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## Frequent coppicing deteriorates the conservation status of black alder forests in the Po plain (northern Italy)

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(Article begins on next page)



# UNIVERSITÀ DEGLI STUDI DI TORINO

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1	Frequent coppicing deteriorates the conservation status of black alder forests in the Po
2	plain (northern Italy)
3	
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14	Highlights
15	• Alluvial forests with black alder are a priority conservation habitat in Europe
16	• We assess whether frequent coppicing is compatible with their conservation
17	• We measured the effect of time since coppicing on forest structure and diversity
18	• Coppicing simplified vertical stand structure and the herbaceous layer for 20-30 years
19	• Non-native species were significantly more abundant in recently coppiced areas.
20	

<sup>&</sup>lt;sup>1</sup> GV wrote the paper and carried out statistical analyses, FM and GC carried out field sampling and forest structure analyses, MFe designed and coordinated the research and carried out imagery analyses, MFr carried out soil sampling and analyses, RM provided input for study design, interpretation and discussions, and ML carried out phytosociological analyses.

### 21 Abstract

22 Alluvial forests with black alder are a priority conservation habitat in Europe. In the Po plain, 23 black alder is traditionally managed by coppicing with frequent rotations. This study aims to 24 ascertain whether such management is compatible with habitat conservation, by measuring 25 the effect of time since coppicing on forest structure and plant species composition across 26 different layers. 27 We compared the effects of three treatments, each thrice replicated: recent (10-20 years), 28 medium (20-30 years) and old coppice (>40 years). In all nine stands we measured basal area, 29 tree and regeneration density, mean tree diameter and height, dominance by alder, species 30 richness, Shannon diversity, and the number of ruderal and non-native species. Significant 31 differences in dendrometric variables, species richness, diversity, and percent cover by 32 chorotype were assessed for treatment effects by two-way ANOVA. 33 Frequently coppiced stands had a lower basal area, mean tree size, and volume, a more 34 simplified vertical structure, a lower cover of the herbaceous layer and higher bare soil cover 35 due to harvesting disturbance, a significantly lower cover by typical woodland *Fraxinetalia* 36 species, and a significantly higher frequency and cover of non-native species. 37 Our study showed that frequent coppicing worsened the conservation status of black alder 38 forests in the study area, simplified stand structure, deteriorated species composition, and 39 increased the spread of non-native and ruderal plant species. Such negative effects persisted 40 even 20-30 years after cutting. We recommend amending the current legislation and 41 introducing mandatory Implications Assessment procedures everywhere alder forests are 42 susceptible to be impacted in a similarly negative way. 43

Keywords: coppice, floodplain forests, forest management, Habitat Directive, non-native
species, plant diversity, understory

47

### 1. Introduction

48

Black alder (*Alnus glutinosa* (L.) Gaertn.) is a tree species of riparian and water-logged
habitats that is naturally widespread from mid-Scandinavia to southern Europe (Kajba et al.
2003). It forms pure stands on periodically submerged sites, while it mixes with ash
(*Fraxinus excelsior* L.), maples (*Acer pseudoplatanus* L. and *Acer platanoides* L.) and oaks
(mostly *Quercus robur* L.) on riverside and plateau sites (Dethioux 1974), where its
intolerance to shading and lower groundwater tables reduce its ability to compete (Claessens
et al. 2010).

56

57 Black alder grows between sea level and 1,300 on the Alps (Shaw et al. 2014). It is largely 58 indifferent to soil parent material, but it requires precipitation above 510 mm per year and 59 high water saturation (McVean 1953), and a high degree of atmospheric humidity throughout 60 its reproductive cycle. When the water-table sinks below the surface during summer, tree 61 growth increases but seedlings may suffer from drought (McVean 1953). The tree is able to 62 fix atmospheric nitrogen in symbiotic root nodules (Bond et al. 1954), and its litter increases 63 nitrogen and phosphorous content of the soil (Moiroud 1991, Giardina et al. 1995). The 64 species has a maximum lifespan of 100 to 160 years (Claessens et al. 2010). It reaches sexual 65 maturity at age 3-30, when it starts producing seeds with mast pulses every 3-4 years 66 (Dethioux 1974). Seeds are dispersed by water or wind (up to 150 m: McVean 1955, but 67 usually within 30 m: Funk 1990). However, regeneration occurs mostly from vegetative reproduction, e.g. in linear flood populations (Koop 1987, Deiller et al. 2003). Regeneration 68 69 from seed is usually scattered and it occurs under favorable establishment conditions, e.g., on 70 low-lying alluvial land or on former meadows (Douda et al. 2009). Seedlings require a higher

light intensity than those of larger-seeded trees (McVean 1956); it was found that natural
regeneration of black alder is not possible under the canopy of a mature stand (Tapper 1993),
except in openings larger than 0.1 ha (Claessens et al. 2010). The regeneration of black alder
also depends on the frequency and intensity of disturbance (e.g. browsers, floods, or forest
harvesting) (Pokorný et al. 2000; Wolf et al. 2004), and on the abundance of herbs that may
compete with the seedlings (McVean 1956).

77

78 Due to their specific hydrological regime and rare occurrence, black alder forests and carrs 79 are considered an endangered forest community in Europe (Ellenberg 1996). Alluvial forests 80 with black alder and ash are a priority habitat of Community interest listed in the Annex I of 81 the Habitats Directive 92/43/EEC as 91E0\* – Alluvial forests with Alnus glutinosa and 82 Fraxinus excelsior (Alno-Padion, Alnion incanae, Salicion albae). These forests are highly 83 important for the conservation of a great number of typical woodland and floodplain plant 84 species (Claessens 2003), particularly when interspersed in an agricultural matrix. Despite 85 being often small and fragmented (Schnitzler 1994), black alder forests are often 86 characterized by a high richness in herbaceous species (Brown et al. 1997). Beyond plant 87 diversity, black alder forests support other ecosystem services as well, such as water filtration and purification in waterlogged soils (Peterjohn and Correll 1984), flood control and 88 89 riverbank stabilization (Piégay et al. 2003).

90

91 Currently, these forests represent less than 1% of the forest cover in most European countries 92 (Claessens et al. 2010) due to both land use changes such as conversion to non-native tree 93 plantations or agricultural land, or to environmental changes related to human activities, e.g. 94 land draining, impact of industrial areas, negative selection in favor of more valuable timber 95 species such as oak and ash, and the introduction of non-native species (EEA 2012). For

96 these reasons, the conservation status of 91E0\* habitat is currently "unfavourable inadequate"
97 or "unfavourable bad" (Kremer et al. 2015).

99 In the Po plain, black alder is traditionally managed by coppicing, with rotations of 10 to 30 100 years due to the fact that the potential for vegetative regeneration from stumps declines at 60-101 80 years of age (Kapustinskaite 1960). Private ownership usually prevails in floodplain 102 forests, with the consequence of creating a mosaic of small but intense and frequent cuttings, 103 which can deteriorate habitat conservation and spatial continuity. Moreover, floodplain 104 forests are highly vulnerable to plant invasions due to the frequent and intense natural 105 disturbances, to their linear nature which facilitates long-distance species dispersal, and to 106 intensive human pressure (Richardson et al. 2007). In this perspective, the question arises 107 whether such management is compatible with habitat conservation. 108 109 This study aims to ascertain the effect of time since coppicing on the conservation status of 110 black alder stands, as measured by (a) forest composition, structure, and biomass, and (b) 111 species composition and naturalness of the herbaceous layer, e.g., the relative frequency of 112 ruderal and non-native herbaceous species. 113 114 2. Study area 115 116 The study was conducted in the Natura 2000 site "IT1110021 - Laghi di Ivrea" (Figure 1), a 1600-ha Site of Community Importance (SCI) at the center of the 500-km<sup>2</sup> Ivrea Morainic 117 118 Amphitheater (IMA). Mean annual temperature and annual precipitation are 12.5°C and 1002 119 mm, respectively (years 1921-2000) (Andreone et al. 2001). The bedrock is a juxtapositions 120 of three metamorphic units (eclogitic micaschists, basic granulites and vulcanites) as a result

121 of uplift and underplating during the Tertiary Alpine orogenesis (Johnson 1973). The 122 Morainic Amphitheater was constructed between the Pleistocene and the Last Glacial 123 Maximum (Carraro et al. 1974). Thereafter, small lakes formed in the gaps between 124 secondary moraines, but most later evolved into peat bogs or were artificially drained. Such 125 low-elevation sites are characterized today by Endoaquepts or Haplosaprists soils (Piazzi et 126 al. 2007). The latter is predominant in peat and raised bogs, where the sapric organic material 127 has an extremely slow hydraulic conductivity and C/N ratios may be as high as 45. 128 129 The site hosts 11 habitat types of the EU Habitats Directive (1992/43/EEC Annex I), among 130 which the priority habitat 91E0\* covers 59 hectares. A total of 32 plant and animal species of 131 the EU Nature Directives (1992/43/EEC and 2009/147/EC Annex II) (Natura 2000 Network 132 Viewer 2016). Anthropic pressure has caused the number of plant species to decline from 179 133 to 160 species between 1950 and 2005; at least 12 non-native plant species were reported in 134 the area so far (Minuzzo et al. 2005; Lonati et al. 2014). 135 136 Forests are mostly owned by small private owners. Between January 2012 and June 2015, 40 silvicultural treatments were authorized across 8 hectares of 91E0\* forests inside the site; 137 10% of this area was treated by thinning, 30% by coppicing, and 60% by contemporary 138 139 cutting of the coppice and high forest layers (Regione Piemonte 2016a). 140 141 3. Methods 142 143 We designed the study as a chronosequence of stands coppiced in three different times: recent 144 (10-20 years, TR1), medium (20-30 years, TR2) and old coppicing (>40 years, TR0). To do 145 so, we preliminarily assigned one of such treatments to all forest stands classified as 91E0\*

146 habitats (according to Andreone et al. 2001) within the study area, based on the analysis of 147 repeated aerial images (years 1954, 1975, 1979, 1994-1996, 1998-1999, 2007, 2009). The 148 images were orthorectified and georeferenced, then visually classified into forested / 149 nonforested categories, and differentiated to obtain age ranges for each forest stand. Age 150 classes were subsequently confirmed by field surveys and exploratory increment core 151 sampling. Only stands belonging to the association Carici remotae-Fraxinetum Koch ex 152 Faber 1926 (alliance Alnion incanae Pawłowski in Pawłowski and Wallisch 1928) and 153 already existing in year 1954 were considered for further analysis, i.e., waterlogged stands of 154 the alliance Alnion glutinosae Malcuit 1929 and secondary stands on former non-forested 155 land were filtered out.

156

Following superposition to cadastrial stand maps, we identified three independent study areas where all three elements of the chronosequence could be found in stands less than 100 m apart from one another, in order to minimize site differences between treatments and counter pseudoreplication. The only three areas where this condition was met in all the SCI are indicated in Figure 1. A total of nine stands (i.e., 3 study areas x 3 treatments) were selected for analysis; stands were at a constant elevation (about 240 m a.s.l.) and had a mean size of  $1120 \text{ m}^2$ .

164

In spring 2015, in each stand we randomly established a circular sampling plot (radius = 10
m) where we recorded species, frequency, diameter at breast heigh (dbh), origin (seed or
sucker) and height of all adult trees with dbh >=7.5 cm. We also recorded species, frequency,
origin, and height of all juvenile trees (dbh <7.5 cm) in a concentric 6-m radius circular plot.</li>
From plot data we computed common descriptors of stand structure (species composition,
number of trees per hectare, basal area, quadratic mean diameter, average and top height,

percent trees originated from seed) and compared them across treatments by Mann-Whitneytest.

173

174 Within each sampling plot we randomly established five understory subplots (radius = 2 m) 175 where we visually assessed percent cover of upper tree (height >15 m), lower tree (height 176 between 5 and 15 m), upper shrub (height between 1.3 and 5 m), lower shrub, herbaceous and 177 bare soil layers, and assigned cover-abundance scores (Braun-Blanquet 1932) to all plant 178 species by each layer. At the center of each understory subplot, we measured canopy cover 179 by taking a hemispherical photograph at 1m height above the ground. Hemispherical 180 photographs were shot in Nikon .NEF format at 400 ISO with a 6 Megapixel Nikon D70S 181 equipped with a Samyang 8mm f/3.5 aspherical IF MC Fisheye Lens set at shutter priority 182 (time = 1/500 s). Canopy cover was determined by averaging the ratio of white to total image 183 pixels obtained by each of four global thresholding algorithms (Otsu 1979; Huang and Wang 184 1995; Yen et al. 1995; Li and Tam 1998) for the Fiji image analysis software (Schindelin et 185 al. 2012), applied to the blue band of each photograph. 186 187 In order to check for the absence of significant edaphic differences, three topsoil samples 188 were extracted from the center of each understory subplot at a depth of 0-10 cm. All samples 189 were air-dried and sieved (< 2 mm). Total carbon (corresponding to total organic carbon, 190 TOC, thanks to the absence of carbonates) and nitrogen (TN) were analyzed by dry 191 combustion with a CN elemental analyzer (CE Instruments NA2100, Rodano, Italy).

192

193 For each understory subplot we computed species richness (total number of species), the

194 Shannon diversity index, and the number of non-native species based on chorotype

195 (according to Celesti-Grapow et al. 2009). We associated a phytosociological optimum

196 (according to Aeschimann et al. 2004) to each vascular plant species at the class level, 197 including all subordinated syntaxa (Lonati et al. 2013; Orlandi et al. 2016; Pittarello et al. 198 2016) (Online Resource 1), and computed the number of species belonging to the following 199 phytosociological groups, listed in order of increasing conservation value: ruderals (classes 200 Artemisietea vulgaris and Bidentetetea tripartitae), tall herbs (classes Filpendulo-201 Convolvuletea and Molinio-Arrhenatheretea), shrubs (classes Crataego-Prunetea, 202 Franguletea and Salicetea purpureae), and European alder forests (order Fraxinetalia). After converting cover-abundance data to mid-percent values ('+' in 0.3%; '1' in 2.8%; '2a' in 203 10.0%; '2b' in 20.5%; '3' in 38.0%; '4' in 63.0%; '5' in 88.0%) (Tasser and Tappeiner 204 205 2005), we computed total percent cover of non-native species and of each phytosociological 206 group for each subplot. 207 208 All variables were log- (for continuous data) or arcsin- transformed (for percent data) to 209 ensure homoskedasticity and normality of error distributions, and assessed for differences as 210 a function of treatment by 2-way ANOVA with study area as a random factor and using 211 Tukey's Honest Significant Difference (HSD) post-hoc test. All analyses were carried out in 212 the SPSS 20.0 statistical environment (IBM Corporation 2011). 213 214 4. Results 215 216 All stands exhibited high tree densities (1000 to 2300 trees per hectare), with a large 217 variability within treatments but a thinning trend as time since coppicing increased (Figure 2). 218 Conversely, basal area, mean diameter, and volume increase with time since coppicing, up to

219 yields of 450 m<sup>3</sup> ha<sup>-1</sup> in stands harvested >40 years before sampling. Seedling density was

220 highly variable, between 260 and 7000 per hectare, mostly originated from seed and

predominantly by ash (64%), but decreased with increasing stand age (Figure 3). Canopy
cover declined with increasing stand age (83%, 78%, and 74% respectively in TR1, TR2 and
TR0, p <0.01 with mixed-model ANOVA), possibly due to a structural change from a dense</li>
coppice to a high-forest with larger but sparser trees.

225

226 In all treatments, alder occupied preferentially the dominant vegetation layer, while ash was 227 found in all layers. However, frequent coppicing simplified vertical stand structure. Relative 228 to TR1 and TR2, old coppices showed a differentiation in two distinct tree layers, the upper 229 dominated by alder, and the lower by ash (Table 1). In both medium and recently coppiced 230 stands, dominant trees were still competing with each other within the same vegetation layer 231 (<15 m height), even after 20-30 years. Moreover, recently coppiced stands (TR1) had a 232 lower herbaceous and higher bare soil cover, likely due to the use of machines to transport 233 harvested woods in the stand (as tracks observed in the field clearly showed).

234

Soils were rich in organic carbon and showed presence of gley. TOC%, TN%, and C/N ratio
varied in the range of 4.5 to 19.3, 0.36 to 1.59, and 11.4 to 14.7, respectively, but they did not
differ significantly between treatments (Table 2). Therefore, we ruled out topographic or
edaphic effects in determining understory species composition.

239

Recently coppiced stands (TR1) had a higher total species richness, and TR1 and TR2 a
higher Shannon diversity, than undisturbed stands (Table 3). However, this did not result in
an increased naturalness of plant species composition. In fact, the number of *Fraxinetalia*species (*Carex remota, Carex brizoides, Carex pendula, Impatiens noli-tangere, Ribes rubrum, Equisetum telmateja*) remained substantially low (3 species per plot) and unchanged

along the chronosequence, while their cover decreased significantly from 80% in TR0, to65% in TR2, and 49% in TR1.

247

248 Conversely, the frequency and cover of non-native species were significantly higher in both 249 medium and recently coppiced stands than in TR0 (Table 3). All 10 non-native species 250 sampled in the study areas (Acer negundo, Acer palmatum, Juglans regia, Ligustrum sinensis, 251 Paulownia tomentosa, Robinia pseudoacacia, Lonicera japonica, Parthenocissus 252 quinquefolia, Duchesnea indica, Oxalis fontana) were found only in the two latter stages of 253 the chronosequence (Appendix 1). These areas also showed a significantly higher number or 254 ruderal nitrophilous species (Urtica dioica, Alliaria petiolata, Geum urbanum, Geranium 255 robertianum, Oplismenus undulatifolius, Galeopsis pubescens, Polygonum hydropiper), 256 which lower the biodiversity value of alder stands and whose cover was relatively high (10%) 257 even 20-30 years after treatment. Also the number and cover of species more typically 258 occurring in mown and fertilized meadows or in hydrophylous herb communities (Poa 259 trivialis, Rubus caesius, Humulus lupulus, Filipendula ulmaria, Lythrum salicaria), rather 260 than in alder forests, were higher in recently and medium vs. old coppices (36% and 41% in 261 TR1 and TR2, 5% in TR0). The same was true also for shrub species (Viburnum opulus, 262 Cornus sanguinea, Corylus avellana, Crataegus monogyna, Euonymus europaeus, Ligustrum 263 vulgare, Frangula alnus, Salix cinerea), as they can take advantage from higher light levels 264 in recently opened gaps. Shrub cover was still high (41%) after 20-30 years from coppicing, 265 showing the magnitude and long lasting legacy of silvicultural treatments on stand structure 266 and light conditions.

267

268 **5.** Discussion

### 270 5.1 Forest structure and dynamics

The yield of black alder in Europe at age 80 is between 500 and 1000 m<sup>3</sup> ha<sup>-1</sup> (Lockow 1995; Sopp 1974). Old coppice stands included in this study approached the lower end of this range. The average yield of black alder in the forest district where the study was carried out was 148 m<sup>3</sup> ha<sup>-1</sup> (Bertani et al. 2003), which is indicative of the relative rarity of undisturbed mature alder forests in the area. Contrary to mountain areas, management of coppices in lowland sites of the Po plain is still quite active, due to their high accessibility and the predominantly private ownership.

278

279 In our study, seedling density decreased with increasing stand age. This confirms that alder 280 regeneration by seed is difficult in closed stands, as seedlings are very intolerant of shading 281 and rank herbaceous vegetation (Latham and Blackstock 1998). Herbaceous competitors (e.g. 282 Carex spp.) were more abundant soon after coppicing (see below) and may have prevented 283 alder seedling development (McVean 1956). Should alder seedlings take advantage of the 284 higher light levels from canopy opening, establishment would decline soon after coppicing 285 (Ash and Barkham 1976). At the same time, intense coppicing with few or no standards 286 favored ash regeneration, both from seed and from faster sprout growth. Ash seedlings and 287 sprouts may grow very slowly if in shade for several years, but are able to respond with rapid 288 growth when gaps appear in the canopy (Merton 1970; Tapper 1992, 1993; Gatsuk et al. 289 1980). Once in the canopy, ash is able to overgrow and suppress alder through competition 290 for light (Tapper 1996).

291

292 Finally, biomass, mean tree size, and complexity of vertical structure (i.e., number of tree
293 layers) declined with increasing coppicing frequency. These effects persisted up to 30 years

after treatment, and may result in a loss of habitats and lower spatial heterogeneity of
resources (Motta et al. 2015) that can hamper diversity and ecosystem stability. A more
diverse array of tree sizes and the existence of multiple vertical layers instead may provide a
greater number of potential ecological niches for a wide array of relevant forest biota, such as
invertabrates, birds, lichens (e.g. Müller et al. 2005; Czeszczewik and Walankiewicz 2006;
Nascimbene et al. 2013; Negro et al. 2015).

300

### 301 5.2 Effects of coppice management on habitat conservation

Many herb species typically occurring in woodland communities are perennials and can persist throughout the coppice cycle. Under a coppice regime, the relatively constant and saturated set of niches that is found in high forests is replaced by a variety of others, which are filled by species capable of survival in a relatively wider range of ecological circumstances (van der Werf 1991). Sometimes, the greatest threat to plant diversity is the abandonment of coppicing, whereby many open-habitat species are slowly outshadowed under the ever denser growing crowns (Baeten et al. 2009; Negro et al. 2015).

309

310 However, when the conservation target is the forest habitat, as in the case of black alder 311 remnants, too frequent coppicing can be detrimental. The first agent of damage is harvesting 312 machinery, which may induce compaction, rutting and stripping, irreversibly reduce soil 313 fertility and drainage, and alter species composition, especially on loamy soils (Decocq et al. 314 2005). Altered drainage may then produce habitat deterioration for other species of 315 conservation interest such as aquatic invertebrates, insects and amphibians (e.g., Pelobates 316 fuscus insubricus: Andreone et al. 2004). Second, frequent coppicing alters light and moisture 317 regimes, which facilitates ruderal, nitrophilous, tall herbs and non-native species (Funk et al. 2008) that can outcompete woodland species of interest. The post-coppicing communities 318

319 analyzed by this study contained species adapted to many types of canopy and soil 320 disturbance and relatively high light levels such as ruderal species, nitrophilous species (e.g., 321 Poa trivialis, Urtica dioica, Rubus caesius) (Honnay et al. 1999; De Keersmaeker et al. 322 2004), tall herbs, and shrubs. In undisturbed stands, germination of these species is prevented 323 by low light levels and the abundant leaf litter (Sydes and Grime 1981). But if this vegetation 324 component increases due to the high coppicing frequency, it may progressively hamper 325 specialist woodland herbs by competitive exclusion (Hipps et al. 2005). Canopy removal 326 during coppicing may increase the amount of solar radiation reaching the soil, the quantity 327 and composition of light, the temperature, humidity, evaporation and mineralization rates. 328 Seeds of many open-habitat species require a high temperature regime for germination and it 329 is apparent that this occurs only when there is no vegetation present to cast shade, such as in 330 the first two years after coppicing. Moreover, the decrease of C and N in recently coppiced 331 stands might indicate an effect of coppicing frequency on organic matter recycling and 332 mineralization. Soils of undisturbed forests have been previously found to be richer in C and 333 N than in intensely harvested ones (Johnson and Curtis 2001; Finér et al. 2003), including in 334 alder carr (Honnay et al. 1999; Verheyen et al. 1999; Dzwonko 2001; Falkengren-Grerup et 335 al. 2006; Orczewska 2009). This may result from a more prolonged accumulation of litter 336 that, in the case of alder, has a high N content and rate of decomposition (Karkanis 1975; 337 Pereira et al. 1998). However, also the opposite was found, i.e., a higher N availability in 338 regularly harvested stands, due to a higher organic matter turnover rate (Covington 1981; 339 Kimmins 1987; Koerner et al. 1997; Keersmaeker et al. 2004). Canopy removal increases soil 340 temperature (Carlson and Groot 1997; Pennock and Kessel 1997; Hashimoto and Suzuki 341 2004), decreases soil water content (Ma et al. 2013) and shortens saturation periods, therefore 342 promoting a faster mineralization in harvested stands. Such different findings may depend on

regional climate (Yin et al. 1989), soil fertility, former agricultural use (Compton and Boone
2000), its duration, and time since abandonment.

345

The picture is further complicated by invasion of non-native species. Changes in the 346 347 hydrologic cycle of humid forests, induced e.g. by upstream water use, canopy cover 348 changes, or climate change, can further facilitate invasion by species adapted to drier 349 conditions (Huston 2004). Moreover, floodplain forests remnants can be more vulnerable to 350 plant invasions due to their small size and high perimeter-to-area ratio, which facilitates 351 species colonization from the margins and makes any canopy disturbance a potential threat to 352 native species. Among the non-native species found by this study in alder stands, some have 353 a well-documented high degree of invasiveness, e.g., Acer negundo L., Robinia pseudoacacia 354 L., Lonicera japonica Thunb. (Regione Piemonte 2015a), while for some others this is less 355 certain. However, also the latter ones have been previously reported in other areas of 356 Piedmont region, indicating that their naturalization potential and invasiveness are likely 357 higher than expected, e.g., Ligustrum sinense (Lonati et al. 2014; Soldano et al. 2015) or 358 Paulownia tomentosa (Selvaggi 2014). Once established, non-native species can persist due 359 to their high regeneration potential (also by vegetative reproduction), high degree of adaptation to disturbances, allelopathy, and influx of non-native seeds in the soil seed bank 360 361 (Lorenzo et al. 2013; Gioria and Pyšek 2015).

362

For all these reasons, we recommend amending the current legislation and introducing
mandatory Implications Assessment procedures everywhere alder forests are susceptible to be
impacted by coppicing in the negative ways suggested by our analyses. Should coppice need
to be maintained, best silvicultural practices should include higher retention of living and
dead biomass, longer rotations (Della Rocca et al. 2014), and cessation of all drainage

activities, as a high water level inhibits the vigorous growth of expansive, nutrient-demanding
species and reduces the competitive exclusion of woodland flora by such herbs. Promoting
shadier conditions in the forest floor may also limit the expansion of ruderal and non-native
plant species, and facilitate the immigration and establishment of typical woodland herbs
(Orczewska 2009).

373

374

### 6. Conclusion

375

376 Many species and habitats associated with natural forested floodplains have disappeared from 377 most of Europe. Current silvicultural practices, especially in small private woodlots, may 378 increase pressures on biodiversity and ecosystem conservation. Our study showed that 379 frequent coppicing has negative consequences for the conservation of black alder forests in 380 the western part of the Po Plain (northern Italy), inducing a simplification of stand structure, a 381 deterioration of species composition, and the spread of non-native plant species. Such 382 negative effects persisted even 20-30 years after cutting, suggesting that the cumulative 383 effects of repeated frequent coppicing would be characterized by even harsher impacts.

384

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392	(https://www.change.org/p/salviamo-la-ricerca-italiana) for the increase of research funding
393	in Italy to the levels requested by the EU Lisbon strategy.
394	
395	Conflict of Interest
396	The authors declare that they have no conflict of interest.
397	
398	Appendices
399	Additional supporting information in the online version of this article (see "Supplementary
400	Material") contains the following: ESM_1 - List of native and non-native species,
401	phytosociological optimum (according to Aeschimann et al. 2004), species frequency (%),
402	and minimum and maximum cover (cover-abundance scores according to Braun-Blanquet
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### 650 Tables

**Table 1.** Percent cover (mean  $\pm$  standard error) of different vegetation layers (all species,653black alder and ash) by treatment. Different letters indicate significant differences between654treatments at p < 0.10 (ANOVA with Tukey's HSD test).</td>

% cover of vegetation layer	TR1 (10-15 years)		TR2 (20-30 years)		TR0 (>40 years)	
Upper tree	0 ± 0.0	а	0 ± 0.0	а	82 ± 2.3	b
Lower tree	83 ± 1.8	b	$87\pm1.2$	b	$19\pm1.7$	а
Upper shrub	$21\pm4.2$	b	$16 \pm 2.1$	ab	$10 \pm 1.4$	a
Lower shrub	$41\pm3.1$	с	$26\pm2.4$	b	$14 \pm 1.4$	a
Herbaceous	$78\pm3.4$	a	$93\pm0.9$	b	$91 \pm 1.1$	b
Bare soil	$22\pm3.4$	b	$7\pm0.9$	а	$9\pm1.1$	а
	Alnus	glutinos	a			
Total	$45 \pm 3.0$	а	$61 \pm 4.5$	b	$78 \pm 3.3$	с
Upper tree	$0\pm0.0$	a	$0\pm0.0$	а	$78\pm3.3$	b
Lower tree	$45\pm3.0$	b	$61\pm4.5$	c	$0\pm0.0$	a
Upper shrub	$0\pm0.3$	а	$0\pm0.0$	а	$0\pm0.0$	а
Lower shrub	-		-		-	
	Fraxinu	s excelsi	or			
Total	66 ± 5.5	b	55 ± 6.7	ab	48 ± 3.7	а
Upper tree	$0\pm0.0$	a	$0\pm0.0$	а	3 ± 1.6	b
Lower tree	$44\pm4.7$	b	$30\pm5.0$	а	21 ± 2.2	а
Upper shrub	$18 \pm 3.5$	а	$16 \pm 2.2$	а	11 ± 1.4	а
Lower shrub	4 ± 1.2	а	$9\pm1.6$	b	$12 \pm 1.3$	b

**Table 2.** Soil data by treatment in the study area (means  $\pm$  standard error). Different letters661indicate significant differences between treatments at p < 0.10 (ANOVA with Tukey's HSD</td>662test).

Variable	TR1 (8-10 years)	TR2 (20-30 years)	TR0 (>40 years)
TN%	$0.8 \pm 0.08$ a	$0.9 \pm 0.07$ a	$1.0 \pm 0.07$ a
TOC%	$10.5 \pm 1.05$ a	$11.8 \pm 0.84$ a	$13.0 \pm 0.81$ a
C/N	$12.6 \pm 0.24$ a	$12.8 \pm 0.17$ a	$12.9 \pm 0.2$ a

**Table 3.** Diversity, richness, and cover of functional groups (mean  $\pm$  standard error) by675treatment. Different letters indicate significant differences between treatments at p <0.05</td>676(ANOVA with Tukey's HSD test).

% cover of vegetation layer	TR1 (10-15 years)	)	TR2 (20-30 years)	)	TR0 (>40 years	s)
Total richness	$15.4\pm0.50$	b	$12.7 \pm 0.67$	a	11.3 ± 0.69	a
Shannon index (H)	$3.0\pm0.06$	с	$2.7\pm0.09$	b	$2.1\pm0.05$	а
	Spe	ecies nu	mber			
Non-native species	$1.7\pm0.27$	с	$0.9\pm0.19$	b	$0.0\pm0.00$	а
Ruderal species	$1.3 \pm 0.12$	b	$1.1\pm0.19$	b	$0.5 \pm 0.13$	а
Hydrophilous tall herb species	$2.3\pm0.21$	b	$3.1\pm0.22$	b	$2.1\pm0.09$	a
Shrub species	$2.8 \pm 0.31$	b	$1.4\pm0.21$	а	$1.9\pm0.36$	а
Fraxinetalia species	$3.0 \pm 0.22$	а	$2.9\pm0.19$	a	$3.0\pm0.24$	a
	Ре	ercent co	over			
Non-native species	$15 \pm 2.0$	с	4 ± 1.2	b	$0\pm0.0$	а
Ruderal species	$15 \pm 2.4$	с	$10 \pm 2.4$	b	$0\pm0.2$	a
Tall herb species	$36 \pm 4.6$	b	$41\pm4.6$	b	$5\pm1.0$	a
Shrub species	$16 \pm 2.4$	b	6 ± 1.4	a	$3\pm0.9$	a
Fraxinetalia species	$49 \pm 3.4$	а	$65\pm3.3$	b	$80 \pm 2.9$	c

682	Figure	captions

**Fig. 1** Location of the study areas in Piedmont, Italy (left: blue – Special Protection Areas,

685 red – Sites of Community Importance, green – Ramsar sites) and within the Site of

686 Community importance "Laghi di Ivrea" (right).

687

**Fig. 2** Stand structural variables in the study areas by treatment. TR0: old (>40 years), TR1:

689 medium (10-20 years), TR2: recent coppice (20-30 years). Different letters indicate

690 significant differences between treatments at p < 0.10 (Mann-Whitney test).

691

692 Fig. 3 Regeneration (individuals per hectare) in the study areas by treatment. TR0: old (>40

693 years), TR1: medium (10-20 years), TR2: recent coppice (20-30 years). Different letters

indicate significant differences between treatments at p < 0.10 (Mann-Whitney test).

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Figure 2 Click here to download high resolution image



