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7	SCALE DEPENDENT EFFECTS OF COPPICING ON THE SPECIES POOL
8	OF LATE-SUCCESSIONAL BEECH FOREST
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32	Running head: Scale dependent effects of coppicing on diversity
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38 39 40 41 42 43 **Abstract** 44 Question: We study the effect of traditional coppicing and its abandonment on community 45 structure and plant diversity at various scales. The aim is to compare active and abandoned beech 46 coppices in terms of: (i) structural features (ii) total, understory and overstory plant diversity across 47 a range of spatial scales (iii) species richness of beech forest specialist species. 48 Location: Monti Sibillini, Central Apennines, Italy. 49 Methods: We applied a multiscale approach working in parallel at fine scale (individual 50 management units, 0.5-1 ha), and at coarser scale (the whole forest patch). Two forests patches 51 were selected according to management regimes (active, 1-40 years, and abandoned coppicing, >40 52 years) in each of the two study areas (Bolognola and Ussita). Individual management units were 53 sampled with 400mq plots; each forest patch was represented by 20 plots arranged in grids. 54 Results: We found significant differences in structural features between coppiced and abandoned 55 forests. At plot scale, only the number of beech forest specialist species was significantly higher in 56 abandoned plots. At the forest patch scale, the Bolognola actively coppied forests hosts many more 57 species than the abandoned ones, but an opposite result was found at Ussita. The abandoned forests 58 show a constant higher beech forest specialist species richness at Ussita, while in Bolognola we 59 found just the opposite. The managed forests host more overstory species than abandoned ones in 60 both areas. Surprisingly the total beech forest specialist species richness was higher in the coppiced 61 plots with respect to the abandoned ones. The species assemblages were more similar between 62 patches having different management regimes within the same area, than between patches having 63 the same management regime across different areas. 64 **Conclusions:** This material provides a novel contribution to the study of species diversity patterns 65 in this forest system, suggesting the importance of a multiple scale approach in forest diversity 66 studies. The beech forest specialist species can largely persist in a heterogeneous coppice landscape, 67 where abandoned stands are mixed with stands under a regular coppicing. The results can link the 68 existing knowledge of beech coppice diversity between landscape- and plot-scale, helping to 69 address new conservation planning.

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Keywords: Scale-dependence; landscape; alpha diversity; beta diversity; Central Apennines; forest

73 stand structure

Introduction

75 In Mediterranean and temperate Europe, coppicing still represents a widespread method of forest resource exploitation (Peterken 1981; Rackham 2008). This management system relies on the 76 77 resprouting capacity of some tree species after coppicing. Some of the trees are not cut during 78 coppicing (the so called "standards") and serve as a seed sources and forest floor protection. These 79 standards are left in the coppice with density dependent planned on different variables. Then, 80 standards are cut on a longer rotation. The rotation period of cutting in Mediterranean coppices is 81 about 15-30 years. Traditionally, coppice systems provide firewood and charcoal but are also 82 grazing areas for livestock (Peterken 1981; Amorini & Fabbio 2001).

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84 In Italy, about 3.9 million ha of forests were managed with coppicing (Ciancio et al. 2006). In the 85 central-Apennines the woodlands dominated by Fagus sylvatica L. were largely coppiced until the 86 last century (Pavari 1999). In the last decades, in parallel with socio-economic changes (e.g. 87 abandonment of the countryside, see Antrop 2004; Agnoletti 2007; Rackham 2008), the coppicing 88 significantly declined in Mediterranean Europe (Barbero et al. 1990; Evans & Barkham 1992; 89 Coppini & Hermanin 2007), and a significant demise of this forest management is also documented 90 for temperate Europe (Van Calster et al. 2007; Hedl et al. 2010). This has made the process of 91 coppicing abandonment a very relevant ecological process in most of southern and Mediterranean 92 Europe.

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94 In the abandonment of coppiced oak woodlands, Debussche et al. (2001) observed a cover decrease 95 in lower height classes of woody stems and an increase in cover and height of the uppermost woody 96 layer, resulting into a simplified vertical structure due to the competition for light. Similarly, a 97 pronounced dominant canopy cover shift was observed from shrubs to trees by Van Calster et al. 98 (2008) in a Fagus sylvatica dominated high forest. The regeneration process after logging of a 99 coppiced forest often causes a decrease of plant diversity at the stand scale (Debussche et al. 2001; 100 Mason & MacDonald 2002; Decocq et al. 2004; Bartha et al. 2008). In a mixed forest of Central 101 Europe (mainly oak-hornbeam forest), Kopecky et al. (2013) demonstrated that the dominant 102 process after abandonment of coppicing was the non-random extinction of light-demanding species, 103 leading to an impoverished, temporally nested plant community structure.

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The consequences of coppice abandonment is in line with a worldwide simplification of forest structure and function (McKinney & Lockwood 1999; Ares et al. 2010), generating a progressive taxonomic impoverishment and homogenization of the woodland vegetation (Keith et al. 2009). In light to these evidences, ecological studies need to explore these issues because plant species assemblages change rapidly under human impact (Hedl et al. 2010; Kopecky et al. 2013).

Although the herbaceous layer contains more than 80% of the total plant species richness, this layer is often an under-considered component of forest ecosystems (Gilliam 2007). However, its effects on tree productivity and long-term stability of the ecosystem have been demonstrated, as well as its influence on the composition, microbiological processes and nutrient cycling of the organic layer (Clement 2001; Gilliam 2007; Salemaa et al. 2008). Herbaceous species also contribute significantly to biodiversity and particularly important for this are those plants which are late successional beech forest species (beech forest "specialist species"). These species usually characterize the forest mature phases and ancient woodlands (Gilliam 2007; Bartha et al. 2008), adapted to the relatively stable environmental conditions of forests (Hermy et al. 1999). The role of forest management in preserving the *specialist species* pool at different spatial scales is still poorly evaluated (Bartha et al. 2008). These species can be used as indicators of biodiversity, and in this respect are important functional components for their habitat and resource specificity (Gillam 2007). Recent studies performed in beech forests of the Italian Apennines (Bartha et al. 2008; Campetella et al. 2011) demonstrated that, at the stand scale, the abundance contribution of the beech forest *specialist species* doubled after about 40–60 years after coppicing.

Species diversity depends considerably on spatial scale (Podani et al. 1993; Palmer & White 1994; Crawley & Harral 2001). Diversity studies usually target coarse (landscapes – several km²) or small (plots - m²) scales (Brockway 1998; Chiarucci et al. 2008a; Bartha et al. 2008) and therefore "in-between" scales are often unexplored. This is a significant gap because management practices are often carried out at intermediate scales and this is likely to have an important effect on species diversity. Moreover, forest fragmentation represents a widespread process (Wulder et al. 2009; Albuquerque & Rueda 2010), and consequently species diversity analysis at the forest patch level is of crucial importance. Although plant biodiversity has become a key element of Central European forest management policy, there is still a limited number of studies examining the influence of coppice abandonment on biodiversity (Kopecky et al. 2013) and very few of them used large dataset (Battles et al. 2001). Information on Mediterranean parts of Europe is largely missing and a study which compares coppice forest units to abandoned ones can contribute to disentangle the ecological processes behind the forest coppice abandonment.

The present study investigates plant species diversity in relation to coppice abandonment, on a range of spatial scales, from the plot to the whole stand. Our overall objective was to compare the structure and diversity of coppiced and abandoned beech forests both at the plot (20 m x 20 m) and the entire forest patch (minimum size of 80 ha) scale in the Italian Apennine, by using a wellbalanced probabilistic sampling design. Our specific research questions were to compare active and abandoned beech coppices in terms of: (i) structural features (ii) total, understory and overstory plant diversity across a range of spatial scales (iii) species richness of beech forest *specialist* species.

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Materials and methods

- 152 Study area
- 153 The study area is located in the central-Apennines, in the Monti Sibillini National Park (southern
- Marche Region, Italy). The area belongs to the bio-climatic 'warm' Fagus forest zone (800-1000 m
- to 1500 m a.s.l., as proposed in the central Appenines by Piovesan et al. (2005), where the mean
- annual precipitation ranges between 1100 and 1400 mm and the mean annual temperature is
- between 8 and 12°C (Biondi & Baldoni 1995). The bedrock is mainly Mesozoic and Tertiary
- 158 limestone. The soils (Skeleti-Calcaric Phaeozems and Calcari-Humic Leptosols) are weakly
- structured, shallow and have low water holding capacity, with medium-high content of carbonates
- 160 (10-40%) loamy texture, and pH ranging from neutral to sub-alkaline (ASSAM 2006). According to
- the phytosociological classification, these beech forests belong to two associations of the alliance
- 162 Geranio striati-Fagion, i.e. the Cardamino kitaibelii-Fagetum sylvaticae and the Lathyro veneti-
- 163 Fagetum sylvaticae.

- 165 Sampling design
- 166 The study was based on a well-balanced sampling design. Several criteria were used to select forest
- patches differing only in current management regime, namely "active coppice" or "abandoned
- 168 coppice", but maintaining all the other ecological factors as constant as possible. To do this, a forest
- management map was used (IPLA Instituto per la Piante da Legno e l'Ambiente 2001). Four
- 170 forest patches were selected in two different areas, with management regimes (active and
- abandoned coppicing) in each area. Each forest patch was located on limestone bedrock on NW-N-
- NE facing slopes, and had a minimum size of 80 ha, considered enough to represent the mean forest
- patch size occurring in the region. The two abandoned coppice patches mostly derived from the
- abandonment of coppices or by the coppice conversion into high forest. One pair of management
- 175 regime ("active" / "abandoned") was in the Bolognola municipality, the other in the Ussita
- 176 municipality.
- 177 To represent the plant species composition and structural attributes at the plot and forest patch
- scales without sampling bias (Chiarucci 2007), a probabilistic sampling design was realized by
- placing 20 m x 20 m plots in the nodes of a grid with cells of 200 m x 200 m, oriented in N-S and E-
- 180 W directions (Figure 1). This distance was chosen to sample the highest level of heterogeneity,
- since the single size of units coppied in a single event is usually between 0.5-2 ha (Canullo et al.

- 182 2009). The grid was virtually overlaid on each of our selected forest patches starting by a random
- point. The south east corner of each plot was positioned at each grid node, after its location by
- 184 Garmin Vista Etrex GPS.
- 185 To minimize edge effects, plots were placed at a minimum distance of 20 m from roads or forest
- edges. Plots falling into non-Fagus facies were placed elsewhere or rejected. Since the resulting
- number of plots in each forest patch was between 20 and 23, they were standardized to 20 by
- 188 removing those plots located in most marginal sites. This resulted into a sample made of 20 plots in
- each of the four forest patches, two for each area and each 2 management regimes (active and
- 190 abandoned).
- 191 In each plot, all vascular plant species were recorded and their cover visually assessed. For
- structural data, in each plot the cover of emergent trees (individuals with crown emerging over the
- dominant tree layer), dominant trees (trees composing the crown dominant layer), subordinate trees
- 194 (trees > 5 m, with crown totally or partially dominated from the upper canopies), shrubs (woody
- individuals < 5 m), herbs, total plant cover, deadwood cover (i.e. coarse woody debris, stumps),
- 196 litter and bare soil were also visually assessed. The first four variables mentioned above are
- 197 considered as vertical physiognomic forest layers. Each individual plant with DBH > 2 cm was
- 198 censused, taxonomically identified and grouped into one physiognomic forest layer. In each plot all
- 199 the individuals released as "standards" were also recorded and their height (we used Vertex
- 200 hypsometer, Haglof Inc., Madison, Miss.) and DBH were measured. Field works were carried out
- 201 from June to August 2008.
- 202 Beech forest specialist species (Bartha et al. 2008) were classified considering their common habitat
- 203 preference, taking into account their similar phytocoenological role (Borhidi 1995; Nagaike et al.
- 204 2003), as determined by regional synecology and their actual role in the local floras (assigned from
- Pignatti 1982, 1998; Pignatti et al. 2004; Ballelli et al. 2005). The complete list of the beech forest
- 206 specialist species identified in this study is provided in the Supplementary material 1.
- 208 Statistical analysis

- 209 The diversity of woody species at the plot scale was quantified by using the Shannon index (H') on
- 210 the basis of the number of individuals with DBH > 2 cm.
- 211 The effects of management and forest patches on the structural variables (number of standards,
- 212 standard diameter, standard height, emergent trees density, dominant trees density, subordinated
- 213 trees density, shrubs density, total woody density, total vegetation cover, herb layer cover,
- 214 deadwood cover) and diversity variables (vascular species richness, overstory species richness,
- 215 understory species richness, beech forest specialist species richness, Shannon index H') was
- analyzed by nested ANOVA statistics (Linear Mixed Models), with management as a fixed factor
- 217 and forest patches nested within the management type. A correction for multiple testing was carried

out to avoid inflating the significance of individual tests ($p \le 0.001$).

219 We addressed the problem of the spatial autocorrelation of error, as this can result in spurious 220 relationships (type I error). For this purpose, we tested for autocorrelation of the models error by 221 using a randomization test. Namely, we plotted the experimental variogram of the models' residuals 222 and produced a 'random envelope' by permuting the coordinates of residual values 999 times while 223 re-computing the variogram at each step. The maximum and minimum value of the semi-variance 224 obtained for each lag provided the 'random envelope'. If the observed variogram values were not 225 within such envelope, this was taken as evidence of autocorrelation and therefore we fitted models 226 with exponentially or spherically correlated spatial error to account for spatial non-independence of 227 the residuals.

- We pooled plots belonging to different forest patches to obtain the diversity measures at the scale of the entire forest. The species richness (total-, understory-, overstory-) was compared by sample (plot) rarefaction curves (Gotelli & Colwell 2001), obtained by using the analytical formula (Kobayashi 1974; Chiarucci et al. 2008b).
- 232 To test the variability in community composition among sites and between management regimes, 233 diversity was multiplicatively partitioned (Baselga 2010) according to the multi-level sampling 234 hierarchy: plot, forest patch, municipality, forest management type. Forest patches were defined on basis of the two management regimes and were tested for difference by comparing variation in 235 236 species composition between patches within the same municipality and, between patches within the 237 same forest management type that belonged to different municipalities. To partition the total 238 diversity of vascular plants, we calculated different measures of forest diversity (Table 1). The beta-239 diversity was calculated according to the Whittaker multiplicative formula, beta = gamma/alpha 240 (Whittaker 1972; Baselga 2010). All the analyses were performed in R environmental (R Core 241 Team 2014), using the "vegan" package (Oksanen et al. 2015).

243 Results

- 244 Forest stand structure at the plot scale
- A total of 5.446 individuals of woody species with DBH > 2 cm were censused. As expected, the mean standards height was significantly higher in the abandoned coppices than in active ones (Table 2), while the density of emergent trees and shrubs were significantly higher in active coppices. As an effect of nested forest areas, standards density, standards diameter and standards height were higher in the abandoned ones (Table 2). Interestingly, total woody individuals density with DBH > 2 cm showed no significant difference among the two management types (Table 2).
- Results for the variables describing the overstory and understory layers evidenced (Table 2) that the two management regimes had similar plant cover (almost 90%, on average). The analysis of the spatial autocorrelation of error shows that all the observed variogram values occurred within the

254 random envelope, supporting the absence of autocorrelation among the different sampling units in

the adopted sampling design (see Supplementary material 2).

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- 257 Diversity at the plot scale
- A total of 291 vascular plant species were found (more than the 10 % of the flora of Marche region,
- see Conti et al 2005) in the 80 plots. The mean species richness of vascular plants, the Shannon
- 260 index of diversity of woody individuals and the species richness of understory plants did not differ
- between the two management types and the two areas. On the other hand, the species richness of
- 262 overstory species resulted statistically different between treatments and areas, being higher in active
- 263 coppices and in the Bolognola area (Table 3). The mean species richness of beech forest specialists
- 264 was significantly higher in the abandoned plots as compared to the forest still actively managed
- 265 (Table 3).

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- 267 Diversity at forest patch scale
- 268 The rarefaction curves for the 20 plots in each forest patch (Figure 2) do not suggest an asymptotic
- accumulation of total species richness. At Bolognola, the actively coppied forest hosts many more
- 270 species than the abandoned one, while an opposite pattern, even if with a lower difference, was
- found at Ussita. The pattern of overstory species was similar in both areas, with the managed forests
- 272 hosting many more species than abandoned ones, while the patterns of understory species showed
- 273 similar difference as the total species. The patterns of beech forest *specialist species* was rather
- 274 complex and divergent between the two areas. At Ussita, abandoned coppices had constantly higher
- species richness of beech specialists at all the sample sizes (from 1 to 20 plots), while at Bolognola
- abandoned coppices had higher species richness of beech specialists only at smaller sample sizes
- 277 (up to 10 plots) and managed coppices had higher species richness at larger sample sizes.

- 279 Diversity partitioning
- 280 Multiplicative partitioning of beta diversity into hierarchical levels showed discordant results in the
- 281 two study areas in term of number of species and species composition. In fact, at the plot scale, the
- 282 number of species was higher in abandoned patch than actively coppied forest in Ussita, but it was
- 283 higher in actively coppied forest than in abandoned forest in Bolognola. On the other hand the
- variability in species composition within each patch was higher within coppied forest in Bolognola
- but little higher within abandoned forest in Ussita (Table 4).
- 286 At patch level, although there were more species in coppied patches than abandoned ones, the
- variability in species composition between patches within the same municipality was slightly lower
- 288 than the variability measured between patches within the same forest management type (Table 5).
- 289 In other words, species assemblage were more similar between patches having different forest

- 290 management regimes within the same area, than between patches having the same forest
- 291 management regime across different areas.
- 292 The same trend was shown by considering differences in beta diversity at the scale of forest
- 293 management type and at the scale of municipality. Beta diversity was slightly higher between the
- 294 two municipalities than between the two forest management types (Table 6).

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Discussion

- 297 Forest structure
- 298 The results of this paper support the hypothesis that management intensity influences forest
- 299 structure as already reported in a number of studies (Nagaike et al. 2003; Decocq et al. 2004;
- 300 Sullivan et al. 2009). The density of emergent trees was found to be higher in actively coppiced
- 301 forests compared to abandoned ones, as it was expected in a coppice management-system. In
- recently coppiced stands, the released "standards" are the only plant physiognomically recognizable
- 303 as trees and they widely contribute to generate this layer. Later, due to a shift of tree layers driven
- 304 by competition for light (Rademacher et al. 2004), the formerly emergent standard trees merge into
- 305 the newly growth dominant layer. The higher density of subordinated trees in the abandoned forests,
- 306 indicates the appearance of a layer formed by those trees that remained dominated during the
- 307 competition for light.
- 308 Other studies indicate that during the process of coppice regeneration, forest structural
- 309 heterogeneity decreases, with upper forest layers becoming dominant (Debussche et al. 2001) and
- 310 shrubs declining (Van Calster et al. 2008). The "loss" of shrubs means a type of simplification
- 311 despite of the higher subordinated tree density. The various regenerative phases of the various
- 312 management units forming forest patches in the Monti Sibillini result in a complex and
- 313 heterogeneous picture of structural characteristics of those forests. The lack of differences between
- actively managed and abandoned coppices in the total density of woody overstory species (DBH
- >2cm), may reflect the fact that abandonment is still relatively recent.
- 316 The persistence of a woody species in coppiced forests is connected to its resprouting ability (Bond
- 317 & Midgley 2001) and this is the biological trait on which coppicing is based in the whole
- 318 Mediterranean forests for centuries (Montes et al. 2004; Pinto-Correia and Vos 2004). As a legacy
- 319 from past land use, these results are probably due to the common origin of the compared forest
- patches (i.e. also the abandoned forest patches included previously active coppiced stands).

- 322 Forest species diversity
- 323 At the plot scale, actively managed and abandoned coppice stands do not significantly differ neither
- 324 in total plants species richness nor in under- and overstory species richness. Our results seem to
- 325 contradict several studies indicating a decrease in richness with maturing coppices at the plot scale

(Debussche et al. 2001; Manson & Macdonald 2002; Decocg et al. 2004; Gondard & Romane 2005; Bartha et al. 2008). A shift from species-rich assemblages of a relatively open and low-nutrient forest to an impoverished plant assemblage of a closed-canopy abandoned forest was also reported by Kopecky et al. (2013). Even in the nearby Siena region, plant richness at the plot scale has been found to decrease in more mature forests, and even if this result was not specifically focused on the abandonment of coppices the basic mechanism was that (Amici et al. 2013). The explanation for this contradiction lies in the small size of coppied management units studied and in the large scale heterogeneity in the successional age of the forests. At the same time, the multi scale probabilistic sampling here performed in four ecologically homogeneous forests, each characterized by an actual precise management orientation, played an important role to understand the effects of scale in the resulting species diversity patterns. In fact, the multi scale approach here adopted offers a totally different view with respect to the observations made at a single spatial scale (e.g., a plot) and a single regenerative forest phase. The small size of the management units (i.e. forest parcels coppied by local owners: 1.28-1.53 ha $\pm 2.03 - 3.46$, Canullo et al. 2009) in the investigated forest patches, makes different regeneration stages co-occurring at the same time, favoring considerable heterogeneity. Consequently, the variability of species richness at plot scale can hide real differences in diversity of the entire forest. This calls for further studies focusing on a range of spatial scales, above the simple plot scale (Podani at al. 1993; Gotelli & Colwell 2001; Chiarucci & Bonini 2005). On the other hand, the results of this study evidenced contrasting results at larger scale with the forest patch generated by still active coppicing in Bolognola being more complex and including diverse set of patches which appears to promote plant species richness and composition due to spatial turnover, even in the case of the beech *specialist species*. Both forest patches located in Ussita resulted less complex (lower beta diversity with respect to the Bolognola ones), with a reduced gamma diversity (50 species less), but with a higher species richness in the abandoned forest patch for all the considered plant groups (total, herb layer and beech forest *specialist species*).

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While it was not expected to find a higher beech forest *specialist species* richness in coppiced patches because of the contribution of species preferring open habitat (i.e. non-forest-species, Bartha et al. 2008), which are favored by coppicing and disturbance, it is somewhat surprising to find a higher species richness of beech forest specialist at larger scales (thus a larger species pool) in the active coppice Bolognola with respect to the abandoned coppice. The difference was really minimal (33 vs. 29) but it suggests the possibility that a rich species pool of beech forest specialists can also be maintained in coppices under active management. This outcome seems to be not in line with other literature outcomes, but it should be noticed that most of the previous results were achieved at a plot or stand scale, without an upscaling as it has done here. For example, Nagaike et al. (2003) and Decocq et al. (2004), found a decrease of forest *specialist species* after silvicultural

362 treatments at the plot scale. Furthermore, Bartha et al. (2008) report a linear positive regression for 363 beech specialist species with coppice age. Our finding on beech forest specialist species could be explained by the fact that when larger scales are considered these actively managed coppices 364 represent very heterogeneous landscapes, due to the contemporary presence of different 365 366 regeneration stages in a rather fine scale patchiness. This can maintain a heterogeneous structure of 367 the forests and consequently a broader spectrum of ecological niches is available for plants, 368 including many of those plants which are linked to mature beech forest stages. In fact, Bartha et al. 369 (2008) reported a remnant of beech forest specialist species also at young stages. Mechanistic 370 insights of a former study on the functional traits of understory forest species in the same area 371 (Campetella et al. 2011) pointed out that beech forest specialist species may meet their habitat 372 requirements already at a successional age of about 14 years after coppicing. Thus, the change of 373 scale perspective leads to the view that despite a reduction of beech forest *specialist species* can be 374 observed at the plot scale, when the focus is on a larger scale this is not valid any more. Practically, 375 the species richness of this group of species is decreased at the alpha diversity level (plot scale) but 376 not at the gamma diversity level (species pool), and this results into a higher beta diversity, i.e. a 377 more heterogeneous distribution in the landscape. In practical terms, these results can be explained 378 by the presence of older coppice stands randomly occurring within the active coppice forest patches 379 in the Bolognola context. There the active forest patch could be considered as a mosaic of 380 contiguous plots in which we can find early cuts with older coppice stands. This is depending on the 381 type of fragmented property, private in most of cases, and to the fact that, during the last decades, 382 due to lack of active management different coppices have been almost abandoned or managed with 383 lower frequency (Amorini & Fabbio 2001; Ciancio et al. 2006; Canullo et al. 2009). This helps to 384 generate a more complex mosaic of forest stands likely to maintain a larger pool of specialist 385 species (with narrow niches). 386 On the other hand, the smaller pool of beech forest specialist species in abandoned patch of 387 Bolognola (accumulation curve reduced the increasing rate at an effort of 14-16 sampled plots) is probably due to the homogeneity of the forest stands, characterized by a more closed-canopy and a 388 389 consequent impoverished flora, more oriented on shade adapted species (i.e. lower availability of 390 light for the understory species, including the beech forest specialist ones; see also Rogers et al. 391 2008; Keith et al. 2009). 392 The higher overstory species richness in the coppied plots is the only convergent result between 393 the two Municipality contexts (areas), which can be explained by a higher availability of woody 394 species niches provided by the coppicing disturbance on the canopy. In fact, also the multiplicative 395 partitioning of beta diversity emphasized that Municipalities (areas) provide a more important effect 396 on the species diversity distribution with respect to the management itself (i.e. beta diversity higher

between the two municipalities than between the two forest management types). This is another

unexpected result (distance between forest stands is a stronger driver than forest management in influencing species composition), probably produced by the significant effects of different forest practices that are implemented by the "local woodcutters" for the maintenance of coppicing. Despite the same management system in both municipalities, the adoption of different criteria in the selection of coppiced parcels (influencing also their spatial distribution) and of standards in terms of number and distribution (as our results demonstrated: 12.65 in Ussita, and 7.35 in Bolognola in 400 m²), can be relevant to support the obtained result in species composition similarity.

405 These results must be inspected more deeply in the future, trying to shed light onto the socio-406 economic and ecological causes of the observed habitat mosaic. The marginalization of forest areas 407 in the Apennines calls for conservation action plans to sustain the considerable biodiversity 408 heritage. Scale, in this respect, is an important factor. We have contributed to show that statements 409 about the effect of forest management on diversity should always define the spatial scale they refer 410 to, as measures considering only plot-level data might result in misleading conclusions. Our results 411 emphasize that analyses should extend toward to the landscape scale in order to better inform and 412 assist conservation management planning.

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- 414 Practical consequences for forest management and ecosystem function
- Despite the local relevance of the study and the limitation in addressing general conclusions at broader geographical scales, we consider the present study as an important contribution in understanding the influence of the management type on species diversity.

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Coppice is an important component of the traditional multifunctional management of forests in the Mediterranean area (Pinto Correia and Vos 2004), but still a fervent debate is active to evaluate the future and the sustainability of such forest management (Ciancio et al. 2006). The importance of coppice management, as a driver of the forest landscape diversification, is reported by different authors (Decoq et al. 2004; Coppini and Hermanin 2007; Kopecky et al. 2013; Müllerová et al. 2015).

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426 Several studies have demonstrated that coppies with standards under a regular felling regime have 427 some negative effects, such as soil erosion, changes in bulk density or soil porosity, changes of pH levels towards soil acidification, depletion of the soil organic matter and nutrient pools (Dyck and 428 429 Skinner 1990; Alban and Perala 1990; Staaf and Olsson 1994; Corona et al. 1996; Worrel and 430 Hampson 1997). In the long term, these changes have been associated with losses of plant species 431 diversity as well (Gilliam and Roberts 1995), so care should be taken in deeming coppicing 'sustainable'. According to our results, however, the coppice with standards in the study area seems 432 433 to be sustainable in terms of the preservation of species diversity on a larger spatial scale. In fact,

(beech forest specialist species) can largely persist in a coppice landscape, if this is managed 435 ensuring a certain heterogeneity with relatively small coppiced stands and a high variability of 436 437 coppice age. 438 The small abandoned patches within the coppice landscape can play a crucial role as propagulum sources and their importance should not be discounted (Gimona et al. 2011). To improve its 439 440 sustainability, coppicing should be planned in a larger scale management, in which the coppice stands are identified as parts of a heterogeneous mosaic, where more developed abandoned stands 441 442 are mixed with others under a regular felling regime. Such a solution would be likely to have environmental benefits (e.g. in terms of ecosystem services) and provide a useful diversification of 443 444 wood production and of forest work and its related professional skill (Coppini and Hermanin 2007), 445 hence enhancing the social sustainability of this management system. This suggestion is also 446 consistent with Kopecky et al. (2013) finding that too large a shift to closed-canopy forest management endangers the diversity of European lowland forests. According to our results, trade-447 offs between open and closed-canopy species might be less than expected at the landscape scale, as, 448 in the studied system, even the forest specialists pool might be maintained in a coppiced landscape 449 450 mosaic. This introduces more interesting reflections about the biodiversity consequences of forest 451 452 management at intermediate spatial scales. For such reasons, our results link the existing knowledge 453 of beech coppice diversity from landscape- and plot-scale and can help to address new management 454 issues and conservation planning on local territorial scale. In the uncertain but certainly variable future, due to, for example, climatic and land use changes (Millar et al. 2007), forest management 455 456 needs to be more resilient (i.e. active adaptive management). In this context, understanding how to 457 promote biodiversity, given its insurance value, is likely to become more important to help sustain 458 desired states of dynamic ecosystem in the face of uncertainty and surprise (e.g Hobbs et al. 2006).

the present study demonstrates that even the group of species more linked to mature forest stages

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Fig. 1. Sampling design applied in the Beech forest patches of the study area near Bolognola and Ussita.

Fig. 2. Species rarefaction curves for total species richness, overstory species richness, understory richness and beech forest specialist richness, in each forest patch and according to the relative municipalities (BOL=Bolognola; USS=Ussita). Vertical lines are confidence intervals. Empty triangles=coppiced forest, black triangles=abandoned forest.

Figures

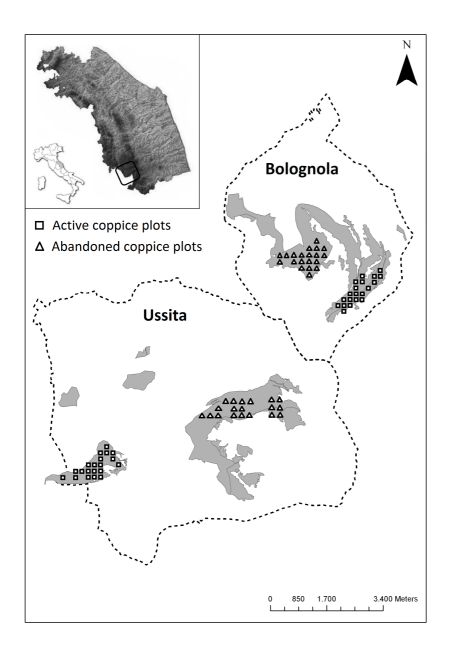
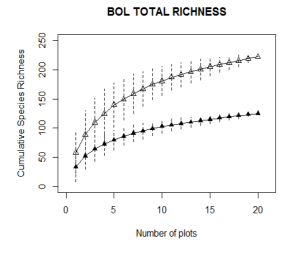
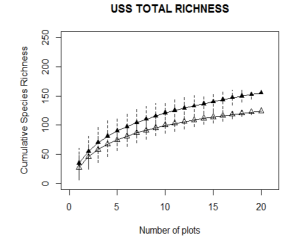
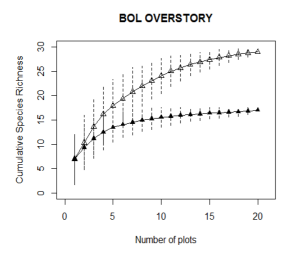
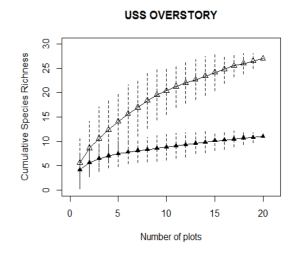


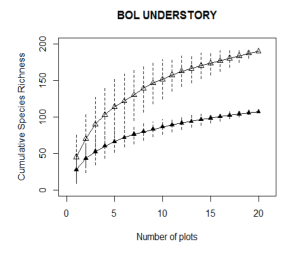
Figure 1 -

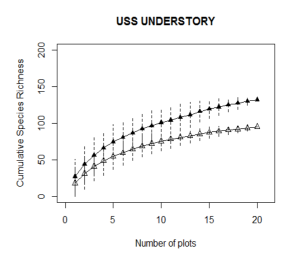


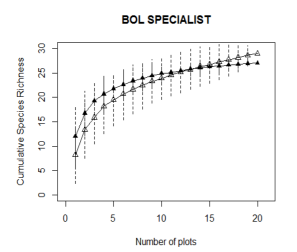


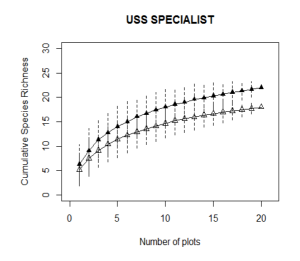












Tables

Table 1 - Partitioning of the total diversity of vascular plants according to the multi-level sampling hierarchy: plot, forest patch, municipality, forest management type.

	Sampling hierarchy	Measurement	Municipalities	Forest management types
	Plot	Mean n. of species per plot	Same	Same
	Patch	Mean n. of species per patch	Same	Different
Alpha	Patch	Mean n. of species per patch	Different	Same
Атриа	Patch	Mean n. of species per patch	Different	Different
	Forest management type	Mean n. of species per management type	Different	Same
	Municipality	Mean n. of species per municipality	Same	Different
	Plot	Beta diversity among the plots	Same	Same
	Patch	Beta diversity among the patches	Same	Different
Beta	Patch	Beta diversity among the patches	Different	Same
Deta	Patch	Beta diversity among the patches	Different	Different
	Forest management type	Beta diversity between forest management types	Different	Same
	Municipality	Beta diversity between municipalities	Same	Different
	Plot	Cumulative n. of species recorded by the plots	Same	Same
	Patch	Cumulative n. of species recorded by the patches	Same	Different
Gamma	Patch	Cumulative n. of species recorded by the patches	Different	Same
Gamma	Patch	Cumulative n. of species recorded by the patches	Different	Different
	Forest management type	Cumulative n. of species recorded by the forest management types	Different	Same
	Municipality	Cumulative n. of species recorded by the municipalities	Same	Different

Table 2 - Mean values of selected variables characterizing released standards, woody individuals (DBH>5cm) and stand layers in the forest patches according to management type (plot scale 20 m x 20 m).

Management	Active Coppice		Abandoned			
Forest patch (Municipalities)	Bol (n=20)	Uss (n=20)	Bol (n=20)	Uss (n=20)	Sign. of Management	Sign. of Area
Mean standards number	7.35ab (±4.17)	11.15abc (±6.27)	6.47a (±5.13)	12.65c (±6.76)	0.759	0.000
Mean standards diameter	20.09a (±6.68)	23.63c (±11.21)	29.36b (±10.59)	20.60a (±8.09)	0.078	0.000
Mean standards height	12.41a (±3.07)	12.55a (±3.60)	20.06b (±4.59)	11.58c (±3.90)	0.000	0.000
Number of emergent trees	4.80a (±3.60)	4.20a (±6.04)	1.95b (±2.28)	1.25b (±1.20)	0.001	0.451
Number of dominant trees	41.20a (±33.20)	26.25a (±14.42)	29.00a (±10.36)	36.6a (±24.86)	0.859	0.987
Number of subordinate trees	20.15a (±22.58)	24.25ab (±18.39)	26.95a (±15.44)	23.55ab (±20.34)	0.482	0.483
Number of shrubs	14.80a (±21.96)	8.65ab (±10.03)	1.60b (±2.72)	2.55b (±2.96)	0.000	0.627
Total woody individuals	80.95a (±52.06)	60.41a (±31.69)	59.51a (±21.94)	63.95a (±38.86)	0.223	0.747
Total vegetation cover (%)	91.35a (±6.53)	88.20a (±5.40)	87.55a (±5.85)	85.5a (±10.08)	0.046	0.158
Herb layer cover (%)	22.20a (±17.60)	6.24b (±11.55)	12.86ab (±11.49)	15.48ab (±14.81)	0.986	0.306
Deadwood cover (%)	2.95a (±2.19)	2.97a (±1.62)	8.25b (±5.40)	3.25a (±2.22)	0.000	0.000

Significant differences (p<0.001) generated both by the management and the nested forest patches were assessed according to ANOVA statistics (Linear Mixed Models). Significant differences between single forest patches are indicated by letters (post hoc Bonferroni). Standard deviation values are reported in parenthesis. (Bol=Bolognola Municipality; Uss=Ussita Municipality; n=number of relevees)

Table 3 - Mean values of species richness and Shannon index (H') characterizing the forest patches according to management type (plot scale $20 \text{ m} \times 20 \text{ m}$).

Management	Active Coppice		Abandoned			
Forest patch (Municipalities)	Bol (n=20)	Uss (n=20)	Bol (n=20)	Uss (n=20)	Sign. of Management	Sign. of Area
Mean vascular species n.	55.05a (±17.08)	28.35b (±10.66)	34.95b (±12.60)	35.30b (±13.00)	0.068	0.023
Mean vascular overstory species n.	7.15a (±2.79)	5.70ab (±2.77)	5.85ab (±2.41)	3.90b (±1.70)	0.006	0.002
Mean vascular understory species n.	47.90a (±17.37)	22.65b (±10.22)	29.10b (±11.13)	31.35b (±12.73)	0.151	0.163
Mean specialist species n.	8.80a (±3.12)	5.15c (±1.69)	12.40b (±3.29)	6.25c (±2.14)	0.000	0.000
Shannon (H') woody individuals	1.09a (±0.45)	0.96a (±0.55)	0.93a (±0.54)	0.74a (±0.53)	0.109	0.216

Significant differences (p<0.001) generated both by the management and the nested forest patches were assessed according to ANOVA statistics (Linear Mixed Models). Significant differences between single forest patches are indicated by letters (post hoc Bonferroni). Standard deviation values are reported in parenthesis. (Bol=Bolognola Municipality; Uss=Ussita Municipality; n=number of relevees)

Table 4 - diversity partitioning between plots within each forest patch.

Municipality	Forest patch management type	Gamma	Mean alpha plot	Beta plot
USSITA (USS)	abandoned	155	35.25	4.40
	coppiced	124	28.35	4.37
BOLOGNOLA (BOL)	abandoned	125	34.95	3.58
	coppiced	222	55.05	4.03

Table 5 - diversity partitioning between patches within the same municipality (a), the same forest patch management type (b), between different municipalities and forest management types (c).

124.50

188.50

1.39

1.41

(a)			
PATCH (within the same municipality)	Gamma	Mean alpha patch	Beta patches
USS-coppiced/USS-abandoned	192	139.50	1.38
BOL-coppiced/BOL-abandoned	239	173.50	1.38
(b)			
PATCH (within the same forest management type)	Gamma	Mean alpha patch	Beta patches
BOL-coppiced/USS-coppiced	246	173	1.42
BOL-abandoned/USS-abandoned	196	140	1.40
(c)			
PATCH (between different municipalities and forest	Gamma	Mean alpha patch	Beta patches

BOL-abandoned/USS-coppiced

USS-abandoned/BOL-coppiced

Table 6 - diversity partitioning between forest management types (a) and between municipalities (b).

(a)

Total Richness	Mean alpha forest management type	Beta forest management type
291	221	1.32
(b)		
Total Richness	Mean alpha municipality	Beta municipality
291	215.5	1.35