dec. 2015. The final version is available at DOI 10.1007/11356-015-5862-z

# Atmospheric pollutants in peri-urban forests of *Quercus ilex*: evidence of pollution abatement and threats for vegetation.

- 3 Héctor García-Gomez<sup>1</sup>, Laura Aguillaume<sup>2</sup>, Sheila Izquieta-Rojano<sup>4</sup>, Fernando Valiño<sup>1</sup>, Anna Àvila<sup>3</sup>,
- 4 David Elustondo<sup>4</sup>, Jesús M. Santamaría<sup>4</sup>, AndrésAlastuey<sup>5</sup>, Héctor Calvete-Sogo<sup>1</sup>, Ignacio González-
- 5 Fernández<sup>1</sup>, Rocío Alonso<sup>1</sup>
- <sup>6</sup> <sup>1</sup>Ecotoxicology of Air Pollution, CIEMAT, Av. Complutense 40, Ed.70, 28040 Madrid, Spain.
- 7 <sup>2</sup>CREAF, Campus de Bellaterra (UAB), Edifici C, 08193Cerdanyola del Vallès, Spain.
- 8 <sup>3</sup>Universitat Autònoma de Barcelona (UAB), Campus de Bellaterra, 08193 Cerdanyola del Vallès, Spain.
- 9 <sup>4</sup>LICA, Universidad de Navarra, C. Irunlarrea 1, 31009 Pamplona, Spain.
- <sup>5</sup> Institute of Environmental Assessment and Water Research (IDAEA-CSIC), C. Jordi Girona 18-26, 08034
- 11 Barcelona, Spain.

### 12 Keywords

13 Atmospheric pollution;nitrogen; ozone; aerosols; ecosystem services; Mediterranean vegetation.

### 14 Abstract

- 15 Peri-urban vegetation is generally accepted as a significant remover of atmospheric pollutants, but it could
- 16 also be threatened by these compounds, with origin in both urban and non-urban areas. To characterize the
- 17 seasonal and geographical variation of pollutant concentrations and to improve the empirical understanding
- 18 of the influence of Mediterranean broadleaf evergreen forests on air quality, four forests of *Quercus ilex*
- 19 (three peri-urban and one remote) were monitored in different areas in Spain. Concentrations of nitrogen
- dioxide (NO<sub>2</sub>), ammonia (NH<sub>3</sub>), nitric acid (HNO<sub>3</sub>) and ozone (O<sub>3</sub>) were measured during two years in
- 21 open areas and inside the forests and aerosols  $(PM_{10})$  were monitored in open areas during one year. Ozone 22 was the only air pollutant expected to have direct phytotoxic effects on vegetation according to current 23 thresholds for the protection of vegetation. The concentrations of N compounds were not high enough to
- directly affect vegetation but could be contributing through atmospheric N deposition to the eutrophization
   of these ecosystems. Peri-urban forests of *Quercus ilex* showed a significant below-canopy reduction of
   gaseous concentrations (particularly NH<sub>3</sub>, with a mean reduction of 29–38%), which indicated the
   feasibility of these forests to provide an ecosystem service of air quality improvement. Well-designed
   monitoring programs are needed to further investigate air quality improvement by peri-urban ecosystems
- 29 while assessing the threat that air pollution can pose to vegetation.
- 30

31

#### 32 Acknowledgements

This research was funded by the Spanish project EDEN (CGL2009-13188-C03-02), by the project from Autonomous Government of Madrid AGRISOST-CM (P2013/ABI-2717), and by the European Projects ECLAIRE (FP7-ENV-2011/282910) and Life RESPIRA (LIFE13 ENV/ES/000417). This study was also supported by the Ministry of Agriculture, Food and Environment (Resolución 15398, BOE n° 230). The authors would like to acknowledge the Department of Environment (DGQA) of the Autonomous Government of Catalonia for performing the active monitoring of air pollutants at LC ("MSY" station from GAW/ACTRIS monitoring networks).

#### 1 **1. Introduction**

2 The continuous growth of urban population has turned air quality into one of the main environmental concerns worldwide. Current urban development needs to consider designs and 3 4 strategies that minimize atmospheric pollution to improve well-being and human health. In the 5 last years, particular attention has been paid to investigate the role of urban and peri-urban 6 vegetation in improving air quality. Vegetation can remove air pollutants via dry deposition, through interception in the canopy surfaces, and via absorption of gases through the stomata. In 7 8 particular, urban and peri-urban vegetation has been proposed as a method to reduce air pollutants 9 such as ozone, nitrogen oxides and particulate matter (Alonso et al. 2011; Kroeger et al. 2014; 10 Nowak et al. 2014; Sgrigna et al. 2015). On the other hand, air pollution can affect these forests, 11 impairing their capacity to provide ecosystem services.

12 Peri-urban areas are transition zones between the denser urban core and the rural hinterland, 13 where natural habitats can be exposed to intermediate concentrations of pollutants linked to both 14 urban and rural activities. Among the most common gaseous pollutants, nitrogen oxides (NO<sub>2</sub>, 15 NO) reach peri-urban areas transported from human agglomerations and highways where they are 16 produced as a result of combustion processes. Nitrogen oxides are in turn precursors for the 17 formation of photochemical oxidants such as ozone (O<sub>3</sub>) and nitric acid (HNO<sub>3</sub>). Ozone is one of 18 the most important and pervasive air pollutants currently affecting vegetation (Kroeger et al. 19 2014). This pollutant is particularly important in the Mediterranean region, where the highest 20 concentrations in Europe are registered (EEA 2013). Ozone levels are usually greater in peri-21 urban and rural areas than in busy urban centres, due to its rapid destruction by reacting with the 22 NO emitted in the cities (The Royal Society 2008). Nitric acid is one of the main components of 23 photochemical smog, together with ozone, and with a similar spatial distribution (Bytnerowicz et 24 al. 1999). In contrast, ammonia ( $NH_3$ ) is mainly emitted from agricultural and livestock activities 25 in rural areas. Ammonia and nitric acid can quickly react with each other, or with other 26 atmospheric gases, to formsecondary inorganic aerosols (SIA), that can represent an important 27 fraction of the particulate matter (PM) concentration measured at regional background stations 28 (EEA 2013). Although atmospheric N pollutant levels are usually not high enough to directly 29 damage vegetation, atmospheric N deposition can contribute to both eutrophication and 30 acidification of ecosystems, which is a bigger problem than the direct exposure to these compounds (Dise et al. 2011; EEA 2013). Atmospheric N deposition can be particularly 31 32 important in peri-urban areas that are receiving contributions of N compounds from both urban and agricultural activities. In fact, Mediterranean forests and mountain scrublands close to 33 34 Barcelona and Madrid cities have been reported to be threatened by N deposition (García-Gómez 35 et al. 2014).

Air pollutant gases and particles are removed from the atmosphere through both wet and dry 1 2 deposition. In Mediterranean environments, atmospheric deposition can be dominated by dry deposition, which can represent up to 50-95% of the total deposition in Mediterranean forests 3 4 (Bytnerowicz and Fenn 1996). In this sense, urban and peri-urban vegetation, through increasing dry deposition, can represent a good strategy to improve air quality, particularly in this region. 5 Dry deposition to vegetation is a function of multiple factors, such as air concentration, chemical 6 7 properties of the depositing species, atmospheric turbulence, moisture and reactivity of receptor 8 surfaces, and vegetation structure and activity (Fowler et al. 2009).

9 Measuring pollutant concentrations outside and within peri-urban forests can provide an insight 10 into the role of vegetation in removing air pollutants (Cavanagh et al. 2009; Setälä et al. 2013; Grundström and Pleijel 2014). Although urban vegetation is accepted as an efficient remover of 11 12 air pollutants, most of the studies are based on large-scale modelling (e.g. Nowak et al. 2014) or 13 laboratory studies (e.g. Chaparro-Suárez et al. 2011), but there are few empirical evidences of the 14 reduction in pollutant concentrations inside urban forested areas (Cavanagh et al. 2009; 15 Grundström and Pleijel 2014). Besides, atmospheric pollution represents a risk for the urban and 16 peri-urban vegetation and should be monitored, particularly in forest potentially withstanding 17 other stressful conditions. Interestingly, NH<sub>3</sub> and HNO<sub>3</sub> concentrations are scarcely measured in 18 the main air-quality networks, despite being major drivers of atmospheric N dry deposition to 19 vegetation (Bytnerowicz et al. 2010).

20 In order to study tropospheric O<sub>3</sub>, gaseous N compounds, and suspended PM in peri-urban forests 21 in Spain, three peri-urban forests of holm oak (Quercus ilex L.) were selected near to three cities 22 in Spain with increasing population and with different influences of traffic and agricultural pollution sources (based on their distances to highways, percentage of agricultural land use and 23 24 presence of livestock). Another holm oak forest site, far from anthropogenic emissions of air pollutants, was established for comparison. Holm oak is an evergreen broadleaf tree species 25 26 representative of the Mediterranean Basin and it is present over a wide range of environments in 27 the region, from cold semi-arid to temperate humid bioclimates. This study was enclosed in the 28 EDEN project (Effects of nitrogen deposition in Mediterranean evergreen holm oak forests), 29 whose main goal was to determine and characterize the nitrogen inputs to holm oak forests in the 30 Iberian Peninsula and the effects in the nitrogen biogeochemical cycle. In the present study, air 31 quality measurements from EDEN project are presented and discussed, with the following 32 objectives: 1) to analyse the main air pollutants that could be affecting holm oak forests close to 33 cities, 2) to characterize air pollutant temporal and geographical variation, and 3) to compare air pollutant concentrations outside and inside the forest to improve the empirical understanding of 34 35 the influence of vegetation on air quality.

#### 1 2. Material and methods

#### 2 2.1. Study sites

Three holm-oak (Ouercus ilex) forests were selected in the vicinity of three cities in Spain with 3 4 increasing population (Fig. 1, Table 1). The Can Balasc (CB) site is placed in a forest located in a 5 natural protected area 4 km away from Barcelona with acidic soils and Mediterranean sub-humid 6 climate. The Tres Cantos site (TC) is a forest located in a natural protected area at 9 km from 7 Madrid, growing on acidic sandy soil with Mediterranean semi-arid climate. The Carrascal site 8 (CA) is located in an agricultural area close to Pamplona (15 km), with calcareous soil and 9 Mediterranean humid climate, and it is the most agricultural-influenced among the three peri-10 urban forests. The canopy in all the sites is dominated by Quercus ilex, mixed with Q. humilisin 11 CB. In the case of TC, vegetation was historically managed as a traditional dehesa (a savannah-12 like agrosilvopastoral system) of Q. ilex, but the low management intensity during the last decades has allowed vegetation to grow as a moderately open forest. An additional holm oak 13 14 forest was selected as a non-urban reference in La Castanya (LC), a long-term biogeochemical 15 study site in a protected mountainous area (Parc Natural del Montseny), situated 40 km away from Barcelona (Fig. 1) and is included in the GAW/ACTRIS monitoring networks ("MSY" 16 station). This site presents moderately acidic soils and montane Mediterranean climate and it is 17 18 relatively sheltered from the surrounding lowland sources of atmospheric pollutants (Hereter and 19 Sánchez 1999). The description of the sites was complemented with land use cover and livestock 20 density data obtained from the Corine Land Cover 2006 of the European Environment Agency 21 (http://www.eea.europa.eu/data-and-maps/data/corine-land-cover-2006-raster-3) and from the 22 Spanish National Statistic Institute (http://www.ine.es) respectively (Table 1). ArcGIS software (version 9.2; Environmental Systems Research Institute Inc., Redlands, CA, USA) was employed 23 to summarize these data using a buffer of 25 km radius around the sampling sites. Meteorological 24 variables were monitored in CB, TC and LC sites, and data from the closest meteorological 25 26 station were collected for the CA site.

**27 2.2.** 

#### 2.2. Air pollution monitoring

Atmospheric concentrations of ozone (O<sub>3</sub>), ammonia (NH<sub>3</sub>), nitrogen dioxide (NO<sub>2</sub>) and nitric 28 29 acid vapour (HNO<sub>3</sub>) were monitored during two years using passive samplers. In every location, 30 two plots were installed: an open-field plot (O) and a below-canopy plot (F -forest plot). Open 31 and below-canopy plots were selected in order to maintain the same orientation, exposure and 32 elevation. Two replicate samplers per gaseous species were exposed at 2 m height in each plot. 33 Gases were measured during two-week-long periods between February 2011 and February 2013; 34 except O<sub>3</sub> in CA, where the sampling survey was only extended until April 2012. Exceptionally, 35 some sampling periods (3% of the total monitoring time) lasted approximately four weeks. In 1 these cases, the same result has been used for the two corresponding regular sampling periods..

During every exposure period, unexposed samplers were used as blanks for each site and type of
passive sampler. After collection, all samples were kept refrigerated (4° C) in darkness until they
were analysed in the laboratory.

5 Tube-type samplers (Radiello®) were used to measure atmospheric concentrations of NH<sub>3</sub>, NO<sub>2</sub> 6 and  $O_3$ . Laboratory analyses were performed according to Radiello's specifications (Fondazione 7 Salvatore Maugeri, 2006). Atmospheric concentrations of  $HNO_3$  were measured by means of 8 badge-type samplers manufactured following Bytnerowicz et al. (2005). In CA, Passam® passive 9 samplers and methods were employed during the second year for monitoring NO<sub>2</sub> after checking 10 their comparability with Radiello<sup>®</sup>. For these sampling periods, correction proposed by Plaisance (2011) was applied to avoid biases caused by high wind speeds. The variability of the duplicate 11 12 passive samplers for each air pollutant averaged from 7% for O<sub>3</sub> to 28% for HNO<sub>3</sub>.

Additionally, concentration of  $O_3$  and nitrogen oxides (NO and NO<sub>2</sub>) were continuously 13 monitored in open-field locations in LC and TC sites with active monitors (in LC: MCV<sup>®</sup> 48AV 14 and Thermo Scientific<sup>®</sup> 42i-TL, respectively; in TC:ML<sup>®</sup> 9810B and ML<sup>®</sup> 9841, respectively). 15 Simultaneous measurements with passive samplers and active monitors were used to estimate 16 17 mean experimental sampling rates, which were applied to calculate atmospheric concentrations. 18 The experimental sampling rates obtained in LC were employed in CB and CA calculations as 19 well, after checking the similarity with concentrations registered at the closest air quality 20 monitoring stations.

Using the data from the active monitors, accumulated  $O_3$  exposure was calculated as AOT40, which is the accumulated amount of hourly  $O_3$  concentrations over the threshold value of 40 nl l<sup>-1</sup>. Following the Ambient Air Quality Directive 2008/50/EC, AOT40 was calculated for the period May–July with the hourly mean values from 8 to 20 hours. Additionally, following the recommendations from the Convention on Long-range Transboundary Air Pollution (CLRTAP 2011), AOT40 was calculated for the entire year (the growing season for *Q. ilex*) during daylight hours.

#### 28 2.3. Particulate matter sampling

Particulate matter with diameter up to 10  $\mu$ m (PM<sub>10</sub>) was collected with 150 mm quartz microfibre filters (2500 QAO-UP, Pall Life Sciences) using high volume samplers installed in openfield plots of TC, CA and LC sites (Digitel<sup>®</sup> DH80 in LC -MSY monitoring station; MCV<sup>®</sup> CAV-A/mb in TC and CA). Samples were collected from February 2012 to February 2013 once a week,using a flow of 30 m<sup>3</sup> h<sup>-1</sup> during 24-h periods. The day of the week for PM<sub>10</sub> collection changed weekly. The concentration was gravimetrically determined and main secondary inorganic aerosols (SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup>) were water-extracted and analysed by ion 1 chromatography. For statistical comparison purposes with gaseous pollutant concentrations,  $PM_{10}$ 2 data were grouped and averaged in accordance to passive sampling periods (except for the 3 comparison of the natural dust events with the rest of the samples).

#### 4 2.4. Statistical analysis

5 Non-parametric statistics was selected for this study because most of the variables did not show a 6 normal distribution according to Shapiro-Wilk test and normal probability plots. Differences 7 among seasons or sites were analysed using the Kruskal-Wallis test; when significant differences 8 were found, differences between pairs of sites were assessed with the Mann-Whitney U test. 9 Correlation between variables was tested with the Spearman rank order correlation coefficient. 10 Differences in pollutant concentration between O and F plots were analysed by applying the 11 Wilcoxon matched pair test to the entire sampling period. The temporal variability is described in 12 this study by the coefficient of variation (CV = standard deviation / mean) of the two-week concentrations for the entire study period. The variability of the duplicate passive samplers for 13 each air pollutant is also described by their respective CV. In this work, seasons were considered 14 15 as periods of three consecutive months, beginning on 1<sup>st</sup> January. Statistica software (version 12; StatSoft, Tulsa, OK) was used for statistical analysis. Alfa level was set at 0.05. 16

17

#### 18 **3. Results**

#### 19 **3.1.** Temporal and spatial patterns of gaseous pollutants

Seasonal and annual pollutant concentrations and differences among sites are described below
based on concentrations in the O plots (Fig. 2; Table 2).

The annual mean of atmospheric NO<sub>2</sub> concentration ranged from 4.3  $\mu$ g m<sup>-3</sup> in LC to 16.2  $\mu$ g m<sup>-3</sup> in CB (Table 2). The highest two-week concentration reached 39.3 and 37.1  $\mu$ g m<sup>-3</sup> registered in CB and TC respectively during the winter 2012 (Supplement, S1). On average for the four sites, temporal variability of NO<sub>2</sub> concentration was 53%. Levels of NO<sub>2</sub> tended to peak during the coldest seasons (autumn and winter). Significant seasonal differences were detected in the sites closest to the big cities of Barcelona and Madrid (CB and TC). LC experienced the lowest concentrations and the lowest inter-seasonal variability (Fig. 2).

Atmospheric NH<sub>3</sub> concentration (Table 2) was the highest in CA (2.5  $\mu$ g m<sup>-3</sup>) and the lowest in TC and LC (0.7  $\mu$ g m<sup>-3</sup>). The maximum two-week value (5.3  $\mu$ g m<sup>-3</sup>)was recorded in CA during late winter (Supplement, S2). The temporal variability showed a mean of 55% across sites. A consistent seasonal pattern was found in TC, where NH<sub>3</sub> concentration increased during spring and summer and decreased during autumn and winter (Fig. 2; Supplement, S2). LC showed a similar seasonal pattern but differences were not statistically significant (p = 0.06). On the contrary, in CB and CA, the highest seasonal concentrations occurred in winter.

1 The concentration of  $HNO_3$  tended to be higher in the sites closest to the Mediterranean coast 2 (CB and LC), but differences among sites were not statistically significant (Table 2). The maximum two-week concentrations found in CB and LC (14.5 and 13.9 µg m<sup>-3</sup> in summer of 3 2012, respectively) were twice the maximum values found in TC and CA (Supplement, S3). The 4 5 temporal variability in HNO<sub>3</sub> concentration was higher than the variability found for the other air pollutants, with an average value of 110%. A general seasonal pattern was detected in HNO<sub>3</sub> 6 7 concentrations, with higher values during spring and summer and lower values in autumn and 8 winter (Fig. 2).

9 The annual mean of atmospheric  $O_3$  concentrations (Table 2) were significantly lower in the sites closest to the big cities of Barcelona and Madrid (57.0  $\mu$ g m<sup>-3</sup> in CB and 69.1  $\mu$ g m<sup>-3</sup> in TC) than 10 in the more rural ones (77.4  $\mu$ g m<sup>-3</sup> and 78.2  $\mu$ g m<sup>-3</sup> in CA and LC, respectively). Ozone was the 11 air pollutant showing the smallest temporal variability with a mean value of 32%. All sites 12 13 showed similar seasonal patterns with higher  $O_3$  concentration during spring and summer than in 14 autumn and winter (Fig. 2). Ozone exposure accumulated during May-July expressed as AOT40 15 ranged from 3.9 ppm h in CA in 2011 to 28.3 ppm h in TC in 2012 (Table 3). When accumulating 16 O<sub>3</sub> exposure throughout the growing season, AOT40 values ranged from 8.2 ppm h in CA in 17 2011 to 49.6 ppm h in TC in 2012 (Table 3).

#### 18 **3.2.** Temporal and spatial patterns of particulate matter

19 The concentration of PM<sub>10</sub> was higher in CA and TC than in LC (Table 2), although differences 20 were only significant between CA and LC, which showed the lowest annual concentration (18.0 21  $\mu$ g m<sup>-3</sup>). Temporal variability in PM<sub>10</sub> concentrations was 50% on average for the three sites. 22 Significant seasonal variations were found in TC and LC, with the highest  $PM_{10}$  concentrations 23 registered in summer and the lowest in autumn (Fig. 3A). Saharan dust events represented 10% of 24 the total amount of samples, and occurred more frequently during the summer season. In the three sites, the highest 24h-concentrations of  $PM_{10}$  (up to 126.4 µg m<sup>-3</sup>) were collected during these 25 26 natural dust events, generally doubling the levels found in the rest of the samples (Fig. 3B).

27 Regarding SIA composition, no differences among sites were found in particulate ammonium 28  $(NH_4^+)$ , while particulate nitrate  $(NO_3^-)$  was significantly the highest in CA (Table 2). Apparently, Saharan dust intrusions did not affect the  $NH_4^+$  and  $NO_3^-$  concentration in  $PM_{10}$  (data not shown). 29 The atmospheric concentration of both water-soluble nitrogen aerosols showed a marked 30 31 seasonality, with higher values detected in winter than in the rest of seasons (Figs. 3C and 3D). 32 However, only for NO<sub>3</sub><sup>-</sup> in CA and LC, these differences were statistically significant. Gaseous 33 nitrogen forms generally predominated over the particulate forms, particularly in spring and 34 summer (Figs. 3E and 3F). However,  $NO_3^-$  clearly predominated over HNO<sub>3</sub> during winter in TC and CA and during autumn in LC, and  $NH_4^+$  predominated over  $NH_3$  during winter in TC. 35

Additionally, no seasonal variations were recorded in ammonium gas/particle ratio in CA (Fig. 3F).

## 3 3.3. Differences in gaseous pollutant concentrations between open-field and below-canopy 4 plots

5 Below-canopy concentrations of gaseous pollutants were, in general, smaller than levels found in 6 the open-field plots (Fig. 4). These differences were more remarkable for  $NH_3$ , which showed an 7 annual mean concentration in F plots 40% lower than in the O plots in average for the four sites 8 (56% in LC, and 29–38% in the peri-urban forests). In the case of NO<sub>2</sub>, differences were not 9 significant in CB, while the concentrations were significantly lower in the F plots in the rest of sites (41% in CA, 13% in TC and 6% in LC). For HNO<sub>3</sub>, the reduction detected inside the forest 10 was significant in TC and CA, showing average concentrations 11-13% lower in the F plot 11 compared to the O plot. Ozone concentrations were significantly lower inside the forests in TC 12 13 and LC (annual mean difference of 7% and 5%, respectively).

14 The reduction of air pollutant concentrations inside the forest showed few evident seasonal 15 patterns. Nitrogen dioxide experienced the highest decrease in concentrations below-canopy 16 (Supplement, S1) during autumn and winter in TC and CA (none and 34% on average for both 17 seasons, respectively), while in LC this difference was larger in spring (18%). The differences in 18 NH<sub>3</sub> levels were consistent most of the time (31% on average; Supplement, S2), although smaller 19 during the summer in the three peri-urban forests. Regarding  $HNO_3$  (Supplement, S3), differences 20 between forest and open plots were slightly higher during spring and autumn in TC and CA (24% 21 in both sites, averaged for both seasons). The reduction of O<sub>3</sub> concentrations inside the forest 22 resulted slightly larger during summer and autumn (8% in TC and 7% in LC, averaged for both 23 seasons; Supplement, S4).

#### 24 3.4. Correlation analysis of pollutant concentrations and meteorology

25 Atmospheric concentrations of  $NO_2$  were poorly correlated with meteorological variables, with the exception of TC site, where NO<sub>2</sub> levels were negatively correlated to temperature, daily solar 26 radiation and wind speed, and positively correlated to relative humidity. In the rest of sites, NO<sub>2</sub> 27 concentrations were negatively correlated with precipitation in CB and LC, and with wind speed 28 29 in CA (Table 4). In the case of NH<sub>3</sub> concentrations, no correlation was found in CA. In the other 30 sites, relative humidity was negatively correlated to NH<sub>3</sub> concentration, while temperature and daily solar radiation were positively correlated in TC and LC, and negatively in CB. 31 Concentrations of HNO<sub>3</sub> and O<sub>3</sub> were positively correlated with temperature and daily solar 32 33 radiation, and negatively with relative humidity in all sites. Besides, HNO3 and O3 concentrations

1 showed a positive correlation with wind speed in TC and CA, and a negative correlation with

2 precipitation in TC (Table 4).

The concentrations of  $PM_{10}$  were negatively correlated with precipitation in TC and CA and positively with solar radiation and temperature in TC and LC. In TC,  $PM_{10}$  was also negatively correlated with humidity. Besides,  $PM_{10}$  was negatively correlated with wind speed in LC. Particulate nitrate was negatively related to temperature and solar radiation only in CA.  $NH_4^+$ concentrations did not show important correlations with meteorological variables. Particulate  $SO_4^{2-}$  was positively correlated to temperature and solar radiation and negatively with wind speed only in LC (Table 4).

10 No significant correlations among gaseous pollutant were found in CA. In the other sites,  $O_3$  and HNO<sub>3</sub> concentrations were positively correlated (Table 4). In TC, O<sub>3</sub> was also negatively 11 12 correlated to NO<sub>2</sub> and NH<sub>3</sub> was positively correlated to O<sub>3</sub> and HNO<sub>3</sub>. Particulate NH<sub>4</sub><sup>+</sup> concentration was correlated with particulate  $NO_3^-$  in the three sites, and with  $SO_4^{2-}$  in CA and 13 LC. However, NH<sub>4</sub><sup>+</sup> was not correlated with NH<sub>3</sub> in any of the sites. Particulate nitrate was 14 15 positively related to NO<sub>2</sub> in TC and CA, and negatively correlated with HNO<sub>3</sub> only in CA (Table 4). Ammonia and HNO<sub>3</sub> concentrations were positively correlated to PM<sub>10</sub> in TC and LC. Finally, 16 scarce significant correlations with meteorological variables were found for the below-canopy 17 18 reductions of atmospheric pollutant concentrations (data not shown).

19

#### 20 4. Discussion

#### 21 4.1. Air pollution affecting peri-urban forests

22 The annual mean of atmospheric NO<sub>2</sub> concentrations decreased from CB to LC (from 16.2 to 4.3  $\mu$ g m<sup>-3</sup>), indicating an order of influence of urban and traffic emissions (CB > TC  $\ge$  CA > LC). 23 24 The levels of NO<sub>2</sub> in the three peri-urban forests (CB, TC and CA) were in the range of values 25 recorded in suburban background monitoring stations in 2012 (AirBase v8 dataset; EEA 2014). 26 Therefore, suburban stations might be considered representative of NO<sub>2</sub> concentration registered in peri-urban forests. Concentrations of NO<sub>2</sub> in the three peri-urban forests followed the expected 27 seasonal pattern of monitoring stations influenced by urban emissions, with highest values 28 29 recorded during autumn and winter. This seasonal pattern is associated with increasing emissions 30 due to urban combustion for heating purposes and with the lower photochemical intensity during the cold season (Karanasiou et al. 2014). The decrease of NO<sub>2</sub> with wind speed in TC and CA 31 32 pointed to a higher influence of local sources rather than regional contribution. Similar results 33 have been reported in other Mediterranean urban sites (Karanasiou et al. 2014). An analogous response would be expected at CB, but the higher urban density around the site and the lower 34 wind speed (annual mean of 0.8 m s<sup>-1</sup>) could be impairing pollutant dispersion. The forest site in 35

LC was more representative of background NO<sub>2</sub> concentrations, since the annual mean was close to the average value of  $3.7-3.5 \ \mu g \ m^{-3}$  recorded in background stations in Spain in 2011 and 2012 respectively (MAGRAMA 2014). Moreover, NO<sub>2</sub> concentrations in LC did not show clear seasonal variations, demonstrating the lack of influence of urban emissions. After adding the estimated NO concentration (from the active monitors), none of the sites are expected to reach the critical level for the protection of vegetation (30  $\mu g \ m^{-3}$ , as annual mean) established in the European Air Quality Directive.

8 The annual mean of NH<sub>3</sub> concentrations in CB, TC and LC were low and similar to the levels recorded in Spanish background stations (0.9 µg m<sup>-3</sup> in 2012; Hjellbrekke 2014). These values 9 10 were lower than concentrations measured in urban backgrounds of their respective closest cities (1.7 µg m<sup>-3</sup> in Madrid and 7.3 µg m<sup>-3</sup> in Barcelona; Reche et al. 2014), and far from levels 11 registered in regions with intensive farming or livestock (up to 60 µg m<sup>-3</sup>; Fowler et al. 1998; 12 Pinho et al. 2012). The higher concentrations found in CA (annual mean of 2.5 µg m<sup>-3</sup>) probably 13 14 is related to the presence of livestock in the nearby area. The seasonal pattern of NH<sub>3</sub> 15 concentrations in TC and LC, with higher values during spring and summer, could be explained 16 by an increasing volatilisation and emission of NH<sub>3</sub> from biological sources under warm 17 conditions. In the case of CB, the highest values recorded in autumn and winter might be related to the emissions of NH<sub>3</sub> from an industrial area 6.5 km west of CB. Concentrations of NH<sub>3</sub> at this 18 site were significantly correlated with west winds (p < 0.01; data not shown), the most frequent 19 20 wind in autumn and winter. The winter maxima NH<sub>3</sub> levels in CA were in agreement with the fertilization practices of cereal crops in the region during this season. Since the annual mean of 21  $NH_3$  concentrations did not exceed the 3  $\mu$ g m<sup>-3</sup> critical level proposed for the protection of higher 22 plants in any of the sites, these forests are not expected to experience relevant ammonia pollution 23 effects (CLRTAP 2011). Moreover, the critical level of 1 µg m<sup>-3</sup> for the protection of lichens and 24 bryophytes (Cape et al. 2009; CLRTAP 2011) was only exceeded in CA. 25

26 No significant differences in HNO<sub>3</sub> annual concentration were detected among the sites included 27 in this study. The concentrations of HNO<sub>3</sub> in the three peri-urban forests were in the range of values found in other peri-urban areas in the Mediterranean region (summer values of 2.8-4.2 µg 28 m<sup>-3</sup>; Danalatos and Glavas 1999) and higher than in urban sites (yearly averaged values of 0.8–1.5 29 µg m<sup>-3</sup>; Anatolaki and Tsitouridou 2007; Tzanis et al. 2009). However, even the highest 30 31 concentrations were below the values reported in forested areas of San Bernardino Mountains in 32 Southern California, where topography, climate and emissions linked tohigh population favour HNO<sub>3</sub> formation (Bytnerowicz and Fenn 1996; Jovan et al. 2012). The typical higher HNO<sub>3</sub> 33 values recorded during spring and summer in the study sites can be explained by the 34 35 photochemical origin of this pollutant (Bytnerowicz et al. 2010; Tzanis et al. 2009). In this sense, 36 positive correlations between solar radiation and HNO<sub>3</sub> concentration were found for all the sites.

The highest levels were found in LC, which must respond to pollutant-transport mechanisms 1 2 rather than to an in-situ formation of HNO<sub>3</sub>, since this is a rural site with low concentration of NO<sub>2</sub> (chemical precursor of HNO<sub>3</sub>). In fact, ageing of air masses over the Iberian Peninsula and 3 4 recirculation along the Mediterranean coast have been reported as processes increasing levels of 5 oxidants, acidic compounds, aerosols and ozone (Escudero et al. 2014; Millán et al. 2002). Although very little information is available on direct effects of HNO<sub>3</sub> on vegetation, the 6 7 concentrations found in this study are much lower than the levels reported for epicuticular 8 damage (Padgett et al. 2009).

9 The annual mean concentration of O<sub>3</sub>increased from CB to LC, following an opposite order of 10 urban influence to the one found for NO2 concentration. A similar behaviour has been described in other studies around cities in the Mediterranean area (Domínguez-López et al. 2014; Escudero 11 12 et al. 2014). CB showed an annual mean similar to values found in 2012 in Spanish suburban 13 areas, while the other sites showed values clearly typical of rural areas (means of 59.0 and  $67.8 \mu g$ 14  $m^{-3}$ , respectively; EEA 2014). Ozone concentrations in the peri-urban forests showed the typical 15 seasonal variations with higher levels during spring and summer, responding to the sum of the 16 hemispheric-scale spring maximum, the increased photochemical production and transport 17 processes, as well as the above mentioned ageing of air masses and recirculation (Cristofanelli 18 and Bonasoni 2009; Millán et al. 2002). In fact, ozone concentrations were significantly 19 correlated with temperature and solar radiation. Besides, the emission of biogenic volatile organic 20 compounds (BVOCs) by vegetation is known to be correlated with temperature, and can 21 exacerbate photochemical reactivity, and thus  $O_3$  formation (Calfapietra et al. 2013). All the 22 calculated AOT40 values were above the concentration-based  $O_3$  critical level proposed by the 23 CLRTAP for protecting forest trees (5 ppm h for the growing season; CLRTAP 2011). The 24 threshold levels for the protection of vegetation established in the European Directive 2008/50/EC 25 (9 ppm h for the period May–July) were also overreached, with the exception of CB site in 2011. 26 Moreover, experimental values of AOT40 similar to those found in this study have been proved to 27 cause a decrease of growth in seedlings of Q. ilex (Alonso et al. 2014; Gerosa et al. 2015).

In the two peri-urban forests with aerosol measurements (TC and CA), the annual mean 28 concentrations of PM<sub>10</sub> were close to the urban background levels measured in Spanish big cities 29 in 2012 (mean of 26 µg m<sup>-3</sup>; MAGRAMA, 2014), and well above the values measured in Spanish 30 background stations (12.9 µg m<sup>-3</sup>; Hjellbrekke 2014). On the other hand, concentrations of 31 particulate NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup> were similar to the national background levels in TC (1.2  $\mu$ g NO<sub>3</sub><sup>-</sup> m<sup>-3</sup>, 32 and 0.4  $\mu$ g NH<sub>4</sub><sup>+</sup> m<sup>-3</sup>; Hjellbrekke 2014), but almost double in CA. The increased concentration of 33 34  $NO_3^-$  and  $NH_4^+$  in CA could respond to the elevated  $NH_3$  concentration caused by agricultural 35 activities, which, combined with the low temperatures, facilitates the formation and stability of 36 ammonium nitrate (NH4NO3). Moreover, at this site, NO3<sup>-</sup> and HNO3showed a negative

correlation, suggesting the existence of conversion of one into the other. The seasonality in PM<sub>10</sub> 1 2 is in agreement with previous studies that attributed the higher summer concentrations to low 3 precipitation, high resuspension, photochemical oxidation and higher frequency of Saharan dust 4 outbreaks (Escudero et al. 2005; Querol et al. 2008; Rodríguez et al. 2002). Interestingly, the natural events of Saharan dust did not modify  $NO_3^-$  and  $NH_4^+$  concentrations. The seasonality 5 observed on particulate N compounds was more related with the thermal instability of NH<sub>4</sub>NO<sub>3</sub>, 6 7 pointing out the importance of temperature-dependent processes within the SIA in the 8 Mediterranean region (Querol et al. 2008; Pey et al. 2009). Gaseous HNO<sub>3</sub> and NH<sub>3</sub> predominated 9 over particulate forms most of the year but aerosol fraction was important mainly during winter. 10 This seasonal variation in gas/aerosol ratios may have implications for N dry deposition 11 estimations and, therefore, should be further investigated. Little information is available on direct 12 effects of particles on vegetation and no threshold of aerosol concentration has been defined yet 13 for the protection of vegetation.

14 According to the established thresholds and the available scientific evidences, the results indicate 15 that O<sub>3</sub> is the only air pollutant considered in this work which is expected to have direct 16 phytotoxic effects on vegetation. The concentrations of N compounds seemed to be not high 17 enough to directly affect vegetation but could be contributing through atmospheric N deposition 18 to the eutrophization of these ecosystems. Moreover, although evergreen broadleaf Mediterranean 19 woody species are assumed to be tolerant to air pollution due to their sclerophyllic adaptations, 20 recent publications suggest that the addition and interaction of different stress factors (O<sub>3</sub>, N 21 deposition, drought) can be affecting the growth of the trees (Alonso et al. 2014; Gerosa et al. 22 2015) and accompanying pastures (Calvete-Sogo et al. 2014). Thus, monitoring of nitrogen 23 compounds such as NH<sub>3</sub> and HNO<sub>3</sub> should be incorporated into air quality monitoring networks.

#### 24 4.2. Below-canopy reduction of atmospheric pollutant concentrations

25 Air pollutant concentrations measured outside and inside the forest (O and F plots) were 26 compared to analyse the influence of vegetation in air quality. In general, the pollutants 27 considered showed lower concentrations inside the forests. Below-canopy reduction of  $NO_2$ 28 concentration in our study sites ranged from none in CB, to 41% in CA. This high reduction 29 detected in CA could be enhanced by the location of the sampling plots, which were at the same distance, but on the opposite sides of a highway. As a result, the O and F plots were located 30 31 downwind and upwind from the highway, respectively, in relation to predominant winds 32 (Supplement, Figure S5). Statistically significant reductions of  $NO_2$  concentrations inside holm 33 oak forests were found in TC and LC, with averaged values of 13% and 6%, respectively. These reductions are comparable to (Grundström and Pleijel 2014) or higher than (Harris and Manning 34 35 2010; Setälä et al. 2013) values reported in similar empirical studies with deciduous forest

species. The larger differences in NO<sub>2</sub> levels in LC were detected during spring, the time when 1 2 holm oak forests usually show higher stomatal conductance (Alonso et al. 2008). Other authors 3 have reported that  $NO_2$  deposition onto forest canopy is governed by plant stomatal aperture 4 (Chaparro-Suárez et al. 2011; Sparks 2009). This behaviour was not observed in TC and CA, where the highest reductions were found during autumn and winter, suggesting that other 5 atmospheric and biogeochemical interactions could be implicated and need further research. In 6 7 this sense, the lack of below-canopy reduction in CB could not be explained by meteorological 8 variables or different pollutant exposure. Other authors have suggested that NO emissions from 9 forest soil in areas with high O<sub>3</sub> levels, could result in the formation of NO<sub>2</sub> below the canopy 10 (Harris and Manning, 2010; Fowler, 2002), diminishing the difference of NO<sub>2</sub> concentrations 11 between outside and inside the canopy. Since dry deposition of atmospheric pollutants depends 12 on multiple factors such as micrometeorology, spatial heterogeneity, plant structure and 13 physiology, and biochemical interaction, further research is needed to clarify the influence of 14 vegetation on air quality.

15 Below-canopy concentrations of NH<sub>3</sub> were on average 40% lower than in the open field, 16 suggesting that holm oak forests act as sinks of ammonia. This difference was relatively higher in 17 the most natural forest (56% in LC) than in the peri-urban ones (29–38%). Since  $NH_3$  stomatal 18 fluxes are bi-directional, emission or deposition of NH<sub>3</sub> will occur depending on ecosystem N-19 status, stomatal conductance, and the ratio between atmospheric and canopy  $NH_3$  concentration 20 (Behera et al. 2013; Fowler et al. 2009). The below-canopy reductions of  $NH_3$  were consistent 21 throughout most of the year, but smaller during the summer, a period of low plant physiological 22 activity in this type of forest. These results indicate a certain regulation of NH<sub>3</sub> fluxes by stomatal 23 uptake. However,  $NH_3$  canopy retention was not the highest in spring, when plants usually 24 experience maximum stomatal conductance, thus other mechanisms must affect the overall 25 ammonia retention by the canopy in autumn and winter. Among other major drivers of 26 atmospheric  $NH_3$  deposition into the canopy, leaf area density, and leaf surface wetness and 27 acidity can enhance the deposition onto the cuticles and epiphytic communities (Geiser et al. 28 2010; Massad et al. 2010).

29 The differences in HNO<sub>3</sub> concentration between O and F plots were only significantly detected in 30 TC and CA, with reductions of 11-13% on annual average. Among the N gaseous pollutants, 31 HNO<sub>3</sub> is supposed to have the highest surface deposition velocity due to its highly reactive and 32 soluble nature, which should lead to large rates of deposition onto leaf surfaces (Fowler et al. 33 2009). However, the rates of bellow-canopy HNO<sub>3</sub> reduction are similar to those of NO<sub>2</sub> in TC and LC, and lower than those of NH<sub>3</sub>. No clear seasonal patterns were found in the below-canopy 34 35 reduction of HNO<sub>3</sub> concentrations that could indicate the main processes involved in HNO<sub>3</sub> dry 36 deposition in these forests.

In regards to O<sub>3</sub> concentrations, urban and peri-urban vegetation has been proposed as a strategy 1 2 to absorb O<sub>3</sub> and diminish atmospheric concentrations (Alonso et al. 2011; Kroeger et al. 2014). In our study, O<sub>3</sub> levels were significantly reduced inside the forests in TC and LC with an average 3 4 decrease of 5–7%. The largest below-canopy reduction of  $O_3$  concentration occurred in summer 5 and autumn, suggesting that stomatal uptake was not the only process involved in this decline, 6 since stomatal conductance in usually low during the summer in these forests due to drought 7 stress. Actually, non-stomatal O<sub>3</sub> deposition in holm oak forests has been reported to account up 8 to ca. 60 % of the total ozone flux (Fares et al. 2014). Surface wetness of the canopy and other 9 forest surfaces can enhance non-stomatal deposition of O<sub>3</sub> (Altimir et al. 2006). This process 10 could explain the higher reductions of  $O_3$  detected during autumn, the wettest season in all the 11 sites. Besides, increased BVOCs emissions linked to high temperatures during the summer could be favouring the photochemical production of O<sub>3</sub> (Calfapietra et al. 2013). This formation of O<sub>3</sub> 12 13 should be more apparent in the open-field plots due to their higher insolation, increasing the 14 difference in O<sub>3</sub> concentrations between O and F plots during this season.

15

#### 16 **5.** Conclusions

17 Peri-urban forests are exposed to air pollutants coming from both urban and rural activities. Ozone concentrations around Spanish cities are high enough to directly impact peri-urban 18 19 vegetation. The concentrations of N compounds would no directly threat vegetation, but could be 20 contributing, through atmospheric N deposition, to the eutrophization of these ecosystems. 21 Besides, the interaction of different stress factors (O<sub>3</sub>, N deposition, drought) could be affecting 22 plant growth and ecosystem functioning. On the other hand, peri-urban forests of Quercus ilex 23 have proved to experience a significant below-canopy reduction of pollutant concentrations, 24 particularly of NH<sub>3</sub>, but also of NO<sub>2</sub>, HNO<sub>3</sub> and O<sub>3</sub>. These results provide scientific evidence of 25 the ability of these ecosystems to improve air quality in urban agglomerations, but further 26 research is still needed to quantify the relevance of this ecosystem service. The high variability 27 found in this study across sites and seasons points that processes and environmental factors 28 involved in air pollution removal must be characterized in order to manage these forest for 29 improving air quality. Well-designed monitoring programs of urban and peri-urban forests could accomplish both objectives of further investigate air quality improvement while assessing the 30 31 threat that air pollution can pose to vegetation.

#### 1 References

- 2 Alonso R, Elvira S, González-Fernández I, et al (2014) Drought stress does not protect Quercus 3 *ilex* L. from ozone effects: Results from a comparative study of two subspecies differing in 4 ozone sensitivity. Plant Biol 16:375-384. doi: 10.1111/plb.12073 5 Alonso R, Elvira S, Sanz MJ, et al (2008) Sensitivity analysis of a parameterization of the 6 stomatal component of the DO<sub>3</sub>SE model for *Quercus ilex* to estimate ozone fluxes. Environ 7 Pollut 155:473-480. doi: 10.1016/j.envpol.2008.01.032 8 Alonso R, Vivanco MG, González-Fernández I, et al (2011) Modelling the influence of peri-9 urban trees in the air quality of Madrid region (Spain). Environ Pollut 159:2138-2147. doi: 10 10.1016/j.envpol.2010.12.005 Altimir N, Kolari P, Tuovinen J-P, et al (2006) Foliage surface ozone deposition: a role for 11 12 surface moisture? Biogeosciences Discuss 2:1739-1793. doi: 10.5194/bgd-2-1739-2005 13 Anatolaki C, Tsitouridou R (2007) Atmospheric deposition of nitrogen, sulfur and chloride in Thessaloniki, Greece. Atmos Res 85:413-428. doi: 10.1016/j.atmosres.2007.02.010 14 15 Behera SN, Sharma M, Aneja VP, Balasubramanian R (2013) Ammonia in the atmosphere: A review on emission sources, atmospheric chemistry and deposition on terrestrial bodies. 16 17 Environ SciPollut Res 20:8092-8131. doi: 10.1007/s11356-013-2051-9 Burkhardt J, Kaiser H, Kappen L, Goldbach HE (2001) The possible role of aerosols on stomatal 18 conductivity for water vapour. Basic ApplEcol 2:351-364. doi: 10.1078/1439-1791-00062 19 20 Bytnerowicz A, Fenn ME (1996) nitrogen deposition in California forests : A review. Environ 21 Pollut 92:127–146. 22 Bytnerowicz A, Fraczek W, Schilling S, Alexander D (2010) Spatial and temporal distribution of 23 ambient nitric acid and ammonia in the Athabasca Oil Sands Region, Alberta. J Limnol 24 69:11-21. doi: 10.3274/JL10-69-S1-03 25 Bytnerowicz A, Padgett P, Percy K, et al (1999) Direct effects of nitric acid on forest trees. In: 26 Miller PR (ed) Oxidant Air Pollution Impacts in the Montane Forests of Southern California 27 (Ecological Studies 134). Springer, New York, pp 270–287 28 Bytnerowicz A, Sanz M, Arbaugh M, et al (2005) Passive sampler for monitoring ambient nitric 29 acid (HNO) and nitrous acid (HNO) concentrations. Atmos Environ 39:2655-2660. doi: 30 10.1016/j.atmosenv.2005.01.018 Calfapietra C, Fares S, Manes F, et al (2013) Role of Biogenic Volatile Organic Compounds 31 32 (BVOC) emitted by urban trees on ozone concentration in cities: A review. Environ Pollut
- 33 183:71–80. doi: 10.1016/j.envpol.2013.03.012

1	Calvete-Sogo H, Elvira S, Sanz J, et al (2014) Current ozone levels threaten gross primary
2	production and yield of Mediterranean annual pastures and nitrogen modulates the response.
3	Atmos Environ 95:197–206. doi: 10.1016/j.atmosenv.2014.05.073
4	Cape JN, van der Eerden LJ, Sheppard LJ, et al (2009) Evidence for changing the critical level for
5	ammonia. Environ Pollut 157:1033-7. doi: 10.1016/j.envpol.2008.09.049
6	Cavanagh JAE, Zawar-Reza P, Wilson JG (2009) Spatial attenuation of ambient particulate
7	matter air pollution within an urbanised native forest patch. Urban For Urban Green 8:21-
8	30.doi: 10.1016/j.ufug.2008.10.002
9	Chaparro-Suarez IGG, Meixner FXX, Kesselmeier J (2011) Nitrogen dioxide (NO <sub>2</sub> ) uptake by
10	vegetation controlled by atmospheric concentrations and plant stomatal aperture. Atmos
11	Environ 45:5742–5750. doi: 10.1016/j.atmosenv.2011.07.021
12	CLRTAP (2011) Mapping critical levels for vegetation. In: UNECE Convention on Long-range
13	Transboundary Air Pollution (ed) Manual on Methodologies and Criteria for Modelling and
14	Mapping Critical Loads & Levels and Air Pollution Effects, Risks and Trends. Available at:
15	www.icpmapping.org
16	Cristofanelli P, Bonasoni P (2009) Background ozone in the southern Europe and Mediterranean
17	area: Influence of the transport processes. Environ Pollut 157:1399-1406. doi:
18	10.1016/j.envpol.2008.09.017
19	Danalatos D, Glavas S (1999) Gas phase nitric acid, ammonia and related particulate matter at a
20	Mediterranean coastal site, Patras, Greece.Atmos Environ 33:3417-3425. doi:
21	10.1016/S1352-2310(98)00342-2
22	Dise NB, Ashmore M, Belyazid S, et al (2011) Nitrogen as a threat to European terrestrial
23	biodiversity. In: Sutton MA, et al (eds) The European nitrogen assessment. Sources, effects
24	and policy perspectives. Cambridge University Press, pp 463-494
25	Domínguez-López D, Adame JA, Hernández-Ceballos MA, et al (2014) Spatial and temporal
26	variation of surface ozone, NO and NO2 at urban, suburban, rural and industrial sites in the
27	southwest of the Iberian Peninsula. Environ Monit Assess 186:5337-5351. doi:
28	10.1007/s10661-014-3783-9
29	EEA (2013) Air quality in Europe - 2013 report (EEA Report No 9/2013).
30	EEA (2014) Airbase v8. Available at: http://www.eea.europa.eu/data-and-maps/data/airbase-the-
31	european-air-quality-database-8 (last accessed 15.10.14), European Environmental Agency
32	Escudero M, Castillo S, Querol X, et al (2005) Wet and dry African dust episodes over eastern
33	Spain.J Geophys Res D Atmos 110:1–15.doi: 10.1029/2004JD004731

- 1 Escudero M, Lozano A, Hierro J, et al (2014) Urban influence on increasing ozone concentrations 2 in a characteristic Mediterranean agglomeration. Atmos Environ 99:322-332. doi: 3 10.1016/j.atmosenv.2014.09.061 4 Fares S, Savi F, Muller J, et al (2014) Simultaneous measurements of above and below canopy 5 ozone fluxes help partitioning ozone deposition between its various sinks in a Mediterranean 6 Oak Forest. Agric For Meteorol 198-199:181-191. doi: 10.1016/j.agrformet.2014.08.014 7 Fondazione Salvatore Maugeri (2006) Instruction manual for Radiello sampler. Edition 01/2006. 8 http://www.radiello.com 9 Fowler D, Pilegaard K, Sutton MA, et al (2009) Atmospheric composition change: Ecosystems-10 Atmosphere interactions. Atmos Environ 43:5193-5267. doi: 11 10.1016/j.atmosenv.2009.07.068 Fowler D, Pitcairn CER, Sutton MA, et al (1998) The mass budget of atmospheric ammonia in 12 13 woodland within 1 km of livestock buildings. Environ Pollut 102:343-348. 14 García-Gómez H, Garrido JL, Vivanco MG, et al (2014) Nitrogen deposition in Spain: modeled 15 patterns and threatened habitats within the Natura 2000 network. Sci Total Environ 485-16 486:450-60. doi: 10.1016/j.scitotenv.2014.03.112 17 Geiser LH, Jovan SE, Glavich D, Porter MK (2010) Lichen-based critical loads for atmospheric 18 nitrogen deposition in Western Oregon and Washington Forests, USA. Environ Pollut 158:2412-2421. doi: 10.1016/j.envpol.2010.04.001 19 20 Gerosa G, Fusaro L, Monga R, et al (2015) A flux-based assessment of above and below ground 21 biomass of Holm oak (*Quercus ilex* L.) seedlings after one season of exposure to high ozone 22 concentrations. Atmos Environ 113:41-49. doi: 10.1016/j.atmosenv.2015.04.066 23 Gerosa G, Vitale M, Finco A, et al (2005) Ozone uptake by an evergreen Mediterranean Forest 24 (Quercus ilex) in Italy. Part I: Micrometeorological flux measurements and flux partitioning. 25 Atmos Environ 39:3255-3266. doi: 10.1016/j.atmosenv.2005.01.056 26 Grundström M, Pleijel H (2014) Limited effect of urban tree vegetation on  $NO_2$  and  $O_3$ 27 concentrations a traffic route. Environ Pollut 189:73-6. doi: near 28 10.1016/j.envpol.2014.02.026 29 Harris TB, Manning WJ (2010) Nitrogen dioxide and ozone levels in urban tree canopies. Environ Pollut 158:2384-6. doi: 10.1016/j.envpol.2010.04.007 30 31 Hereter A, Sánchez JR (1999) Experimental Areas of Prades and Montseny. In: Rodà, F., et al. 32 (eds) Ecology of Mediterranean evergreen oak forests. Springer-Verlag Berlin Heidelberg,
- 33 New York, pp 15–28.

1 2 3	Hjellbrekke A-G (2014) Data report 2012. Acidifying and eutrophying compounds and particulate matter (EMEP/CCC, 03/2014). Edited by Norwegian Institute for Air Research (NILU) – Chemical Coordination Center of EMEP (CCC), Oslo,				
4 5	Jovan S, Riddell J, Padgett PE, Nash TH (2012) Eutrophic lichens respond to multiple forms of N: implications for critical levels and critical loads research. EcolAppl 22:1910–22.				
6 7 8	Karanasiou A, Querol X, Alastuey A, et al (2014) Particulate matter and gaseous pollutants in the Mediterranean Basin: Results from the MED-PARTICLES project. Sci Total Environ 488- 489:297–315. doi: 10.1016/j.scitotenv.2014.04.096				
9 10 11	Kroeger T, Escobedo FJ, Hernandez JL, et al (2014) Reforestation as a novel compliance measure in State Implementation Plans for ground-level ozone. ProcNatlAcadSci U S A 52:1–43 doi: 10.1073/pnas.1409785111				
12 13 14	MAGRAMA (2014) Banco Público de Indicadores Ambientales. Edited by Spanish Ministry of Agriculture, Food and Environment (MAGRAMA). Available at http://www.magrama.gob.es				
15 16 17	Massad R-S, Nemitz E, Sutton MA (2010) Review and parameterisation of bi-directiona ammonia exchange between vegetation and the atmosphere. AtmosChem Phys 10:10359-10386. doi: 10.5194/acp-10-10359-2010				
18 19	Millán MM, Sanz MJ, Salvador R, Mantilla E (2002) Atmospheric dynamics and ozone cycles related to nitrogen deposition in the western Mediterranean. 118:167–186.				
20 21 22	Nowak DJ, Hirabayashi S, Bodine A, Greenfield E (2014) Tree and forest effects on air quality and human health in the United States. Environ Pollut 193:119–129. doi: 10.1016/j.envpol.2014.05.028				
23 24 25	Padgett PE, Parry SD, Bytnerowicz A, Heath RL (2009) Image analysis of epicuticular damage to foliage caused by dry deposition of the air pollutant nitric acid. J Environ Monit 11:63–74. doi: 10.1039/b804875d				
26 27	Pey J, Perez N, Castillo S, et al (2009) Geochemistry of regional background aerosols in the Western Mediterranean. Atmos Res 94:422–435.				
28 29 30	Pinho P, Theobald MR, Dias T, et al (2012) Critical loads of nitrogen deposition and critical levels of atmospheric ammonia for semi-natural Mediterranean evergreen woodlands. Biogeosciences 9:1205–1215. doi: 10.5194/bg-9-1205-2012				
31 32	Plaisance H (2011) The effect of the wind velocity on the uptake rates of various diffusive samplers. Int J Environ Anal Chem 91:1341–1352. doi: 10.1080/03067311003782625				

1	Querol X, Alastuey A, Moreno T, et al (2008) Spatial and temporal variations in airborne
2	particulate matter (PM10 and PM2.5) across Spain 1999-2005. Atmos Environ 42:3964-
3	3979.

Reche C, Viana M, Karanasiou A, et al (2014) Urban NH<sub>3</sub> levels and sources in six major Spanish cities. Chemosphere 119C:769-777. doi: 10.1016/j.chemosphere.2014.07.097

Rodríguez S, Querol X, Alastuey A, Mantilla E (2002) Origin of high summer PM10 and TSP concentrations at rural sites in Eastern Spain. Atmos Environ 36:3101-3112. doi: 10.1016/S1352-2310(02)00256-X

Setälä H, Viippola V, Rantalainen A-LL, et al (2013) Does urban vegetation mitigate air pollution in northern conditions? Environ Pollut 183:104–112. doi: 10.1016/j.envpol.2012.11.010

Sgrigna G, Sæbø A, Gawronski S, et al (2015) Particulate Matter deposition on *Ouercus ilex* leaves in an industrial city of central Italy. Environ Pollut 197:187-194. doi: 10.1016/j.envpol.2014.11.030

- Sparks JP (2009) Ecological ramifications of the direct foliar uptake of nitrogen.Oecologia 159:1-13. doi: 10.1007/s00442-008-1188-6
- The Royal Society (2008) Ground-level ozone in the 21st century: future trends, impacts and policy implications. Edited by: Fowler D, et al. R. Soc., London.
- Tzanis C, Varotsos C, Ferm M, et al (2009) Nitric acid and particulate matter measurements at Athens, Greece, in connection with corrosion studies. AtmosChem Phys Discuss 9:14683– 14711. doi: 10.5194/acpd-9-14683-2009

Site code	СВ	ТС	СА	LC
Site name	Can Balasc	<b>Tres Cantos</b>	Carrascal	La Castanya
Province (administrative unit)	Barcelona	Madrid	Navarra	Barcelona
Type of site	Peri-urban	Peri-urban	Peri-urban	Rural
Altitude (m)	255	705	592	696
Longitude	2° 04' 54" E	3° 43' 59" O	1° 38' 40'' O	2° 21' 29'' E
Latitude	41° 25' 47" N	40° 35' 17" N	42° 39' 13" N	41° 46' 47" N
Mean annual temperature (°C) $^{1}$	15.2	14.6	12.3	13.7
Mean annual rainfall $(mm y^{-1})^{1}$	652	348	645	812
Distance to the nearest big city (km)	4	9	15	40
Population of the nearest big city (million inhabitants)	1.6	3.2	0.20	1.6
Distance to the nearest highway (km)	0.15	1.5	0.05	16
Average daily flow in the nearest road (thousand vehicles day <sup>-1</sup> ) $^{2}$	40-50	50-60	20-30	20-30
Agricultural land-use cover <sup>3</sup>	23%	21%	62%	23%
Artificial land-use cover <sup>3</sup>	35%	28%	3.1%	7.6%
Livestock density (LU km <sup>-2</sup> ) <sup>4</sup>	14.5	13.7	26.9	88.8

#### **Table 1.**Characterization of the study sites.

<sup>1</sup>: Mean values calculated for the study period.
<sup>2</sup>: Values for 2012 from the Spanish Ministry of Development (http://www.fomento.gob.es/).
<sup>3, 4</sup>: From the Corine Land Cover 2006 (http://www.eea.europa.eu/data-and-maps/data/corine-

land-cover-2006-raster-3) and the Spanish National Statistic Institute (http://www.ine.es), 

respectively, using a buffer of 25 km radius around the sampling sites. 

1 Table 2.Basic statistics of the monitored pollutant concentrations in open-field plots for

2 the entire monitoring periods.

	SITE	MEAN	MIN. – MAX.	CV
	CB	$16.2 \pm 1.0$ a	5.7 - 39.3	42%
NO <sub>2</sub>	ТС	$11.1 \pm 1.1$ b	3.8 - 37.1	71%
(µg m <sup>-3</sup> )	CA	$10.6\pm0.7\;b$	4.4 - 26.0	45%
	LC	$4.3 \pm 0.3 \text{ c}$	0.8 - 9.4	52%
	СВ	$1.0 \pm 1.0 \text{ b}$	0.3 - 2.6	53%
NH <sub>3</sub>	TC	$0.7 \pm 0.1 \ c$	0.1 - 1.7	60%
(µg m <sup>-3</sup> )	CA	$2.5 \pm 0.2$ a	0.6 - 5.3	47%
	LC	$0.7 \pm 0.1 \ c$	0.1 - 1.7	59%
	СВ	$2.7\pm0.6$	0.0 - 14.5	134%
HNO <sub>3</sub>	TC	$1.5 \pm 0.2$	0.0 - 6.4	73%
(µg m <sup>-3</sup> )	CA	$2.3\pm0.3$	0.3 - 9.7	98%
	LC	$3.3 \pm 0.7$	0.0 - 13.9	134%
	СВ	$57.0 \pm 2.4 \text{ c}$	10.8 - 86.1	30%
<b>O</b> <sub>3</sub>	ТС	$69.1\pm2.9~b$	28.7 - 101.4	30%
(µg m <sup>-3</sup> )	CA	$77.4 \pm 4.7 \text{ a}$	25.3 - 122.3	32%
	LC	$78.2 \pm 3.2$ a	34.9 - 117.3	29%
DM	ТС	$23.0 \pm 3.2$ ab	5.2 - 61.0	67%
$PM_{10}$	CA	$26.9\pm2.6~\mathrm{a}$	6.8 - 49.2	41%
(µg m )	LC	$18.0\pm1.5~b$	4.8 - 32.8	41%
	ТС	$1.3\pm0.4\ b$	0.1 - 8.1	129%
$NO_3$	CA	$2.2 \pm 1.5 \text{ a}$	0.5 - 8.8	99%
(µg m)	LC	$1.1\pm0.2\ b$	0.2 - 4.2	80%
<b></b>	ТС	$0.6 \pm 0.1$	0.2 - 2.7	54%
$\mathbf{NH}_4$ (ug m <sup>-3</sup> )	CA	$0.9\pm0.2$	0.3 - 3.7	97%
(µg m )	LC	$0.5 \pm 0.1$	0.0 - 1.6	71%
SO 2-	ТС	$1.2 \pm 0.2$ b	0.1 - 4.2	70%
$SU_4^-$	CA	$1.9 \pm 0.2$ a	0.8 - 3.7	48%
(µg m )	LC	$1.7 \pm 0.2 \text{ a}$	0.4 - 3.3	52%

3 Mean: arithmetic mean  $\pm$  standard error. Min. – Max.: Minimum and maximum two-

4 week values. CV: coefficient of variation, representing the temporal variability. Different

5 letters indicate significant differences (p < 0.05) between sites. The absence of letters

6 indicates no significant differences.

7

8 9

10

11

12

Table 3. Ozone exposure expressed as AOT40 for years 2011 and 2012, following
 criteria from the Convention on Long-range Transboundary Air Pollution (CLRTAP) and
 the Ambient Air Quality Directive 2008/50/EC.

AOT40 (ppm h)					
SITE	CLRTAP (Jan-Dec)		Directive 2008/50/EC (May–July)		
	2011	2012	2011	2012	
СВ	8.2	18.8	3.7	9.4	
ТС	31.8	49.6	17.4	28.3	
CA	32.6	32.3	15.5	16.5	
LC	27.3	34.9	12.5	18.3	

#### **1 FIGURE CAPTIONS**

2 Fig. 1 Distribution of *Quercus ilex* habitats in Spain, and location of the study sites. LC: La

Castanya (Barcelona); CB: Can Balasc (Barcelona); CA: Carrascal (Navarra); TC: Tres Cantos
(Madrid).

Fig. 2 Seasonal mean concentration of atmospheric pollutants in the open-field (O) plots of the
four study sites and standard error of the mean. Different letters indicate significant differences
amongseasons.

8 Fig. 3 Seasonal mean concentrations of aerosols and standard errors, and ratios of particulate to 9 gaseous pollutants in the three aerosol monitoring sites. A)  $PM_{10}$  concentration; B)  $PM_{10}$ 10 concentration for measurements during Saharan dust events compared with the rest of the 11 samples; C) particulate nitrate concentrations; D) particulate ammonium concentrations; E) 12 concentrations ratios of nitric acid and particulate nitrate, expressed as percentage of the sum of 13 both compounds; F) concentrations ratios of ammonia and particulate ammonium, expressed as 14 percentage of the sum of both compounds. Different letters indicate significant differences 15 between seasons. One outlier value (CA, spring) was removed from the graphs C-F.

Fig. 4 Mean concentration of pollutants in O plots (open field) and F plots (below canopy), and
standard error of the mean. Significance of the Wilcoxon matched pairs test: \*: p < 0.05; \*\*: p <</li>
0.01; \*\*\*: p < 0.001.</li>

- Fig. 1



- 1 Fig. 2







- 1 Fig. 3



- .

- 1 Fig. 4

