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Vegetative regeneration of beech coppices for biomass in Piedmont, NW Italy

This is the author's manuscript

Original Citation:

Availability:

This version is available <http://hdl.handle.net/2318/1657723> since 2018-01-16T14:41:39Z

Published version:

DOI:10.1016/j.biombioe.2017.10.018

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1 Vegetative regeneration of beech coppices for biomass in Piedmont, NW Italy

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11 Abstract

12 Interest in coppices is growing due to the need to replace fossil fuels with renewable energy sources. In
13 Italy, beech covers one million hectares, half of which originated by coppicing. This study tested which
14 factors drive the presence and growth of beech resprouts, with a focus on fertility, cutting intensity, age,
15 and size at time of coppicing.

16 We analyzed 509 stools in 24 stands coppiced between 1 and 26 years before sampling. We fitted
17 Generalized Linear Mixed Models of the probability of sprouting and height of the tallest resprout for
18 each stool as a function of elevation, slope, aspect, bedrock, precipitation, temperature, age at coppicing,
19 time since coppicing, residual shoot density, the sum, average and coefficient of variation of the
20 diameter of cut shoots, and type of stool treatment.

21 Of all harvested stools, 249 (49%) had sprouted with an average of 7.6 resprouts per stool. Height of the
22 tallest resprout on each stool ranged from 3 to 800 cm, mainly as a function of time since coppicing.

23 Resprout mortality was on average 1.4% per plot. Sprouting decreased with decreasing site fertility,
24 increasing precipitation, and increasing size of cut stems. Leaving one or more shoots on the stool after
25 felling produced a high proportion of sprouting stools (82%).

26 Although based on a limited sample, our quantitative analysis of the driving factors of sprouting in
27 beech can be used to support silvicultural decisions in over-mature beech coppices, and to optimize
28 trade-offs between ecosystem services such as biomass production, biodiversity, and hydro-geologic
29 protection.

30 **Keywords:** biomass, *Fagus sylvatica* L., forest management, sprouting

31 Highlights:

- 32 • Forest coppicing allow to replace fossil fuels with renewable energy sources
- 33 • We tested which factors drive the probability and vigor of beech sprouting in the Italian Alps
- 34 • Sprouting decreased with poor soils, higher precipitation and increasing stool size
- 35 • When one or more shoots were left, 82% of the stools sprouted
- 36 • This evidence can be used to sustainably manage beech coppices for biomass

1. Introduction

Coppicing is a silvicultural treatment that takes advantage of the ability of broadleaves to reproduce clonally [1]. When a stem or root is damaged by natural disturbance or cut, changes in hormonal controls prompt root collar buds to generate a new resprout [2]. Such ability varies among species [3] and can be maintained indefinitely or lost with ageing [4].

Since regeneration by coppicing is relatively easy to obtain, and usually grows faster than seedlings thanks to the reserves stored in the root systems of the living stools [5], coppicing has been one of the most common forms of forest management, mostly preferred when seed regeneration is impractical for time, money, or site constraints, and to obtain a fast and steady production of firewood or charcoal [6].

To compensate for the progressive exhaustion of root reserves and consequent mortality of whole stools, and to mitigate the loss of soil and nutrients, clearcut coppicing has often been replaced by sheltered coppicing [7], selection coppicing [8], or maintained under a seed-regenerated overstory.

In Europe, coppicing has historically been more common in southern and eastern countries, e.g., Italy, Spain, Bulgaria, or Serbia, but has recently declined due to depopulation of rural areas and the spreading of fossil fuels, leading to an increase of stored or neglected coppices [9]. To avoid growth stagnation and mechanical instability, steer the forest towards a more "natural" structure, and promote the harvest of larger and more valuable timber, many administrations subsidized the conversion of stored coppices to high forest [1,10,11], while most private-owned coppices were neglected. In recent years, however, the interest in coppices has been rekindled due to the need to replace fossil fuels with renewable energy sources [12-14]. Above being a source of bioenergy, coppices can sustain effectively other ecosystem services such as biodiversity conservation [15-18, but see also 19], protection from natural hazards [20], provision of non-wood forest products [21], or climate change resilience [22].

The key factor in successful coppice management restoration, whether for nature conservation or for economic reasons, is the permanence of the capacity of trees to sprout from their stool. However, there is still very little quantitative information on sprouting ability and its drivers in the main European tree

69 species. For some of them, sprouting success is known to be limited when coppicing is carried out on
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70² older stems [8,23,24]. Beech (*Fagus sylvatica* L.), one of the most widespread and commercially
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71⁴ important forest trees in Europe, is one of such species. In non-managed stands dominated by species of
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72⁷ the genus *Fagus*, the incidence of clonal reproduction is often greater where disturbances are more
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73⁹ severe [25]. Studies on such survival benefits of sprouting in natural systems exist [15,26], but research
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74¹³ exploring the contribution of vegetative reproduction in managed stands are still scarce.

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75¹⁶ Beech coppices are widespread in Southern European mountains, and have been used extensively to
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76¹⁹ provide firewood and charcoal. However, beech has been showed as one of the weakest sprouters among
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77²¹ temperate broadleaves, especially when the temporal span of such sprouting ability is concerned [27]. In
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78²⁴ Italy, beech covers more than one million hectares (10% of total forest cover), half of which originated
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79²⁶ by coppicing [28]. In Alpine regions such as Piedmont (NW Italy) the share of coppices among beech
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80²⁸ forests reaches 90%. Here, the recent regional Forest Management Act prohibited coppicing beech
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81³¹ stands older than 40 years. To support decisions on the ecological and economic sustainability of
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82³³ maintaining coppicing, and to test legal disposition against ecological evidence, we carried out an
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83³⁶ analysis of vegetative regeneration of beech coppices in Piedmont. The aim was to test which factors
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84³⁸ ensure a successful sprouting of beech stools, with a specific attention on fertility, cutting intensity, age
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85⁴¹ and size at the time of coppicing.

86⁴³ 2. Material and methods

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88⁴⁸ We analyzed 24 beech coppice stands in 13 municipalities of Piedmont (Fig. 1). Stands were within an
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89⁵¹ elevation range of 850-1350 m a.s.l.; mean annual temperature and precipitation ranged from 6.4 to 11.0
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90⁵⁴ °C and from 1120 to 2315 mm, respectively (interpolated weather station data for the period 1951-86
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91⁵⁶ [29]). At all sites, soil water regime was classified as udic; soil type and soil nutrient supply were note
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92⁵⁹ measured directly, but site fertility was described by a dummy variable ranging from 1 (poor) to 4 (very
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93 good), following a region-wide forest cover type classification [30]. All stands had been coppiced
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94² between 1 and 26 years before sampling. Data on tree density before and after coppicing were available
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95⁴ from local forest management records; in absence of any other information, we assumed that all stems
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96⁵ had originated as shoots in the original stands. Harvest intensity was defined as the relative change in the
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97⁷ number of shoots before and immediately after coppicing (Table 1). Coppicing was carried out as either
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98⁹ clearcut (one site, harvest intensity =100%), coppice-with-standards (72-95%, 15 sites), or conversion to
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99¹⁰ high forest (61-82%, 8 sites); no selective coppices were present.
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[Figure 1 here]

103²⁴ Stands ranged in size from 0.8 to 4.2 ha, as a result of the extreme fragmentation of privately owned
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104²⁶ forests such as beech coppices in Northern Italy. In each stand we established a circular sampling plot
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105²⁷ (radius = 8-14 m, proportional to tree density) centered on randomly extracted coordinates within the
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106²⁸ stands (making sure that the whole plot area was contained within the stand limits). In each plot we
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107²⁹ recorded slope, aspect, and visually estimated canopy cover. For each stool in the plot, we counted all
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108³⁰ cut and uncut shoots, and all live and dead resprouts (i.e., shoots that we could confidently classify as
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109³¹ having sprouted after the cut), and measured their diameter at stump height, origin (root or stem
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110³² sprouting), and height (Table 1). We estimated the age of the stand at the time of coppicing by averaging
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111³³ tree ring counts from 3-5 exposed stools per plot. We also estimated the age of the resprouts by
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112³⁴ averaging the ring counts from 3-5 increment cores taken from each resprout diameter class.
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113³⁵ We defined as “sprouting” all stools with at least a living resprout at the time of survey, and computed
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114³⁶ resprout mortality as the relative frequency of dead over all resprouts. These figures include mortality
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115³⁷ due to all causes (e.g., competition or natural disturbance), but do not account for resprouts that have
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116³⁸ died and fallen from the stool. The frequency of sprouting stools is therefore a conservative estimate.
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117 We fitted Generalized Linear Mixed Models (GLMMs) of the probability of sprouting (0 or 1, n = 509
118 stools) and height of the tallest resprout (n = 249 stools with live resprouts), which has been often found
119 to correlate with future resprout survival [31]. GLMMs allowed us to accommodate non-independent
120 observations (random variable =plot), non-normality and heterogeneity of the response, by using a
121 binomial and lognormal distribution to model the probability of sprouting and the height of the tallest
122 resprout, respectively. Independent variables included slope, bedrock (granite/acidic or limestone),
123 annual precipitation, mean annual temperature, age at coppicing, time since coppicing, the mean and
124 coefficient of variation of the diameter of cut shoots, and type of stool treatment (total or partial cut)
125 (Table 1). Predictors were filtered for collinearity using variance inflation factors (VIF) thresholds
126 (predictors were excluded if VIF >4). All models were optimized including only significant predictors
127 and successfully scrutinized for overdispersion, normality of random factors, and residual patterns.
128 Goodness-of-fit was assessed by the marginal (fixed factors) and conditional (fixed and random factors)
129 coefficient of determination R^2 for GLMMs [32,33], computed by the function *r.squaredGLMM* in the
130 *MuMIn* package for R [34].

3. Results

131 The stands were coppiced at an age of 35 to 55 years (25th, 50th, 75th percentiles: 40 – 45 – 50 years,
132 respectively). Average shoot density before coppicing was in a range of 969-3550 trees ha⁻² (1524 –
133 2134 – 2383), and harvest intensity was in a range of 61-100% (70% – 79% – 90%). Of all 509
134 harvested stools, 249 (49%) had sprouted a total of 2163 new resprouts (range: 0-100% of sprouting in
135 each plot), with an average of 7.6 resprouts per stool (range: 0.3 - 18.4). Only 5% of them sprouted from
136 roots. Six out of 24 plots experienced resprout mortality (average: 1.4%, range: 1-17%). Except for one
137 recently treated plot where no resprouts existed yet at the time of sampling, the height of the tallest
138 resprout in the plot was significantly correlated with time since coppicing (Pearson's $R = 0.66$, $p < 0.001$).
139 The average age of resprouts was also correlated to time since coppicing ($R = 0.86$, $p < 0.001$) but the two

142 variables did not match perfectly, with a difference of 0-13 years between time since coppicing and
143 average age of resprouts. Current canopy cover ranged from 20 to 100% (average: 55%, correlation with
144 time since coppicing: $R = 0.58$, $p = 0.003$), and tended to saturate after about 15 years from coppicing.
145 (Fig. 2).

146 At the plot level, both the proportion of sprouting stools and maximum resprout height were weakly
147 correlated to decreasing residual shoot density ($R = -0.45$ and -0.33 , $p = 0.03$ and 0.14 , respectively)
148 (Fig. 3). The low significance of the correlation between height and residual density has to do with the
149 fact that age since coppicing is not factored in the analysis, indicating the need for a multiple regression
150 approach. At the stool level, both the number of resprouts per stool and height of tallest resprout
151 decreased with decreasing site fertility (Fig. 4), although the robustness of the correlation is somehow
152 limited by the categorical classification of fertility. However, since fertility and bedrock were highly
153 collinear ($R = 0.85$), we decided to use only the second variable in GLMMs.

154 [Figure 2 here]

155 [Figure 3 here]

156 [Figure 4 here]

157 The model for probability of sprouting explained 41.2% of the total variance in the data (marginal R^2 :
158 26.4% from fixed factors only; dispersion parameter = 0.86). The significant variables were
159 presence/absence of shoots left alive on the stool, mean diameter of cut shoots, age at the time of
160 coppicing, and precipitation (Table 2). Sprouting decreased with increasing size of cut shoots (80% of
161 the those > 40 cm in diameter did not resprout) and increasing precipitation, and increased with
162 increasing age at time of coppicing (Fig. 5); the presence of live shoots on the stool improved the
163 probability of sprouting by 20-25% (Fig. 6). When one or more stools were left on the stool after felling,
164 82% of the stools sprouted.

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[Table 2 here]

[Figure 5 here]

[Figure 6 here]

The model for height of tallest resprout explained 88.9% of the total variance (marginal R^2 : 62.2%). The significant variables were annual precipitation, bedrock, and time since coppicing (Table 3). Age at time of coppicing was, as expected, the dominant driver of maximum resprout height (Fig. 7). However, height also decreased significantly with increasing annual precipitation, and on acidic bedrocks (Fig. 8).

[Table 3 here]

[Figure 7 here]

[Figure 8 here]

4. Discussion

Knowledge about coppicing of European beech stands has a long history in both textbook and operational silviculture [35]. The two main results reported in this paper are not new to the forester community, i.e., that larger shoots are less successful in sprouting due to faster desiccation of the cut stem, and that leaving a shoot on the stool acting as sap sucker keeps the stool tissues vital. However, such knowledge got somewhat obsolete in last decades and very few peer reviewed contributions exist on the topic. Having such popular knowledge confirmed and detailed by the main quantitative results of this paper is of high interest for managers.

Despite the limited sample size and the use of some non-quantitative or partially subjective measurement due to data limitations (e.g., bedrock, fertility, canopy cover), some clear trends emerged.

In managed stands from this study, beech sprouting was influenced by stem diameter, age at the time of

192 coppicing, presence of uncut shoots, time since coppicing, annual precipitation and bedrock. The effect
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193 of size at the time of coppicing has been documented by previous studies [4,36]; small stools are
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194 younger and their resprouts may be more vigorous [37]. This is consistent with anecdotal knowledge of
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195 European foresters [38] and with studies from other species of the *Fagaceae* family, e.g., on North
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196 American oaks, showing a decline in the number of resprouts and a lower height growth with increasing
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197 diameter at the time of coppicing [39,40]. Such behavior may be related to failure of hidden epicormic
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198 buds to develop into new resprouts due to the increasing physical resistance of the bark as diameter
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199 increases with age [41].
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200 All other things being equal, age of the shoot at the time of coppicing had a positive effect at the
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201 individual level, but a negative effect at the stand level; if mean age was <40 years, on average 70% of
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202 stools sprouted, but only 50% if mean age was >50 years. Other European species show a positive effect
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203 of age at time of coppicing on sprouting, e.g., hornbeam [42], as young resprouts can benefit from the
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204 mature root system of a parent tree [43]. The ability of stools to sprout at a later age is the main obstacle
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205 for coppice restoration, as many coppice stands have been either neglected, or converted to high forest
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206 several decades ago. These results are generally consistent with provisions by regional forest regulations
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207 that, in many Italian regions, forbid coppicing of beech beyond 40 years of age.
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208 In our study, more productive sites – those with limestone bedrock rather than acidic – were associated
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209 to an increased probability of sprouting, other factors being equal. Relative to the overall figure (70%),
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210 stands on the most fertile sites (eutrophic beech forest cover type) showed a much higher proportion of
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211 sprouting stools at age <40 years (91%). This suggests the relevance of resource allocation mechanisms,
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212 as abundant non-structural carbohydrates can be preferentially allocated to bud growth rather than shoot
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213 biomass [44], but data on this trait are largely lacking [45].
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214 Regarding the role of climate, high precipitation and moisture may cause the bark to partly fall off from
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215 the stool in shade-tolerant species characterized by a thin bark such as beech, consequently killing or
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216 damaging the buds [31]; in sites where annual precipitation was higher than 1500 mm we observed total
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217 decay of stools as soon as eight years after cutting.

218³ Finally, a very strong effect on sprouting probability was played by the presence of one or more uncut
219⁵ shoots on the stool. This has the advantage of slowing down stool decay and avoid drying out of the cut
220⁸ surface. All else being equal, stools with uncut shoots (even if these had poor growth and form) showed
221¹⁰ a higher probability of sprouting, a higher number of resprouts and a better average resprout growth.

222¹² This result is consistent with earlier evidence from selection coppices, where the number and growth of
223¹⁴ resprouts after a rotation of 20 years was higher than in clearcut coppices or coppices with standards of
224¹⁵ the same age [8]. We therefore suggest that sprouting ability can be improved by retaining at least one
225¹⁷ shoot per stool uncut, especially when restoring coppice in unfavorable situations (poor fertility or late
226¹⁹ age).

227²⁰ Similar variables appeared to influence the growth of resprouts after the cut. Beyond the obvious
228²² influence of time since coppicing, height of the tallest resprout was positively influenced by nutrient
229²⁴ availability (limestone bedrock), and negatively by precipitation. Warmer aspect (south-facing) were
230²⁶ expected to improve resprout growth, especially due to the sensitivity of young resprouts to late frost,
231²⁸ but the effect was not significant (albeit positive).

232³⁰ Another driver of resprout growth was residual shoot density. At the stand level, shading from
233³² increasing stand density markedly affected stool sprouting and growth of the resprouts: the average
234³⁴ proportion of sprouting stools was 69% when residual shoot density was <400 trees per hectare, but only
235³⁶ 32% above that threshold (Fig. 3). Trees that are, or have to be, left standing as a seed source to
236³⁸ compensate for stool exhaustion, also limit the amount of light that reaches the forest floor, and may
237⁴⁰ therefore either favor or hinder the regeneration depending on their density [41].

238⁴² This work, albeit conducted on a limited sample size, confirmed that increasing age and size at the time
239⁴⁴ of coppicing are associated to a decline in sprouting ability of European beech. Over-mature coppices
240⁴⁶ suffer from detrimental physiological changes, a reduction in re-sprouting ability, and increased

241 mechanical failure that substantially decreases their longevity [46]. The recommended maximum age for
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242 coppicing beech is 40 years. However, leaving one shoot uncut on the stool significantly improves
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243 sprouting. All else being equal, residual shoot density, site fertility and climate were also found to play
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244 an important role.
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245 10 11 12 13 246 **5. Conclusions** 14 15

247 We showed how beech sprouting in the western Italian Alps decreases with poor soils, higher
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248 precipitation, and increasing stool size and age. An effective measure to preserve vegetative regeneration
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249 (82% of cases) is to leave one or more shoots uncut.
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250 Such evidence can support silvicultural decisions in overmature beech coppices, which still represent
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251 present an unsolved silvicultural dilemma. When the choice must be made between abandonment, active
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252 conversion into high forest, mixed regeneration systems [47], or restoration of clearcut or selection
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253 coppicing, forest managers should carefully examine stand age, climate, soil, and site characteristics.
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254 Old coppices on poor soils and humid climates exhibit a poor sprouting capacity, and are the first
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255 candidates for conversion to high forest. On the other hand, coppices on more fertile soils that have not
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256 yet reached 40-50 years of age can be targeted for a continuation or restoration of vegetative
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257 regeneration aimed at the provision of sustainable energy wood, provided that one or more live shoots
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258 are left on the stool.
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259 In a warming world, over-mature and neglected coppices are also more vulnerable to climatic stress and
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260 xylem cavitation [48,49]. Therefore, restoration of coppices has the potential to increase forest resilience
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261 in all broadleaves forests at risk from e.g. drought or pests – that put especially large, old trees at risk of
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262 death faster than they can be replaced by seed regeneration – and still sustain key ecosystem services
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263 such as biodiversity, biomass production, and hydrogeologic protection.
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Acknowledgments

The authors support the campaign #ricercaprecaria for the full implementation of the European Charter for Researchers and the acknowledgement of all researchers as workers, and the Change.org petition “Salviamo la ricerca italiana” (<https://www.change.org/p/salviamo-la-ricerca-italiana>) for the increase of research funding in Italy to the levels requested by the EU Lisbon strategy.

Conflict of interest

The authors declare no conflict of interest and compliance with the current laws of the country in which field measurement were performed.

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Figure captions

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Fig. 1 – Distribution of beech in Piedmont, Italy (blue) and beech coppice stands analyzed by this study (red dots)

Fig. 2 – Mean resprout age, maximum resprout height, and current canopy cover as a function of time since coppicing

Fig. 3 - Relationship between residual shoot density and (a) proportion of sprouting stools or (b) maximum resprout height in each plot (n =24)

Fig. 4 - Relationship between site fertility and number of resprouts per stool (a) (n =509) or maximum resprout height (b) (n =249)

Fig. 5 – Marginal effects of significant predictors in the GLMM for the probability of sprouting (binomial regression with log link); in each panel, all other predictors were kept at their mean level

Fig. 6 - Predicted probability of sprouting as a function of mean cut diameter with or without uncut shoots, conditioned on fixed and random effects. Dots represent observed data. Precipitation and age at time of coppicing were set at the average value for all plots

Fig. 7 – Marginal effects of significant predictors in the GLMM for maximum resprout height (lognormal regression with identity link); in each panel, all other predictors were kept at their mean level

1
2 Fig. 8 - Predicted maximum resprout height as a function of time since coppicing and
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4 bedrock, conditioned on fixed and random effects. Dots represent observed data.
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7 Precipitation was set at the average value for all plots
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Tables

Table 1 – Definition and units of variables used in this study

Variable (units)	Description
<i>Descriptive</i>	
Tree density before coppicing (ha^{-1})	From local forest management records
Harvest intensity (%)	$100 - (\text{Residual shoot density} / \text{Tree density before coppicing})$
Resprouts (ha^{-1})	Shoots that we could confidently classify as having sprouted after the cut
Number of stools (ha^{-1})	Only including those that were cut
Proportion of sprouting stools (%)	Stools with live or dead resprouts / Number of stools (per plot)
Number of resprouts per stool (-)	Resprouts / Number of stools
Residual shoot density (ha^{-1})	From local forest management records; includes all shoots but no resprouts
Fertility (ordinal 1 to 4)	From forest cover type map
Resprout diameter (cm)	Measured at stump height
Resprout origin (binary)	Root or shoot
Resprout height (cm)	Measured from stump height
Resprout status (binary)	Dead or alive
Mean resprout age (years)	Average of ring counts from 3-5 increment cores taken from each resprout diameter class
Resprout mortality rate (%)	Relative frequency of dead resprouts (e.g., by competition or natural disturbance), not including those that had died and fallen from the stool
<i>GLMMs - Dependent</i>	
Probability of sprouting (0-1)	On each stool
Height of the tallest resprout (cm)	On each stool
<i>GLMMs - Independent</i>	
Age at time of coppicing (years)	Average of tree ring counts from 3-5 exposed stools per plot
Time since coppicing (years)	From documental records
Bedrock (categorical)	Limestone or granite
Annual precipitation (mm)	From regional climate database
Mean annual temperature ($^{\circ}\text{C}$)	From regional climate database
Type of stool treatment (binary)	Total or partial cut
Sum of the diameter of cut shoots (cm)	
Average diameter of cut shoots (cm)	
Coefficient of variation of the diameter of cut shoots (0-1)	

Table 2 - GLMM for probability of sprouting - summary of standardized coefficients.

Dispersion =0.86. Marginal R^2 : 0.264. Conditional R^2 : 0.412. St.dev. of random effects: 1.09.

N =479 after deletion of missing cases

	β	Std. Error	p-value
intercept	-0.57	0.29	0.047
Type of stool treatment: partial	1.86	0.35	<0.001
average cut diameter	-0.37	0.15	0.012
age at coppicing	0.75	0.29	0.009
precipitation	-1.01	0.28	<0.001

Table 3 - GLMM for height of tallest resprout - summary of standardized regression

coefficients. Marginal R^2 : 0.622. Conditional R^2 : 0.888. St.dev. of random effects: 0.66. N

=242 after deletion of missing cases.

	β	Std. Error	p-value
intercept	3.89	0.23	<0.001
time since coppicing	1.03	0.17	<0.001
precipitation	-0.40	0.21	0.071
bedrock: limestone	0.93	0.43	0.046

Figure 1
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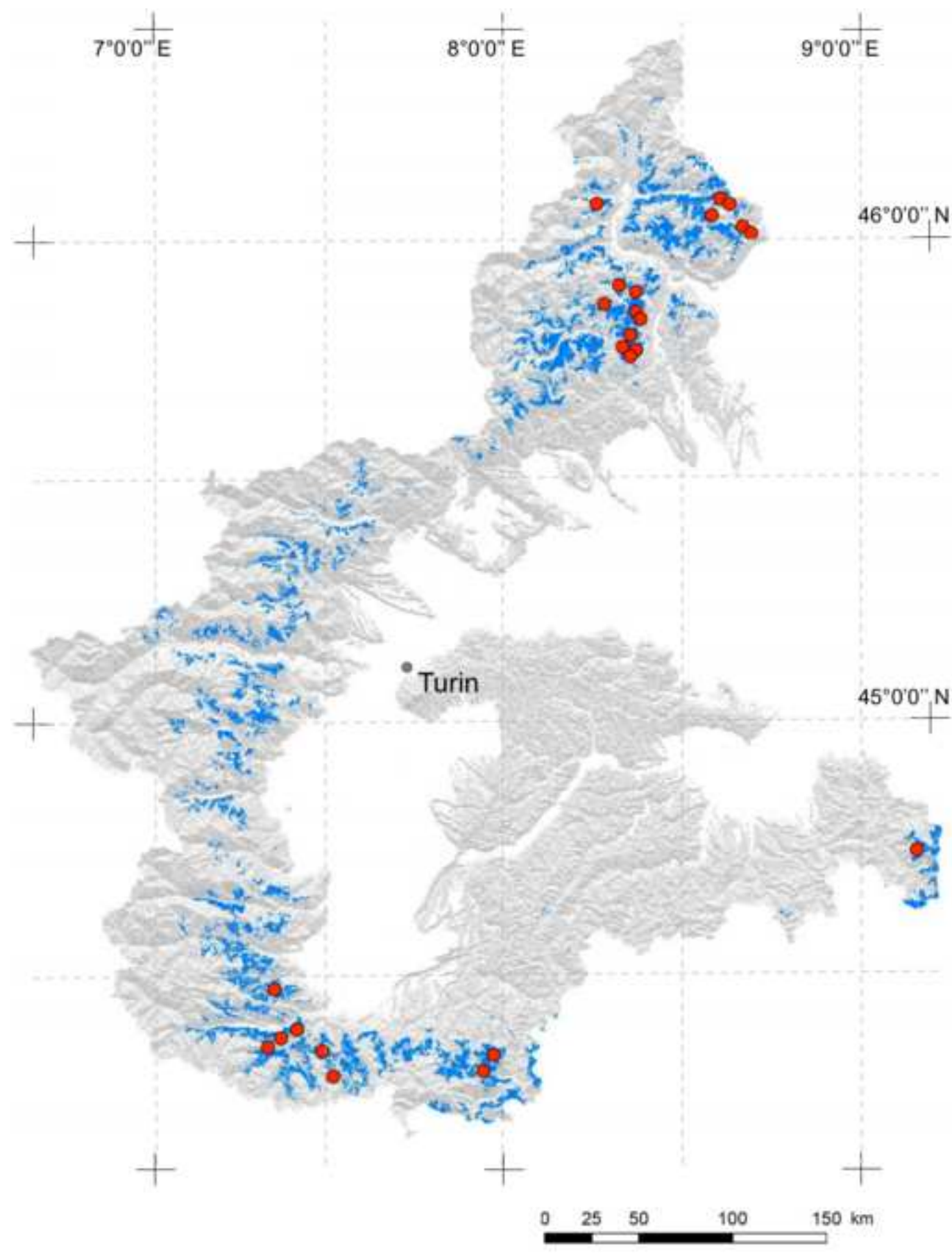


Figure 2

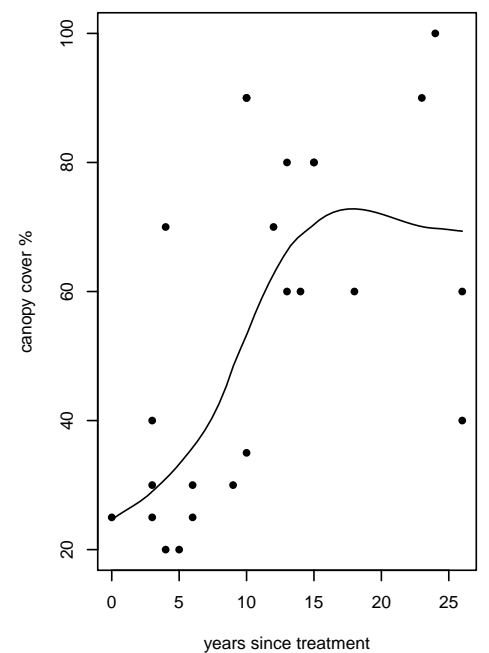
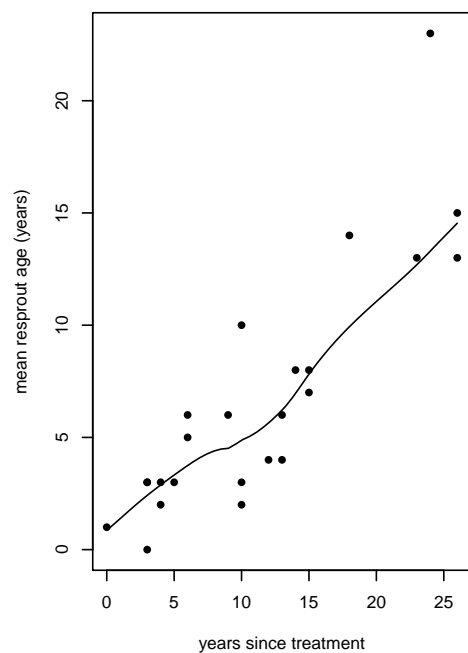
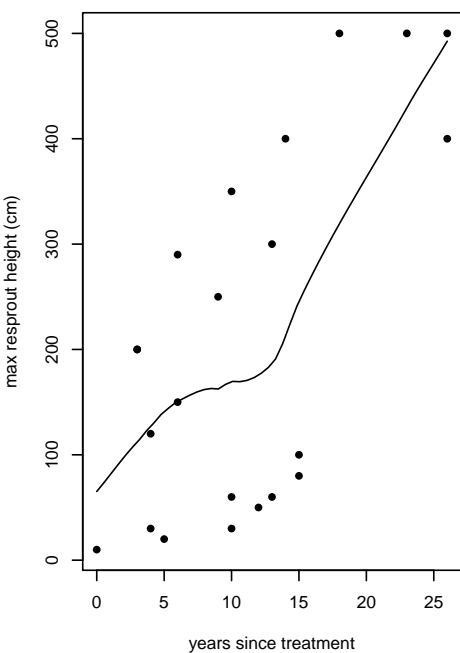


Figure 3

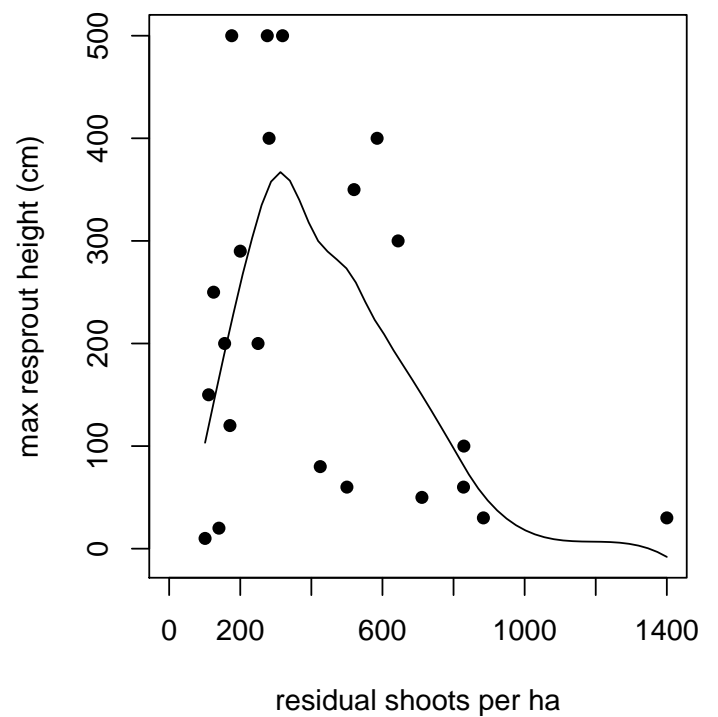
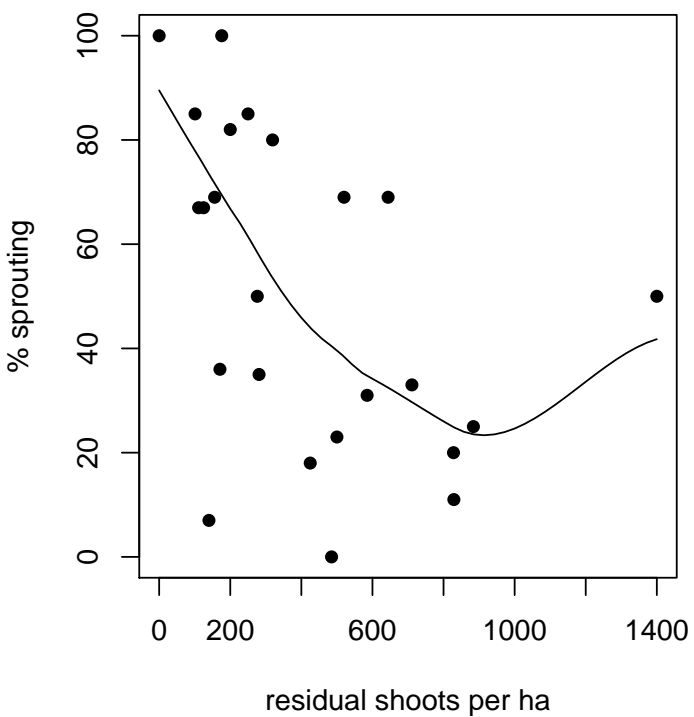


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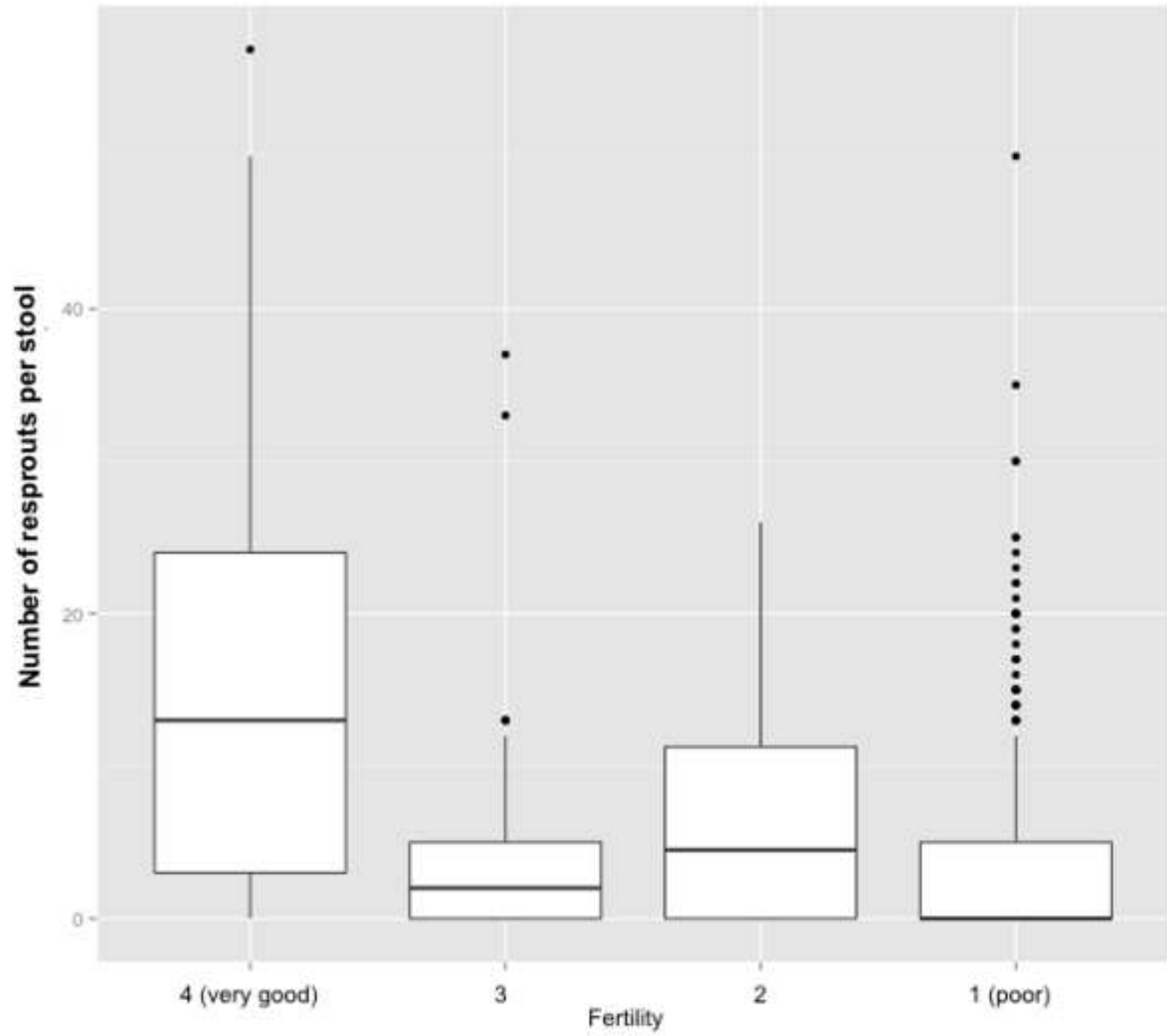


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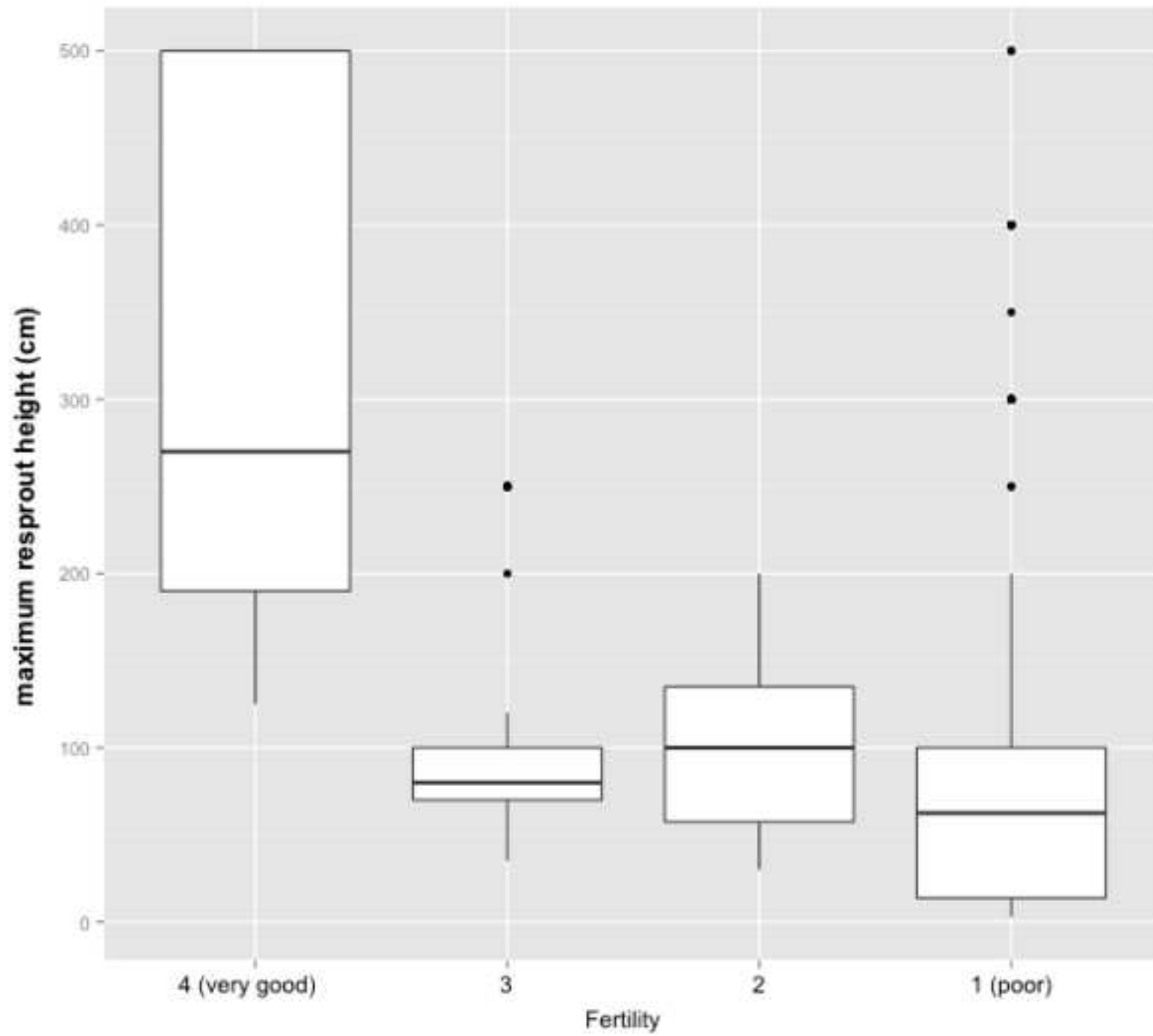


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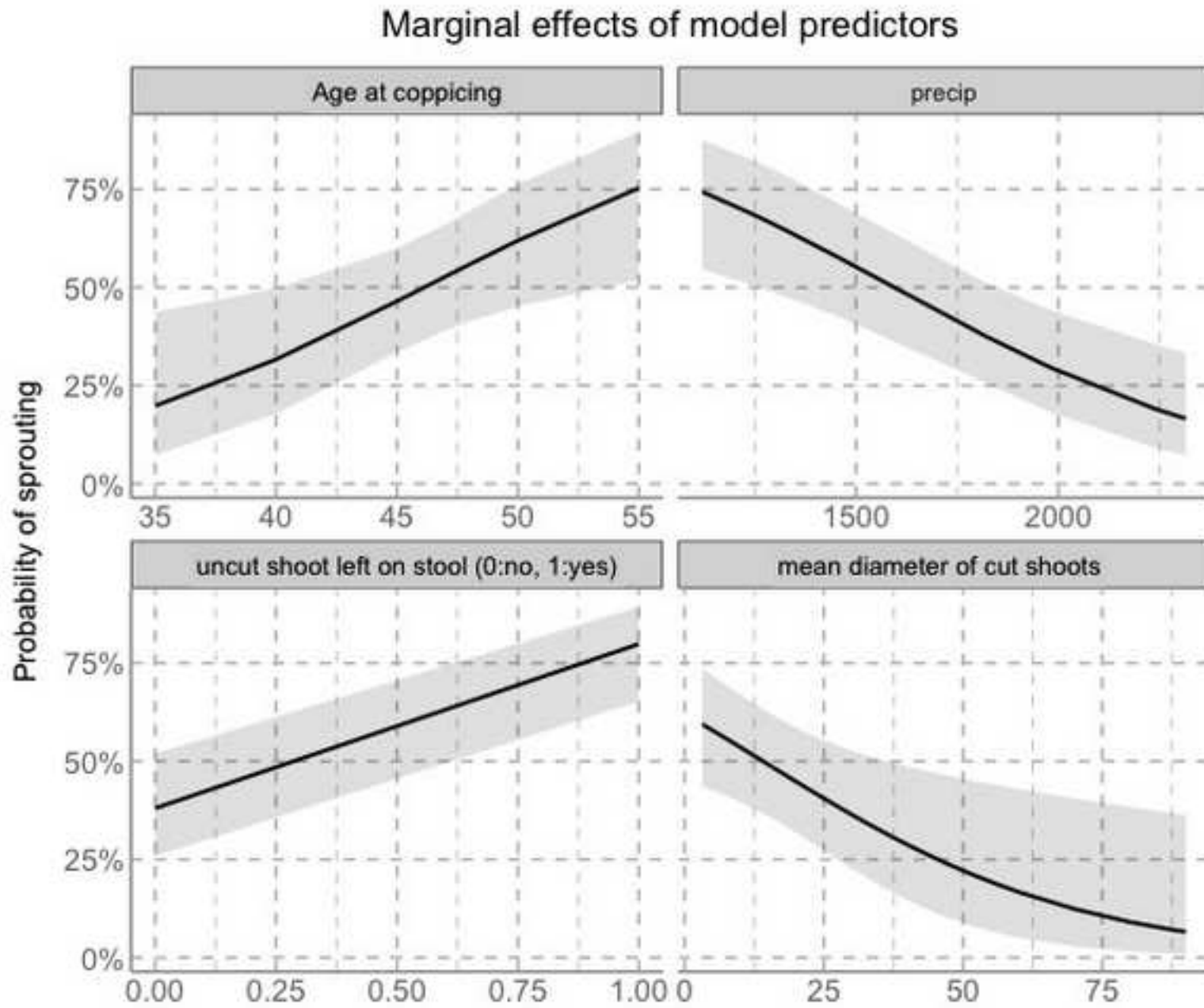


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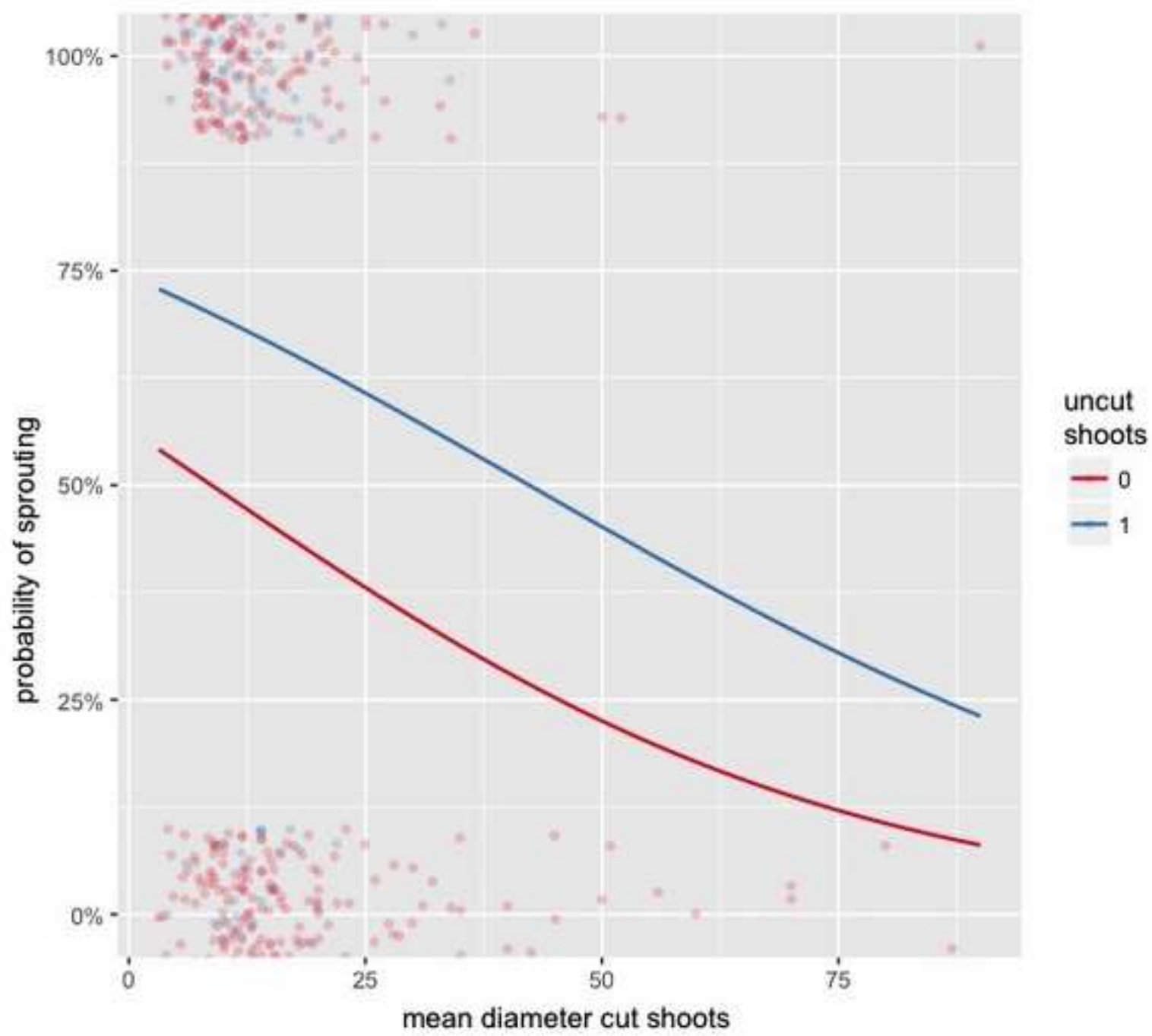


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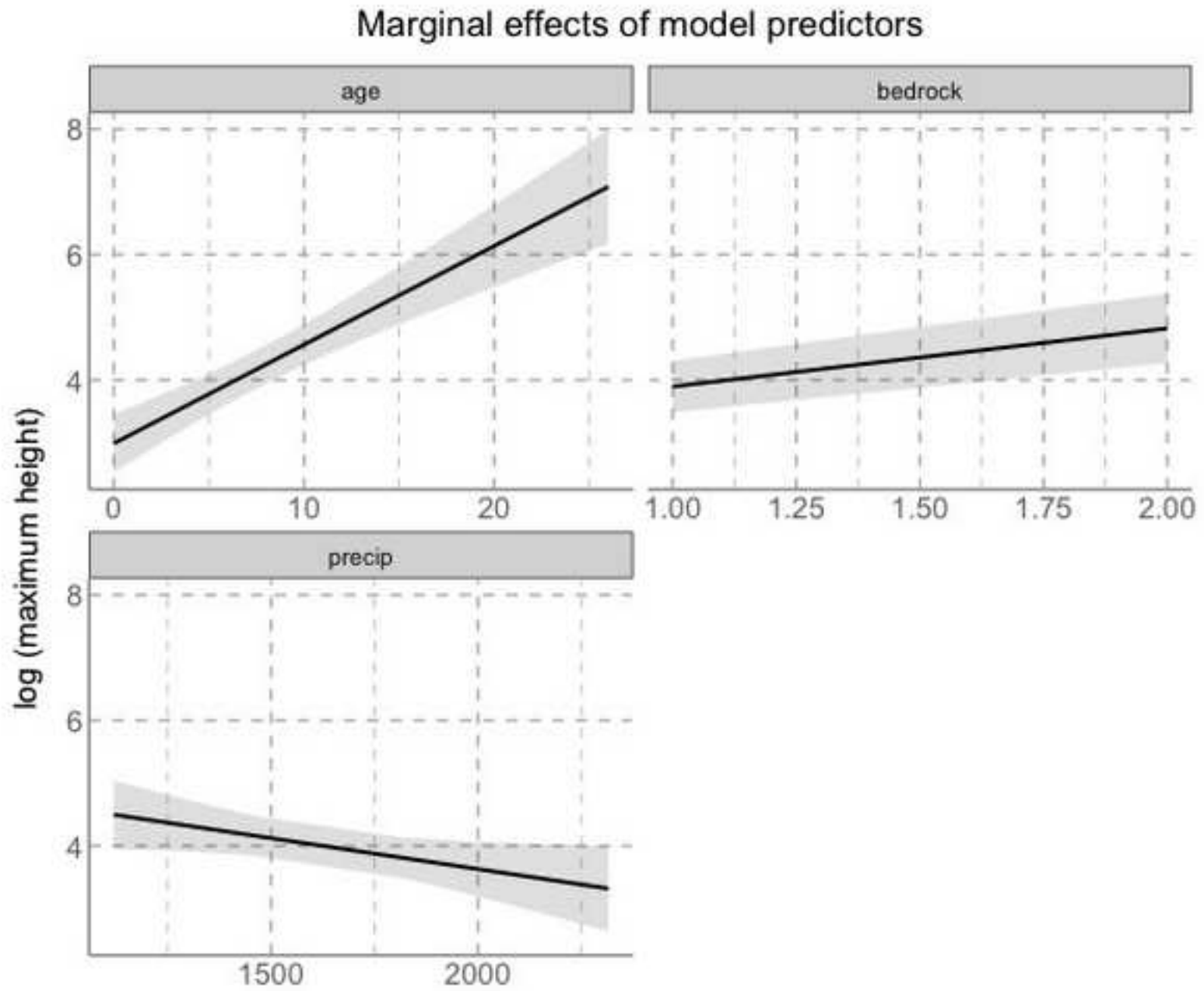


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