

1	Title: Soil	nutrients ar	nd microbial	biomass in	three	contrasting	Mediterranean	forests
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21 ABSTRACT

- 22 Aims: The extent to which the spatial and temporal patterns of soil microbial and available nutrient pools hold
- 23 across different Mediterranean forest types is unclear impeding the generalization needed to consolidate our

24 understanding on Mediterranean ecosystems functioning.

- 25 Methods: We explored the response of soil microbial, total, organic and inorganic extractable nutrient pools (C,
- 26 N and P) to common sources of variability, namely habitat (tree cover), soil depth and season (summer drought),
- 27 in three contrasting Mediterranean forest types: a *Quercus ilex* open woodland, a mixed *Q. suber* and *Q.*
- 28 canariensis woodland and a Pinus sylvestris forest.
- 29 Results: Soil microbial and available nutrient pools were larger beneath tree cover than in open areas in both oak
- 30 woodlands whereas the opposite trend was found in the pine forest. The greatest differences in soil properties
- 31 between habitat types were found in the open woodland. Season (drought effect) was the main driver of
- 32 variability in the pine forest and was related to a loss of microbial nutrients (up to 75% loss of N_{mic} and P_{mic}) and
- 33 an increase in microbial ratios (C_{mic}/N_{mic}, C_{mic}/P_{mic}) from Spring to Summer in all sites. Nutrient pools
- 34 consistently decreased with soil depth, with microbial C, N and P in the top soil being up to 208%, 215% and
- 35 274% larger than in the deeper soil respectively.
- 36 Conclusions: Similar patterns of variation emerged in relation to season and soil depth across the three forest
- 37 types whereas the direction and magnitude of the habitat (tree cover) effect was site-dependent, possibly related
- 38 to the differences in tree species composition and forest structure, and thus in the quality and distribution of the
- 39 litter input.
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- 41 Keywords: soil fertility, plant-soil interactions, soil carbon, nitrogen, phosphorus
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44 INTRODUCTION

Soil microorganisms, in their role in organic matter decomposition, have the capacity to both mineralize and immobilize nutrients (Singh et al. 1989) thereby influencing soil nutrient availability and plant growth (Lambers et al. 1998). Spatial and temporal changes in soil microbial biomass may determine the patterns of availability of limiting nutrients such nitrogen (N) and phosphorus (P), thus having profound influence on plant communities and ecosystem functioning (Ettema and Wardle 2002; Gallardo and Schlesinger 1994; Sardans et al. 2005; van der Putten et al. 2009).

51 Spatial and temporal variations of soil microbial biomass and activity are related to different biotic and 52 abiotic factors that modulate the temperature, moisture conditions and substrate quality and availability. For 53 instance, vegetation composition and structure control the spatial distribution, quality and quantity of nutrients 54 inputs via litter and root exudates (Aponte et al. 2011; Huang et al. 2013; Prescott and Grayston 2013; Ushio et 55 al. 2010). Soil nutrients and microbial activity usually decrease as soil depth increase due to a decline in the 56 quality and quantity of organic matter (Gaudinski et al. 2000; Xiang et al. 2008). Seasonal changes in 57 temperature, water and substrate availability also have a large impact on soil microbial activity and nutrient 58 cycling (Corre et al. 2002; Quilchano and Marañón 2002; Schmidt et al. 1999). In highly seasonal ecosystems, 59 such as Mediterranean forest, the effects imposed by seasonal variations, in particular associated to the summer 60 drought, are especially important for ecosystem functioning (Aponte et al. 2010b; Marañón-Jiménez et al. 2011; 61 Matías et al. 2011).

62 Many studies have described soil nutrient heterogeneity in Mediterranean forests; however, most of them have 63 been conducted at local spatial scales, focused on a single forest type (Barcenas-Moreno et al. 2011; Carreira et 64 al. 1994; Gallardo 2003; García et al. 2006; Maltez-Mouro et al. 2005; Monokrousos et al. 2004). The detection 65 of general patterns across different forest types is necessary to fully understand microbial biomass and nutrients 66 dynamics and their consequences for plant community. At the same time, the emergence of site-dependent 67 effects will be of interest from a modelling and management perspective, to properly determine nutrient pools at 68 wide geographical scales including a mosaic of forest types. Patterns of microbial biomass and nutrient 69 heterogeneity across different forest types have been largely investigated in temperate, boreal and tropical forest 70 (e.g. Hackl et al. 2005; Lindo and Visser 2003; Liu et al. 2012; Zhong and Makeschin 2006), while remain far 71 less studied in Mediterranean forest (but see García et al. 2002; Goberna et al. 2006). This coordinated study 72 addressed this knowledge gap and aimed to evaluate whether the effects of main sources of variability, namely 73 habitat (i.e. tree cover), soil depth and season, in the soil nutrients and microbial C, N and P pools could be

74 generalised across three contrasting Mediterranean forests: a Quercus ilex open woodland, a mixed Q. suber and 75 Q. canariensis woodland and a Pinus sylvestris forest. While this study builds upon previous knowledge on soil 76 nutrient heterogeneity at local scales (Aponte et al. 2010b; Matías et al. 2011), it focuses on the comparison 77 among forests with different structure and species composition, thus taking a step forward towards 78 understanding general patterns of soil microbial responses to biotic and abiotic environmental drivers. 79 Explicitly, we aimed to answer the following questions: 1) Is there a common pattern across the three forests in 80 relation to the tree effect, soil depth and seasonal drought?; 2) Are the interactions between the effects of tree 81 cover, soil depth and season (summer drought) similar across forest types?; 3) What is the quantitative 82 importance of the studied factors (tree effect, soil depth and seasonal drought) on the soil and microbial 83 variables in each forest type? 4) Do the relationships between microbial and soil chemical properties hold when 84 examined across forest types?

85

86 METHODS

87 Study areas

88 The study was conducted in three different Mediterranean forest types: a mixed woodland of Quercus suber L. 89 (evergreen) and Q. canariensis Willd. (deciduous) in Los Alcornocales Natural Park in the extreme south, near 90 the Strait of Gibraltar, an open woodland dominated by the sclerophyllous *Quercus ilex* subsp. ballota L. and 91 eventually mixed with other Quercus species (Q. suber, Q. pyrenaica Willd., Q. faginea Lam.) in Sierra de 92 Cardeña and Montoro Natural Park (Cardeña), in the south mainland, and a forest mainly comprised of Pinus 93 sylvestris L. interspersed with Q. ilex subsp. ballota in Sierra Nevada National Park in the southeast of Spain 94 (Fig. 1). In all three forest types, the main tree species are intermingled with open areas covered by sparse 95 herbaceous vegetation. The study sites vary in altitude, climate and soil conditions (Table 1). The general 96 climate of the three sites is Mediterranean-type, characterized by hot and dry summers, and cold and wet winters 97 with most rainfall occurring from October to May. The sites in Cardeña and Sierra Nevada experience more 98 extreme temperatures due to their continental and altitudinal locations (respectively), while temperatures in 99 Alcornocales site are milder due to the lower elevation and proximity to the Mediterranean Sea and Atlantic 100 Ocean. Mean annual rainfall follows a rising gradient from Cardeña to Alcornocales (Table 1). The sites in 101 Alcornocales and Cardeña stand on a bedrock of sandstone and granite, both producing acidic sandy soils. On 102 the contrary the site in Sierra Nevada stands on limestone, which gives rise to basic loamy soils. Cambisols 103 dominated in Alcornocales and regosols in Cardeña (nomenclature follows WRB 2006), indicating a greater

soil development i.e. soil depth, structure, water holding capacity and chemical fertility in the former than thelater.

106

107 Experimental design

108 At each forest site 10-20 replicates (depending on the site, Table 1) of two main habitat types were identified 109 within a stand: beneath the canopy of the dominant tree species (Q. suber and Q. canariensis in Alcornocales, 110 O. ilex in Cardeña and P. sylvestris in Sierra Nevada), and in open areas with bare soil or sparse herbaceous 111 cover and no tree cover. These habitat types will be referred as 'Tree' and 'Open' respectively hereafter. At 112 each replicate point, four soil cores (0-16 cm) were extracted using an auger after removing the litter layer, 113 divided between 'Top soil' (0-8 cm) and 'Deeper soil' (8-16 cm) and homogenized within the same depth to 114 obtain a composite soil sample per habitat type replicate and depth. Soil samples were taken in Spring (May-115 June) and Summer (August-September) 2007, coinciding with the moment of maximum soil biological activity 116 and maximum water stress in soil, respectively. In total 400 soil samples were taken corresponding to 10-20 117 replicates (Table 1) of 2 habitat types x 2 soil depths x 2 seasons x 3 forest sites. Litter, i.e. dead plant material 118 relatively undecomposed standing on the ground, was collected once in all sampling points using a 10 x 10 cm 119 quadrat (in Sierra Nevada) or a 30 x 30 cm quadrat (in Alcornocales and Cardeña). Litter samples were oven-120 dried at 60°C for 72 h and weighted.

121

122 Laboratory analyses

Soil samples were brought to the laboratory in an ice-box, fresh-sieved at 2 mm removing stones, roots and other recognizable plant parts and stored at 4°C for analyses. Water content was determined on a subsample as the difference in weight between fresh and oven dried (105°C) soil.

126 Microbial C, N and P were estimated in fresh soils using a chloroform fumigation-extraction procedure

127 (Brookes et al. 1985; Brookes et al. 1982; Vance et al. 1987). Dissolved organic C (DOC) and N (DON) and

128 inorganic P (P_{inorg}) were determined in non-fumigated and chloroform fumigated soil subsamples (24h).

129 Dissolved C and N were extracted with 0.5M K₂SO₄, and their concentration was determined using a Shimadzu

- 130 TOC-V CSH analyzer. Inorganic P was extracted with either 0.025N HCl+0.03N NH₄F (Bray Kurtz 1 method
- 131 (Bray and Kurtz 1945) for the acidic soils of Alcornocales and Cardeña) or 0.5M NaHCO₃ (Olsen method
- 132 (Olsen et al. 1954) for the basic soils of Sierra Nevada) and its concentration was determined by colorimetry

using the ascorbic acid-molybdenum blue method (Sparks 1996). Microbial C (C_{mic}), N (N_{mic}) and P (P_{mic}) were estimated as the difference in DOC, DON and P between fumigated and non-fumigated samples.

135Inorganic nitrogen (N_{inorg}) was extracted from non-fumigated soils using 2M KCl and the extracts were136analyzed for NH_4^+ and NO_3^- by the Kjeldhal method (Bremner and Keeney 1965). Soil total C (C_{tot}) and N (N_{tot})137were determined on oven dried soils by combustion at 850°C (Leco TruSpec autoanalyzer) and total inorganic C138(C_{inorg}) was measured by acidification with HClO4 in a TIC analyzer (UIC CM-5014). The difference between139 C_{tot} and C_{inorg} gave the total organic C (C_{org}).

140

141 Data analysis

142 Differences among habitat, soil depth and season were analyzed using repeated measurement ANOVAs with 143 season as a within-group effect and habitat and depth as between-group effects. Forest site was also included in 144 the analysis to test for potential interactions with the studied factors. Variables were transformed (log, arcsin) 145 when necessary to meet normality assumptions. To control the type I error inflation resulted from repeated tests, 146 the false discovery rate (FDR), i.e. the expected proportion of tests erroneously declared as significant, was 147 controlled at 5% using a step-up procedure (Benjamini and Hochberg 1995; García 2003). The percentage of 148 the total variance explained by the studied factors (habitat, soil depth and season) was calculated for each 149 variable and site using a repeated measurement ANOVA with no interactions. Patterns in pairwise Pearson's 150 correlations between microbial and soil nutrient factions were explored using correlation network analysis (R 151 package igraph, Csardi and Nepusz 2006). Multivariate relationships between variables were analysed using 152 Principal Component Analysis (PCA). The 'Broken stick' method (King and Jackson 1999) was used to select 153 significant components. Habitat, soil depth and season were included in the PCA as supplementary variables, 154 i.e. these factors did not participate in the analysis, but were projected on the multivariate space generated by the 155 PCA for the purpose of interpretation.

156

157 RESULTS

158 Overall, the study forests differed in all the analyzed soil and microbial properties (Table 2 and 3). Cardeña was

159 the least fertile site while Alcornocales had the largest fraction of microbial nutrients (from 3 to 6-fold the

160 values of the other sites) and the largest pool of total and dissolved C and N and organic C (Fig. 2 and 3). Sierra

- 161 Nevada showed the highest inorganic N and P values (~2-fold to 8-fold the values of the other sites), the highest
- 162 C_{mic}/N_{mic} ratio (2-fold) and the largest litter pool (~6-fold) (Fig. 3). The ratios of nutrients retained in the

- 163 microbial biomass vs. the pool of available nutrients (N_{mic}/N_{inorg} and P_{mic}/P_{inorg}) as well as the fraction of soil
- 164 organic carbon and total nitrogen in the microbial biomass (C_{mic}/C_{org} and N_{mic}/N_{tot}) were the highest in

165 Alcornocales and Cardeña and the lowest in Sierra Nevada (Online Resource 1).

166

167 Effect of habitat

168 Soil parameters differed significantly between the two habitat types in all forest sites (Table 3, Fig. 2 and 3). 169 However, the magnitude and direction of those differences varied across sites, as the interaction Site \times Habitat 170 was significant for most of the variables (Table 3). In both oak woodlands, Alcornocales and Cardeña, the 171 nutrient pools (microbial, dissolved organic and inorganic) tended to be larger beneath tree canopy than in open 172 areas with the exception of nitrate that showed the opposite trend (Fig. 2 and 3). Greater concentrations of 173 ammonium (156% in both sites), phosphate (120% in Alcornocales and 182% in Cardeña), and microbial 174 nutrients (123 and 166% for C_{mic} ; 126 and 227% for N_{mic} ; 215 and 175% for P_{mic}) were found beneath tree cover 175 than in open areas. Mean soil organic carbon was also greater beneath tree cover than in open areas in Cardeña 176 (1.7% vs. 0.97%; P<0.0001) and Alcornocales (4.1% vs. 3.6%, not significant difference). A different pattern 177 was observed in the pine forest (Sierra Nevada) where most of the soil nutrient pools were similar between the 178 two habitats or even decreased beneath trees as it occurred with N_{mic} and inorganic N and P (Fig. 2 and 3). 179 Organic and inorganic C also decreased significantly from open areas (3.1%, and 2.85% respectively) to 180 beneath pine tree cover (2.9% and 0.93% respectively). Nevertheless, the amount of litter was larger beneath 181 tree canopy than in open areas in all sites, being the values larger in Sierra Nevada than in the other two forests 182 (Table 1). There were no habitat differences in the fractions of microbial values relative to soil pools 183 $(N_{mic}/N_{inorg}, P_{mic}/P_{inorg}, C_{mic}/C_{org} and N_{mic}/N_{tot}; data not shown).$

184

185 *Effect of soil depth*

In general all variables measured showed a consistent pattern with soil depth in the three forest sites, with values decreasing from Top soil to Deeper soil (Fig. 2 and 3). However, there was a significant Site × Depth interaction (Table 3) due to the lack of statistical significance of soil depth for many variables in Cardeña (C_{mic} , P_{mic} , DON, NH₄ and P_{inorg}) (Fig. 2 and 3).

Microbial C, N and P in Top soil were higher than in Deeper soil with the largest variations found in
Alcornocales (208, 215 and 274% respectively) and the smallest changes found in Cardeña (128, 155 and 119%
(Fig. 2). On average across sites, the pool of inorganic N and P, DOC, DON and C_{org} was 133% (site mean

range 110 – 146%), 155% (129 – 177%), 142% (117-172%), 140% (112-159%) and 118% (114-120%) higher in Top soil than in Deeper soil respectively. As with microbial pools, variation was the least in Cardeña (Fig. 2 and 3). Microbial ratios C_{mic}/C_{org} and N_{mic}/N_{tot} showed the largest decrease with soil depth in Alcornocales but remained constant in Cardeña. The ratio of microbial biomass nutrients (C_{mic}/N_{mic} and C_{mic}/P_{mic}) showed no significant variation from Top soil to Deeper soil in any site. The only exception was found for soils in open areas in Cardeña where C_{mic}/P_{mic} increased with soil depth from 55 to 143, as evidenced by a significant Site × Habitat × Depth interaction (Table 3).

200

201 Effect of season

202 Soil microbial fractions and nutrient pools varied significantly with the season. However, the seasonal patterns 203 of variation were site-dependent as indicated by a significant Site × Season (Table 3). Seasonal variations were 204 stronger in Sierra Nevada and Alcornocales whereas Cardeña showed the lowest variability between seasons 205 (Fig. 2 and 3). In general, microbial pools were larger in Spring than in Summer, particularly for N_{mic} and P_{mic} 206 which values were on average 237% and 258% higher in Spring (Fig 2). The fraction of microbial C and N 207 relative to soil total pools (Cmic/Corg and Nmic/Ntot) decreased from Spring to Summer in Sierra Nevada but not in 208 Cardeña. Microbial ratios (Cmic/Nmic, Cmic/Pmic) increased from Spring to Summer in all sites revealing a larger 209 loss of N_{mic} and P_{mic} as compared to C_{mic}.

210 The seasonal variability of N_{mic}, P_{mic}, C_{org}, DOC and DON was larger in soils beneath tree canopy 211 whereas the variation was subdued in the open habitats (Season × Habitat interaction, Table 3). We also found a 212 strong and significant Site \times Season interaction for N_{inorg} and P_{inorg} (Table 3), which was due to opposite seasonal 213 changes across forest types. For example, the pool of available inorganic nutrients (ammonium and phosphate) 214 as well as DON increased from Spring to Summer in Cardeña and Sierra Nevada, whereas the values decreased 215 in Alcornocales (Fig. 3). Despite the discrepancies in the seasonal dynamics of P_{inorg}, the proportion of P_{mic} 216 relative to P_{inorg} was higher in Spring than in Summer in all sites (data not shown). The observed seasonal 217 patterns were similar at the two soil depths.

218

219 Variance partitioning among habitat, soil depth and season

As shown in the partition of variance (Fig. 4) and the principal component analysis (Fig. 5) the main drivers

221 of variability differed between sites. Soil depth and season accounted for the largest part of the variability

222 observed in the microbial and soil nutrient pools in Alcornocales and Sierra Nevada. For instance, in

Alcornocales soil depth explained 50, 38 and 30% of the variation of microbial C, N and P respectively and

season explained 55 and 39% of the variation of N_{inorg} and P_{inorg}. In Sierra Nevada season was the main driver of

225 microbial variability accounting for 26, 58 and 66% of the variation of microbial C, N and P. In contrast, the

variability of soil biotic and abiotic properties in Cardeña was mainly driven by habitat type, which explained

- 227 10, 16 and 4% of microbial C, N and P variation respectively and 9 and 12% of N_{inorg} and P_{inorg} variation.
- 228

229 Relations between microbial pools and soil properties

230 Soil microbial C, N and P were significantly correlated among them in all sites (Fig. 5 and Online Resource 1). 231 Microbial C and N were consistently and strongly coupled (r >0.76 in all sites), whereas P_{mic} was more weakly 232 but still significantly related with C_{mic} (from r = 0.28 in Cardeña to r = 0.69 in Sierra Nevada) and N_{mic} (from r 233 = 0.30 in Cardeña to r = 0.82 in Sierra Nevada). Microbial C, N and P were positively related to most of the 234 measured soil properties in each site (Fig 5). The strongest correlations were found with C_{org} , N_{tot} and soil 235 moisture reflecting microbial biomass dependence on substrate and water availability. Microbial C also showed 236 a significant correlation with P_{inorg} in all sites (r ~0.36). The relationship between litter and C_{org} (r_C) and N_{tot} (r_N) 237 varied across forest sites, being positive in Cardeña ($r_c = 0.43$, $r_N = 0.35$, P < 0.0001, seasons and depths 238 pooled), positive in Top soil in Alcornocales ($r_c = 0.28$, $r_N = 0.27$, P < 0.05; not significant in Deeper soil) and 239 negative in Sierra Nevada ($r_c = -0.16$, P < 0.06; $r_N = -0.36$, P < 0.0001). The correlation network was the 240 strongest in Alcornocales, i.e. there was a tight coupling between most variables, and the weakest in Cardeña 241 (Online Resource 1).

242 The multivariate analyses (PCAs) showed similar patterns of covariation among the nutrient pools for 243 all sites (Fig 5). Two main significant gradients (axes) emerged for each PCA from the analysis based on the 244 'Broken --stick' method (King and Jackson 1999). For all sites the first axis was strongly correlated to microbial 245 C, N and P, total N and organic C. In Alcornocales and Sierra Nevada the firts axis was also positively related to 246 soil moisture and negatively related to Cmic/Nmic. The separation of samples along the main axis and the analysis 247 of the supplementary variables indicated that both season and soil depth imposed a similar degree of variability 248 in Alcornocales whereas season was the main driver of variation in Sierra Nevada, which is agreement with our 249 variance partitioning analysis. In Cardeña the first axis was related to litter amount, but not to soil moisture, and 250 separated the samples by habitat type. The second axis in all PCAs was related to the availability of inorganic 251 nutrients (N and/or P), which covariation with other variables was inconsistent across forest types. Higher 252 microbial ratios (Cmic/Nmic) were consistently associated to lower soil moisture and Summer samples in all sites.

The relationship between litter abundance and microbial and total nutrient pools was positive in Alcornocalesand Cardeña, but negative in Sierra Nevada.

Variables covaried similarly when all three sites were combined in a single PCA (Online Resource 2): the first axis accounted for 34% of the variability and was strongly correlated to most nutrient pools (microbial, dissolved organic and total) and soil moisture. The second axis accounted for 27% of the variability and was mostly related to inorganic N and P. The two axes clearly separated between forest sites, with Cardeña at the poorest end of both axes and Alcornocales and Sierra Nevada at the richest end of the first and second axes, respectively.

261

262 DISCUSSION

263 Overall, the three sources of variability considered (habitat, soil depth and season) had significant effects on the 264 soil microbial pools and nutrient concentrations in the studied forests. However the direction and magnitude of 265 these effects varied across forest types and with the soil parameter examined.

266 The expected positive effect of tree canopy on soil and microbial nutrients was confirmed for the two oak 267 woodlands (Cardeña and Alcornocales) but not for the pine forest (Sierra Nevada), where the soil and microbial 268 nutrients pools were smaller beneath tree canopy than in open areas. The inconsistency of the habitat effect 269 could be attributed to the forests' distinct species composition. Trees generate species-specific effects on soil 270 conditions through multiple pathways, such as changing microclimatic conditions or via leaf and root litter input 271 or root exudates (Alameda et al. 2012; Aponte et al. 2013; Aponte et al. 2011; Malchair and Carnol 2009). Tree 272 species changes in soil abiotic properties might in turn affect soil biota (Aponte et al. 2013; Aponte et al. 2010a; 273 Prescott and Grayston 2013). In particular, tree-mediated changes in soil acidity and in the amount and quality 274 of substrate are known to affect microbial communities size and composition (Lucas-Borja et al. 2012; Sagova-275 Mareckova et al. 2011; Thoms et al. 2010). In Sierra Nevada soil acidity was higher beneath pine cover than in 276 open areas, as evidenced by their distinct pH (7.7 vs. 8.1) and C_{inorg} (0.93% vs. 2.85% respectively), while clay 277 content was lower (18.5 vs. 21.6%). Litter biomass was 15 times greater (8594 vs. 559 g m⁻²) and the amount 278 and quality of the substrate (Core, DOC, Ntot, Core/Ntot) were significantly lower beneath tree cover (Pinus) than 279 in open areas, in agreement with the negative correlation observed between litter and soil C_{org} and N_{tot} . 280 Meanwhile, the opposite was found in the two oak forest sites, i.e substrate quality was higher beneath tree 281 cover (Quercus) than in open areas, and it was positively related to litter biomass, thus sustaining the 282 counteracting patterns observed for microbial nutrients. In accordance with our results, previous studies on the

283 effects of tree species on soils have related the lower soil nutrient and microbial values found beneath pine 284 cover, compared to other broadleaves tree species (including *Quercus*), with the poorer quality of the pine litter, 285 and thus to its lower decomposition rate and nutrient release, and its capacity to acidify soils (Augusto et al. 286 2002; Rutigliano et al. 2004; Smolander and Kitunen 2002; Ste-Marie et al. 2007). Nonetheless, the observed 287 differences in soil and microbial nutrients between habitat types should not be solely attributed to vegetation 288 cover. Other soil physicochemical properties, such as soil depth, structure and texture, which may be the 289 underlying reason for the distinct cover type, can also control microbial development (Hassink 1994). 290 Interestingly our results also revealed a difference in the magnitude of the positive tree-effect on soil 291 nutrients between the two oak woodlands, Cardeña and Alcornocales. These two sites significantly differed in 292 their soil type and nutrient content: Cardeña sited over Regosols, i.e. weakly developed soils with a low organic 293 matter content and water holding capacity (WRB 2006) (Table 1). In contrast, soils in Alcornocales were 294 cambisols (also known as Brown forest soils, WRB 2006), they were well structured and presented a thick 295 humic horizon (15-20cm beneath tree canopy; Garcia et al, unpublished data), and a relatively high soil organic 296 matter content (11% in 0-25cm upper soil, Polo 2006). Mean site Corg was greater in Alcornocales (3.9%) than 297 in in Cardeña (1.3 %), clay content was 7 times higher in the former (36%) than in the later (5%), and CEC 298 (cation exchange capacity) was two-fold in Alcornocales than in Cardeña (Table 1), all of which supported the 299 distinct soil fertility and microbial nutrient levels observed in both forest sites (Table 2). These two sites also 300 differed in their stand structure, with a lower tree density in Cardeña than in Alcornocales (131vs. 219 stems ha 301 ¹). It is possible that the interaction between their distinct soil types and stand structure could be determining 302 why habitat type was the main driver of variability of soil microbial properties in Cardeña but it was of lesser 303 importance in Alcornocales. The intensity of tree effects on soil properties is modulated by the spatial 304 distribution of tree canopies (Bennett et al. 2009; Ushio et al. 2010). It is well-known that oak trees in 305 Mediterranean savannah-like systems (dehesas) generate islands of fertility beneath their canopies where the 306 leaf litter and root exudates accumulate and build up the soil organic matter that sustain microbial biomass and 307 nutrient cycling (Alameda et al. 2012; Gallardo 2003). In sparse forests, such as Cardeña, trees are scattered in a 308 matrix of open areas and their footprints on soil fertility are expected to be more intense beneath the canopy. In 309 contrast, a more diffuse footprint occurs in dense forests where open areas are intermingled in a matrix of trees. 310 This is consistent with C_{org} and C_{mic} being greater beneath tree cover than in open areas by a factor of 1.7 and 1.7 311 in Cardeña and a factor of 1.1 and 1.2 in Alcornocales respectively. In addition, the small concentrations of

312 substrate (Core, DOC, Ntot, DON) in Cardeña could be a limiting factor for microbial biomass and a tree-313 mediated increase in its availability would render a larger boost of microbial growth than in more fertile sites. 314 Microbial C, N and P showed a common and seasonal pattern, with values decreasing from Spring to 315 Summer in response to summer drought. This response was the weakest in Cardeña, where changes were not 316 significant. Seasonal variation was larger for N_{mic} and P_{mic} than for C_{mic}, rendering a shift in the microbial ratios, 317 as evidenced by the multivariate analyses. The change in C_{mic}/N_{mic} was the largest in Sierra Nevada (from 9 in 318 Spring to 34 in Summer), where a decrease in C_{mic}/C_{org} , a proxy for microbial C assimilation efficiency 319 (Sparling 1992), was also observed. Soil microorganisms in Mediterranean ecosystems have adapted to 320 withstand the seasonal variation in water availability and temperature that define the Mediterranean-type climate 321 (Goberna et al. 2007). Seasonality, in particular the summer drought, may influence microbial biomass directly 322 by inducing microbial metabolic responses to changes in soil moisture and temperature (Chen et al. 2003; 323 Jensen et al. 2003), or indirectly by influencing plant productivity, organic matter release and C diffusion in soil, 324 and hence substrate availability (Rey et al. 2002; Xiang et al. 2008). The high microbial values found in Spring 325 may reflect favourable environmental conditions and more labile substrates derived from roots or from materials 326 incorporated into the soil whereas the decrease in Summer might indicate a loss in the total number of 327 organisms. This is consistent with previous work conducted in the same forest stand in Alcornocales, which 328 showed higher soil enzyme activity during the rainy season than in summer (Quilchano and Marañón 2002). On 329 the other hand, summer increases in microbial ratios can be related to an increasing proportion of fungi vs. 330 bacteria (Jensen et al. 2003), since fungi have a higher carbon to nitrogen ratio (Cmic/Nmic) (related to the their 331 lower efficiency, Cleveland and Liptzin 2007); and are more drought-tolerant than bacteria(Wilkinson et al. 332 2002). In addition at low water potentials, fungi are able to increase their cytoplasmic C (thus further increasing 333 C_{mic}/N_{mic} and C_{mic}/P_{mic}) to reduce osmotic pressure and maintain hydration (Schimel et al. 2007). We propose 334 that the loss of N_{mic} and P_{mic} as compared to C_{mic} in all sites could be explained by a net decrease in the size of 335 the microbial biomass, driven by lower substrate (Corg) and water availability, together with an increase in the 336 proportional abundance of fungi. However, neither microbial activity nor community composition indicators 337 were measured in our study, thus the underlying mechanisms for the observed seasonal changes remain unclear. 338 Although the microbial pool showed a common trend affected by the summer drought, we observed 339 significant discrepancies on the seasonal dynamics of the available pools. In Alcornocales, nutrient availability 340 was higher in Spring, whereas the opposite was found for the other two sites. Net nutrient pools size is the result 341 of the nutrient release through mineralization, nutrient immobilization and uptake by microorganisms and

342 plants. The rates of N mineralization and nitrification can be more influenced by soil type and soil organic 343 matter quality than by changes in temperature, and the effect of temperature on the rate of P mineralization can 344 vary among soil types (Nadelhoffer et al. 1991). The more severe summer drought in Cardeña and Sierra 345 Nevada might reduce plant uptake capacity (Kozlowski and Pallardi 2002), and increase the proportion of 346 nutrients in the soil when compared to Alcornocales. In a climate change study conducted in the same forest site 347 in Sierra Nevada, Matías et al. (2011) observed that under a dry scenario (30% summer rainfall reduction) soil 348 available nutrients increased and plant and microbial nutrient pools decreased. Thus the contrasting seasonal 349 patterns observed could be the result of different interacting factors such as the activity rates of soil 350 microorganisms, the substrate availability and accessibility, the soil acidity and texture, and the plant nutrient 351 uptake.

352 The effect of soil depth, i.e. decreasing soil and microbial nutrient content from Top soil to Deeper soil, was 353 similar in all forest types. However the magnitude of the change varied among forests, with Cardeña showing 354 the smallest changes. This effect of soil depth has been previously reported for Mediterranean and other forest 355 types (Aponte et al. 2010b; Raubuch and Joergensen 2002; Ross et al. 1996; Wang et al. 2004), the main causes 356 being a decrease in the labile C pools and an increase in the concentration of recalcitrant compounds (Fierer et 357 al. 2003; Goberna et al. 2006). In our study C_{org} decreased with soil depth in all sites, the largest change 358 observed in Alcornocales (from 4.6% to 3.2%) and the smallest one in Cardeña (from 1.5% to 1.2%). The 359 stronger vertical development of cambisol soils in Alcornocales, as evidenced by the deeper soil layer having a 360 significantly lower amount (Corg) and quality (Corg/Ntot) of carbon compounds than the top soil, explains the 361 larger variability of soil properties associated to soil depth observed in this site. This is consistent with the 362 changes observed in microbial C and N values related to soil total pools (Cmic/Corg and Nmic/Ntot) in 363 Alcornocales, which are an indicator of a lower efficiency of the microbial biomass to assimilate C and N 364 possibly due to a higher proportion of the soil organic matter being highly recalcitrant (Sparling 1992). 365 Meanwhile, the dominance of shallower and more weakly developed soils in Cardeña (i.e. regosols) underpin 366 the low importance of soil depth as a driver of soil and microbial nutrient content. 367 The size of the microbial pool fell within the ranges observed in other Mediterranean forests (Gallardo et al. 368 2000; Goberna et al. 2006) although it differed significantly between sites. It was not within the scope of this 369 study to investigate the overall differences between forest types, but in general differences among the studied 370 forest types were probably underpinned by the variation in soil types, the amount and quality of soil organic

371 matter, the soil texture and water content, all of them factors constraining the size of the soil microbial biomass

(Bohlen et al. 2001; Nielsen et al. 2009). For example, clay content was the highest in Alcornocales (site mean
of 35% *vs.* 8% in Cardeña). Clay content is positively related to microbial biomass and soil organic carbon
because it protects microbial biomass from predation by creating refuge microsites. Furthermore, it increases
soil organic matter stabilization and soil water retention thus enhancing soil conditions for microbial
development (Insam et al. 1989; Sparling 1992).

377

378 Conclusions

Our findings revealed that across three contrasting Mediterranean forest types with significant differences in soil abiotic conditions, the microbial nutrient pools showed a consistent response in relation to soil depth and seasonal (drought effect) variability, which is indeed mirrored in many other ecosystems at a global scale. In contrast, the direction and magnitude of the variability associated to habitat (tree effect) varied among forest types suggesting a higher complexity in the biotic interactions between the aboveground and belowground components of these ecosystems. Few consistent interactions between factors (tree effect, soil depth and seasonal drought) were observed across forest types.

Microbial and soil chemical properties showed similar patterns of covariation in all sites, with microbial biomass responding to variations in the amount and quality of soil organic carbon and soil moisture. Thereby, the quantitative importance of the three studied factors on soil microbial nutrients varied across site, being the most important factor in each case that one which alleviated limitations and imposed the largest variability in substrate and water availability. As such, differences in forest structure and species composition between forest types would underpin the observed inconsistent tree effect on soil microbial properties, since they are related to the amount, quality and spatial and temporal distribution of the resources available to soil microorganisms.

393

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- 598 599

600 FIGURE CAPTIONS

601

602 **Fig. 1.** Location of the studied forest sites in the Iberian Peninsula.

Fig. 2. Microbial and soil nutrient fractions in Alcornocales (A), Cardeña (C) and Sierra Nevada (SN).

604 Differences between the levels of each factor are as indicated by repeated measurement ANOVAs with season

605 as a within-group effect and site, habitat and depth as between-group effects followed by Tukey's posthoc

606 comparisons (* P < 0.05; ** P < 0.01; *** P < 0.001). Abbreviations are: C_{mic}, microbial C; N_{mic}, microbial N;

607 P_{mic}, microbial P; DOC, dissolved organic C; DON, dissolved organic N.

608 Fig. 3. Soil inorganic nutrient fractions and organic carbon in Alcornocales (A), Cardeña (C) and Sierra Nevada

609 (SN). Differences between the levels of each factor are as indicated by repeated measurement ANOVAs with

610 season as a within-group effect and site, habitat and depth as between-group effects followed by Tukey's

611 posthoc comparisons (* P < 0.05; ** P < 0.01; *** P < 0.001). Abbreviations are: P_{inorg}, inorganic available P;

612 C_{org}, organic C.

613 Fig. 4. Percentage of the total variance explained by each of the studied factors, habitat (tree effect), soil depth

and season, for each variable in each site. C_{mic}, N_{mic}, P_{mic}: microbial C, N and P, respectively; Corg: organic C;

615 P_{inorg}: inorganic P; N_{inorg}: inorganic N; DOC, DON: dissolved organic C and N, respectively; C_{mic}/N_{mic}, C_{mic}/P_{mic}:

616 ratios between respective variables; Overall: mean across all variables.

- 617 **Fig. 5**. PCA ordination plot showing the distribution of Spring and Summer values of each study sites. C_{mic},
- 618 N_{mic}, P_{mic}: microbial C, N and P, respectively; P_{inorg}: inorganic P; DOC, DON: dissolved organic C and N,

619 respectively; N_{tot}: total N; C_{org}: organic C; C_{mic}/P_{mic}, C_{mic}/N_{mic}, N_{mic}/N_{inorg}, P_{mic}/P_{inorg}: ratios between respective

620 variables. Depth, season and habitat (in grey) are supplementary variables included as passive in the analysis.

621

623 TABLES

- Table 1. Characteristics of the studied forest sites. Values are site means (± standard deviation, when
- 625 provided) and habitat means in square brackets [Open; Tree].
- 626

	Alcornocales	Cardeña	Sierra Nevada		
Coordinates	36°31' N, 5°34' W	38° 15´ N, 4° 21´ W	37°05' N, 3°28'W		
Altitude (m a.s.l.)	545	750	1650		
Soil					
Bedrock	sandstone	granite	limestone		
pН	acidic	acidic	basic		
	$[6.34; 6.07]^{a}$	5.4 ^b	$[8.1; 7.7]^{c}$		
Soil type	cambisol	regosol	regosol, cambisol		
Texture	sandy	sandy	loamy		
Sand (%)	[44; 49] ^a	[80; 79] ^d	[22; 19] ^c		
Clay (%)	[39; 33] ^a	$[4.8; 4.5]^{d}$	[29; 37] ^c		
CEC (meq 100g ⁻¹)	[23.1; 19.7] ^a	[7.8; 8.6] ^d	[14.7; 18.5] ^c		
Litter (g m^{-2})	[45±38; 936±350]	$[282\pm 259; 1200\pm 875]$	[559±362; 8594±5543]		
Climate					
Temperature (°C)					
mean annual	15.5	15.3	12.1		
mean min	9.1	7.3	-1.1		
mean max	23.6	25.3	29.2		
Rainfall (mm)					
annual	1117	752	811		
spring	259	151	206		
summer	28	39	43		
Vegetation					
Tree density (stems ha ⁻¹)	219	131	787		
Basal area $(m^2 ha^{-1})$	24	13	-		
Experimental design (n)	Open (10)	Open (19)	Open (16)		
_ _ · · ·	Q. suber/ Q . canariensis (20)	Q. ilex (19)	P. sylvestris (16)		

^a Values determined in 0-25 cm deep soil samples (Polo 2006)

^b Mean value for regosols in the region (0-15cmGil Torres et al. 2003)

^c Values determined in 0-16 cm deep soil samples (Matías et al, unpublished data)

630 ^d Values determined in 2-14 cm deep soil samples (Alameda et al. 2012/ Alameda et al.,

631 unpublished results).

Table 2. Mean (±SE) values of the measured soil variables across habitat, season and soil depth by

634 site. Letters indicate differences between sites (P < 0.05).

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	Alcornocales	Alcornocales			Sierra Nevada		
$C_{mic} (mg kg^{-1})$	378 ± 18	а	58 ± 4	b	63 ± 3	b	
$N_{mic} (mg kg^{-1})$	46 ± 3	а	17 ± 1	b	6.9 ± 0.5	с	
$P_{mic} (mg kg^{-1})$	7.9 ± 0.6	а	1.0 ± 0.1	b	4.0 ± 0.3	с	
$C_{\text{org}}(\%)$	3.9 ± 0.1	а	1.3 ± 0.1	b	3.0 ± 0.1	с	
P_{inorg} (mg kg ⁻¹)	2.7 ± 0.2	а	1.3 ± 0.1	b	3.4 ± 0.2	с	
NH_4 (mg kg ⁻¹)	8.1 ± 0.6	а	2.6 ± 0.2	b	19 ± 1	с	
$NO_3(mg kg^{-1})$	2.5 ± 0.2	а	0.7 ± 0.1	b	7.9 ± 0.4	с	
DOC (mg kg ⁻¹)	168 ± 7	а	117 ± 8	b	41 ± 3	с	
DON (mg kg ⁻¹)	27 ± 1	а	6.7 ± 0.3	b	3.9 ± 0.2	с	
N _{tot} (%)	0.28 ± 0.01	а	0.09 ± 0.0	b	0.22 ± 0.01	с	
C_{mic}/N_{mic}	9.3 ± 0.3	а	7.2 ± 0.7	b	21 ± 4	с	
C_{mic}/P_{mic}	108 ± 18	а	145 ± 28	а	23 ± 2	b	
Moisture Spring (%)	22 ± 1	а	9.3 ± 0.6	b	13 ± 1	с	
Moisture Summer (%)	11.0 ± 0.4	а	2.7 ± 0.2	b	3.4 ± 0.2	с	

Table 3. Repeated measurements ANOVA for the studied soil properties. F and *P*-values for between effects (forest site, habitat and soil depth), within effect (season) and three way interactions are presented. Significant effects are marked with asterisks (* P < 0.05; ** P < 0.01; *** P < 0.001). C_{mic}, N_{mic}, P_{mic}: microbial C, N and P, respectively; C_{org}: organic C; N_{inorg}: inorganic N (NH₄ + NO₃); P_{inorg}: inorganic P; DOC, DON: dissolved organic C and N, respectively.

	C _{mic}	N _{mic}	P _{mic}	C _{org}	P _{inorg}	N _{inorg}	DOC	DON	C_{mic}/N_{mic}	$C_{mic}\!/P_{mic}$	Moisture
Effect	F P	F P	F P	F P	F F	F P	F P	F P	F P	F P	F P
Site	342***	368***	170.4***	11.2***	75.6***	619***	303.3***	590.2***	65.6***	126.9***	265.4***
Habitat	12.1***	7.79**	14.9***	15.4***	3.83	12.5***	39.2***	16.6***	1.13	5.82*	7.96**
Depth	41.5***	69.2***	63.5***	6.93**	31.1***	18.3***	45.3***	37.6***	1.73	10.9**	0.93
Site×Habitat	9.7***	22.3***	13***	7.55**	6.61**	7.36***	3.15	9.12***	6.79***	18.2***	1.71
Site×Depth	8.1***	3.67*	22.2***	3.96*	3.25	1.87	6.2**	4.76*	0.1	4.86**	8.93***
Habitat×Depth	0.97	2.14	0.01	0.01	0.02	0.07	2.54	3.52	0.19	8.68**	0.91
Site×Habitat×Depth	2.54	1.37	0.47	4.05*	2.67	2.58	3.17	1.35	0.22	7.93***	0.9
Season	8.83**	116.8***	154***	0.14	5.65	0.97	144.9***	11.2***	92.1***	36.4***	1283***
Season×Site	23.3***	27.2***	31.7***	45.8***	105***	193***	21.2***	22.9***	12.3***	0.40	13.6***
Season×Habitat	0.05	4.41*	11.4***	6.42*	0.04	5.11*	10.9***	8.13**	6.27*	7.67**	0.06
Season×Depth	0.04	0.09	0.5	1.54	5.32*	4.27	0.52	0.75	0.89	0.00	13.4***
Season×Site×Habitat	0.18	0.02	2.45	3.36*	7.48***	3.17	8.67***	5.95**	0.7	0.65	1.45
Season×Site×Depth	0.28	1.7	0.24	1.46	2.32	3.79	8.61***	5.98**	0.03	0.03	6.61**
Season×Habitat×Depth	0.08	0.27	3.21	0.84	0.63	0.51	2.97	0.45	0	1.26	0.38

Site: Alcornocales, Cardeña and Sierra Nevada

Habitat: Tree and Open;

Depth: Top soil (0-8cm) and Deeper soil (8-16cm);

Season: Spring and Summer .

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Fig. 1.





С SN

А

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Variance explained (%)

Fig 5.









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