

The closer the better? Relative influence of forest continuity and distance to water on the taxonomic and functional structure of riparian plant communities along headwater streams

Philippe Janssen, Richard Chevalier, André Evette, Frédéric Archaux

▶ To cite this version:

Philippe Janssen, Richard Chevalier, André Evette, Frédéric Archaux. The closer the better? Relative influence of forest continuity and distance to water on the taxonomic and functional structure of riparian plant communities along headwater streams. Applied Vegetation Science, 2021, 24 (2), pp.e12591. 10.1111/avsc.12591. hal-03322975

HAL Id: hal-03322975 https://hal.inrae.fr/hal-03322975

Submitted on 20 Aug 2021 $\,$

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

1	The closer the better? Relative influence of forest continuity and distance to water on the taxonomic and
2	functional structure of riparian plant communities along headwater streams
3	
4	Philippe Janssen ¹ *, Richard Chevalier ² , André Evette ¹ , Frédéric Archaux ²
5	
6	¹ Univ. Grenoble Alpes, INRAE, LESSEM, F-38402 St-Martin-d'Hères, France
7	² INRAE, EFNO, F-45290 Nogent-sur-Vernisson, France
8	
9	* Corresponding author, email philippe.janssen@inrae.fr, phone +33 476762879
10	
11	Email addresses of other authors:
12	R. Chevalier (<u>richard.chevalier@inrae.fr</u>), A. Evette (<u>andre.evette@inrae.fr</u>), F. Archaux
13	(frederic.archaux@inrae.fr)
14	
15	Running head: high value of ancient forests near stream
16	
17	Funding information
18	Financial support was provided by the Conseil Départemental de l'Yonne (Auxerre, 89, Convention Conseil
19	Départemental de l'Yonne - Irstea n° 2016_01509 relating to the actions carried out within the framework of

20 the Protection of Sensitive Natural Spaces) and INRAE.

21 ABSTRACT

22 **Questions:** (1) Is forest temporal continuity an important factor in structuring the riparian plant communities

along small alluvial valleys? (2) To what extent can distance from the stream modulate the effect of forest

24 continuity and provide a better understanding of the taxonomic and functional structure of these

25 communities?

26 **Location:** Ouanne watershed, France.

27 Methods: We used a paired samples design, involving stands close and far from the stream, in ancient or

recent riparian forests, in which we studied the vegetation. We performed analyses of species richness and

29 cover, as well as mean values of species preferences and functional traits, and measures of species

30 composition, distinguishing between the understory and overstory.

31 **Results:** Strong differences in richness and mean values, as well as in composition, were found between

32 ancient and recent forests. Ancient forests were characterized by species with a dominant stress-tolerant

33 strategy, whereas recent forests were dominated by hygrophilous and nitrophilous species with a dominant

34 competitive strategy. Furthermore, we found that ancient and recent stands close to the stream encompassed

a larger richness of species, than ancient and recent riparian stands far from the stream, a pattern mostly

driven by the greater co-occurrence of small ruderal species with rapid leaves turnover but also of ancient

37 forest species.

38 **Conclusions:** In small riparian forest stands along headwater streams, we have shown that past human

39 activities leave still detectable traces in contemporary plant communities. In addition, we pointed out that

40 colonization by ancient indicator species was faster in recent forests closer to the stream. This underscores the

41 need to protect remaining ancient riparian forests that are well connected to the stream, even if they are small,

42 and suggests that bank reforestation as well as restoration of hydrological connectivity on rivers fragmented by

43 artificial barriers may be an effective way to increase forest biodiversity in riparian zones.

44

Keywords: ancient forests, connectivity, conservation, creek, distance to stream, forest continuity, functional
traits, legacy effect, riparian vegetation.

47 INTRODUCTION

Historical ecology has demonstrated the legacy effects that past land uses can have on the properties of 48 49 contemporary forest ecosystems and associated biodiversity (Peterken & Game 1984; Hermy et al. 1999; De 50 Frenne et al. 2011). Changes in land cover over time can often be deduced from historical land-use maps, such 51 as the Ordnance Survey Map in Great Britain, the Val der Maelen Map in Belgium or the État-Major Map in 52 France. With these maps, it is possible to determine precisely whether a site has been occupied by forest at different times and thus to characterize the so-called 'forest continuity', also referred as 'forest ancientness' or 53 'ancient woodland' (Flinn & Vellend 2005; Goldberg et al. 2007; Hermy & Verheyen 2007). More precisely, this 54 55 characterization makes it possible to distinguish between ancient forests, which have been forested 56 continuously for centuries, and recent forests, i.e. newly established forests resulting from spontaneous 57 afforestation on former agricultural land or artificial plantations (also known as 'second growth' or 'post-58 agricultural forests'). Forest continuity thus refers to the maintenance of forest cover over time, irrespective of 59 stand maturity and management type. It influences forest biodiversity through two mechanisms, a dispersal 60 limitation, related to increased exposure time to colonization, and a recruitment limitation, related to changes 61 in soil properties and competitive interactions (Flinn & Vellend 2005; Hermy & Verheyen 2007). 62 Research that has linked biodiversity to the temporal continuity of forest cover has mainly studied the 63 response of vascular herbaceous plants in lowland temperate forests, particularly in northwestern Europe (De 64 Frenne et al. 2011). Numerous studies have shown that the composition of plant species differed between 65 ancient and recent forests and several regional lists of ancient forest plant species have been developed (e.g. 66 Hermy et al. 1999; Matuszkiewicz et al. 2013; Bergès et al. 2016). The response of other taxa - insects (e.g. 67 Assmann 1999), lichens (e.g. Janssen et al. 2019), fungi (e.g. Spake et al. 2016) – to forest continuity has also 68 been studied, and despite variations in the magnitude and significance of the effects, have overall highlighted 69 the importance of ancient forests for conservation. However, although some studies have been carried out in 70 areas other than temperate lowland forests, as is the case in mountain and Mediterranean areas (Sciama et al. 71 2009; Janssen et al. 2018; Abadie et al. 2018), several forest types have been largely ignored in historical 72 ecology (Bergès & Dupouey 2021). This is especially the case for riparian forests, i.e., forests at the interface 73 between freshwater and terrestrial ecosystems, usually established on riverbanks. 74 Riparian forests, although representing only 1% of the European continental surface (Weissteiner et al. 2016),

host a unique species pool and are of critical concern for biodiversity conservation (González et al. 2017). This

76 strong ecological interest is, however, threatened by numerous anthropogenic pressures affecting riparian 77 vegetation either directly, e.g. by clearing vegetation, or indirectly, e.g. by altering the flow regime through 78 dams (González et al. 2017). Although the extent of riparian forests has increased sharply since the 1950s in 79 western Europe, due to the natural afforestation of stabilized floodplains and to large plantations of hybrid 80 poplars (Liébault & Piégay 2002; Archaux & Martin 2009), a large part of the natural riparian vegetation has 81 already been lost, degraded or fragmented. There is thus an urgent need to preserve existing intact riparian 82 vegetation and better assess the conservation value of the remaining ancient riparian forests. This is 83 particularly true in headwater streams, where the harvesting of forest stands bordering creeks may threaten 84 the quality of downstream ecosystems (Richardson & Danehy 2007; Suurkuukka et al. 2014).

85 Here, we combined a taxonomic and functional approach, to test whether forest continuity influences the 86 structure of plant communities in small remnant riparian forest stands along creeks. This double approach 87 allows both a focus on conservation issues, studying variations in species richness and identity, and a better 88 understanding of the underlying ecological processes captured by functional traits (i.e. Westoby's Leaf-Height-89 Seed), species preferences for habitat conditions (i.e. Ellenberg's indicator values) and ecological strategies 90 deployed along disturbance and stress gradients (i.e. Grime's CSR). Based upon a dedicated sampling design in 91 which we selected ancient and recent riparian forest stands, either in close proximity to stream or far from the 92 bank, we more precisely tested whether the distance to the water changed the taxonomic and functional 93 response of plant communities to forest continuity. This has been motivated by the fact that the dispersion of 94 plant along streams is to a large part dependent on floods (Nilsson et al. 2010; Bourgeois et al. 2016), which 95 should not only favour the arrival of a higher density of propagules but also a better recruitment of species 96 (Glaeser & Wulf 2009). However, although a few studies have shown differences in the species composition of 97 ancient and recent forests in large alluvial valleys (Verheyen et al. 2003; Glaeser & Wulf 2009; Douda 2010; 98 Chevalier et al. 2014), no studies have yet focused on forest fragments bordering creeks in headwater systems and none have tested the interaction between forest continuity and distance from the stream. In order to fill 99 100 this knowledge gap, we addressed the following two questions: (1) Is temporal continuity of forest cover an important factor in the taxonomic and functional structuring of riparian plant communities in forest stands 101 102 along headwater streams? (2) To what extent can distance from the stream modulate the effect of forest 103 continuity and provide a better understanding of the taxonomic and functional structure of riparian plant 104 communities?

105

106 MATERIALS & METHODS

107 Study area and experimental design

108 The study was carried out along the headwater area of the Ouanne creek (size of study area watershed = 950 109 km^2 , stream width = 2-6 m, mean annual discharge at Toucy gauging station = 0.976 m^3/s) and its tributaries in 110 the center of France (Figure 1). These small alluvial valleys are under the influence of punctual and irregular 111 floods that cause disturbances of small spatial extent, i.e. often limited to riparian stands directly in contact 112 with the stream. They are characterized by a temperate climate with oceanic influences (mean annual temperatures of 11.6°C and annual precipitations ranging between 700 and 800 mm) and a geological context 113 114 dominated by acidic silts and flint clays. To limit variation in environmental conditions, the study area was 115 restricted to the alluvial part of streams, made up of modern alluvial deposits and colonized by forests 116 dominated by European alder (Alnus glutinosa), field maple (Acer campestre), European hornbeam (Carpinus 117 betulus) and European ash (Fraxinus excelsior). The surrounding landscape is mainly composed of agricultural areas, consisting of vast fields of crops and pastures, and wooded areas more or less connected to each other. 118 119 During the summers of 2018 and 2019, we sampled 24 sites in ancient (n = 12) or recent (n = 12) riparian 120 forests. For each site, we studied the vegetation in a pair of stands (n stands = 48), one directly in contact with 121 the stream, referred to as "close to stream" (ancient = 12, recent = 12), the other remote from the stream, 122 referred to as "far from stream" (ancient = 12, recent = 12) (Figure 1). Each pair was established on the same 123 stream segment and positioned in small adjacent riparian forest fragments, averaging 1.46 ha in size. The 124 distinction between stand close and far from the stream was made by establishing a buffer zone of 5 to 10 m 125 between the boundaries of the two stands where vegetation was surveyed (Figure 1). Forest continuity was 126 characterized by crossing digitized and geo-referenced 1:40,000 État-Major maps of France, charted in 1837 for 127 our study area, with 1:10,000 up-to-date forest maps (BD Forêt® V2) in a Geographic Information System (QGIS Development Team 2015). Forest cover overlapping in both maps was considered to indicate ancient forests 128 129 (i.e. with a continuity of forest cover well above 180 years), while current forest cover overlapping with crops or meadows in the État-Major maps was considered to indicate recent forests (i.e. with a continuity of forest 130 cover well below 180 years). To confirm the continuity of the forest cover since the middle of the 19th century, 131 132 we used a series of aerial photographs taken in 1950 at and around each site. Combined with an inspection of 133 the most recent aerial photographs available, this approach allowed us to prospect for polygons occupied by

forests with a natural appearance, i.e. avoiding poplar groves or harvested plots. Finally, to avoid biases due to
low accuracy of vegetation mapping from historical land use maps, isolated polygons < 0.25 ha and < 20 m wide
were eliminated.

137 Vegetation survey

At the scale of the stand (i.e. ancient-close, ancient-far, recent-close, recent-far), vegetation surveys were realized following the Braun–Blanquet abundance-dominance methodology. Within a 200-m² (20 m x 10 m) rectangular plot, parallel to the stream, a complete inventory was done in June/July, either in 2018 (n = 26) or in 2019 (n = 22). All the vascular plants (pteridophytes and phanerogams) were recorded by considering two layers, that of the overstory (i.e. tree and shrub layers \geq 2 m height) and that of the understory (i.e. herbaceous layer < 2 m height). To limit biases associated with observer effect, all surveys were performed by the same experienced botanist (RC), assisted by colleagues, considering an average search effort of 30 min per plot.

145 Stand and soil data

146 At the scale of the site (ancient forests, n pairs = 12; recent forests, n pairs = 12), stand and soil data were 147 measured only in the stand far from the stream. This was motivated by the fact that the overstory composition 148 and structure of these riparian forests were homogeneous between stands close and far from the stream. 149 Stand attributes were characterized using a series of nested circular subplots: a 4-m-radius subplot to measure 150 trees with a diameter at breast height (DBH, at 1.3 m standard height) \geq 5 cm, a 6-m-radius subplot for DBH \geq 151 10 cm, a 9-m-radius subplot for DBH \ge 25 cm and a 15-m-radius subplot for DBH \ge 45 cm. For each tree, 152 species, diameter and vitality state (alive, dying, dead), were recorded. Soil properties were characterized using 153 a soil corer in the stand center to describe the soil texture classes at 20-cm depth, based on the Jamagne's 154 (1967) texture triangle, and estimated the percentage of coarse elements ($\emptyset > 2$ mm) in the first 50 cm of soil. 155 Humus forms, i.e. mull, moder, mor and their subdivisions, were described based on three sampling points 156 located 2 m from the plot center in opposite directions. Finally, the coordinates of the stand center and the 157 elevation difference with the stream water level were measured using a GPS device and an inclinometer 158 respectively. 159 From the field measurements, we calculated a set of variables to compare environmental conditions between 160 ancient and recent forests. Stand variables (basal area of living trees and of dead trees, number of 5-cm

diameter classes, canopy cover percentage), were standardized to a per hectare basis and used to compare
overstory characteristics. Soil variables (percentage of sand/silt/clay and of coarse elements) were derived

either from direct measurements or from the conversion of texture classes to percentage using the central
value of the texture triangle. Humus forms were converted to a quantitative assessment using the humus index
proposed by Ponge et al. (2002). These soil variables were used to highlight possible legacy effects of past
human activities on edaphic conditions or pre-existing differences in environmental conditions. Spatial (latitude
and longitude) and topographic (altitude and elevation from water) variables were used to control for possible
biases in the distribution of sites within the study area.

169 Plant preference and trait data

170 To analyze how the ecological preferences and strategies of plants differed according to forest continuity and distance from the stream, we collected data for each species from three sources of information. Ellenberg's 171 172 indicator values (Ellenberg et al. 1992) for light (L), nutrient availability (N) and soil moisture (F) were extracted 173 from the Baseflor database (Julve 1998) and used to characterize variations in species habitat preference. 174 Among the 194 species encountered, these values were unavailable for three species with a single occurrence 175 in our data set (i.e. singleton species) and for five taxa identified at the genus level. Grime's values (Grime 176 1977) for respectively Competitors, Stress-tolerators and Ruderals, i.e. ternary coordinates, were extracted 177 from the list of species available in Pierce et al. (2017) and used to characterize variations in the strategyenvironment relationships of riparian plants. For missing values (n = 54), data were either completed by using 178 179 ecological information on closely related species, i.e. mean value for the same genus (n = 43), or discarded 180 (genus level taxa, n = 4; singleton species, n = 2; others, n = 2). Westoby's values for specific leaf area (SLA; leaf 181 area per dry mass), plant height at maturity and seed mass were extracted from the LEDA database (Kleyer et 182 al. 2008) and used to characterize variations in resource acquisition, competitive and dispersal abilities. Missing 183 values were restricted to ten taxa (genus level taxa, n = 4; singleton species, n = 4; others, n = 2). 184 At the scale of each forest stand (n = 48), we then computed the mean values weighted by the relative cover of 185 each species bearing each value, i.e. community-weighted means (CWMs) (FD package, Laliberté et al. 2014). Since herbaceous species are the most sensitive to forest continuity (Gilliam 2007) and tree species can be 186 187 shaped by forestry practices, CWMs were calculated only for herbaceous understory species. Finally, to analyze how ancient forest species differed according to forest continuity and distance from the 188 189 stream, we used three local species lists of plants (Hermy et al. 1999; Dupouey et al. 2002; Bergès et al. 2016) 190 and considered as ancient forest indicator a species that appears at least twice in these lists.

191 Statistical analysis

To determine whether environmental parameters varied between ancient and recent forests (n = 24), we used
the Mann-Whitney U-test. For proportional data (i.e. sand, silt, clay, soil coarse elements and canopy cover),
logit transformation was applied prior to the calculation.

195 To determine whether species richness and cover, as well as the mean values of species' preferences and traits, 196 were influenced by forest continuity, distance to stream and their interaction, we used linear mixed models 197 (LMMs) in which "Sites" (n = 24) was included as a random effect (Ime4 package, Bates et al. 2015). This 198 random effect is designed to control for the non-independence of observations made in paired stands close to 199 or far from the stream. For taxonomic measures, we used the total richness and cover of forest overstory and 200 understory species as well as the richness and cover of ancient forest indicator species as dependent variables 201 (Appendix S1). For mean values, we used as dependent variables CWMs related to species preferences (light, 202 nutrient availability and soil moisture), ecological strategies (competitors, stress-tolerators and ruderals) and 203 functional strategies of plants (SLA, height and seed mass) (Appendix S1).

204 To determine whether species composition was influenced by forest continuity, distance to the stream and 205 their interaction, we used multivariate generalized linear models (GLMs) (mvabund package, Wang et al. 2012). 206 We fitted the full model, testing the interaction between forest continuity and distance of the stream to each 207 species that was present in more than 5% of the stands, using presence-absence data, with a binomial 208 distribution and summed the likelihood ratio statistics across the univariate responses to estimate their 209 multivariate response. Because overstory composition can influence the recruitment of ancient-forest species 210 (Thomaes et al. 2014) and because tree composition can change within a few decades during riparian 211 succession (Schnitzler 1995; Fierke & Kauffman 2005; Janssen et al. 2020), we fitted one multivariate GLM for 212 the understory and one for the overstory. The significance of the factors 'forest continuity' and 'distance to 213 stream' in the multivariate GLM was assessed using an analysis of variance with the PIT-trap method and 999 214 bootstrap resamples (Warton et al. 2017). This method has the advantage of using a resampling scheme that 215 ensures that p-values are approximately correct when the independence assumption is not satisfied, as is the 216 case of our paired sampling design. To determine which herbaceous and tree species best contribute to the 217 overall model deviation, we extracted univariate test statistics and adjusted p-values for each species, using the 218 Holm's step-down procedure to correct for multiple testing (for details see, Wang et al. 2012). Finally, to 219 provide a graphical representation of the two main factors and their interaction, we used a canonical analysis

- 220 of principal coordinates (CAP, Anderson & Willis 2003) with the Jaccard distance community matrix (vegan
- 221 package, Oksanen et al. 2013).
- Analyses were performed with R version 4.0.0 (R Core Team 2020).
- 223
- 224 RESULTS
- 225 Variations in environmental conditions between ancient and recent riparian forests
- 226 Mann–Whitney U-test revealed that only the humus index varied significantly with forest continuity, being
- 227 more acidic in ancient forests than in recent forests. All others environmental variables did not vary with forest
- continuity (Appendix S2), indicating that ecological conditions were comparable between ancient and recent
- 229 forests in the studied area.
- 230 Effect of forest continuity and distance to stream on species richness and cover and mean indicator and traits
- 231 values
- A total of 194 vascular plant species have been inventoried on the 48 forest stands, of which 34 species have
- been identified as ancient forest indicator species. At stand scale, overstory richness ranged from 4 to 13
- species (mean \pm SD = 8 \pm 2.31), understory richness ranged from 14 to 58 species (mean \pm SD = 36 \pm 11.9), while
- ancient plant richness ranged from 3 to 18 species (mean \pm SD = 8 \pm 3.77).
- 236 LMMs revealed no significant interaction between forest continuity and distance to stream (Table 1). The
- 237 richness and cover of ancient forest species and the mean Grime value for stress-tolerators were significantly
- higher in ancient forest stands, while the richness and cover of overstory vegetation, the mean Ellenberg value
- for soil moisture and nutrient availability, as well as the mean Grime value for competitors were significantly
- 240 higher in recent forest stands. Also, the richness of understory vegetation, the richness and cover of ancient
- forest species, the mean Ellenberg value for nutrient availability, the mean Grime value for ruderals and the
- 242 mean value for SLA were significantly higher close to the stream, while the mean Grime value for stress-
- tolerators and the mean value for plant height were significantly higher far to the stream.

244 Effect of forest continuity and distance to stream on the species composition of overstory and understory

- 245 The multivariate GLM for understory vegetation showed that species composition varied significantly with
- forest continuity (Deviance = 259.8, p = 0.001) and distance to the stream (Deviance = 179.5, p = 0.007), but
- not with their interaction term (Deviance = 75.9, p = 0.617). Based on the deviation explained by factors, most
- of the variation in species composition was related to differences between ancient and recent forests, as

249 shown by the first CAP-axis (Figure 2-A). The second axis was more evidently related to the distance to the 250 stream. Univariate tests for each species showed that few of them contributed significantly to the multivariate 251 significant effect (Table 2). Among the species that captured the largest amount of the deviance explained by 252 forest continuity, Angelica sylvestris and Filipendula ulmaria were more frequent in recent forests, while 253 Milium effusum and Carex sylvatica were more frequent in ancient forests. For the distance to the stream, 254 species that captured the largest amount of the deviance, i.e. Veronica montana, Stellaria holostea, Elymus 255 caninus and Phalaris arundinacea, were all more frequent in stands close to the stream (Appendix S3). 256 The multivariate GLMs for overstory vegetation showed that species composition varied significantly with forest continuity (Deviance = 99.3, p = 0.001) but neither with distance to the stream (Deviance = 19.9, p =257 258 0.643), nor with the interaction term (Deviance = 15.6, p = 0.719). The strong forest continuity effect is well 259 represented by the first CAP-axis (Figure 2-B), which shows clear difference in species composition between 260 ancient and recent forests. Univariate tests for each species showed that few of them contributed to the 261 multivariate significant effect. Among the trees/shrubs that captured the largest amount of the deviance 262 explained by forest continuity, Alnus glutinosa, Sambucus nigra and Corylus avellana were more frequent in 263 recent forests, while Carpinus betulus was more frequent in ancient forests (Appendix S4).

264

265 DISCUSSION

266 In accordance with the large amount of previous studies in European temperate forests (Hermy et al. 1999; De 267 Frenne et al. 2011; Matuszkiewicz et al. 2013; Bergès et al. 2016), we found that past land uses leave still 268 detectable traces in contemporary riparian plant communities. This highlights that even in narrow strips of 269 riparian forests along small streams, and in addition to the numerous environmental factors that structure 270 riparian communities, it is important to consider forest continuity to better understand biodiversity patterns 271 (Verheyen et al. 2003; Glaeser & Wulf 2009; Douda 2010). In addition, we showed that plant species richness, 272 including that of ancient forest indicator species, was higher in stands near the stream, and that this effect was 273 found both in ancient and recent forests. This suggests that the restoration of the hydrological connectivity on 274 rivers fragmented by artificial barriers (e.g. weirs and bed sills) could be an effective way to increase forest biodiversity (Glaeser & Wulf 2009). Overall, by showing that ancient riparian forest stands, even if small in size, 275 276 supported distinct assemblages of plant species, our results highlight their strong ecological interest and the 277 need to better conserve the remaining fragments in agricultural landscapes.

278 Forest continuity strongly influences riparian communities of headwater streams

279 Our results showed that the richness and cover of overstory vegetation was higher in recent riparian forest 280 stands than in ancient ones. This may be related to the maintenance of shrubs in early successional stages 281 (Douda 2010), as confirmed by univariate analyses for each species (Appendix S4), when forest gradually colonized former agricultural lands (Bergès & Dupouey 2021). This may also be related to a higher level of 282 283 available resources in recent forests (Hermy et al. 1999; Herault & Honnay 2005; Sciama et al. 2009), which 284 should favor a higher density of individuals and benefit a larger diversity of species (i.e. species-energy 285 hypothesis, Wright 1983). Indeed, Ellenberg's values support the view that nutrient availability was greater in 286 recent than in ancient forests, due to legacy effects of past agricultural practices or pre-existing differences in 287 soil properties, thus conditioning past land uses. Our results also showed that the richness and cover of ancient 288 forest species were significantly higher in ancient forests (Hermy & Verheyen 2007). This confirms the validity 289 of previously established lists (Hermy et al. 1999; Dupouey et al. 2002; Bergès et al. 2016), even in the riparian 290 context (Douda 2010; Chevalier et al. 2014), and, as ancient forest species represent a guild of specialists 291 sensitive to habitat loss and fragmentation (Bergès & Dupouey 2021), it pleads for priority conservation of 292 ancient forest stands, even of small size, along the banks of headwater streams. Specifically, our results 293 showed that the Grime's ecological strategies were different between ancient and recent forests, dominated 294 by stress-tolerators in the first case and by competitors in the second. Those results are consistent with 295 previous ones (Hermy et al. 1999; Hermy & Verheyen 2007; Douda 2010) and reflect a filtering of plants species 296 in relation to available environmental conditions, less limited by resources and leading to greater competitive 297 interactions between vigorous species in recent forests. However, due to overall more fertile habitat 298 conditions, it has been shown that the recruitment of ancient forest species was faster in recent forests in 299 alluvial valleys than in non-alluvial contexts (Dumortier et al. 2002; Verheyen et al. 2003). Our results tend to 300 confirm this, since of the 34 ancient forest species recorded in our study area, 26 were inventoried in recent 301 forests. This indicates that habitat conditions in recent riparian forests are globally favorable to the recruitment 302 of ancient forest species and suggests that restoration actions promoting the reforestation of riverbanks could 303 benefit forest biodiversity (McClain et al. 2011) as well as that of streams (Suurkuukka et al. 2014). Regarding 304 species composition, we found significant differences between ancient and recent forest stands. For the 305 understory, these results are consistent with those reported in temperate forests (De Frenne et al. 2011; 306 Matuszkiewicz et al. 2013; Bergès et al. 2016), including riparian forests (Verheyen et al. 2003; Glaeser & Wulf

307 2009; Douda 2010). As such, among the herbaceous species that predominantly occurred in ancient forests, 308 numerous were already listed as ancient forest indicator species, e.g. Milium effusum, Carex sylvatica or Vinca 309 minor (Hermy et al. 1999; Dupouey et al. 2002; Bergès et al. 2016). Interestingly, our results also emphasized 310 plants indicative of recent forests, which are usually found in wetlands (Douda 2010), e.g. Angelica sylvestris, 311 Caltha palustris or Filipendula ulmaria. Beyond showing that understory composition responds consistently to 312 increasing soil moisture in recent forests, as expressed by Ellenberg values, our results highlight that recent 313 forest species captured most of the explained deviance. This suggests recent forest species have a higher 314 indicative power than ancient forest species and thus, that relevant lists of recent forest plants (Bergès et al. 2016) may be a complementary tool for identifying the continuity of forest cover in areas lacking historical 315 316 land-use maps or where their accuracy is too coarse. Our results also highlighted differences in the composition 317 of overstory. This can be attributed to different maturation stages between ancient forests, dominated by the 318 post-pioneer species Carpinus betulus, and recent forests, dominated by the pioneer species Alnus glutinosa. 319 Associated with changes in humus forms, these results reveal the difficulty of disentangling the effects of forest 320 continuity from those of forest succession. Indeed, it has been shown that changes in canopy composition 321 (Thomaes et al. 2014) and maturity stage (Janssen et al. 2018) of stands can induce a shift in herbaceous 322 communities in ancient forests. Thus, although the difference in successional stage may partially explain the 323 understory changes between ancient and recent forests, the fact that all other environmental variables did not 324 vary with forest continuity supports the idea of a strong effect left by past human activities in the studied 325 alluvial valleys.

326 Stream proximity enhances the conservation value of riparian forests

327 Total understory and ancient forest species richness was higher in stands located near the water. On the one 328 hand, and linked to the predictions of the species-energy hypothesis (Wright 1983), this difference can be 329 attributed to more fertile habitat conditions, as expressed by the increase in the mean Ellenberg value for 330 nutrient availability near the stream. Indeed, in our predominantly agricultural study area, it is possible that 331 nutrient leaching into streams preferentially enriches the best-connected adjacent riparian zones. On the other hand, this effect can be attributed to more frequent and intense floods in the stands closest to the stream. This 332 333 assumption is supported by the fact that not only the richness of ruderals but also the mean value of specific 334 leaf area increased in stands close to the stream, while the mean value of plant height decreased. Given that 335 ruderal species are fast-growing with an annual life cycle (Grime 1977), that high SLA values indicate the

336 dominance of "acquisitive" species with rapid leaf turnover, and that smaller species indicate a shorter 337 potential growth duration between disturbances (Westoby 1998), our results provide strong evidence that 338 near-stream herbaceous communities are adapted to a more disturbed environment. Another important point 339 is related to the potential of colonization, which is known to increase along flooded areas due to the addition of hydrochorous species to the local species pool (Jansson et al. 2005; Nilsson et al. 2010). However, contrary 340 341 to expectations, the mean value of seed mass did not increase with distance to the stream. Although seed 342 traits have been demonstrated to explain long-distance dispersal patterns (Nilsson et al. 2010; Bourgeois et al. 343 2016), local patterns of seed deposition may be less dependent on seed morphology, especially along creeks. 344 Indeed, in headwater streams, it has been shown that non-hydrochores can successfully disperse by water flow 345 (Honnay et al. 2001) but also that propagules can be caught and carried away by floating debris (Nilsson et al. 346 2010). We therefore infer that the positive effect of stream proximity on richness patterns was mainly due to 347 flooding which increased both disturbances and colonization opportunities in ancient and recent forest stands. 348 Specifically, the fact that almost all of the recent forests studied were a short distance downstream from 349 ancient forests may explain why they received more ancient forest species than stands far from the stream 350 (Honnay et al. 2001).

351 Regarding species composition, significant differences were found for understory vegetation but not for 352 overstory vegetation between stands close and far from the stream. These results thus confirm that riparian 353 forest stands belonged to the same successional stage in each pair of stands but also that distance to water 354 strongly structures herbaceous communities. Indeed, on riverbanks, the distance to the stream is most often 355 related to an elevation gradient and, by extension, to the flood gradient, well known for its great importance in 356 the structuring of riparian communities (Naiman & Decamps 1997; Poff et al. 1997). As such, numerous studies 357 have documented strong shift in species and traits composition from the bottom to the top of the riverbank 358 (Kyle & Leishman 2009; McCoy-Sulentic et al. 2017). Thus, although no differences in Ellenberg values for soil 359 moisture were found between stands close and far from the stream, in the riparian forests studied, we infer 360 that compositional changes were related to spatial variations in flood frequency and intensity as well as in nutrient availability. In stands close to the stream, flooding, by periodically disturbing understory vegetation 361 362 but also by providing nutrients, may have favored the establishment of species requiring fertile environments 363 (e.g. Alliaria petiolata, Elymus caninus, Phalaris arundinacea) and/or ruderals (e.g. Stellaria holostea, Veronica 364 montana, Viola reichenbachiana).

365

366 CONCLUSION

- 367 Using a dedicated sampling design, we demonstrated that stand position relative to the stream and forest 368 continuity strongly shaped riparian plant communities, even for small wooded patches in headwater streams. 369 Specifically, we show that in fertile and humid riparian forests, the colonization credit usually documented in 370 non-alluvial settings appears to resolve more rapidly, at least for the recent forests closest to the stream. These 371 results complement well the few previous studies carried out in large alluvial valleys (Verheyen et al. 2003; 372 Herault & Honnay 2005; Glaeser & Wulf 2009; Douda 2010) and give weight to the need to increase the 373 conservation of the remaining ancient riparian forests. Moreover, by showing that recent forests established 374 along streams were rapidly colonized by ancient forest species, our results suggest that riverbank reforestation 375 actions could be an effective way to increase the forest biodiversity. In Europe's predominantly agricultural 376 landscapes, this knowledge provides complementary arguments to encourage the conservation and restoration 377 of ancient forests. This is particularly the case along headwater streams, which account for up to 80% of the 378 total length of streams in watersheds around the world (Gomi et al. 2002). 379
- 380 ACKNOWLEDGEMENTS
- 381 We thank Nadia BARUCH and Antoine DELBERGUE of the CRPF Bourgogne-Franche-Comté, Marie GARNIER of 382 the IGN and Noémie BERTRAND, Adélie CHEVALIER, Barthélémy DESSANGES and Hilaire MARTIN of INRAE for 383 their help in the field and in data acquisition.

384

385 AUTHOR CONTRIBUTIONS

R.C. conceived the research idea and collected data; P.J. performed statistical analyses; P.J., wrote the first
 draft of the paper; all authors discussed the results and commented on the manuscript.

388

389 DATA ACCESSIBILITY

390 The research data supporting this publication are provided in the online supporting information.

391

392 **REFERENCES**

- Abadie, J., Avon, C., Dupouey, J.-L., Lopez, J.-M., Tatoni, T., & Bergès, L. 2018. Land use legacies on forest
 understory vegetation and soils in the Mediterranean region: Should we use historical maps or in situ
 land use remnants? *Forest Ecology and Management* 427: 17–25.
- Anderson, M.J., & Willis, T.J. 2003. Canonical analysis of principal coordinates: a useful method of constrained
 ordination for ecology. *Ecology* 84: 511–525.
- Archaux, F., & Martin, H. 2009. Hybrid poplar plantations in a floodplain have balanced impacts on farmland
 and woodland birds. *Forest Ecology and Management* 257: 1474–1479.
- Assmann, T. 1999. The ground beetle fauna of ancient and recent woodlands in the lowlands of north-west
 Germany (Coleoptera, Carabidae). *Biodiversity and Conservation* 8: 1499–1517.
- Bates, D., Mächler, M., Bolker, B., & Walker, S. 2015. Fitting Linear Mixed-Effects Models Using Ime4. *Journal of Statistical Software* 67:.
- Bergès, L., Avon, C., Arnaudet, L., Archaux, F., Chauchard, S., & Dupouey, J.-L. 2016. Past landscape explains
 forest periphery-to-core gradient of understorey plant communities in a reforestation context.
 Diversity and Distributions 22: 3–16.
- Bergès, L., & Dupouey, J. 2021. Historical ecology and ancient forests: Progress, conservation issues and
 scientific prospects, with some examples from the French case. *Journal of Vegetation Science* 32:
 e12846.
- Bourgeois, B., González, E., Vanasse, A., Aubin, I., & Poulin, M. 2016. Spatial processes structuring riparian plant
 communities in agroecosystems: implications for restoration. *Ecological Applications* 26: 2103–2115.
- Chevalier, R., Archaux, F., Berthelot, A., Carnnot-Milard, L., Duprez, M., Gaudin, S., & Villemez, A. 2014. Le
 concept de forêt ancienne s'applique-t-il aux peuplerais cultivées? Test de pertinence avec la flore des
 vallées de Champagne. *Revue Forestière Française* LXV: 375–388.
- 415 De Frenne, P., Baeten, L., Graae, B.J., Brunet, J., Wulf, M., Orczewska, A., Kolb, A., Jansen, I., Jamoneau, A.,
 416 Jacquemyn, H., Hermy, M., Diekmann, M., De Schrijver, A., De Sanctis, M., Decocq, G., Cousins, S.A.O.,
 417 & Verheyen, K. 2011. Interregional variation in the floristic recovery of post-agricultural forests:
 418 Recovery of post-agricultural forest. *Journal of Ecology* 99: 600–609.
- Douda, J. 2010. The role of landscape configuration in plant composition of floodplain forests across different
 physiographic areas. *Journal of Vegetation Science* 21: 1110–1124.
- Dumortier, M., Butaye, J., Jacquemyn, H., Van Camp, N., Lust, N., & Hermy, M. 2002. Predicting vascular plant
 species richness of fragmented forests in agricultural landscapes in central Belgium. *Forest Ecology and Management* 158: 85–102.
- 424 Dupouey, J.-L., Sciama, D., Koerner, W., Dambrine, É., & Rameau, J.-C. 2002. La végétation des forêts anciennes.
 425 *Revue Forestière Française* LIV: 521–532.
- Ellenberg, H., Weber, H.E., Düll, R., Wirth, V., Werner, W., & Paulißen, D. 1992. Zeigerwerte von Pflanzen in
 Mitteleuropa. Scripta geobotanica 18: 1–248.
- Fierke, M.K., & Kauffman, J.B. 2005. Structural dynamics of riparian forests along a black cottonwood
 successional gradient. *Forest Ecology and Management* 215: 149–162.
- Flinn, K.M., & Vellend, M. 2005. Recovery of forest plant communities in post-agricultural landscapes. *Frontiers in Ecology and the Environment* 3: 243–250.
- Gilliam, F.S. 2007. The ecological significance of the herbaceous layer in temperate forest ecosystems.
 Bioscience 57: 845–858.

- 434 Glaeser, J., & Wulf, M. 2009. Effects of water regime and habitat continuity on the plant species composition of 435 floodplain forests. *Journal of Vegetation Science* 20: 37–48.
- Goldberg, E., Kirby, K., Hall, J., & Latham, J. 2007. The ancient woodland concept as a practical conservation
 tool in Great Britain. *Journal of Nature Conservation* 15: 109–119.
- Gomi, T., Sidle, R.C., & Richardson, J.S. 2002. Understanding processes and downstream linkages of headwater
 systems. *BioScience* 52: 905.
- González, E., Felipe-Lucia, M.R., Bourgeois, B., Boz, B., Nilsson, C., Palmer, G., & Sher, A.A. 2017. Integrative
 conservation of riparian zones. *Biological Conservation* 211: 20–29.
- Grime, J.P. 1977. Evidence for the existence of three primary strategies in plants and its relevance to ecological
 and evolutionary theory. *The American Naturalist* 111: 1169–1194.
- Herault, B., & Honnay, O. 2005. The relative importance of local, regional and historical factors determining the
 distribution of plants in fragmented riverine forests: an emergent group approach. *Journal of Biogeography* 32: 2069–2081.
- Hermy, M., Honnay, O., Firbank, L., Grashof-Bokdam, C., & Lawesson, J.E. 1999. An ecological comparison
 between ancient and other forest plant species of Europe, and the implications for forest
 conservation. *Biological Conservation* 91: 9–22.
- Hermy, M., & Verheyen, K. 2007. Legacies of the past in the present-day forest biodiversity: a review of past
 land-use effects on forest plant species composition and diversity. *Ecological Research* 22: 361–371.
- Honnay, O., Verhaeghe, W., & Hermy, M. 2001. Plant community assembly along dendritic networks of small
 forest streams. *Ecology* 82: 1691–1702.
- 454 Jamagne, M. 1967. Bases et techniques d'une cartographie des sols. *Annales agronomiques* 18: n° hors-série.
- Janssen, P., Bec, S., Fuhr, M., Taberlet, P., Brun, J.-J., & Bouget, C. 2018. Present conditions may mediate the
 legacy effect of past land-use changes on species richness and composition of above-and below ground assemblages. *Journal of Ecology* 106: 306–318.
- Janssen, P., Fuhr, M., & Bouget, C. 2019. Beyond forest habitat qualities: Climate and tree characteristics as the
 major drivers of epiphytic macrolichen assemblages in temperate mountains. *Journal of Vegetation Science* 30: 42–54.
- Janssen, P., Stella, J.C., Piégay, H., Räpple, B., Pont, B., Faton, J.-M., Cornelissen, J.H.C., & Evette, A. 2020.
 Divergence of riparian forest composition and functional traits from natural succession along a
 degraded river with multiple stressor legacies. *Science of The Total Environment* 721: 137730.
- Jansson, R., Zinko, U., Merritt, D.M., & Nilsson, C. 2005. Hydrochory increases riparian plant species richness: a
 comparison between a free-flowing and a regulated river. *Journal of Ecology* 93: 1094–1103.
- Julve, P. 1998. Baseflor. Index botanique, écologique et chorologique de la flore de France. Version : 2017.
 http://perso.wanadoo.fr/philippe.julve/catminat.htm.
- Kleyer, M., Bekker, R.M., Knevel, I.C., Bakker, J.P., Thompson, K., Sonnenschein, M., Poschlod, P., van
 Groenendael, J.M., Klimeš, L., Klimešová, J., Klotz, S., Rusch, G.M., Hermy, M., Adriaens, D., Boedeltje,
 G., Bossuyt, B., Dannemann, A., Endels, P., Götzenberger, L., Hodgson, J.G., Jackel, A.-K., Kühn, I.,
 Kunzmann, D., Ozinga, W.A., Römermann, C., Stadler, M., Schlegelmilch, J., Steendam, H.J.,
 Tackenberg, O., Wilmann, B., Cornelissen, J.H.C., Eriksson, O., Garnier, E., & Peco, B. 2008. The LEDA
 Traitbase: a database of life-history traits of the Northwest European flora. *Journal of Ecology* 96:
 1266–1274.

- Kyle, G., & Leishman, M.R. 2009. Plant functional trait variation in relation to riparian geomorphology: The
 importance of disturbance. *Austral Ecology* 34: 793–804.
- Laliberté, E., Legendre, P., & Shipley, B. 2014. FD: measuring functional diversity from multiple traits, and other
 tools for functional ecology. R package version 1.0-12.
- Liébault, F., & Piégay, H. 2002. Causes of 20th century channel narrowing in mountain and piedmont rivers of
 southeastern France. *Earth Surface Processes and Landforms* 27: 425–444.
- 481 Matuszkiewicz, J.M., Kowalska, A., Kozłowska, A., Roo-Zielińska, E., & Solon, J. 2013. Differences in plant 482 species composition, richness and community structure in ancient and post-agricultural pine forests in
 483 central Poland. *Forest Ecology and Management* 310: 567–576.
- 484 McClain, C.D., Holl, K.D., & Wood, D.M. 2011. Successional models as guides for restoration of riparian forest
 485 understory. *Restoration Ecology* 19: 280–289.
- McCoy-Sulentic, M.E., Kolb, T.E., Merritt, D.M., Palmquist, E., Ralston, B.E., Sarr, D.A., & Shafroth, P.B. 2017.
 Changes in community-level riparian plant traits over inundation gradients, Colorado River, Grand
 Canyon. *Wetlands* 37: 635–646.
- Naiman, R.J., & Decamps, H. 1997. The ecology of interfaces: riparian zones. *Annual review of Ecology and Systematics* 28: 621–658.
- 491 Nilsson, C., Brown, R.L., Jansson, R., & Merritt, D.M. 2010. The role of hydrochory in structuring riparian and
 492 wetland vegetation. *Biological Reviews* 85: 837–858.
- Oksanen, J., Blanchet, G.F., Kindt, R., Legendre, P., Minchin, P.R., O'Hara, R.B., Simpson, G.L., Solymos, P.,
 Stevens, M.H.H., & Wagner, H. 2013. *vegan: Community Ecology Package. R package version 2.0-10.*
- Peterken, G.F., & Game, M. 1984. Historical factors affecting the number and distribution of vascular plant
 species in the woodlands of central Lincolnshire. *Journal of Ecology* 72: 155–182.
- Pierce, S., Negreiros, D., Cerabolini, B.E.L., Kattge, J., Díaz, S., Kleyer, M., Shipley, B., Wright, S.J., Soudzilovskaia,
 N.A., Onipchenko, V.G., van Bodegom, P.M., Frenette-Dussault, C., Weiher, E., Pinho, B.X., Cornelissen,
 J.H.C., Grime, J.P., Thompson, K., Hunt, R., Wilson, P.J., Buffa, G., Nyakunga, O.C., Reich, P.B.,
 Caccianiga, M., Mangili, F., Ceriani, R.M., Luzzaro, A., Brusa, G., Siefert, A., Barbosa, N.P.U., Chapin,
 F.S., Cornwell, W.K., Fang, J., Fernandes, G.W., Garnier, E., Le Stradic, S., Peñuelas, J., Melo, F.P.L.,
 Slaviero, A., Tabarelli, M., & Tampucci, D. 2017. A global method for calculating plant CSR ecological
 strategies applied across biomes world-wide. *Functional Ecology* 31: 444–457.
- Poff, N.L., Allan, J.D., Bain, M.B., Karr, J.R., Prestegaard, K.L., Richter, B.D., Sparks, R.E., & Stromberg, J.C. 1997.
 The natural flow regime. *BioScience* 47: 769–784.
- Ponge, J.-F., Chevalier, R., & Loussot, P. 2002. Humus Index: an integrated tool for the assessment of forest
 floor and topsoil properties. *Soil Science Society of America Journal* 66: 1996–2001.
- 508QGIS Development Team. 2015. QGIS Geographic Information System. Open Source Geospatial Foundation509Project.
- R Core Team. 2020. *R: A language and environment for statistical computing.* R Foundation for Statistical
 Computing, Vienna, Austria.
- Richardson, J.S., & Danehy, R.J. 2007. A synthesis of the ecology of headwater streams and their riparian zones
 in temperate forests. *Forest Science* 53: 131–147.
- Schnitzler, A. 1995. Successional status of trees in gallery forest along the river Rhine. *Journal of Vegetation Science* 6: 479–486.

- Sciama, D., Augusto, L., Dupouey, J.-L., Gonzalez, M., & Domínguez, C.M. 2009. Floristic and ecological
 differences between recent and ancient forests growing on non-acidic soils. *Forest Ecology and Management* 258: 600–608.
- Spake, R., van der Linde, S., Newton, A.C., Suz, L.M., Bidartondo, M.I., & Doncaster, C.P. 2016. Similar
 biodiversity of ectomycorrhizal fungi in set-aside plantations and ancient old-growth broadleaved
 forests. *Biological Conservation* 194: 71–79.
- Suurkuukka, H., Virtanen, R., Suorsa, V., Soininen, J., Paasivirta, L., & Muotka, T. 2014. Woodland key habitats
 and stream biodiversity: Does small-scale terrestrial conservation enhance the protection of stream
 biota? *Biological Conservation* 170: 10–19.
- Thomaes, A., De Keersmaeker, L., Verschelde, P., Vandekerkhove, K., & Verheyen, K. 2014. Tree species
 determine the colonisation success of forest herbs in post-agricultural forests: Results from a 9 yr
 introduction experiment. *Biological Conservation* 169: 238–247.
- Verheyen, K., Bossuyt, B., Honnay, O., & Hermy, M. 2003. Herbaceous plant community structure of ancient
 and recent forests in two contrasting forest types. *Basic and Applied Ecology* 4: 537–546.
- Wang, Y., Naumann, U., Wright, S.T., & Warton, D.I. 2012. mvabund an R package for model-based analysis of
 multivariate abundance data. *Methods in Ecology and Evolution* 3: 471–474.
- Warton, D.I., Thibaut, L., & Wang, Y.A. 2017. The PIT-trap—A "model-free" bootstrap procedure for inference
 about regression models with discrete, multivariate responses. *PloS one* 12: e0181790.
- Weissteiner, C., Ickerott, M., Ott, H., Probeck, M., Ramminger, G., Clerici, N., Dufourmont, H., & de Sousa, A.
 2016. Europe's Green Arteries—A Continental Dataset of Riparian Zones. *Remote Sensing* 8: 925.
- 536 Westoby, M. 1998. A leaf-height-seed (LHS) plant ecology strategy scheme. *Plant and Soil* 199: 213–227.
- 537 Wright, D.H. 1983. Species-energy theory: an extension of species-area theory. *Oikos* 41: 496–506.
- 538

539 SUPPORTING INFORMATION

- 540 Appendix S1: Correlation matrix of dependent variables
- 541 Appendix S2: Variations of environmental variables in relation to forest continuity
- 542 Appendix S3: Univariate tests for understory species
- 543 Appendix S4: Univariate tests for overstory species
- 544 Appendix S5: Raw data: environmental variables
- 545 Appendix S6: Raw data: species by sites matrix

Table 1. Mean (±SD) of species richness and species cover, as well as of the community-weighted means of trait value, in relation to forest continuity (ancient vs recent) and

547	distance to stream	(close vs far) along the	Ouanne creek and its tributaries,	France (p-value based on lir	near mixed models).
-----	--------------------	--------------------------	-----------------------------------	------------------------------	---------------------

		Ancient	forests	Recent forests		Forest	Distance	Interaction
	Variable	close	far	close	far	continuity	to stream	Interaction
		Mean (±SD)	Mean (±SD)	Mean (±SD)	Mean (±SD)	p value	p value	p value
Richness and cover	Overstory richness	6.83 (±1.70)	6.33 (±1.92)	9.42 (±1.31)	8.50 (±2.78)	0.001	0.197	0.700
	Overstory cover	86.79 (±20.59)	76.62 (±19.99)	91.48 (±17.07)	97.62 (±20.03)	0.027	0.721	0.154
	Understory richness	39.25 (±10.66)	28.42 (±10.03)	43.67 (±7.76)	34.92 (±13.87)	0.161	<0.001	0.654
	Understory cover	118.05 (±33.02)	109.66 (±35.68)	141.57 (±28.00)	126.32 (±35.38)	0.087	0.134	0.656
	Ancient forest richness	11.25 (±3.52)	8.42 (±4.12)	8.58 (±2.87)	5.42 (±2.11)	0.026	<0.001	0.778
	Ancient forest cover	42.38 (±21.53)	28.02 (±15.23)	28.02 (±22.21)	15.38 (±11.57)	0.037	0.004	0.842
Mean value (CWM)	Soil moisture	5.35 (±0.35)	5.19 (±0.17)	5.74 (±0.53)	5.81 (±0.46)	0.002	0.557	0.157
	Light	4.61 (±0.40)	4.66 (±0.38)	4.84 (±0.32)	4.95 (±0.38)	0.066	0.242	0.662
	Nutrient availability	5.25 (±0.82)	4.76 (±1.23)	6.37 (±0.71)	5.82 (±0.99)	0.003	0.021	0.897
	Competitors	29.12 (±5.89)	26.95 (±5.45)	35.94 (±6.42)	35.81 (±5.06)	0.001	0.369	0.426
	Stress-tolerators	27.70 (±8.83)	36.67 (±16.17)	16.61 (±5.98)	22.68 (±10.86)	0.003	0.007	0.569
	Ruderals	43.18 (±9.19)	36.38 (±14.35)	47.45 (±6.96)	41.52 (±11.32)	0.246	0.004	0.829
	Specific leaf area	27.82 (±5.80)	23.86 (±6.13)	31.99 (±4.03)	27.77 (±6.32)	0.067	<0.001	0.892
	Plant height	0.70 (±0.23)	0.94 (±0.36)	0.54 (±0.17)	0.77 (±0.30)	0.094	0.001	0.953
	Seed mass	2.87 (±1.25)	4.10 (±2.49)	2.82 (±1.47)	2.77 (±0.94)	0.163	0.221	0.188

- 549 **Table 2.** Top-ranking species (n = 10) among 92 herbaceous species that explained most of the variation in species composition between ancient/recent forests and stands
- 550 far/close to the stream (Test = univariate test statistics from the multivariate GLM; p = p-values adjusted). To facilitate interpretation, species frequency in either
- ancient/recent forests or far/close stands is provided (* = ancient forest indicator species).

Така	Forest continuity		Ancient	Decent	Tava	Distance to stream		For	Class
IdXd	Test	р	Ancient	Recent	IdXd	Test	р	Fdf	close
Angelica sylvestris	14.736	0.015	4.2%	29.2%	Veronica montana*	13.801	0.019	10.4%	35.4%
Filipendula ulmaria	13.023	0.023	8.3%	33.3%	Stellaria holostea*	12.566	0.045	2.1%	22.9%
Milium effusum*	10.759	0.104	20.8%	2.1%	Elymus caninus*	11.740	0.053	12.5%	35.4%
Carex sylvatica*	9.200	0.153	22.9%	4.2%	Phalaris arundinacea	9.305	0.209	0.0%	12.5%
Caltha palustris	9.178	0.195	0.0%	12.5%	Veronica chamaedrys	9.161	0.245	2.1%	18.8%
Carex elongata	9.178	0.219	0.0%	12.5%	Atriplex prostrata	7.544	0.480	0.0%	10.4%
Vinca minor*	9.178	0.228	12.5%	0.0%	Viola reichenbachiana*	7.057	0.531	14.6%	33.3%
Glechoma hederacea	8.268	0.294	25.0%	43.8%	Geum urbanum	6.579	0.572	31.3%	45.8%
Scrophularia auriculata	5.965	0.596	2.1%	14.6%	Alliaria petiolata	6.261	0.684	18.8%	35.4%
Urtica dioica	5.842	0.768	22.9%	39.6%	Myosoton aquaticum	6.019	0.741	0.0%	8.3%

553 **Figure 1.** Location of the study area in France (A), distribution of sampled riparian forests along the Ouanne

creek and its tributaries (B), form of the sampling design used to survey plant communities in relation to their

distance to the stream (C) and flow regime variations over the last 10 years at the Toucy gauging station (D,



dashed green line = interannual mean flow; dashed red line = 2-year return flood).

Figure 2. Constrained canonical analysis of principal coordinates of riparian understory (A) and overstory (B) communities, for each studied sites along the Ouanne creek and its tributaries, France. To facilitate graphical interpretation, the centroids for the interaction between forest continuity and distance to the stream are provided. In addition, the centroids in principle coordinate space of the most influential (i.e. with the highest cumulative deviance) understory and overstory species are provided (* = ancient forest indicator species, see also Appendix S3 and S4).

