_{2.5}, ranging from 136 µg m⁻³ in VI to 202 µg m⁻³ in
341 RO. This period was not included in the present study.

342

343 On a monthly basis, each ion exhibited a typical seasonality and similar seasonal trends were
344 generally observed in all the territory. Figure 2 reports the mass concentration time series of
345 the three SIA components, while seasonal average levels for all ions are shown as Figure SI2.
346 Results for SIA ions show that both the concentrations and the daily variations of SIA at the
347 four sites in the flatter areas of the Po Valley (VI, VE, PD and RO) are quite similar and are
348 in line with results observed at urban sites in other nearby regions (Table SI2). Low SIA
349 concentrations were recorded at all sites in June, and in October in the Alpine valley (BL),
350 the SIA components were extremely low as well. Nitrate concentrations in PM are inversely
351 related to the ambient temperature: they are higher in the colder months, mainly because
352 ammonium nitrate tends to volatilise at temperatures above 20°C (Schaap et al., 2004a;
353 Vecchi et al., 2009). This is observed all over Europe (e.g., Allen et al., 1989; Schaap et al.,
354 2004b; Revuelta et al., 2012). Sulphate presents a peculiar bimodal seasonality, with two
355 maxima in August and February. A peak in the warmest period is commonly recorded in
356 Europe (e.g., Revuelta et al., 2012) and is probably due to the increased photochemical
357 activity favouring the oxidation of SO₂ via hydroxyl radical reaction (Stockwell and Calvert,
358 1983; Khoder, 2002; Seinfeld and Pandis, 2006), whereas the peak in February may be
359 associated with aqueous phase oxidation. Ammonium concentrations tend to parallel those of
360 nitrate and sulphate. Calcium shows no evident seasonality. However slightly higher levels
361 were recorded in August and winter. Potassium, a known tracer of biomass combustions,

362 (e.g., Puxbaum et al., 2007; Saarnio et al., 2010) presents an evident seasonality with higher
363 concentrations in the coldest period. Wood (i.e. logs, briquettes, chips and pellet) is becoming
364 a popular renewable alternative to natural gas in Northern Italy (Pastorello et al., 2011) and
365 the increasing emissions from its use for domestic heating can be considered the most
366 plausible source. Chloride has a seasonal behavior similar to potassium. Its presence in PM
367 can derive from various sources, i.e. sea-salt, biomass burning, resuspension of road deicing
368 salts, coal combustion and various industrial processes. The marine origin can be probably
369 excluded as no significant gradients of concentration are observed from the stations close to
370 the coast (VE) to the more continental ones (VI and PD). Therefore, biomass burning and the
371 resuspension of road salt are probably the most important sources.

372 Seasonal trends of gaseous pollutants are also given in Figure SI2. Nitrogen oxides increased
373 during the cold season due to changes in mixing depths and emission rates, while ozone
374 reached the highest levels in the warmest period due to its photochemistry. Sulfur dioxide
375 showed no clear seasonal trends, but reached the highest levels in VE during the warmest
376 period (June-August).

377

378 **3.3 Spatial Variations**

379 Starting from the evidence that PM_{2.5} and most ions have quite similar seasonal trends at all
380 the sites, an inter-site comparison of the annual concentrations was conducted for each ion.
381 Since the data were not distributed normally, the nonparametric Kruskal–Wallis one-way
382 analysis of variance was used. This test is based on the rank of each sample instead of its
383 value and the null hypothesis assumes that the central values of the groups (medians) are
384 equal, and is rejected for $p < 0.05$. Thus, the post hoc Dunn's test was applied to identify
385 which sites are significantly different from the others. Results generally show that PM_{2.5},
386 nitrate, sulphate and ammonium in BL and TV are significantly ($p < 0.05$) different from the

387 other sites and concentrations increased from North (mountain) to South (lowland) and from
388 East (coastal) to West (more continental). On the other hand, K^+ and Ca^{2+} levels are not
389 significantly different and their concentrations are therefore uniform in all of the Veneto
390 region. These results show that biomass burning, which has been identified as a major source
391 of potassium, and the re-suspension of mineral dust and soil, which is the major source of
392 calcium, are quasi-uniformly distributed throughout the region.

393

394 An indirect quantification of differences in concentrations among the sites was carried out by
395 regressing $PM_{2.5}$ mass concentration and nitrate+sulphate (expressed as $neq\ m^{-3}$) among
396 pairs of sites (intercept forced to zero). Results are provided in Figures SI3 and SI4,
397 respectively. Results for both $PM_{2.5}$ and nitrate+sulphate show that sites located in the main
398 Po Valley (VI, VE, PD and RO) have regression slopes around 1 (0.84–1.17) and high
399 coefficients of determination ($R^2 > 0.8$), which indicate good agreement between
400 concentrations. On the contrary, slopes (range 1.26–1.42) and R^2 (≤ 0.2) between BL and
401 sites in the main Po Valley indicate a very poor agreement. TV has an intermediate behavior
402 with sites in the main Po Valley: it presents a moderate relationship (R^2 0.6–0.8), but high
403 slopes (1.2–1.8).

404

405 The spatial and temporal relationships among the sites for $PM_{2.5}$ and ionic species were
406 further investigated by using correlation analysis. A preliminary inter-site correlation analysis
407 among the $PM_{2.5}$ concentrations for the whole year (365 days) was conducted. The $PM_{2.5}$
408 distributions were tested for normality by applying the Shapiro-Wilk's tests and the normality
409 assumption at $p < 0.05$ was not met. A Box-Cox transformation of the dataset was therefore
410 made. The resultant transformed data were normally distributed and Pearson's correlation
411 analysis was run. Results (Table 3) generally show significant correlations ($p < 0.01$, $r > 0.8$)

412 among all the sites, with the exception of BL, which appears slightly less correlated ($r \approx 0.7$)
413 with the others. $PM_{2.5}$ exhibits a similar temporal trend in all the cities even if these are
414 located in different territories of the region. It is evident that the processes of emission,
415 accumulation and removal are quite similar in the six cities.

416

417 However, the correlation analysis for the full dataset may be affected by the marked
418 seasonality of the variables, with the result that the correlation links variables with similar
419 seasonal trends and not sites with simultaneous daily variations. This problem was solved by
420 monthly-averaging the original data: the monthly means were subtracted from each daily
421 value in each selected period. This normalisation procedure had also the advantage of
422 generating variables that were quasi-normally distributed. The correlation matrices of the
423 monthly-averaged data are reported in Table 3 and show that the $PM_{2.5}$ is still strongly
424 correlated at all the sites located in the lowland area, while the mountain site (BL) is less
425 correlated. Sulphate has usually significantly ($p < 0.05$) positive relationships for all pairs of
426 sites, indicating that it has a similar (synchronous) behavior in the whole region. Highly ($r >$
427 0.75) significant correlations are also found for $PM_{2.5}$, nitrate and ammonium, except at BL
428 which appears to be uncorrelated with the other sites. Potassium is very well correlated in the
429 central part of the region (VI, VE, PD), while significant but weak correlations are found in
430 TV and RO, and BL is uncorrelated with the other sites. Calcium shows few inter-site
431 correlations (Table SI3).

432

433 This analysis generally shows that $PM_{2.5}$, potassium, nitrate, sulphate and ammonium follow
434 a similar day-to-day trend at all sites throughout the region, in particular in the lowland
435 territory, and confirms that both the emission sources and the accumulation/removal

436 processes in the region are similar. A similar finding was also recently reported for the levels
437 of PM₁₀-bound polycyclic aromatic hydrocarbons at 21 sites in Veneto (Masiol et al., 2013).

438

439 **4. DISCUSSION**

440 The SIA mass is generally calculated as the simple sum of ammonium, sulphate and nitrate or
441 is derived from the results of source apportionment approaches. Nevertheless, its prediction is
442 not straightforward because the ion generation, transport, aging or removal in the particle-
443 phase strongly depends on weather conditions, but also on the presence of precursor gases
444 and oxidant species (mainly hydroxyl radical, hydrogen peroxide and ozone). Basically, SIA
445 generation is a two-step process, in which the gaseous precursors SO₂ and NO_x undergo
446 photochemical and heterogeneous thermal oxidation to form sulfuric and nitric acids,
447 respectively. Subsequently, the acids are neutralised by ammonia, and in the case of
448 ammonium nitrate, partitioned according to thermodynamic equilibria, mostly determined by
449 temperature and relative humidity (Baek et al., 2004; Seinfeld and Pandis, 2006; Allen et al.,
450 1989). Reactions with other ions may also form mixed salts. Using the experimental data
451 obtained in this study, some preliminary conclusions regarding the SIA are drawn.

452

453 **4.1 Sulfur and Nitrogen Oxidation Ratios**

454 The degree of atmospheric conversion of gaseous precursors, SO₂ and NO₂, to sulphate and
455 nitrate, respectively, can be indirectly assessed by means of the sulfur (SOR) and nitrogen
456 (NOR) oxidation ratios:

$$\text{SOR} = \frac{n\text{-nssSO}_4^{2-}}{n\text{-nssSO}_4^{2-} + n\text{SO}_2}$$

$$\text{NOR} = \frac{n\text{-NO}_3^-}{n\text{-NO}_3^- + n\text{NO}_2}$$

457 where the n units are in moles m^{-3} and nss-SO_4^{2-} is the non-sea-salt sulphate calculated as
458 $[\text{SO}_4^{2-}] - 0.25 \cdot [\text{Na}^+]$. The SOR and NOR have been used by many authors (e.g., Khoder, 2002;
459 Bencs et al., 2008; Behera and Sharma, 2010) to describe the degree of ageing of the air
460 mass. The results appear as Table SI4, alongside those of other similar studies for
461 comparison. Annually, the average SOR varied from 0.4 (VE) to 0.6 (PD) suggesting a high
462 degree of oxidation of SO_2 in the atmosphere, while the annual average NOR ranged between
463 0.04 (BL) and 0.1 (PD). SOR shows no clear spatial variation and generally its seasonal
464 concentrations follow those of sulphate. However, it is important to point out that the
465 minimum SOR is reached at VE in the warmest period. This is probably due to the highest
466 concentrations of SO_2 in summer caused by: (1) the peak of energy production of a coal-fired
467 power plant meeting the demand for air conditioning; (2) the presence of higher shipping
468 traffic using the cruise harbour. This assumption is also supported by the emission inventory
469 for 2010 (ISPRA, 2014) showing that the Venice province has the highest production of SO_2
470 (4586 Mg y^{-1}), followed by Padova (1324 Mg y^{-1}). About 71% of the emissions in VE are
471 attributed to combustion in energy and transformation industries. Spatially, NOR seems to
472 increase slightly from North to South and from the coast to the mainland.

473

474 **4.2 Ammonia Availability and Neutralisation Ratio**

475 Ammonia is known to neutralise sulfuric acid irreversibly, and then nitric acid. In addition,
476 hydrochloric acid may react with gaseous ammonia to form ammonium chloride aerosol.
477 However, in thermodynamic equilibrium conditions ammonium chloride is reported to be 2-3
478 times more volatile than ammonium nitrate (Stelson and Seinfeld, 1982; Pio et al., 1992) and
479 its formation occurs later. It is well known that in low ammonia conditions, NH_3 acts as the
480 main limiting factor for SIA generation (Erisman and Schaap, 2004). On the other hand, in
481 case of high NH_3 availability, ammonium nitrate formation is principally limited by the

482 availability of nitric acid. These conditions are important in agricultural areas because
483 livestock farming and the use of soil fertilizers are primary sources of atmospheric NH₃
484 (Galloway et al., 2004; Sutton et al., 2008). Recent modeling simulations on a continental
485 scale (Wichink Kruit et al., 2012) have reported that ammonia levels in the Po Valley are
486 among the highest in Europe (range 4–10 µg m⁻³). This is also confirmed by satellite
487 observations (Clarisse et al., 2009) indicating the Po valley as one of the most evident
488 hotspots for NH₃ at a global scale. The 2010 Italian emission inventories (ISPRA, 2014)
489 reported that ~50.2·10³ Mg of NH₃ are emitted annually in Veneto, most of which is from
490 agriculture (48.9·10³ Mg), followed by road transport (0.7·10³ Mg). Because SO₂ emissions
491 have been sharply reduced in the last decades in most developed countries, including Italy
492 (Manktelow et al., 2007; Hamed et al., 2010), more NH₃ is available for the formation of
493 ammonium nitrate (Bauer et al., 2007; Pye et al., 2009). Recent data indicated that in Veneto
494 SO₂ concentrations are generally < 8 µg m⁻³, i.e. below the EU lower threshold (ARPAV,
495 2013).

496

497 Reactions of gaseous acids with other particles (e.g., sea salt, crustal dust, anthropogenic) can
498 form secondary salts, mainly replacing Cl⁻ with sulphate and nitrate, or forming salts with
499 Na⁺, K⁺, Mg²⁺ or Ca²⁺. For example, sulphate and nitrate may affect the hygroscopic
500 behaviour of mineral dust (Shi et al., 2008) and may form nitrate-containing particles mainly
501 in the coarse mode (Pakkanen et al., 1996; Metzger et al., 2006).

502

503 However, in this study, the masses of Na⁺, Mg²⁺, Ca²⁺ and Cl⁻ were low, if compared to
504 NH₄⁺, NO₃⁻ and SO₄²⁻ and therefore their contribution to salts in PM_{2.5} can be assumed to be
505 negligible. From a linear regression analysis between ammonium and the sum of nitrate and
506 sulphate (expressed as neq m⁻³) significant coefficients of determination (R² varying from

507 0.94 in BL and 0.99 in VI, VE and RO), almost unitary slopes (from 0.83 in BL 1.06 in VI,
508 VE and RO) and very low intercepts were obtained for all the sites. The scatterplots are
509 reported as Figure SI5. They also reveal that the relationships are constantly linear in all the
510 seasons, even if the mass contributions of each ion varied greatly during the year.

511

512 The neutralisation ratio (NR) (Bencs et al., 2008), also called acidity ratio (Engelhart et al.,
513 2011), expresses the degree of neutralisation of sulphate and nitrate by ammonium
514 (concentrations are in equivalents) and was used to describe the aerosol acidity:

$$\text{NR} = \frac{[\text{NH}_4^+]}{[\text{SO}_4^{2-}] + [\text{NO}_3^-]}$$

515 Figure 3 shows the NR time series and permits some inferences: (i) on an annual basis,
516 average NRs were equal to 1 within the analytical variability, or slightly less: 0.8 in BL, TV,
517 PD and 0.9 in VI, VE, RO; (ii) the lowest NRs were recorded in spring, while they were
518 almost constant in the remaining months at all the sites; (iii) both the concentrations and the
519 daily variations of SIA at the 4 sites in the Po valley had similar trends; (iv) NR variability in
520 August and February, i.e. in the warmest and coldest months of the year, respectively, was
521 small, while strong daily changes were recorded in April and October. It is unclear if this
522 trend is linked mainly to a discontinuity of the sources (e.g., domestic heating switching off
523 and on), to weather factors controlling the SIA generation, or to external transport effects.

524

525 To investigate the extent of neutralisation of the SIA in more detail, NR was plotted against
526 the ammonium concentration (Figure 4a). Results show that for all the sites: when
527 concentrations of NH_4^+ exceed $\sim 150 \text{ neq m}^{-3}$, the NR appears to be constant around 1 and
528 SIA is likely to be composed of ammonium nitrate and ammonium sulphates; for lower levels
529 of ammonia, the variability of NR increases and, generally, the ratio becomes smaller. These

530 results confirm that ammonia may effectively act as a limiting agent for SIA and suggest that
531 during ammonia-limiting conditions, sulfuric and nitric acids may react with other particles to
532 form salts. This assumption can be further confirmed by plotting the NR versus ionic balance
533 (ratio between the sum of all analysed cations and anions) (Figure 4b). The graph clearly
534 shows that most of samples are set in the 4th quadrant, a region where the relative lack of
535 ammonium ($NR < 1$) corresponds to an excess of cations (cations > anions), i.e. nitrate and
536 sulphate are potentially combined with other cations than ammonium. Figure 4b also shows
537 that no samples are plotted in the opposite quadrant (2nd), demonstrating that on days with an
538 excess of ammonium ($NR > 1$) no excess anions are present, thus showing the absence of other
539 inorganic salts of ammonium, such as NH_4Cl . A few samples mainly pertaining to the
540 mountain site (BL) are scattered in the 1st and 3rd quadrants: samples in the 1st quadrant are
541 characterised by an excess of ammonium and a positive ionic balance, i.e. an excess of
542 positive charges probably neutralised by organic acids, not measured in this study. Samples
543 in the 3rd quadrant were almost all collected in April and a possible explanation is that the
544 lack of positive charges may be balanced by H^+ (which was not measured), resulting in acid
545 aerosol.

546

547 **4.3 Potential Contribution of Long-Range Transport**

548 The analysis of the back-trajectories was used to give some insight into the potential
549 contribution of long-range aerosol transport upon the Veneto region. As known from the
550 literature, the use of trajectories has some limitations in accuracy for various reasons (e.g.,
551 Stohl et al., 1998). However, taking into account the range of associated uncertainties, the use
552 of some trajectory statistical methods is recognised as very useful to investigate potential
553 source areas (Kabashnikov et al., 2011; Abdalmogith and Harrison, 2005).

554 For the purpose of this study, the variability of back-trajectories was tested using different
555 starting heights and hours: errors associated with a single trajectory were reduced by
556 simulating four trajectories for each sampling day (at 6, 12, 18, 24 local time). The cluster
557 analysis was applied to all the 4-days back-trajectories computed and for the each site, i.e. 4
558 trajectories every day, which have been merged with daily data. In fact, this expedient
559 allowed the spread of daily chemical data over 4 trajectories and thus can account for days
560 that may have changes in trajectories within 24-h. The number of extracted clusters was
561 carefully evaluated by analysing the change in the total spatial variance and the best
562 compromise was 5 clusters for all the sites. Results show that all sites present similar mean
563 trajectories (Figure 5) named (1) Western Europe, (2) Mediterranean, (3) local, (4) Northern
564 Europe and (5) Eastern Europe. Statistics for chemical composition data in each cluster are
565 presented as boxplots in Figures 5 and SI6. The number of trajectories grouped in each
566 cluster generally differ among BL, TV and other sites (Table SI5). The reason is linked to the
567 topography of the territory: BL is located in an alpine valley, TV is at the border of Alps,
568 whereas other sites are located in flatter areas of the Po Valley. As a consequence, results for
569 BL and TV differ from the other sites with results sometimes showing opposite trends.
570 Generally, $PM_{2.5}$, nitrate and K^+ show similar results, with concentrations higher for cluster 3
571 in BL and TV and for clusters 1, 4 and 5 for the remaining sites. Sulphate in BL and TV
572 appears to have higher concentrations when air masses are associated with clusters 1 and 5,
573 whereas it is associated with clusters 3 and 5 at the other sites. Calcium and chloride show
574 only small differences. Sites BL and TV show a different behaviour with respect to $PM_{2.5}$.
575 The highest concentrations are associated with trajectory 3, which for the other sites shows
576 the lowest $PM_{2.5}$. This effect is probably the result of trapping of lower level emissions at BL
577 and TV in the slow moving northerly air. With regard to high sulphate and nitrates, these
578 sites behave rather similarly to the others as a results of regional influences.

579

580 On the other hand, the analysis of the potential effects of long-range transport on a regional
581 scale through the CWT model returned very similar results at all sites and clearly indicate
582 some predominant source areas for potential transboundary transport of PM_{2.5} and some ions
583 (Figures 6a and 6b). In particular, a wide area spanning across Eastern and Central Europe
584 and Northern Italy is identified as a main potential source of all species. Similar results have
585 also been obtained from a previous study conducted at a site near VE during 2009-2010
586 (Squizzato et al., 2014). In addition, other minor source areas are also identified: an area in
587 Central Italy which roughly coincides with the heavily populated areas of Rome and Naples
588 as a source of PM_{2.5} nitrate, and an area in North Africa, which may be linked to Saharan dust
589 outbreaks. CWT also shows that air masses passing over continental Europe are responsible
590 for the highest NR and SOR, while this effect is less evident for NOR (Figure SI7). If NR,
591 SOR and NOR are taken as indicative of the aging of air masses (generally highest values of
592 oxidation ratios and NR values close to 1 are expected in aged air masses) these results stress
593 that transboundary transport from continental Europe may have an important impact on levels
594 of secondary species in the Po Valley. The lower values of NOR than SOR probably reflect
595 the higher local emissions of NO_x compared to SO₂.

596

597 **4.4 Analysis of Single Episodes**

598 Three episodes of high SIA concentrations occurred during the campaign (Figure 2): (1) 15th
599 to 21st October, (2) 13th to 17th February and (3) 17th to 22nd February. Despite all sites
600 showed covariant daily variations in the levels of nitrate, some differences during those
601 episodes were identified. A further analysis of single back-trajectories was thus performed to
602 explain why some sites are generally experiencing much lower concentrations than others.
603 Figure 7 shows the single back-trajectories associated with the daily concentration of SIA. In

604 the first and third episodes, it is evident that all sites show similar daily air mass pathways
605 from the Mediterranean and Central Europe, respectively. However, only VI, VE, PD and RO
606 show similar daily variations and levels associated with single trajectories, whereas TV had a
607 similar daily variation, but significantly lower concentrations. There will be a number of
608 reasons explaining this result: (i) data indicate that transboundary transport of polluted air
609 masses may have a higher impact over the Eastern Po Valley; (ii) the cluster and CWT
610 analyses both indicate Central Europe as a major source area of ammonium nitrate aerosol;
611 (iii) results suggest the topography may influence the local impact of long-range transport: a
612 general homogeneity in the SIA levels is often recorded in the flat area of the valley, while
613 the Alpine chain may act as a barrier for the dispersion of pollutants at ground-level.

614

615 The results for the second episode are quite different. Despite all sites show similar air mass
616 histories, the levels of SIA were higher in RO and VI. As the differences cannot be explained
617 by differing air mass origins, it can be concluded that ammonium nitrate generation may also
618 occur locally as a consequence of oxidation of locally emitted NO_x .

619

620 In conclusion, these results indicate that SIA pollution may be sensitive to both long-range
621 transport and local generation processes. Due to the relatively short period investigated in this
622 study (60 days over one year), there is a limit to the conclusions which may be drawn.

623 However, as a few events such as those considered in detail can have a considerable effect
624 upon the annual mean $\text{PM}_{2.5}$ concentration, the different characteristics and effects of long-
625 range or local SIA episodes should be investigated in more detail over a longer period, by
626 collecting a large number of samples.

627

628

629 **5. CONCLUSIONS**

630 This study is the first one investigating the spatial and temporal properties of secondary
631 inorganic aerosol in a large area of the Po Valley using simultaneous experimental
632 measurements at multiple receptor-sites. The statistical processing of the data shows that
633 $PM_{2.5}$ and individual ions to have very similar concentrations across all urban sites and to be
634 very well correlated throughout the region, even though the sampling stations are located in
635 different cities and in an area $\sim 18.4 \cdot 10^3 \text{ km}^2$ -wide. Therefore, it can be concluded that the PM
636 pollution and the relative amount of SIA in the Veneto is quasi-uniformly distributed
637 throughout the region and the formation and removal processes affecting all sites are quite
638 similar. Moreover, a comparison with previous studies conducted in other nearby regions of
639 NE Italy indicates quite constant levels, seasonal trends and speciation of SIA over a wide
640 area of the Po Valley. The main results can be summarised as follows:

641

- 642 • Annually, water soluble inorganic ions account from 30% to 41% of the total $PM_{2.5}$
643 mass concentrations and the most abundant ion is nitrate (36%–47%), followed by
644 sulphate (22%–29%), ammonium (17%–21%) and potassium (3%–5%).
- 645 • Each ion exhibits a characteristic seasonality and similar seasonal trends are generally
646 recorded over the entire study area.
- 647 • $PM_{2.5}$, nitrate, sulphate and ammonium in BL and TV are significantly different from
648 other sites and generally levels of analysed pollutants increased from North (mountain)
649 to South (lowland) and from East (coastal) to West (more continental). In contrast, K^+
650 and Ca^{2+} show weak spatial gradients.
- 651 • Potassium, nitrate, sulphate and ammonium also show similar daily trends throughout
652 the region, in particular in the lowland territory, and confirm that both the sources and
653 the accumulation/removal processes in the region are similar.

- 654 • The neutralisation ratio and the ionic balance were jointly investigated to provide
655 information about the processes affecting SIA and the interactions between the
656 secondary ions and other particles. Results confirm the probable formation of
657 secondary salts with potassium, sodium and calcium.
- 658 • The application of trajectory-based methods (cluster and CWT analyses) was useful to
659 identify potential source areas leading to increases in $PM_{2.5}$ and ions concentrations
660 across the region. Results showed that higher concentrations of all analysed species are
661 mainly associated with air masses originating in a widespread area located in the
662 Eastern-Central Europe. Central Italy and Northern Africa are also identified as
663 possible source areas particularly for $PM_{2.5}$ and K^+ .
- 664 • The analysis of three single episodes of high ammonium nitrate levels indicate that both
665 long-range transport and local formation processes may lead to high SIA levels during
666 colder months. Those events have a large potential for raising the annual average levels
667 of $PM_{2.5}$ and should be investigated in more detail.

668

669 As a final remark, this study concluded that SIA pollution has similar and concurrent effects
670 over the entire study area and probably in the whole Po Valley. Findings clearly indicate that
671 any action to mitigate the $PM_{2.5}$ pollution to meet the present target and the future air quality
672 standards in Veneto must be taken concurrently in the entire region and well beyond its
673 boundaries.

674

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680

681 **DISCLAIMER**

682 This study was not financially supported by any public or private institution. We would like
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960 **TABLE LEGENDS**

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962

963 **Table 1.** Characteristics of the selected sampling sites and the number of analysed
964 samples.

965

966 **Table 2.** Annual average concentrations of analysed pollutants. A full list of results
967 including monthly average concentrations is provided as supplementary
968 material Table SI1.

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970 **Table 3.** Inter-site correlation matrices. Upper-left: box-cox transformed PM_{2.5} dataset
971 for the whole year (365 day); other matrices are calculated on the selected
972 periods (60 days) and data were monthly normalized. Only significant ($p <$
973 0.05) correlations are shown; correlations significant ($p <$ 0.01) are bold faced.
974 Correlation matrices for all analysed compounds is provided in Table SI2.

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980 **FIGURE LEGENDS**

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983 **Figure 1.** Map of selected sites (a; left) and annual average percentages of analysed ions
984 on Σ WSII (b; right).

985

986 **Figure 2.** Time series of sulphate, nitrate and ammonium in the six sites.

987

988 **Figure 3.** Time series of neutralisation ratio (NR) in the six sites.

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990 **Figure 4.** Scatterplots of a) ammonium vs NR and b) ionic balance vs NR. Samples
991 collected in six sites are coloured differently.

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993 **Figure 5.** Results of the back-trajectory clustering (upper) and distributions of PM_{2.5} and
994 ion concentrations for each identified cluster (bottom). Results for remaining
995 ions are provided as Supplementary Information Figure SI6.

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997 **Figure 6a.** CWT analysis for PM_{2.5}, nitrate and sulphate. Concentrations are expressed as
998 $\mu\text{g m}^{-3}$.

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1000 **Figure 6b.** CWT analysis for chloride, potassium and calcium. Concentrations are
1001 expressed as $\mu\text{g m}^{-3}$.

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1003 **Figure 7.** Single back-trajectories during three high-nitrate concentration events.

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1005

1006 **Table 1.** Characteristics of the selected sampling sites and the number of analysed samples.

	Municipality	Latitude	Longitude	Alt (m)	Site characteristics	Other automatic measurements
BL	Belluno	46.143 N	12.218 E	401	Public park, residential-commercial area	SO ₂ ; O ₃ ; NO ₂ ; NO; NO _x ; CO; Benzene; PM ₁₀ (gravimetric); PM ₁₀ (BAMs)
TV	Conegliano	45.890 N	12.307 E	72	Residential area	SO ₂ ; O ₃ ; NO ₂ ; NO; NO _x ; CO; PM ₁₀ (gravimetric)
VI	Vicenza	45.560 N	11.539 E	36	Residential area	NO ₂ ; NO; NO _x ; PM ₁₀ (gravimetric); PAHs
PD	Padova	45.371 N	11.841 E	13	Residential area	SO ₂ ; O ₃ ; NO ₂ ; NO; NO _x ; CO; Benzene; PM ₁₀ (gravimetric); PAHs
VE	Venice-Mestre	45.498 N	12.261 E	1	Public park, residential area	SO ₂ ; O ₃ ; NO ₂ ; NO; NO _x ; CO; Benzene; PM ₁₀ (gravimetric); PAHs
RO	Rovigo	45.074 N	11.782 E	7	Residential-commercial area	SO ₂ ; TSP (gravimetric); O ₃ ; NO ₂ ; NO; NO _x ; CO; PM ₁₀ (gravimetric)

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1019 **Table 2.** Annual average concentrations of analysed pollutants. A full list of results including
 1020 monthly average concentrations is provided as supplementary material Table SII.
 1021

		BL	TV	VI	VE	PD	RO
PM_{2.5}	μg m ⁻³	17	20	28	25	29	27
Na⁺	μg m ⁻³	0.14	0.31	0.15	0.16	0.47	0.23
NH₄⁺	μg m ⁻³	0.9	1.1	2.3	1.9	2	2.3
K⁺	μg m ⁻³	0.28	0.29	0.31	0.38	0.39	0.3
Ca²⁺	μg m ⁻³	0.11	0.15	0.15	0.15	0.16	0.15
Cl⁻	μg m ⁻³	0.12	0.12	0.19	0.17	0.19	0.24
NO₃⁻	μg m ⁻³	1.8	2.4	5	3.6	4.6	5.2
SO₄²⁻	μg m ⁻³	1.5	1.7	2.4	2.6	2.4	2.6
SIA	μg m ⁻³	4.2	5.2	9.7	8.1	9	10.2
SIA	%	23	25	32	30	29	35
ΣWSII	μg m ⁻³	5.2	6.5	10.8	9.2	10.7	11.4
ΣWSII	%	30	34	36	35	38	41
NO	μg m ⁻³	15	12	24	22	27	26
NO₂	μg m ⁻³	23	27	33	32	37	36
NO_x	μg m ⁻³	45	45	70	65	79	76
O₃	μg m ⁻³	49	47	48	49	61	46
SO₂	μg m ⁻³	1.1	—	—	2.8	1	2.5

1022

Table 3. Inter-site correlation matrices. Upper-left: box-cox transformed PM_{2.5} dataset for the whole year (365 day); other matrices are calculated on the selected periods (60 days) and data were monthly normalized. Only significant ($p < 0.05$) correlations are shown; correlations significant ($p < 0.01$) are bold faced. Correlation matrices for all analysed compounds is provided in Table SI2.

PM _{2.5} (Whole year)	BL	TV	VI	VE	PD	RO	PM _{2.5} (Monthly norm.)	BL	TV	VI	VE	PD	RO
BL	1						BL	1					
TV	0.74	1					TV	0.33	1				
VI	0.75	0.86	1				VI	0.26	0.84	1			
VE	0.75	0.82	0.86	1			VE		0.84	0.89	1		
PD	0.74	0.83	0.89	0.94	1		PD	0.29	0.85	0.89	0.87	1	
RO	0.71	0.82	0.88	0.9	0.93	1	RO	0.26	0.8	0.81	0.83	0.95	1
NO ₃ ⁻ (Monthly norm.)	BL	TV	VI	VE	PD	RO	SO ₄ ²⁻ (Monthly norm.)	BL	TV	VI	VE	PD	RO
BL	1						BL	1					
TV		1					TV	0.51	1				
VI		0.84	1				VI	0.39	0.51	1			
VE		0.85	0.95	1			VE	0.53	0.86	0.58	1		
PD		0.87	0.97	0.96	1		PD	0.54	0.74	0.59	0.9	1	
RO		0.79	0.86	0.84	0.92	1	RO	0.39	0.73	0.55	0.83	0.89	1
NH ₄ ⁺ (Monthly norm.)	BL	TV	VI	VE	PD	RO	K ⁺ (Monthly norm.)	BL	TV	VI	VE	PD	RO
BL	1						BL	1					
TV		1					TV		1				
VI	0.26	0.81	1				VI		0.58	1			
VE		0.86	0.92	1			VE		0.64	0.82	1		
PD		0.87	0.94	0.95	1		PD		0.51	0.83	0.77	1	
RO	0.26	0.77	0.85	0.83	0.92	1	RO	0.32	0.52	0.56	0.77	0.77	1

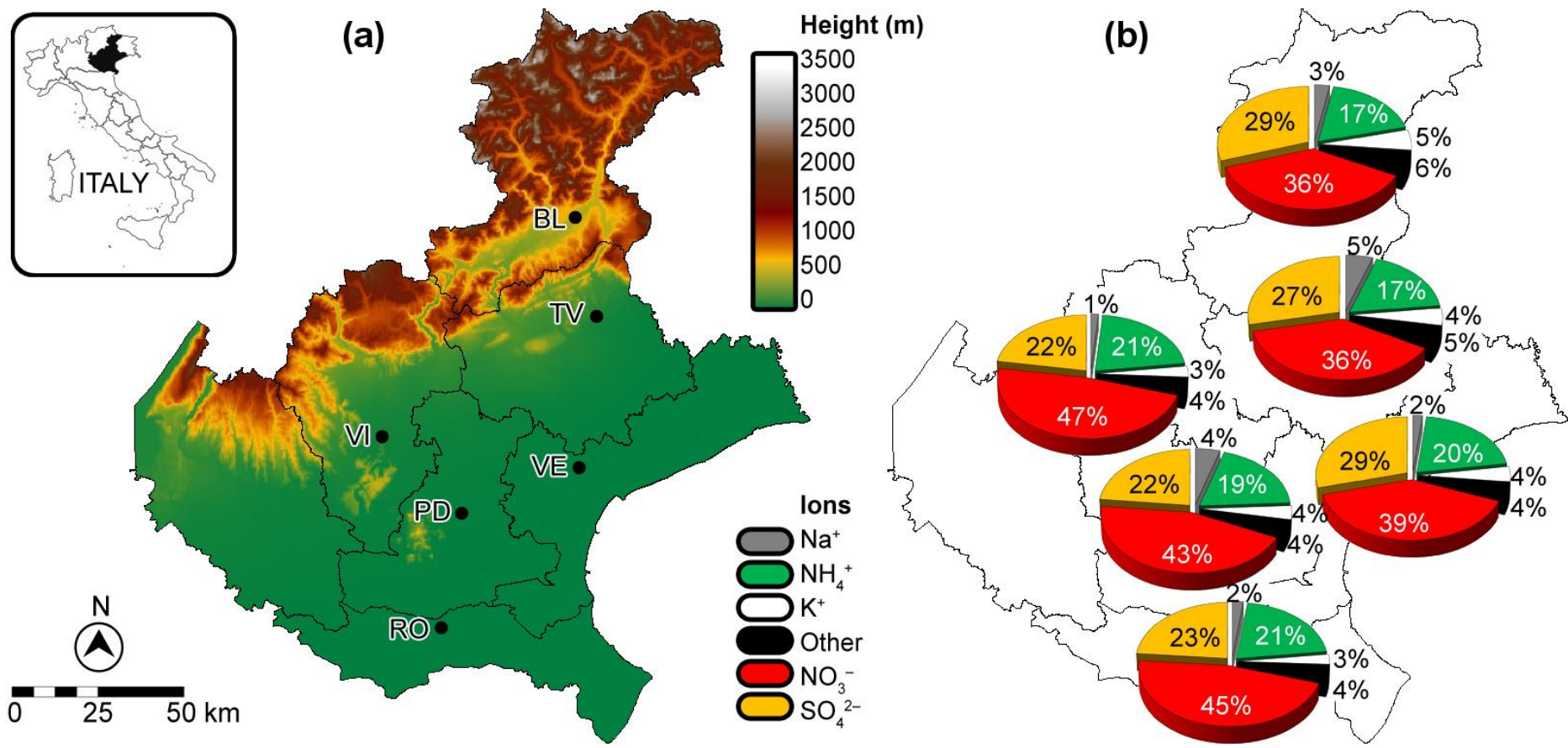


Figure 1. Map of selected sites (a; left) and annual average percentages of analysed ions on Σ WSSII (b; right).

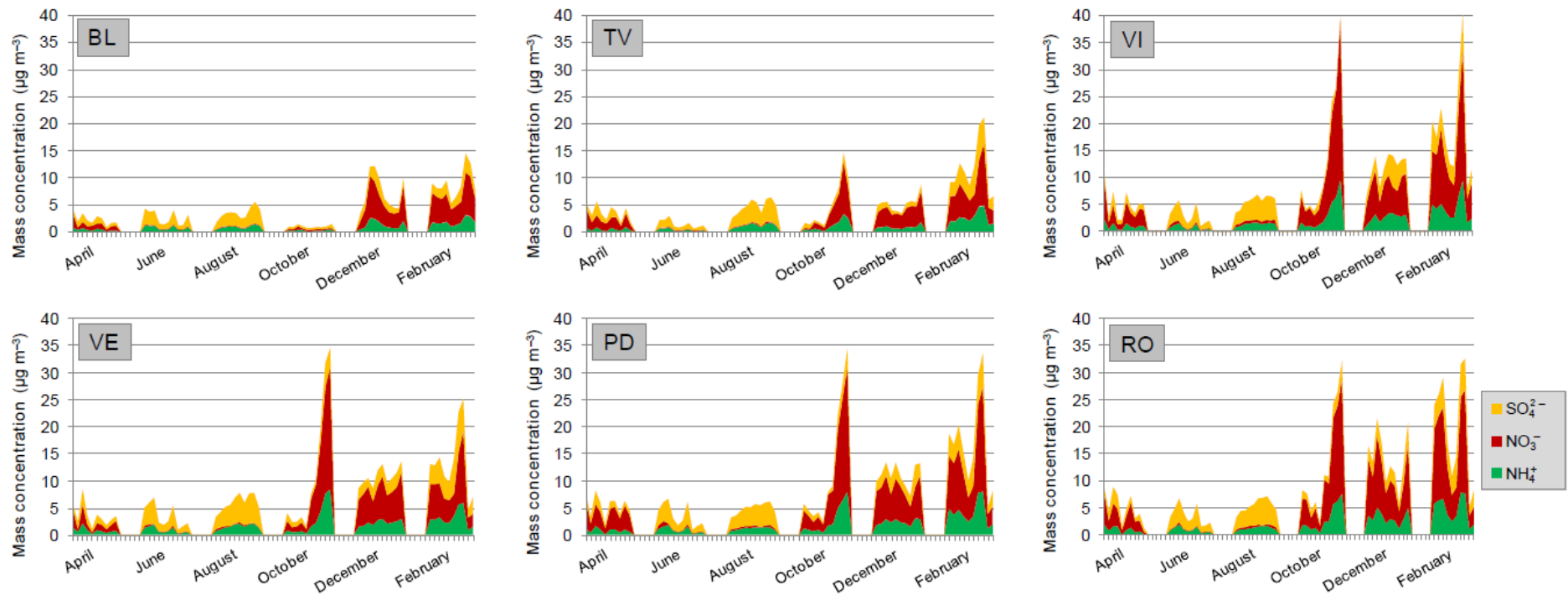


Figure 2. Time series of sulphate, nitrate and ammonium in the six sites.

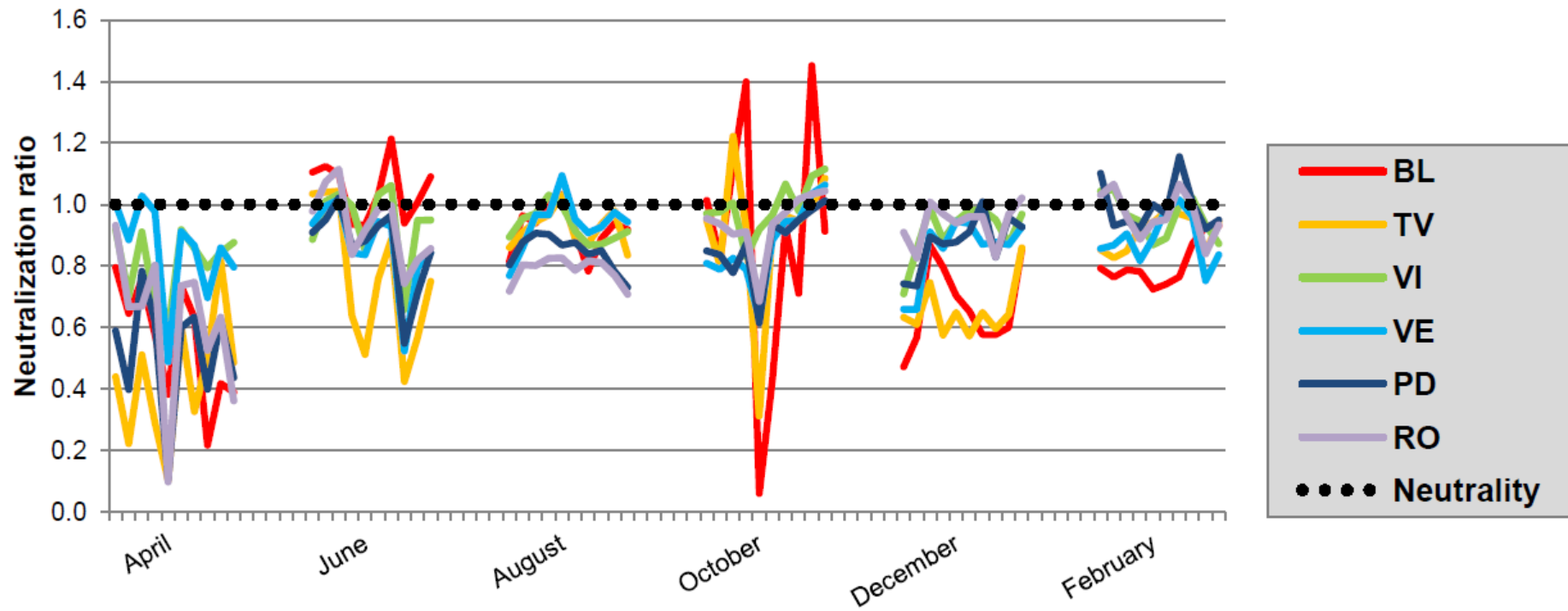


Figure 3. Time series of neutralisation ratio (NR) in the six sites.

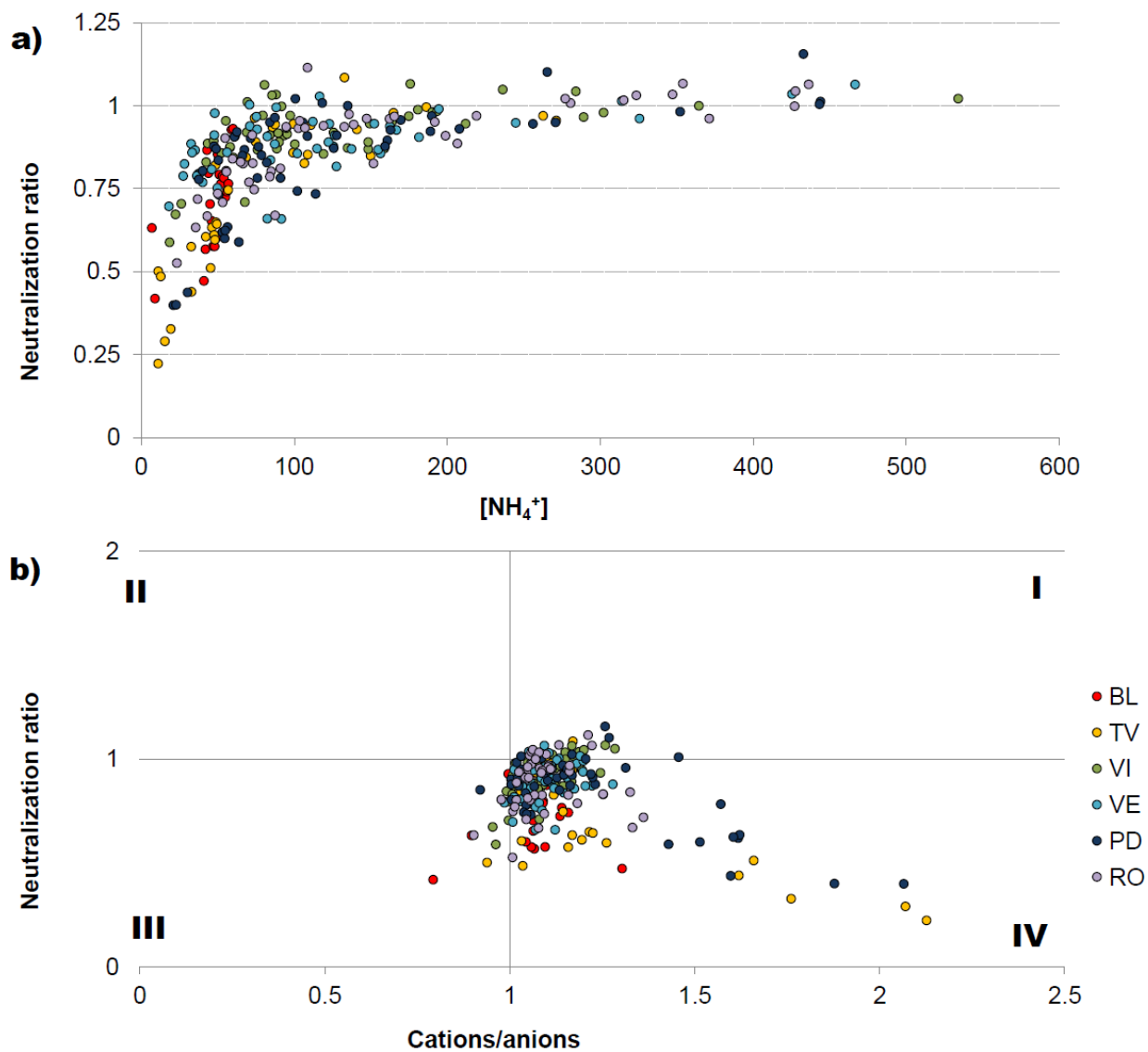


Figure 4. Scatterplots of a) ammonium vs NR and b) ionic balance vs NR. Samples collected in six sites are coloured differently.

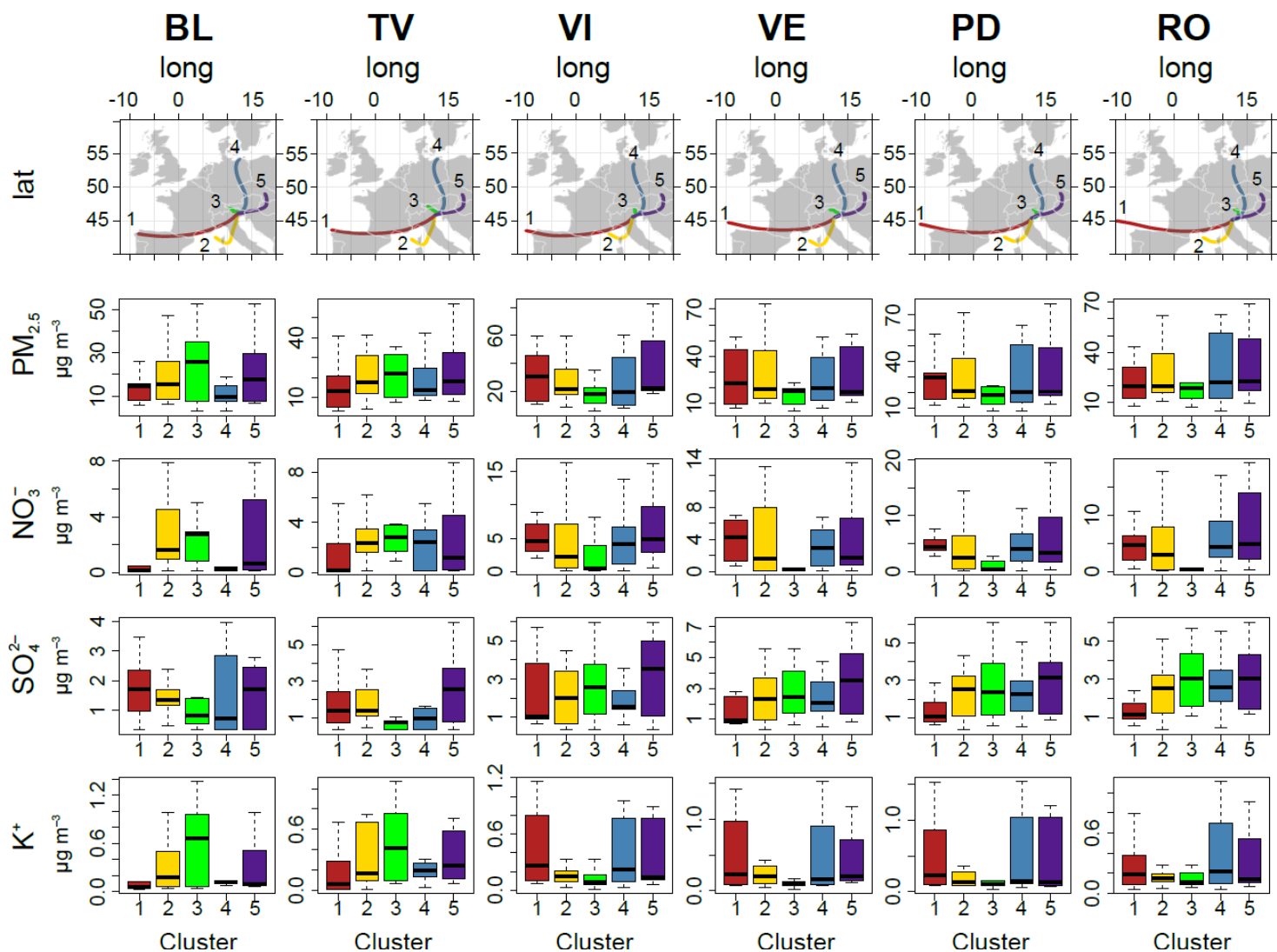


Figure 5. Results of the back-trajectory clustering (upper) and distributions of $PM_{2.5}$ and ion concentrations for each identified cluster (bottom). Results for remaining ions are provided as Supplementary Information Figure SI6.

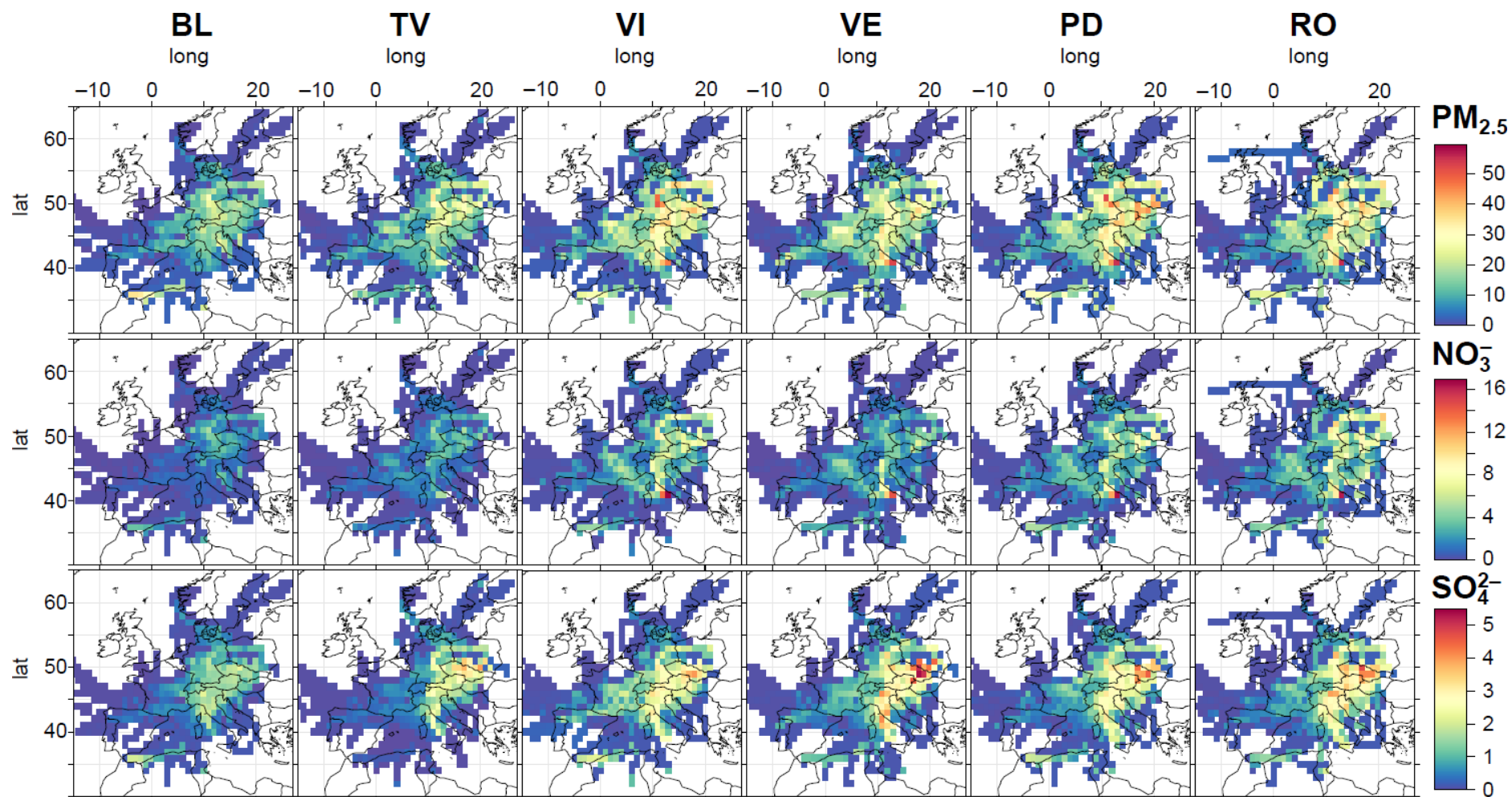


Figure 6a. CWT analysis for $\text{PM}_{2.5}$, nitrate and sulphate. Concentrations are expressed as $\mu\text{g m}^{-3}$.

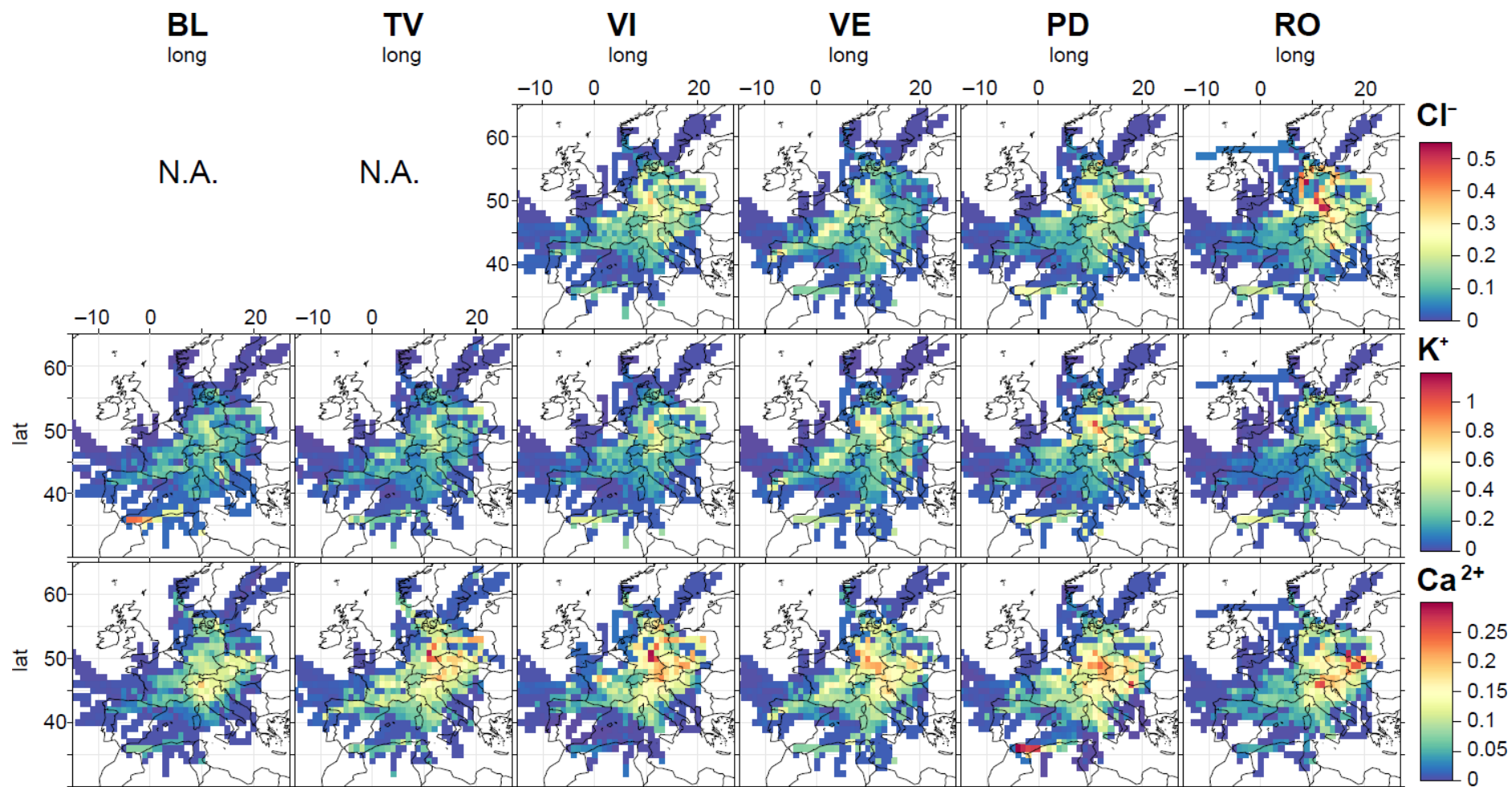


Figure 6b. CWT analysis for chloride, potassium and calcium. Concentrations are expressed as $\mu\text{g m}^{-3}$.

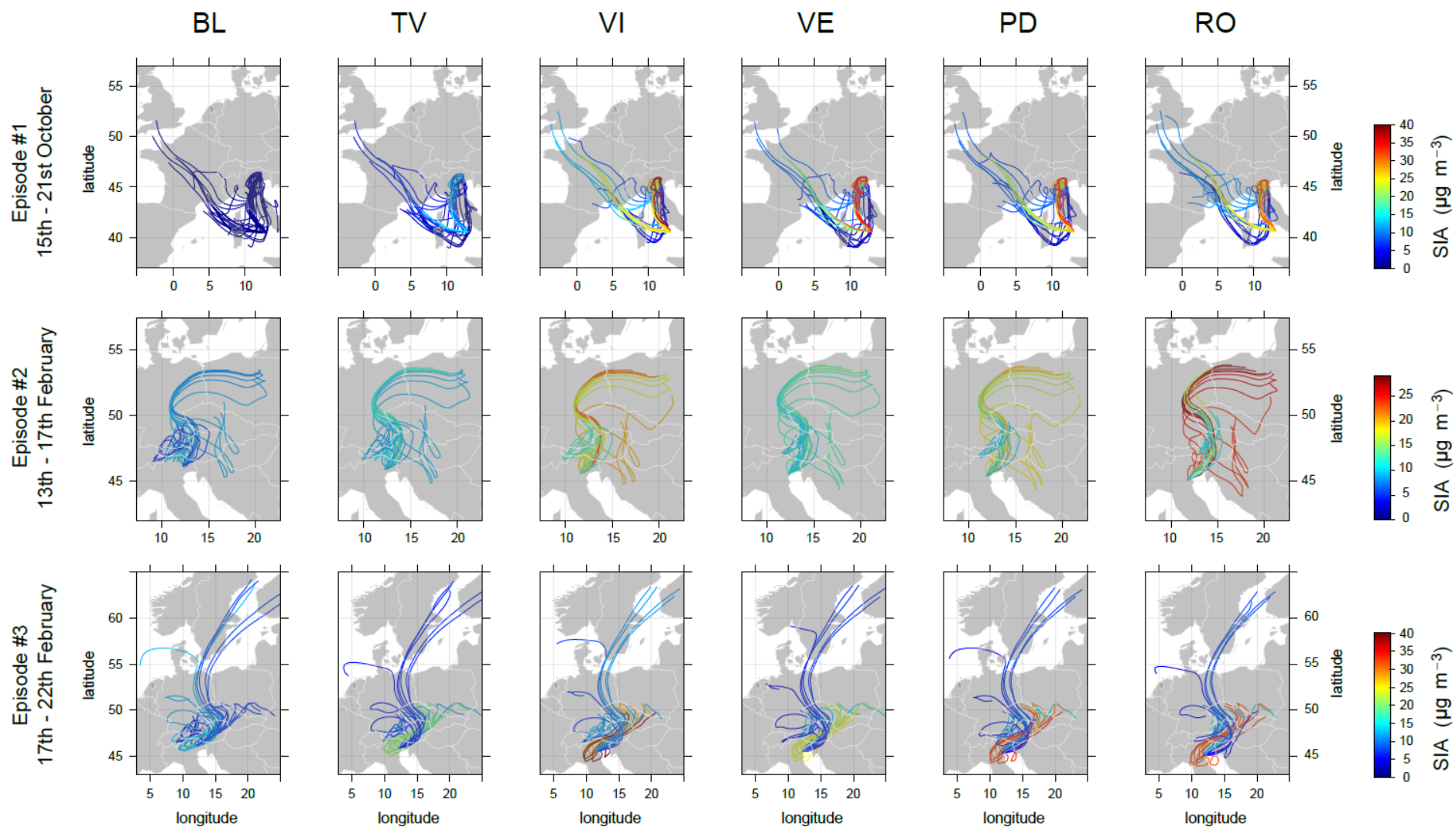


Figure 7. Single back-trajectories during three high-nitrate concentration events.

Graphical abstract

