Ecological impacts of atmospheric pollution and interactions with climate change in terrestrial ecosystems of the Mediterranean Basin: Current research and future directions

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Keywords: Air pollution, Climate change, Coordinated research networks, Environmental monitoring

Mediterranean Basin ecosystems, their unique biodiversity, and the key services they provide are currently at risk due to air pollution and climate change, yet only a limited number of isolated and geographically-restricted studies have addressed this topic, often with contrasting results. Particularities of air pollution in this region include high O3 levels due to high air temperatures and solar radiation, the stability of air masses, and dominance of dry over wet nitrogen deposition. Moreover, the unique abiotic and biotic factors (e.g., climate, vegetation type, relevance of Saharan dust inputs) modulating the response of Mediterranean ecosystems at various spatiotemporal scales make it difficult to understand, and thus predict, the consequences of human activities that cause air pollution in the Mediterranean Basin. Therefore, there is an urgent need to implement coordinated research and experimental platforms along with wider environmental monitoring networks in the region. In particular, a robust deposition monitoring network in conjunction with modelling estimates is crucial, possibly including a set of...
1. Introduction

Human activities and natural processes have shaped each other over ca. eight millennia within Mediterranean Basin ecosystems (Blondel, 2006). This coevolution, together with the heterogeneous orography and geology, the large seasonal and inter-annual climatic variability, the refuge effect during the last glaciations, and the crossroad location between European temperate ecosystems and North African and Asian drylands, has resulted in the high diversification of the flora and fauna that we observe today, making Mediterranean ecosystems a hotspot of biodiversity, but also of vulnerability (Schröter et al., 2005; Blondel, 2006; Phoenix et al., 2006). Moreover, the Mediterranean Basin is one of the world’s largest biodiversity hotspots and the only one within Europe, otherwise dominated by temperate natural and semi-natural grasslands, temperate deciduous forests and boreal conifer forests (Myers et al., 2000). Species-rich ecosystems exclusive to the Mediterranean Basin include Spanish matorrales and garriegas, Portuguese matos, Italian macchias, Greek phryganas, and agrosilvopastoral ecosystems of high natural and economic value such as Spanish dehesas and Portuguese montados (Cowling et al., 1996; Blondel, 2006). However, the biodiversity and other ecosystem services of this region are currently at risk due to human pressures such as climate change, land degradation and air pollution (Schröter et al., 2005; Scarcia-Mugnoza and Matteucci, 2012). Air pollution in the Mediterranean Basin is primarily in the form of particulate matter, nitrogen (N) deposition and tropospheric ozone (O3) (Paoletti, 2006; Ferretti et al., 2014; García-Gómez et al., 2014; Fig. 1). Production of pollutants is mainly associated with industrial activities, construction, vehicle emissions and agricultural practices and, within the European context, is characteristically exacerbated by more frequent droughts and the typical stability of air masses in the region, with important consequences for ecosystem and human health (Millán et al., 2002; Vestreng et al., 2008; Izqueta-Rojano et al., 2016a). This also has important social consequences for the Mediterranean region, where approximately 480 million people live, and where more frequent droughts, extreme climatic events and wildfires will only reinforce the current migrant and humanitarian crisis (Werz and Hoffman, 2016).

Environmental pollution causes and interacts synergistically with climate change (Alonso et al., 2001, 2014; Bytnorowicz et al., 2007; Sardans and Peñuelas, 2013). This is particularly true for seasonally dry regions like the Mediterranean Basin (Baron et al., 2014), but the effects of this interaction on the structure and function of Mediterranean ecosystems are not adequately quantified and, therefore, the consequences are poorly understood (Bobbink et al., 2010; Ochoa-Hueso et al., 2011). Projections for 2100 suggest that mean air temperatures in the Mediterranean Basin region will rise from 2.2 °C to 5.1 °C above 1990 levels and that precipitation will decrease between −4 and −27% (Christensen et al., 2007 and Fig. 2). The sea level is also projected to rise, and a greater frequency and intensity of extreme weather events (e.g., drought, heat waves and floods) are expected (EEA, 2005). These changes will exacerbate the already acute water shortage problem in the region, particularly in drylands (Terray and Boé, 2013; Sicard and Dalstein-Richier, 2015), impairing their functionality and ability to deliver the ecosystem services on which society and economy depend (Bakkenes et al., 2002; Lloret et al., 2004). Functions that will be synergistically impaired by air pollution and climate change include reductions in crop yield and carbon sequestration (Maracchi et al., 2005; Mills and Harmens, 2011; Shindell et al., 2012; Ferretti et al., 2014). In addition, a higher fire risk is attributed to higher temperatures and more frequent droughts coupled with an N-driven increase of grass-derived highly-flammable fine fuel (Pausas and Fernández-Muñoz, 2012).

In the last decades, atmospheric concentrations of major anthropogenic air pollutants such as particulate matter and sulphur dioxide (SO2) have decreased in Southern Europe due to emission control policies and greener technologies (Querol et al., 2014; Barros et al., 2015; Aguillauema et al., 2016; Ávila and Aguillauema, 2017). However, mitigation strategies have not been equally effective with other compounds such as reactive N and tropospheric O3 (Fig. 1; Paoletti, 2006; García-Gómez et al., 2014; Sicard et al., 2016). For example, recent increases in N deposition, particularly dry deposition of NOx, have been detected in North-eastern Spain, where N deposition is estimated in the range of 15–30 kg N ha−1 yr−1 (Ávila and Rodà, 2012; Camarero and Catalan, 2012; Aguillauema et al., 2016). This has been attributed to increased nitrogen oxide (NOx) and ammonia (NH3) emissions and changes in precipitation patterns (Aguillauema et al., 2016). Background O3 pollution is typically high in Mediterranean climates due to the meteorological conditions of the area (Paoletti, 2006) and recent reviews have demonstrated that while O3 in cities has generally increased, no clear trend, or only a slight decrease, has been detected in rural areas (Sicard et al., 2013; Querol et al., 2014); the annual average at rural western Mediterranean sites over the period 2000–2010 was 33 ppb, with a modest trend of −0.22% yr−1 (Sicard et al., 2013). The Mediterranean Basin is also exposed to frequent African dust intrusions, which can naturally increase the level of suspended particulate matter and nutrient deposition, changing the chemical composition of the atmosphere (Escudero et al., 2005; Maricorena and Fornetti, 2013; Ávila and Aguilluema, 2017). This has profound impacts on the biogeochemical cycles of both aquatic and terrestrial ecosystems (Mona et al., 2006), further exacerbating the negative consequences of air pollution and climate change on ecosystem and human health.

In this review, originated as a result of the 1st CAPEReMED (Committee on Air Pollution Effects Research on Mediterranean Ecosystems; http://capermmed.weebly.com/) Conference in Lisbon, Portugal, we (i) summarize the current knowledge about atmospheric pollution trends and effects, and their interactions with climate change, in terrestrial ecosystems of the Mediterranean Basin, (ii) identify research gaps that need to be urgently filled, and (iii) recommend future steps. Due to lack of information for other regions within the Mediterranean Basin, we mainly focused our review on studies carried out in south-western European countries.
Fig. 1. Modelled nitrogen deposition for the Mediterranean region based on the European Monitoring and Evaluation Programme (EMEP) model at 0.1°-0.1° longitude-latitude resolution [EMEP MSC-W chemical transport model [version rv4.7; www.emep.int]]. Modelled N deposition is based on 2013 emissions data. (a) Total N deposition (oxidized + reduced; dry + wet), (b) percentage of dry deposition, (c) percentage of wet deposition, (d) percentage of oxidized deposition and (e) percentage of reduced deposition.
(France, Italy, Portugal and Spain). In contrast, we discuss information generated through a variety of experimental approaches (field manipulation experiments, greenhouse studies, open top chambers [OTCs], observational studies, modelling, etc.) from studies carried out in a wide range of representative natural (e.g., shrublands, grasslands, woodlands and forests) and semi-natural (e.g., montados or dehesas) ecosystems.

2. Measurement and modelling of atmospheric pollution and deposition

Estimating pollutant deposition loadings, particularly dry deposition, still presents important uncertainties and challenges, both in terms of modelling and measurements (Simpson et al., 2014). This is particularly true in studies at small regional scales and in regions with complex topography or under the influence of local emission sources (García-Gómez et al., 2014), which is very often the case in the Mediterranean Basin. Dry deposition in Mediterranean ecosystems can represent the main input of atmospheric N, contributing up to 65–95% of the total deposition (Fig. 1b; Sanz et al., 2002; Avila and Rodà, 2012). For example, wet N deposition at the Levantine border of the Iberian Peninsula can be considered low to moderate (2–7.7 kg N ha⁻¹ yr⁻¹), but total N deposition loads are comparable to more polluted areas in central and northern Europe (10–24 kg N ha⁻¹ yr⁻¹) when dry deposition is included (Avila and Rodà, 2012). Given that dry deposition is important in the Mediterranean Basin but is also difficult to measure, we should ideally combine modelled dry deposition with wet deposition measures from representative monitoring stations. A recent modelling analysis has also highlighted that mountain ecosystems in Spain, where monitoring stations are even scarcer, are frequently exposed to exceedances of empirical critical N loads (García-Gómez et al., 2014, 2017). Moreover, mountain areas of the Mediterranean Basin also frequently register very high O₃ concentrations that are not recorded in air quality monitoring networks (Díaz-de-Quijano et al., 2009; Cristofanelli et al., 2015; Elvira et al., under review). This observation should encourage the inclusion of monitoring stations in mountain areas in air quality networks in the Mediterranean Basin to protect these highly valuable and vulnerable ecosystems (García-Gómez et al., 2017). Another important aspect to be considered in both deposition monitoring networks and model-based estimates is the quantification and characterization of ammonium (NH₄⁺) and the organic N fraction (Jickells et al., 2013; Fowler et al., 2015). Dissolved organic N (DON) can represent a significant component of wet and dry deposition fluxes but it is often overlooked and not routinely assessed (Mace, 2003; Violaki et al., 2010; Im et al., 2013; Izquieta-Rojano and Elustondo, 2017). However, DON fluxes may have significant implications in terms of critical loads, reaching up to 34–56% of the total N deposition (12 kg DON ha⁻¹ yr⁻¹) in Mediterranean agricultural areas (Izquieta-Rojano et al., 2016a). The quantification of temporal trends in air pollution is equally important for evaluating the impact of changing precursor emissions and informing local and regional air quality strategies.

3. Impacts of atmospheric pollution and climate change on natural and semi-natural terrestrial ecosystems

The ecological impacts of air pollution (particularly for N deposition and O₃) on natural and semi-natural ecosystems have been primarily studied in the temperate and boreal regions of Europe and North America and, more recently, in steppe and subtropical areas of China (Paoletti, 2006; Xia and Wan, 2008; Bobbink et al., 2010; Ochoa-Hueso, 2017). In contrast, much less is known for Mediterranean Basin ecosystems, which differ from these better-studied ecosystems in critical aspects that justify their separate consideration, such as their much-higher levels of biodiversity (particularly for plants) and their higher-than-average levels of biologically-relevant spatial and temporal environmental heterogeneity, including the characteristic summer drought period (Cowling et al., 1996; Myers et al., 2000). Most studies on the impacts of atmospheric pollution in terrestrial ecosystems from the
Fig. 3. Examples of terrestrial ecosystems and experimental facilities set up to investigate the effects of air pollution and climate change in the Mediterranean Basin (see Supplementary Table 2 for details): a) Companhia das Lezírias, Samora Correia, Portugal; b) Alambre, Serra da Arrábida, Portugal; c) Herdade da Coitadinha, Barrancos, Portugal; d) Alto de Guarramillas, Madrid, Spain; e) La Higuerauela, Toledo, Spain; f) El Regajal, Madrid, Spain; g) Tres Cantos, Madrid, Spain; h) Capo Caccia, Sardinia, Italy; i) La Castanya, Spain; j) Ozone FACE (Free-Air Controlled Exposure) facility, Florence, Italy; k) Fontblanche, Provence, France.
Mediterranean Basin have been carried out in just a small part of the geographic area (i.e. certain localities in Italy, Portugal and Spain) and have used different experimental design and methodologies (Fig. 3 and Supplementary Table 1). Similarly, instead of taking advantage of the development of statistical methods to integrate responses at the ecosystem level (e.g., structural equation modelling; Eisenhauer et al., 2015), studies have typically focused solely and independently on plants (community or, more frequently, individual species), lichens (community or, again more frequently, individual species) and soil properties (soil biogeochemistry, structure and functioning; Supplementary Table 1). One notable exception to this is NitroMed, a unique network of three comparable N addition experimental sites (Capo Caccia [0 and 30 kg N ha\(^{-1}\) yr\(^{-1}\)], Alambre [0, 40 and 80 kg N ha\(^{-1}\) yr\(^{-1}\)], and El Regajal [0, 10, 20 and 50 kg N ha\(^{-1}\) yr\(^{-1}\)]; see Fig. 3b, f and h) that is currently using common experimental methodology and structural equation modelling to understand the cause-effect mechanisms that determine changes in gas (CO\(_2\)) exchange and litter decomposition and stabilization rates in response to N deposition in semiarid Mediterranean ecosystems (see Ochoa-Hueso and Manrique, 2011 and Dias et al. 2014 for further details on experimental methodologies). Preliminary results suggest that N deposition increases soil N availability and reduces soil pH which, in turn, has an effect on microbial community structure (lower fungi to bacteria ratio) and overall enzymatic activity, direct responsible for reduced litter decomposition and higher stabilization rates (Lo Cascio et al., 2016). Similarly, a new coordinated project is looking at the effects of N addition at realistic doses (10 and 20 kg N ha\(^{-1}\) yr\(^{-1}\)) in conjunction with P. on alpine ecosystems from five National Parks in Spain.

Moreover, most of these studies addressed the impact of one global change driver alone (often increased N availability, mostly the N load, or O\(_3\)) and so comprehensive studies on the interaction between global change drivers (e.g., air pollution and climate change) are few. However, recent studies have described a heterogeneous response of annual pasture species to O\(_3\) and N enrichment, with legumes being highly sensitive to ozone but not N, while grasses and herbs were more tolerant to O\(_3\) and more responsive to N (Calvete-Sogo et al., 2016). Thus the interactive effects of O\(_3\) and N can alter the structure and species composition of Mediterranean annual pastures via changes in the competitive relationships among species (González-Fernández et al. 2013 and references therein; Calvete-Sogo et al., 2014, 2016). Similarly, only a few studies have addressed the impacts on edaphic fauna and above- and below-ground biotic interactions such as mycorrhiza, biological N fixation, herbivory or pollination in ecosystems from the Mediterranean Basin (Supplementary Table 1 and references therein), despite the relevance of ecological interactions to maintain healthy, functional ecosystems (Tylianakis et al., 2008). For example, Ochoa-Hueso et al. (2014a) found that edaphic fungal abundance, particularly collemobolans, increased in response to up to 20 kg N ha\(^{-1}\) yr\(^{-1}\) and then decreased with 50 kg N ha\(^{-1}\) yr\(^{-1}\), whereas 10 kg N ha\(^{-1}\) yr\(^{-1}\) were enough to completely suppress soil microbial N fixation (Ochoa-Hueso et al., 2013a). Another notable exception is Ochoa-Hueso (2016), who showed how even low-N addition levels (10 kg N ha\(^{-1}\) yr\(^{-1}\)) can completely disrupt the tight coupling of the network of ecological interactions in a semiarid ecosystem from central Spain, despite the lack of evident response of most of the individual abiotic and biotic ecosystem constituents evaluated (i.e., soils, microbes, plants and edaphic fauna). Ozone and N soil availability can also alter volatile organic compound (VOC) emissions, and thus biosphere-atmosphere interactions, of some Mediterranean tree and annual pasture species. The consequences of these interactions need to be further studied (Peñuelas et al., 1999; Llusia et al., 2002; Llusia et al., 2014).

Therefore, a more comprehensive and integrative experimental approach is urgently needed to fully capture the real consequences of air pollution in the Mediterranean region (see Fig. 4).

3.1. Sensitivity of Mediterranean forests to air pollution and climate change

Mediterranean forest ecosystems have naturally evolved cross-tolerance to deal with harsh environmental conditions (Paoletti, 2006; Matesanz and Valladares, 2014). However, climate change, N deposition and O\(_3\) are currently threatening Mediterranean forests in unprecedented and complex manners, with consistent stoichiometric responses to increased N deposition (higher leaf N:P ratios; Sardans et al., 2016), but with physiological and growth-related consequences forecasted to vary among the three main tree functional types (i.e., conifers, evergreen broadleaf trees, and deciduous broadleaf trees). As deposition increases, photosynthesis, water use efficiency, and thus growth, often increase in conifers (Leonardi et al., 2012), although under chronic N deposition, other nutrients such as P can become more limiting, counteracting the initial benefits of more N availability (Blanes et al., 2015). Nitrogen deposition could also increase pine mortality rates in response to drought due to a decline of ectomycorrhizal colonization rates, a phenomenon of widespread occurrence in US dryland woodlands (Allen et al., 2010). On the other hand, their low stomatal conductance and their high stomatal sensitivity to vapour pressure deficit and water availability might limit the diffusion of O\(_3\) to the mesophyll (Flexas et al., 2014). Similarly, conservative strategies of water and nutrient-use may also play a key role in allowing conifers to keep a positive balance between assimilation and respiration in response to climate change (Way and Oren, 2010). However, O\(_3\) exposure might be impairing their ability to withstand other environmental stresses such as those triggered by drought, high temperature and solar radiation (Barnes et al., 2000; Alonso et al., 2001).

In contrast, evergreen broadleaf species inhabiting resource-poor ecosystems might be jeopardized by N deposition by shifting biomass partitioning (Cambui et al., 2011) and altering allometric ratios (e.g., leaf area/sap wood or root/leaf biomass), which may have consequences for their ability to deal with water stress, particularly in the context of the characteristic summer drought period and climate change (Martinez-Vilalta et al., 2003; Mereu et al., 2009). Ecophysiological responses to O\(_3\) vary from downregulation of photosystems (Mereu et al., 2009) to reduced stomatal aperture and increased stomatal density (Fusaro et al., 2016) and sluggishness (Paoletti and Grulke, 2005, 2010). However, Mediterranean vegetation usually has efficient antioxidant defences (Nali et al., 2004), which are key factors in O\(_3\) tolerance (Calatayud et al., 2011; Mereu et al., 2011), and is usually known to be more O\(_3\)-tolerant than mesophilic broadleaf trees (Paoletti, 2006). Nevertheless, biomass losses and allocation shifts cannot be excluded, especially as a consequence of synergistic effects of N deposition and drought, although local differentiation may result in significant intraspecific tolerance differences (Alonso et al., 2014; Gerosa et al., 2015).

Responses of deciduous broadleaf species to N deposition may be modulated by water and background nutrient availability (mainly P) but, in general terms, growth is favoured over storage (Ferretti et al., 2014). In contrast, broadleaf tree species are highly sensitive to climate change, particularly to the combination of drought and increased temperature (Lopez-Iglesias et al., 2014), which also suggests relevant interactions between air pollution and climate change. In this direction, De Marco et al. (2014) predicted that crown defoliation will increase in Mediterranean environments due to drought events and higher temperatures by 2030, a
phenomenon that could be exacerbated by excessive N. Deciduous broadleaf species also have lower capacity to tolerate oxidative stress than evergreen broadleaf species due to traits such as thinner leaves and higher stomatal conductance (Calatayud et al., 2010). Gas exchange and antioxidant capacity in deciduous broadleaves are, therefore, generally more affected by high O$_3$ concentrations than in evergreen broadleaves (Bussotti et al., 2014). Based on their levels of visible foliar injury and expert judgement, deciduous broadleaf species range from highly to moderately sensitive species such as Fagus sylvatica and Fraxinus excelsior, respectively (Baumgarten et al., 2000; Tegischer et al., 2002; Gerosa et al., 2003; Deckmyn et al., 2007; Paoletti et al., 2007; Sicard et al., 2016), to O$_3$-tolerant species like some Quercus species (Q. cerris, Q. ilex and Q. petraea; Gerosa et al., 2009; Calatayud et al., 2011; Sicard et al., 2016).

Relatively little is known about the effects of O$_3$ on perennial, perennial and woody understory vegetation of Mediterranean forest ecosystems. Under experimental conditions, some species characteristic of the annual grasslands associated with Q. ilex dehesas have high O$_3$ sensitivity. Interestingly, N fixing legumes, of higher nutritional value, are more O$_3$ sensitive than grasses (Bermejo et al., 2004; Gimeno et al., 2004), particularly in terms of flower and seed production (Sanz et al., 2007), which could affect their competitive fitness and, ultimately, reduce the economic value of the pasture. Nitrogen availability can partially counterbalance O$_3$ effects on aboveground biomass when the levels of O$_3$ are moderate, but O$_3$ exposure reduces the fertilization effect of higher N availability (Calvette-Sogo et al., 2014). Anyhow, given that O$_3$ levels are higher in summer, when herbaceous species are dormant, Mediterranean species that are summer-active such as pines and oaks are more likely to be directly affected by O$_3$ than forbs and grasses. This suggests that the seasonality of O$_3$ concentrations as well as plant phenology and functional type must be considered if we are to fully understand the consequences of air pollution on the highly diverse Mediterranean plant communities. A unique ozone FACE (free air controlled experiment) is now available in the Mediterranean Basin (Fig. 3) to help fill this gap (Paoletti et al., in preparation).

3.2. Role of environmental context in the response of biodiversity and C sequestration

The local abiotic (e.g., climate, soil properties) and biotic (e.g., vegetation type, community attributes, etc.) contexts are known to modulate ecosystem responses to environmental drivers at different temporal and spatial scales (Bardgett et al., 2013). Given that plant biodiversity at the regional (10$^{-6}$ km$^2$) and local (<0.1 ha) scales in Mediterranean ecosystems ranks among the highest in the world (Cowling et al., 1996), this is particularly relevant for the region. Various studies in Mediterranean ecosystems have shown that increased N availability may have a positive (Pinho et al., 2012; Dias et al., 2014), negative (Bonanomi et al., 2006; Bobbink et al., 2010) or even no effect (Dias et al., 2014) on plant species richness, which is probably due to cumulative effects and modulating factors such as the ecosystem type, the initial N status of the system, the dominant form of mineral N in the soil (NH$_4$, NO$_3$), and/or the N form added. Positive effects on species richness, however, have only been observed in areas characterized by strong environmental stress and low nutrient availability (e.g., open arid and semiarid Mediterranean ecosystems) and are often associated with an increase in nitrophytic and weedy species (Bobbink et al., 2010; Pinho et al., 2011; Dias et al., 2014). The presence and density of shrubs, as well as the availability of inorganic phosphorus (P) and other macro and micronutrients, can also modulate the response of the herbaceous vegetation to N addition and plant invasion in semiarid Mediterranean areas (Ochoa-Hueso et al., 2013b; Ochoa-Hueso and Stevens, 2015). For example, Ochoa-Hueso and Manrique (2014) found that N addition increased the nitrophytic element, particularly native crucifers, only when these species were present in the seed bank in relevant densities and there was sufficient P, whereas a closed scrub vegetation is known to be less susceptible to invasion by N-loving species than open shrublands, woodlands and grasslands (Dias et al., 2014). The role of soil nutrient availability, typically lower than in other Mediterranean-type ecosystems such as those from Chile (Cowling et al., 1996), in the ecosystem response to extra N can also be linked to induced nutrient imbalances, particularly N in relation to P, and therefore to an alteration of ecosystem stoichiometry (Ochoa-Hueso et al., 2014b; Sardans et al., 2016).

The behaviour of terrestrial ecosystems as a global C sink or source under increased N deposition or O$_3$ pollution scenarios is currently a research hot-topic and is of paramount importance for the mitigation of climate change (Felzer et al., 2004; Reich et al., 2006; Pereira et al., 2007). Recent studies have suggested that seasonally water-limited ecosystems, such as those typically found in the Mediterranean basin, may have a disproportionately big role in the inter-annual C sink-source dynamics at the global scale due to higher C turnover rates (Poulter et al., 2014); this is attributed to their large inter-annual climatic variability, with unusually wet years contributing to strengthen the terrestrial C sink but where
multiple processes like fire or rapid decomposition could result in a rapid loss of most of the accumulated C. These aspects are, however, still poorly understood in Mediterranean ecosystems, where different studies have reported contrasting results (Ochoa-Hueso et al., 2013a, 2013c; Ferretti et al., 2014). In Mediterranean ecosystems, ecosystem C storage should, therefore, be evaluated in terms of altered abundance and patterns of rainfall (both within and between years) (Pereira et al., 2007), in relation to the levels of N saturation (NO₃⁻) and toxicity (NH₄⁺) in soil (Dias et al., 2014), as well as other site-dependent characteristics such as dominant vegetation, soil type (texture and pH), and stand history and age (Ferretti et al., 2014). Experimental and observational field studies suggest that, at least in the short-term, seasonal and inter-annual dynamics may override any potential effect of atmospheric N pollution, despite potential cumulative negative impacts in the long-term due to an overall decline in ecosystem health (Ochoa-Hueso et al., 2013c; Ferretti et al., 2014).

Although within the Mediterranean Basin there is still a large gap in the knowledge of the impacts of atmospheric pollution and climate change on natural and semi-natural ecosystems, taken together, all the scattered information available suggests the particularly key role of spatial and temporal environmental heterogeneity, biotic interactions, and ecosystem stoichiometry in mediating the ecosystem response to air pollution.

### 3.3. Critical loads and levels

The concepts of critical loads and critical levels were developed within the United Nation Economic Commission for Europe (UNECE) Convention on Long-Range Transboundary Air Pollution (CLRTAP) for assessing the risk of air pollution impacts to ecosystems and defining emission reductions. This tool is commonly used to anticipate negative effects of air pollution and, therefore, to protect ecosystems before the changes become irreversible. The derivation of empirical critical loads for nutrient N is based on experimental activities performed on different vegetation types and they are assigned to habitat classes, while the derivation of NH₃ and NOₓ critical levels is based on the responses of broad vegetation types such as higher plants or lichens and bryophytes. The pan-European critical level for atmospheric NH₃ is currently set at an annual mean of 1 µg m⁻² for lichens and bryophytes and 3 µg m⁻³ for higher plants, while the NOₓ critical level for all vegetation types is an annual mean of 30 µg m⁻³ (CLRTAP, 2011). Although some modelling approaches exist to define critical N loads, the identification of empirical critical loads is recommended for Mediterranean ecosystems due to its particularities such as co-occurrence with other pressures and high seasonality (de Vries et al., 2007; Fenn et al., 2011). Empirical critical loads of N for European-Mediterranean habitats have only been proposed for four ecosystems: (1) Mediterranean xeric grasslands (EUNIS [European Nature Information System] E 1.3), 15–25 kg N ha⁻¹ yr⁻¹; (2) Mediterranean maquis (F5), 20–30 kg N ha⁻¹ yr⁻¹; (3) Mediterranean evergreen (Quercus) woodlands (G 2.1), 10–20 kg N ha⁻¹ yr⁻¹, and (4) Mediterranean Pinus woodlands (G 3.7), 3–15 kg N ha⁻¹ yr⁻¹ (Bobbink and Hettelingh, 2011). However, these critical loads are based on very little information and are thus classified as expert judgement. Similarly, NH₃ critical levels have only been set for Mediterranean evergreen woodlands and dense holm oak forests. Critical levels of atmospheric NH₃ of <1.9 and 2.6 µg m⁻³ have been estimated for evergreen woodlands surrounded by intensive agricultural landscapes (Pinho et al., 2012; Aguillaume, 2015), while for evergreen woodlands under light agricultural influence but strong oceanic influence, the critical level was estimated to be 0.69 µg m⁻³ (Pinho et al., 2014). Nevertheless, the N critical loads and NH₃ critical levels for many European-Mediterranean ecosystems remain unstudied, despite their relevance for protecting relatively undisturbed and oligotrophic ecosystems. Therefore, long-term manipulation experiments across a range of typical Mediterranean terrestrial ecosystems are desperately needed to obtain a more complete set of reliable empirical critical N loads and levels for the Mediterranean Basin (Bobbink et al., 2010; Bobbink and Hettelingh, 2011). Ozone critical levels have also been proposed for the protection of natural vegetation at European level for two vegetation types, forests and semi-natural vegetation (CLRTAP, 2011). The new flux-based O₃ critical levels allow species-specific physiological conditions and O₃ uptake mechanisms to be included considering the particularities of Mediterranean species. Interestingly, multiple studies performed with Mediterranean tree species recommend higher O₃ critical levels for the protection of Mediterranean forests than the values currently accepted (Calatayud et al., 2011; Alonso et al., 2014; Gerosa et al., 2015). The possible definition of different O₃ critical levels for different biogeographical regions or vegetation types is currently under analysis within the Convention (CLRTAP, 2011).

### 3.4. Cryptogams as indicators of the impact of air pollution and climate change

Lichens and bryophytes (i.e., cryptogams), very often used in the definition of critical loads and levels, are important components of the vegetation in Mediterranean ecosystems. These organisms are key drivers of ecosystem properties (soil aggregation and stability) and processes (C and N fixation and nutrient cycling), particularly in the case of biological soil crusts (hereafter biocrusts), a functionally-integrated association of cyanobacteria, protists, fungi, mosses and lichens inhabiting the first millimetres of soil (Cornelissen et al., 2007; Maestre et al., 2011). Cryptogams are usually extremely sensitive to environmental changes and so they often provide early-warning indicators of impacts before any other constituent of the ecosystem, particularly in the case of N (Pardo et al., 2011; Munzi et al., 2012). For example, mosses have been used in N deposition surveys under the ICP-Vegetation framework (Harmens et al., 2014). The results showed that N concentration in mosses can potentially be used as an indicator of total atmospheric N deposition. Similarly, Root et al. (2013) showed that lichens can be a suitable tool for estimating throughfall N deposition in forests. However, the relationship between N deposition and tissue N concentration can also be affected by environmental factors such as local climate and the form of N deposition.

Mosses and lichens have been instrumental to the evaluation of the impacts of global change drivers on temperate and boreal ecosystems (e.g., Arróniz-Crespo et al., 2008), although the number of studies carried out in Mediterranean ecosystems is very limited. Recent studies have, however, reported significant impacts of increased N deposition on Mediterranean biocrust and epiphytic communities. For example, two studies carried out in the Iberian peninsula found higher tissue N content and a shift from N to P limitation in the terricolous moss Tortella squarrosa (=Pleurochaete squarrosa; Ochoa-Hueso and Manrique, 2013; Ochoa-Hueso et al., 2014a). Similarly, an alteration of physiological and chemical responses in lichen transplants (Branquinho et al., 2010; Paoli et al., 2010, 2015) and a shift in epiphytic lichen communities from oligotrophic-dominated to nitrophytic-dominated species have also been reported in Portugal (Pinho et al., 2008, 2009) and Spain (Aguillaume, 2016). Recent studies have also observed a change in the isotopic N composition of mosses due to the impact of N from fuel combustion sources (shift to more positive δ¹⁵N signature) and agricultural activities (shift to more negative δ¹⁵N signature; Delgado et al., 2013; Varela et al., 2013; Izquieta-Rojano et al., 2016b).
Cryptogam traits (e.g., morphology, anatomy, life form) are also strongly connected to water availability. For example, mosses from dry habitats are organized in dense cushions, naturally retaining water by capillarity and dehydrating slowly, whereas mosses from moist habitats have a less dense morphology and require the activation of specific mechanisms to survive during dry periods (Arróniz-Crespo et al., 2011; Cruz de Carvalho et al., 2011, 2012, 2014). Similarly, lichen growth form and photobiont type have been shown to be relevant traits in the response to water availability in Mediterranean areas (Concostrina-Zubiri et al., 2014; Matos et al., 2015). Cryptogam traits related to water availability could, therefore, be equally effective biomarkers to detect climate-induced hydrological changes in Mediterranean ecosystems but the application of biomonitoring techniques using cryptogams in the Mediterranean region may be complicated by the fact that cryptogam species are simultaneously exposed to both severe water restriction and pollution, and some biomarkers (e.g., ecophysiological responses) are similarly affected by both stress factors (Pirintos et al., 2011). Thus, we need to disentangle the multiple environmental drivers (Munzi et al., 2014a), possibly by integrating physiological and ecological data to understand the specific response mechanisms to different ecological parameters and environmental changes (Munzi et al., 2014b).

3.5. Anticipating global tipping points using ecological indicators

The fact that ecosystem responses to air pollution and climate change are very often non-linear may complicate the use of bioindicators in the Mediterranean Basin. Non-linear dynamics often manifest in the form of tipping points, defined as ecosystem thresholds above which a larger-than-expected change happens, shifting ecosystems from one stable state to another stable state (Scheffer and Carpenter, 2003). Due to its climatic peculiarities, tipping points may be particularly relevant for the Mediterranean Basin. One example is the ability of soils to store extra mineral N. Above a certain N deposition value, N-saturated soils will start leaching N down into the soil profile. This excessive N can also accumulate as inorganic N in seasonally dry soils and be leached by surface flows that, as in the case before, will eventually reach and, therefore, pollute aquifers and watercourses (Fenn et al., 2008). Another relevant example is related to increased fire risk due the accumulation of highly flammable leaf litter, particularly from exotic grasses, as a consequence of N deposition; above a certain N deposition threshold the probability of a fire to occur increases exponentially, priming the ecosystem for a state change (Rao et al., 2010).

Despite the potential prevalence of tipping point-like dynamics in Mediterranean ecosystems in response to air pollution and climate change, we are not aware of any vegetation-based tools available to predict ecosystem thresholds in the Mediterranean Basin context. A notable exception is the work by Berdugo et al. (2017), who suggested that changes in the spatial configuration of drylands may be an early-warning indicator of desertification. However, we suggest that if we are to aim for universal indicators of environmental change (i.e., at wide geographical ranges) and to account for the role of the environmental context as a driver (i.e., across ecosystem types), functional trait-based approaches (e.g., functional diversity and community weighted mean trait values [CWM]) should be preferred over other widely used indicators, including species richness (Jolyan and McCune, 2005; Valencia et al., 2015). Functional diversity and CWM are independent of species identity and may be functionally linked to the environmental variable of interest (e.g., oligotrophic species, nitrophytic species, or subordinate species responding to eutrophication, species-specific leaf litter traits, etc.). More research is, however, needed to integrate these concepts (ecological indicators, ecological thresholds and functional diversity) in a meaningful way.

4. Linking functional diversity to the provision of ecosystem services

The universal applicability and ecological relevance of the functional trait diversity concept makes it equally valuable to establish possible connections between global environmental change and the loss of ecosystem services. Ecosystem services that may be impaired by air pollution and climate change and that may be particularly associated with changes in functional diversity include C sequestration, soil fertility and nutrient cycling and pollination, among many others. However, research on the link between functional diversity and ecosystem services is lagging behind in the Mediterranean region where only a few controlled experiments exist (Hector et al., 1999; Pérez-Camacho et al., 2012; Tobner et al., 2014; Verheyen et al., 2016), species trait databases are still incomplete (Gachet et al., 2005; Paula et al., 2009), and field surveys along climatic and air pollution gradients are only recently starting to emerge (De Marco et al., 2015; Sicard et al., 2016).

The few studies available within the Mediterranean Basin context have shown that N deposition has already induced changes in functional diversity of epiphytic lichens along a NH₃ deposition gradient in Mediterranean woodlands, with a drastic increase and decrease of nitrophytic and oligotrophic species, respectively (Pinho et al., 2011). Similarly, a continuous increase of nitrophytic species (plants, lichens, mosses) has been detected in the Iberian Peninsula for the period 1900–2008 using the Global Biodiversity Information Facility (GBIF) database (Ariño et al., 2011). Increased N availability in nutrient-poor ecosystems like Mediterranean maquis can also alter plant functional composition (e.g., higher proportion of short-lived species in relation to summer semi-deciduous and evergreen sclerophylls), leading to changes in litter amount and quality (e.g., higher proportion of evergreen sclerophyll litter from affected shrubs and a general increase in lignin and N content in litter and a decrease in lignin/N ratio) and microbial community (e.g., reduction in biomass and activity), thus affecting nutrient cycling (an ecosystem function) and, therefore, soil fertility (including soil C accumulation, an ecosystem service) (Dias et al., 2010, 2013, 2014). In another study, Concostrina-Zubiri et al. (2016) showed that livestock grazing greatly affected the abundance and functional composition of moss–lichen biocrusts in a Mediterranean agro-silvo-pastoral system, with direct negative consequences on microclimate regulation and other ecosystem processes (CO₂ fixation, habitat provision and soil protection). This also affected the cork-oak regeneration processes, one of the traditional and most economically valuable services in these systems. Given the negative impacts of air pollution on cryptogamic biocrusts, a similar effect of air pollution on the cork-oak regeneration processes mediated by biocrusts might be expected.

5. Common experimental design, data sharing and global networks

The understanding of the ecological impacts of pollution and climate change across the Mediterranean region would improve through coordinated efforts and networks, which could take several forms. One possible approach is the use of large-scale regional surveys on existing pollution gradients representative of the current range of pollution loads (e.g., from big cities and/or extensive agricultural areas to their periphery). This approach was successfully used to survey 153 acid grasslands in ten countries across the Atlantic biogeographic zone of Europe (significantly less biodiverse than their Mediterranean counterparts) (Stevens et al., 2016).
where each partner surveyed sites in their local area according to an agreed protocol. Other networks have been successful using experimental approaches. For example, the Nutrient Network (NutNet) is a global network of over 90 sites following a common experimental protocol for nutrient addition and grazing (Borer et al., 2014). Similarly, the previously presented NitroMed network, originated within the CAPERmed platform, aims at using the same experimental protocols to integrate results from three comparable experiments in semiarid Mediterranean ecosystems. Other experimental networks have not used common experimental protocols, but through coordinated analyses have added value to individual experiments (Phoenix et al., 2012). Coordinated experimental networks (e.g., low-cost N addition experiments) bring many advantages such as the ability to assess the general applicability of results, additional statistical power resulting from well-established and robust statistical methods (e.g., linear mixed effects models, hierarchical Bayesian models, structural equation modelling), and opportunities to explore interactions with other natural and human-caused gradients such as climate, ecosystem and soil type, land use, atmospheric pollution (including O3 gradients), etc. They can also provide support and collaboration for individual scientists. An inventory of the existing sites with manipulation experiments in the Mediterranean Basin would provide added value to the individual sites through the implementation of common protocols and experiments.

In the Mediterranean region, another path to follow may be to build upon existing research and to participate more in already existing large-scale initiatives, in which the Mediterranean research community is not particularly well-represented. For example, interacting with the International Long Term Ecological Research (ILTER) network or with the International Cooperative Programme (ICP), established under the United Nation Economic Commission for Europe (UNECE) “Convention on Long-Range Transboundary Air Pollution” (CLRTAP) that includes several initiatives such as ICP Forest, ICP-Vegetation, and ICP-M, would facilitate the collection of large-scale spatial and temporal data series. Cooperation with other more specific networks like NitroMed (N deposition), ICOS (C cycle), and GLORIA (Alpine environments) would also help to establish a wider and more collaborative research community focused on air pollution impacts in Mediterranean terrestrial ecosystems.

The need of more coordination and investment to better understand the Mediterranean responses to climate change and air pollution has already been acknowledged by several groups of scientists both at the European (e.g. CAPErmed) and global scales (e.g. MEDDECO). These groups not only represent suitable arenas to discuss scientific results, but can also provide leading members able to manage the above-mentioned research and networking activities. However, all the presented approaches require considerable funding and determined political support to foster the exchange of information and best practices across the entire Mediterranean region and, thus, to promote the development of concrete projects and initiatives. In this context, the European Commission, through funding programs like Horizon 2020, could and should have, in our opinion, a pivotal role in supporting research projects (as it happened with the CIRCE project) and to provide the logistic means for transferring the scientific knowledge to the society.

Increasing awareness about the effects of climate change and pollution among stakeholders and society is encouraging the development of several European and Pan-European Programs (e.g. UNECE/ICP, Climate-ADAPT). One important step towards the coordinated action of the Mediterranean-basin countries in relation to Adaptation to climate change was the creation of “The Union for the Mediterranean Climate Change Expert Group” (UMCCEG), a partnership promoting multilateral cooperation between 43 countries (28 EU Member States and 15 Mediterranean countries). These initiatives show that opportunities do exist for countries to make progress. Due to campaigning, and partially because of the considerable losses from extreme weather events in recent years, public awareness in Mediterranean countries about risks associated with climate and air pollution increased. Governments and organisations at the EU level, national and sub-national level, have developed or are in the process of developing adaptation strategies. Therefore, there is an opportunity to make progress by actively engaging actors from all sections of the Mediterranean society.

6. Conclusions and future directions

The comparatively fewer number of studies on the effects of air pollution and its interactions with climate change on terrestrial ecosystems from the Mediterranean Basin is particularly noteworthy considering the high biodiversity, cultural value, and unique characteristics of this region such as high O3 levels, dominance of dry deposition over wet deposition, and long dry periods. Therefore, we emphasize the need to urgently implement common and coordinated research and experimental platforms in the Mediterranean region along with wider and more representative environmental monitoring networks. In particular, a robust connection between N deposition monitoring networks and modelling estimates is crucial. Ideally, monitoring and assessment programs should regularly include a set of common biomonitors such as local and/or transplanted cryptograms to identify local pollutant sources and, thus, help refine pollutant deposition maps (physiological indicators) and to provide early warning indication of potential critical thresholds (community shifts). Only by filling these gaps can the scientific community reach a full understanding of the mechanisms underlying the combined effects of air pollution and climate change in the Mediterranean Basin and, consequently, provide the science-based knowledge necessary for the development of sustainable environmental policies and management techniques and the implementation of effective mitigation and adaptation strategies. Finally, CAPErmed, a bottom-up initiative (from the researchers to the institutions), can be the longed-for catalyst that brings the Mediterranean community together and, therefore, represents an excellent opportunity to make all this happen.

Acknowledgements

This manuscript resulted from the 1st “Committee on Air Pollution Effects Research on Mediterranean Ecosystems” (CAPErmed) Conference celebrated in Lisbon, 2014 (http://capeired.weebly.com/). All attendants to this conference are, therefore, greatly acknowledged. Fundação para a Ciência e Tecnologia (FCT) is acknowledged through the Post Doc grant SFRH/BPD/85419/2012 to Teresa Dias and the Investigador grant to Silvana Munzi. ADM, EP and PS thank the LIFE10 ENV/FR/208 project MOTTLES. EP thanks also the Fondazione Cassa di Risparmio di Firenze (2013/7956). ROH is supported by a Juan de la Cierva Fellowship (JCI-2014-21252) of the Spanish Ministry of Economy and Competitiveness. The authors declare no conflicts of interest.

Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.envpol.2017.04.062.


