Does the strength of facilitation by nurse shrubs depend on grazing resistance of tree saplings?

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Abstract

Facilitation of tree regeneration by nurse shrubs that offer protection against large herbivores is an important driver of wood-pasture dynamics. Here we asked whether the response to facilitation by nurse shrubs depends on the grazing resistance of the protégé saplings. We experimentally tested the protective effects of the thorny \textit{Rosa rubiginosa} on browsing frequency, survival, and biomass change of saplings of two species-groups, presumably differing in grazing resistance: the coniferous \textit{Abies alba} and \textit{Picea abies} and the deciduous \textit{Acer pseudoplatanus} and \textit{Fagus sylvatica} saplings. The saplings were planted under and outside (1.5 m) planted nurse shrubs, under zero, low and high grazing intensity. In total, 1920 young saplings were transplanted to 60 blocks and followed for 1 year.

Although the number of saplings browsed did not differ between species-groups, the coniferous saplings showed lower resistance to cattle browsing (i.e. lower survival and growth rates) than the deciduous saplings. The less resistant coniferous saplings benefited significantly more from nurse shrubs than the more resistant deciduous species in terms of growth of the surviving saplings, but not in terms of overall survival. This was likely due to herbivory on the nurse shrubs causing incidental browsing on protégé saplings and differences in biomass off-take. At high grazing intensity facilitative effects of the nurse shrubs decreased, especially for the coniferous species.

These results have important management implications for the endangered wood-pastures in Western Europe. For a sustainable management and conservation that allows tree recruitment, grazing intensity should remain low to best promote facilitation processes for all tree species, but in particular for the less resistant conifer saplings.

Zusammenfassung

Die Förderung der Baumregeneration durch Ammengebüsche, die einen Schutz gegen große Herbivoren bieten, ist eine wichtige treibende Kraft für die Dynamik von Waldweiden. An dieser Stelle fragten wir uns, ob die Reaktion auf

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Introduction

The importance of facilitation for determining ecosystem functioning and dynamics, plant community structure, and regulating biodiversity is now widely acknowledged (Brooker et al., 2008; Bruno, Stachowicz, & Bertness, 2003; Callaway, Kikodze, Chiboshvili, & Khetsuriani, 2005; Cheng, Wang, Chen, & Wei, 2006; Tirado & Pugnaire, 2005). Facilitative interactions between plants occur when one plant ("nurse plant") modifies, directly or indirectly, the abiotic or biotic environment and consequently increases the performance of a nearby plant ("target plant") (Bertness & Callaway, 1994; Brooker & Callaghan, 1998). Facilitative and competitive interactions generally act simultaneously upon neighbouring plants, and recent studies have focussed on how this balance of plant-plant interactions changes along gradients of environmental severity, including abiotic stress or biotic disturbance (sensu Brooker & Callaghan, 1998; Grime, 1977). In general, the role of facilitation increases with abiotic stress whereas competition decreases (Callaway et al., 2002; Cheng et al., 2006; Choler, Michalet, & Callaway, 2001; Liancourt, Callaway, & Michalet, 2005). However, this relationship is not always linear and is currently being vigorously debated (Maestre, Valladares, & Reynolds, 2006; Michalet et al., 2006; Travis, Brooker, Clark, & Dytham, 2006). Besides abiotic factors, biotic disturbances such as grazing also influence plant interactions. In ecosystems driven by large herbivores, unattractive, toxic or thorny plants were found to have positive indirect (i.e. grazer mediated) effects on palatable herbs, shrubs or trees (Bakker et al., 2004; Callaway et al., 2005; Smit, Béguin, Buttler, & Mueller-Schärer, 2005; Smit, den Ouden, & Müller-Schärer, 2006). Brooker, Scott, Palmer, and Swaine (2006) and Smit, Vandenberghe, den Ouden, and Müller-Schärer (2007) found a hump-backed relationship between facilitation of saplings by a nurse plant and the level of grazing disturbance. The reduced facilitation for saplings under high grazing intensities was related to high net search effort by the herbivores and increased damage to nurse plants. It is known that the strength of facilitation along abiotic stress gradients can differ between target species due to differences in their tolerance to a particular abiotic stress (e.g. shade and moisture) and/or competitive ability (Gómez-Aparicio, Valladares, Zamora, & Quero, 2005; Liancourt et al., 2005). However, responses of facilitation processes to a biotic disturbance gradient, such as a grazing intensity, have thus far been rarely studied for species that differ in their grazing resistance.

Wood-pastures are ecosystems in which large herbivores drive vegetation dynamics through the maintenance of complex assemblages of grassland, shrub and woodland patches (Bakker et al., 2004; Olff et al., 1999; Vera, Bakker, & Olff, 2006). Due to their structural heterogeneity, wood-pastures harbour a great biodiversity, including many rare and endangered plant and animal species (Kirby, Thomas, Key, McLean, & Hodgetts, 1995; Pott & Hüppe, 1991). They also have a high cultural, socio-economic and landscape value...
Most ancient wood-pastures are found in places with a long history of extensive livestock grazing and forest management. However, due to current changes in agricultural practices (Etienne, 1996), only a few remnants of this rich system remain scattered across Europe, e.g., Jura Mountains (CH), New Forest (UK), Borkener Paradies (DE), Junner Koeland (NL). Worldwide, there is interest in combining agriculture and silviculture to conserve and re-establish these endangered systems (Lehmkuhler et al., 2003; Manning, Fischer, & Lindenmayer, 2006). This requires the maintenance of large herbivore densities which are neither too high to allow shrubs or trees to regenerate nor too low for grassland to regenerate (Gillett, 2005; Kirby et al., 1995; Vera et al., 2006).

Facilitation from nurse plants (e.g., the thorny shrub Rosa rubiginosa L.) is an important process for natural tree establishment in wood-pastures as tree saplings frequently die due to cattle activity (Smit et al., 2006), particularly when grazing intensity is high (Vandenbergh, Freléchoux, Moravie, Gadallah, & Buttlar, 2007). The probability of a sapling being browsed and killed as a result depends on grazing intensity (Hester, Mitchell, & Kirby, 1996) and browsing resistance of the species (Hester, Bergman, Iason, & Moen, 2006). Resistance includes appearance (e.g., size or structure), mechanical or chemical defences (e.g., stiff needles, secondary compounds) and specific intrinsic abilities (i.e., tolerance) to survive and regrow after biomass loss under different environmental conditions (Boege & Marquis, 2005; Rosenthal & Kotanen, 1994). In the remaining wood-pastures of the Swiss Jura Mountains, two coniferous (Abies alba Miller and Picea abies L.) and two deciduous (Acer pseudoplatanus L. and Fagus sylvatica L., Karst) tree species co-occur naturally. Saplings of coniferous trees appear affected more severely in their growth and survival by cattle browsing than saplings of deciduous trees (Smit et al., 2006; Vandenbergh et al., 2007). The differences in consumption of these tree saplings tend to decrease with grazing intensity most likely as a result of decreased selectivity (Crawley, 1983; Vandenbergh et al., 2007).

Hence, the hump-backed curve between the level of grazing disturbance and facilitation strength is expected to be higher for less resistant species, especially at low disturbance. We hypothesized that the benefits of facilitation will be stronger for the less resistant coniferous species and that such benefits will change in a group-specific manner as a consequence of changes in grazing intensity. We therefore experimentally tested the group-specific relationships between a facilitative nurse shrub and the aforementioned coniferous and deciduous tree species under varying grazing intensities. Saplings were planted under and outside the canopy of R. rubiginosa in paddocks with three levels of grazing intensity: zero, low, and high. The following question was addressed: do coniferous trees benefit more from facilitation by a nurse shrub than deciduous trees (in terms of browsing frequency, survival and growth), under both low and high grazing intensities?

**Methods and materials**

**Study area**

The experiment was conducted from April 2004 until May 2005 in the Swiss Jura Mountains, at ‘La Petite Ronde’ (Les Verrières, 6°27’35”E, 46°56’18”N, altitude 1125 m a.s.l.). The climate is cold and wet, with a yearly mean temperature of 6.7 °C (±1.67) and yearly mean precipitation of 1646 mm (±424), of which 20% falls as snow (mean±1 SD over 1991–2003). Mean monthly temperature and total precipitation during the grazing season (May–September) of 2004 was 12.1 °C (±4.6) and 673 mm, respectively.

The vegetation is dominated by grasses (76%), mainly by Agrostis capillaris L. and Festuca rubra agg. L. The soils are on Argovian marls (Mühlethaler, 1930), rich in clay (30%), acidic (5.5<pH<6.0), locally hydro-morphic, and belong to the brown-soil class. No fertilizers have been applied since 1986. The mean productivity of the area for 2004 was 389.4 g m⁻².

The study area contains three sites and each site consists of two adjacent paddocks. Each year, the paddocks of each site are simultaneously grazed by cattle between May and October following a rotational grazing system with four grazing periods per year. Different grazing intensities are achieved by varying the size of the paddocks, while maintaining a stable herd size of twenty-four 2-year-old heifers.

**Experimental design**

Each of the three sites contained a large (6.9–8.2 ha) and small (3.5–4.1 ha) paddock, representing, respectively, low (83.6–106.1 LU (i.e. Livestock Units) days ha⁻¹) and high (181.7–204.1 LU days ha⁻¹) grazing intensities (Fig. 1). We fenced an exclosure (i.e. zero grazing intensity) of 10 m × 60 m at the border of each paddock. Sward heights (mean±1 SE) measured before and after four rotations correlated well with grazing intensities (low = 56.7±6.4 mm and high = 90.1±10.7 mm) (see Smit et al., 2007 for more details). In April 2004, five blocks of 10 m × 10 m were evenly distributed in each of the 6 paddocks and the zero grazing intensity exclosures (Fig. 1). In each block, we planted four multi-stemmed R. rubiginosa of 1.5 m in height as thorny nurse shrubs at a distance of 7 m from each other. One shrub each was selected as a nurse plant for one of the four tree species (coniferous: A. alba and P. abies; deciduous: A. pseudoplatanus and F. sylvatica).
deciduous: *A. pseudoplatanus* and *F. sylvatica*). Eight saplings were planted at each shrub: four saplings were planted near the shrub, i.e. at <0.1 m from the stem of the shrub (position “in”) and four saplings were planted at 1.5 m (position “out”) from the shrub, all according to the cardinal directions (N, E, S, W). In total, 1920 young saplings were transplanted to the 60 blocks. Saplings were obtained from a local nursery (Lobsigen, region of Bern, Switzerland). Prior to plantation of saplings in the field, heights and diameters at 1 cm above the root collar were measured (Table 1).

### Measurements

After each of the four grazing periods in 2004 and in May 2005, we recorded, for each sapling, whether it was browsed and whether it was still alive. Saplings without living stem (i.e. brown under the bark) as well as saplings pulled out by cattle were scored as dead. Browsing frequency and survival were calculated for each species in each block at position “in” and “out” of the shrub as the proportion (0, 0.25, 0.50, 0.75 and 1) of the four individuals browsed and still alive. All living saplings (*n* = 1367) were harvested in May 2005, before cattle arrived. Saplings were cut 1 cm above the root collar, oven-dried and weighed. The impact of browsing and nurse shrubs on the growth of surviving saplings was assessed by analysing sapling biomass change (BC) by final harvest. BC was calculated as (aboveground dry mass at harvest) / (predicted aboveground dry mass at planting)−1. BC was averaged for each shrub plant and for each position relative to the shrub plant (BC >1 represents an increase in biomass and 0 < BC < 1 a decrease in biomass). Aboveground dry mass at planting was estimated using regression models. These had been derived from measurements on randomly selected extra saplings (*n* = 40 per species; see Table 1).

### Statistical analyses

We tested for the effects of species-group (coniferous versus deciduous), grazing intensity (zero, low, high) and position (“in”, “out”) and their interactions on final browsing frequency (*n* = 240), final survival (*n* = 360) and BC (*n* = 336; unbalanced design) using a linear mixed-effects (LME) model with restricted maximum likelihood estimation. Site, with grazing intensity nested within site and block nested within grazing intensity, were specified as random effects. The use of LME models is more reliable than conventional ANOVA when analysing unbalanced data (Pinheiro & Bates, 2000).

<table>
<thead>
<tr>
<th>Species</th>
<th>Age (yr)</th>
<th>Height (cm)</th>
<th>Diameter (cm)</th>
<th>Dry mass (g)</th>
<th>log (dry mass)</th>
<th>R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abies</td>
<td>3</td>
<td>11.6 ± 0.1</td>
<td>0.262 ± 0.003</td>
<td>0.61 ± 0.01</td>
<td>0.5402 × log(D² + H) − 0.1619</td>
<td>0.666</td>
</tr>
<tr>
<td>Picea</td>
<td>2</td>
<td>12.5 ± 0.2</td>
<td>0.264 ± 0.004</td>
<td>1.04 ± 0.3</td>
<td>0.7283 × log(D² + H) + 0.0271</td>
<td>0.843</td>
</tr>
<tr>
<td>Acer</td>
<td>1</td>
<td>15.5 ± 0.2</td>
<td>0.292 ± 0.003</td>
<td>0.26 ± 0.01</td>
<td>0.9101 × log(D² + H) − 0.7246</td>
<td>0.955</td>
</tr>
<tr>
<td>Fagus</td>
<td>1</td>
<td>19.1 ± 0.2</td>
<td>0.332 ± 0.004</td>
<td>0.51 ± 0.02</td>
<td>0.8327 × log(D² + H) − 0.593</td>
<td>0.914</td>
</tr>
</tbody>
</table>

Means (± 1 SE) are given for each variable and species. Regression formulas, based on 40 extra saplings per species, were used to estimate aboveground dry mass from the height and diameter measures. For all models R² is given and *p* < 0.0001.

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Fig. 1. Experimental design: one of the three sites, consisting of two adjacent paddocks of varying size (low and high grazing intensity) and their exclosure (zero grazing intensity). Five blocks (represented by numbers) were evenly located in each paddock and exclosure. Each block (10 m x 10 m) consisted of four rose shrubs (grey triangles). One of the four tree species was randomly attributed separately to each shrub. Four saplings (dots) were planted near the shrub (i.e. position “in”) and four saplings were planted at 1.5 m from the shrub (i.e. position “out”).

Table 1. Age, initial height (*H*), diameter (*D*) at 1 cm above root collar and estimated initial aboveground dry mass (*DM*) of the transplanted saplings (*n* = 480 per species)
and the interpretation of the results is more straightforward than those of generalized linear mixed models (GLMMs) (Venables & Ripley, 2002). Assumptions of normality and homoscedasticity were fulfilled. The exclusion-data (zero grazing intensity) were never considered for analysis of the browsing frequency since the saplings within the exclosures were not browsed by cattle. Survival and growth measurements in both exclosures at each site were averaged over the same block numbers (resulting in five measurements for zero intensity at each site). We conducted an additional analysis (LME as described before) for survival and BC excluding the exclusion-data in order to test more precisely the differences between low and high grazing intensity. A posteriori contrasts were tested with Tukey HSD tests.

Furthermore, we tested the effects of browsing status (unbrowsed and browsed) and species-group on BC of individual saplings still alive at harvest under low and high grazing intensities (n = 537; unbalanced design) using the same analysis as described above. BC was log transformed to reduce heteroscedasticity. All data were analysed with R, version 2.3.0. (R Development Core Team, 2006).

Results

Browsing frequency

Differences in browsing frequency between positions “in” and “out” were larger under low grazing intensity for both coniferous and deciduous species (intensity x position, Table 2). Browsing frequency did not differ between species-groups, although the species-group-intensity interaction was marginally significant (Fig. 2, Table 2).

Table 2. Results of the linear mixed-effects models LME with the fixed effects of grazing intensity, species and position and their interactions for browsing frequency (n = 240; exclusion-data not included), survival (n = 360) and biomass change (BC; n = 336)

<table>
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<tr>
<th></th>
<th>Browsing</th>
<th>Survival</th>
<th>BC</th>
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<tbody>
<tr>
<td>Species group</td>
<td></td>
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<tr>
<td>Intensity</td>
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<td>Position</td>
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<td>Spgroup × intensity</td>
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<td>Intensity × position</td>
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<td>Spgroup × intensity × position</td>
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Browsing was evaluated after the last grazing period; survival and BC were measured at harvest (May 2005). The significance of bold interactions changes after removing the exclusion-data from the LME analysis (see Results).

ms: 0.05 < p < 0.1; ns: non-significant result. *p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001.

Fig. 2. The effects of grazing intensity (low and high) and position (in and out) on the browsing frequency (mean proportion ± 1 SE, n = 30) of coniferous and deciduous saplings, after the fourth grazing period. Different letters indicate significantly different means (Tukey post hoc comparisons within each species-group, p < 0.05).
Survival

Differences in survival between both shrub positions were largest under low grazing intensity (intensity x position, Table 2), also when exclosure-data were removed from the data-set. Furthermore, differences in survival between species-groups depended on grazing intensity (species-group x intensity, Table 2): without grazing, survival was similar for both species-groups (86–90%), whereas with grazing, survival (mean ± 1 SE) of coniferous species (46%) was lower than survival of deciduous species (66%) (Fig. 3). The species-group-intensity interaction term was no longer significant ($F_{1, 204} = 1.5, p = ns$) when exclosure-data were not considered in the analysis (bold in Table 2). For both species-groups, survival (mean ± 1 SE) outside shrubs was approximately equal under low and high grazing intensities, with $35 ± 5.3\%$ and $53 ± 6.1\%$ for evergreen...
and deciduous species, respectively. Only the survival of coniferous saplings beneath shrubs was significantly reduced under low intensity compared to zero intensity (Fig. 3).

**Biomass change of saplings alive at harvest**

Differences in BC between positions depended on grazing intensity (intensity × position, Table 2): differences between ‘in’ and ‘out’ were smallest under zero grazing intensity and similar under low and high grazing intensities (Fig. 4). When the exclosure-data were excluded from the analysis, the intensity × position interaction ($F_{1, 180} = 0.1$, $p = \text{ns}$) disappeared (bold in Table 2). However, the species-group × position interaction became significant ($F_{1, 180} = 5.3$, $p < 0.05$): only beneath the shrubs, BC was larger for coniferous saplings than for the deciduous ones (Fig. 4).

Differences in BC between species-groups depended on the browsing status (species-group × browsing status: $F_{1, 503} = 30.1$, $p < 0.0001$): coniferous saplings suffered from larger growth losses after cattle browsing (Fig. 5). Growth reduction due to browsing of the surviving saplings [$1 – \frac{\text{BC}_{\text{browsed saplings}}}{\text{BC}_{\text{unbrowsed saplings}}}^{-1}$] was 51% for coniferous and 27% for deciduous species. 40% of the coniferous and 59% of the deciduous saplings still alive at harvest was browsed.

**Discussion**

**Coniferous and deciduous sapling responses to facilitation**

As expected, the saplings of the coniferous *Abies* and *Picea* were less resistant (i.e. higher mortality and biomass loss) to cattle browsing than the *Acer* and *Fagus* saplings. Since browsing frequency did not differ between species, avoidance could not have been the main mechanism explaining this difference. The lower survival of the coniferous species was presumably, as found previously by Vandenberghe et al. (2007), due to a greater biomass off-take as a consequence of more vulnerable plant architecture (i.e. dense, large horizontal evergreen branches and canopy close to the herbaceous field layer versus less dense, erect branches) rather than higher compensation abilities of deciduous species (Chapin, Schulze, & Mooney, 1990; Hester, Millard, Baillie, & Wendler, 2004). Moreover, the compensatory growth responses of deciduous species did not differ from that of the coniferous species in a simulating browsing experiment using saplings of the same species and size (Vandenberghe, Frele´ choux, & Buttler, 2008).

Although deciduous species were initially on average 5 cm taller, and thus a priori more apparent, cattle affected browsing frequency and survival of coniferous species more rapidly than deciduous species (data not shown). The abundant and attractive fresh foliage of the coniferous species early in the season (Welch, Staines, Scott, French, & Catt, 1991) might therefore explain their lower resistance.

Shrubs indirectly facilitated the saplings by protecting them from cattle. For both species-groups, facilitation in terms of sapling browsing frequency and sapling survival was strongest at low levels of grazing intensity. The reduction in facilitation under high grazing intensities was related to increased damage of the nurse plants (Smit et al., 2007), allowing for incidental browsing of saplings under the shrubs, and it was most likely a consequence of decreased potential for selectivity by grazers (Crawley, 1983). Increased browsing on nurse shrubs will not always lead to a hump-backed response as for some nurse species it may also result in increased spinescence or a different structure leading to better protection for associated tree saplings (Garcı´a & Obeso, 2003). In contrast with the reduced facilitation at high grazing intensity in terms of browsing frequency and sapling survival, facilitation in terms of sapling growth was not reduced at high grazing intensity. This
was due to the fact that surviving saplings did not differ in growth between the low and high grazing intensities in both positions.

We expected that the less resistant coniferous saplings would benefit more from nurse shrubs than would the more resistant deciduous species. Evidence for this hypothesis was found for growth of the surviving saplings, but not for browsing frequency or survival. The surviving coniferous saplings were less browsed (20%) than the deciduous saplings, which resulted in lower overall growth loss beneath shrubs. The surprising lack of species-group response to facilitation in terms of survival is due to the fact that the less resistant coniferous species had an overall lower survival than the deciduous species, both outside and beneath nurse shrubs. These lower survival rates beneath shrubs were not related to differences in browsing frequency but rather to a greater biomass off-take (see above, Fig. 5). Moreover, facilitation of survival under high grazing intensity tended to become weaker for coniferous species than for deciduous species. Consequently, the expected increase in facilitation strength of coniferous species will most probably be reached at grazing intensities lower than the one used in this study. In other words, if shrubs would act as a perfect barrier against cattle browsing (i.e. browsing frequency equals zero), species-group responses to facilitation could become significant.

No significant facilitative or competitive interactions under zero grazing intensity were found. Other studies demonstrate how tree species react differently to modified environmental conditions (e.g. shade, competition from neighbours, nutrients, water, etc.) caused by nurse plants (Dolezal, St’astna, Hara, & Srutek, 2004; Gómez-Aparicio et al., 2005; Gómez-Aparicio, Zamora, Castro, & Hódar, 2008). Such interactions were demonstrated in more severe ecosystems (i.e. drier and warmer) than our high-productive study area. Pagès and Michalel (2003) found that the indirect positive effect of competition release for tree saplings by adult trees in a temperate hardwood forest was outweighed by the direct negative effect of light reduction. We suggest that microsites and resources (e.g. nutrients, light) inside and outside shrubs did not differ sufficiently to provoke large facilitative or competitive interactions. Thus, protection against grazing was the main indirect facilitation mechanism. Our study was based on sapling heights initially ranging between 10 and 20 cm, while other sizes of the target species might change the net outcome of the biotic interactions (Olff et al., 1999). Associational effects will also vary strongly among systems as they depend on the selectivity of the herbivores, heterogeneity and productivity of the grazed patches, and palatability of nurse plants and protégés (Baraza, Zamora, & Hódar, 2006; García & Obeso, 2003; Gómez, Hódar, Zamora, Castro, & Garcia, 2001; Gómez-Aparicio et al., 2008). In conclusion, we found that the less resistant surviving coniferous saplings did benefit more from facilitation in terms of growth than the more resistant deciduous saplings, which is in agreement with our hypothesis. In contrast, the coniferous saplings did not benefit more from facilitation in terms of overall survival due to incidental browsing by herbivores on the nurse plants and protégé sapling browsing underneath (Gómez et al., 2001; Smit et al., 2007). At high grazing intensities, coniferous saplings tended to benefit even less from facilitation.

Implications for wood-pasture systems

Most of the ancient wood-pastures suffered a dramatic decline during the last century due to changes in agricultural practices (Etienne, 1996; Manning et al., 2006). If management deviates from a balance between intensification and extensification, wood-pastures evolve towards either open pastures or closed forests with concomitant loss of biodiversity (Gillet, 2005). Periods where grazing is excluded to allow for regeneration of the tree resource or where grazing is intensified to limit natural reforestation, could only be a short-term management solution before reinstating seasonal or year-round grazing. A more sustainable and natural solution may be applying facilitation of tree recruitment via nurse plants, particularly as extensive grazing is an integral part of long-term management of wood-pastures. This study shows that the strength of facilitation decreases with increasing grazing intensity and in particular for the less resistant conifers. Within the context of conservation and restoration of wood-pastures, this means that grazing intensity needs to be kept low to promote the effectiveness of facilitation for all tree species.

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