



Post-dispersal seed predation in Patagonia temperate forest depends on habitat patchiness and seed species

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Abstract Post-dispersal seed predation is a key process regulating plant population dynamics and community composition. Because food preference (i.e., seed species selection) can interact with habitat features such as vegetation characteristics, integrating both is important for a better understanding of the processes that drive plant community structure. In order to study how forest habitat patchiness and seed species influence post-dispersal seed predation, we monitored seed predation of native common

understory plant species in Patagonia temperate forests. By performing a cafeteria-style experiment, we assessed consumption on the three most common understory seed species, in forest interior and forest gaps. We found that seed predation by rodents differed between habitats and, independently, between seed species. Seed predation was more than $2 \times$ higher in forest gaps than in forest interior, and medium-sized seed species were the least preyed-upon. Although counterintuitive, given that granivores such as rodents usually prefer sheltered habitats to forage, these results highlight the importance of site-specific variables in plant-granivore interactions.

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Introduction

Post-dispersal seed predation is a key process regulating plant population dynamics and community composition (Hulme 1998; Bricker et al. 2010; Maron et al. 2012; Hegstad and Maron 2019). Seed predation can be highly variable and influenced by multiple factors including seed predator abundance and behavior (Orrock et al. 2010), habitat characteristics, and seed traits (Kollman and Buschor 2003; García and Chacoff 2007; Moyano et al. 2019; Dylewski et al. 2020). Theory posts that while foraging, rodents should avoid low-quality food patches in favor of high-quality ones (MacArthur and Pianka 1966) and seek for safe patches while avoiding risky ones (the “foraging dilemma,” McArthur et al. 2014). On the one hand, habitats with complex structure can enhance rodents foraging activity by offering shelter from predators (Morris and Davidson 2000; Kollman and Buschor 2003). For example, habitat variables such as substrate and distance to nearest tree in Mediterranean forests (Fedriani 2005), and grass and shrub cover in temperate northern forests (Kollman and Buschor 2003) have better explained seed predation by rodents than seed phenotypic traits. In temperate forest ecosystems, understory cover and patchiness have shown to be relevant in driving seed predation patterns (Abe et al. 2001; Schnurr et al. 2004). On the other hand, seed traits such as mass (Jansen et al. 2004), size (Dylewski et al. 2020), and volume (Moyano et al. 2019) have explained rodents preference for seeds. Therefore, because food preference (i.e., seed selection) can interact with or overcome habitat features such as vegetation characteristics (Pons and Pausas 2007; Booman et al. 2009; García et al. 2011), integrating both is important for a better understanding of the processes that drive plant community structure (Larios et al. 2017).

Patchiness or forest cover variations are fundamental drivers of diversity and community dynamics in forest ecosystems (e.g., Jackson and Wong 1994; Schnurr et al. 2004; Heinemann et al. 2006; Ushio et al. 2010; Echeverria et al. 2014). Particularly, the regeneration and persistence of tree species in southern temperate forests can depend on forest-clearing

dynamics (Veblen 1985; Bustamante & Armesto 1995; Pollmann 2003). Gutiérrez et al. (2004) found that small-scale disturbances (e.g., tree-fall originated gaps) increased the heterogeneity of the forest floor, producing microsites that favor the coexistence of plants with different regeneration modes. Also, in forest gaps, seeds previous to perturbation or seeds coming from adjacent patches are important for native vegetation to recover (Armesto et al. 2001; Parkes et al. 2003; Guidetti et al. 2016). In this context, it is known that forest cover variation can alter plant-animal interactions such as seed predation (Schnurr et al. 2004; Caccia et al. 2006; Royo and Carson 2008) which can vary among habitats in response to biotic effects (e.g., direct and indirect predator cues; Sivy et al. 2011) or environmental drivers (e.g., vegetation context, Booman et al. 2009; Pons and Pausas 2007; moonlight, Kotler et al. 2010). Therefore, it is reasonable to expect that habitat change alter seed predation patterns (Diaz et al. 1999; García and Chacoff 2007), which in turn can influence forest composition and regeneration (Schreiner et al. 2000; García et al. 2005; Caccia et al. 2006).

In order to understand how forest habitat patchiness (forest interior vs. gaps) and seed species influence post-dispersal seed predation, we monitored seed predation of native common understory plant species in Patagonia temperate forests, by assessing consumption on three different native seed species in forest interior and forest gaps. Understanding how forest habitat heterogeneity affects seed predation is fundamental to understand plant community dynamics in forest ecosystems.

Methods

Study area

Our study was conducted in Llao-Llao Reserve, a 1220 ha area within Nahuel Huapi National Park in Patagonia–Argentina (41° 03′ S, 71° 30′ W), in Autumn 2005. Regional climate is humid in autumn–winter and dry in spring–summer, with 9 °C average annual temperature and 1800 mm average annual precipitation (Cabrera 1976). The native forest vegetation belongs to the Subantarctic biogeographical region (Cabrera 1976), the dominant tree species being the evergreen southern beech (*Nothofagus*

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dombeyi) and cordilleran cypress (*Austrocedrus chilensis*) (Mermoz and Martín 1986). Llao-Llao Reserve has been protected since the 1960s, but it was previously logged in certain areas, and canopy gaps of variable size have been generated by tree falls, giving the forest a patchy distribution (Amico et al. 2008). These gaps present some of the common understory vegetation dominated by the native shrub (*Aristotelia chilensis*) and native bamboo (*Chusquea culeou*) (Mermoz and Martín 1986). The main post-dispersal seed predators in the area are Cricetidae rodents (~ 25 gr.) such as the long-haired grass mouse (*Abrothrix hirta*), long-tailed mouse (*Oligoryzomys longicaudatus*), and olive grass mouse (*A. olivacea*) (Caccia et al. 2006; Nuñez et al. 2008; García et al. 2011). So far, there are no reports of scatter-hoarding rodents, and the authors found no evidence of bird seed predation (no removed soiled or bird excrements around seed depots). As the experiment was during Autumn, invertebrate or insect predation can be negligible.

Cafeteria experiment

In order to study if post-dispersal seed predation varied between forest interior and forest gaps (“habitat”) and if there was a preference for different seeds (“species”), we established a cafeteria-style experiment (Lobo et al. 2009; Pearson et al. 2014; Moyano et al. 2019). We selected six forest gap sites distributed haphazardly inside the Llao-Llao Reserve and six intact native forest interior sites, with gap and forest habitats differenced by the occurrence of tree canopy cover, forest having $> 80\%$ and gaps $< 10\%$ (see Fig. 1a for a schematic representation). As for seeds, we chose the three most common understory native species in these forests (García et al. 2011) and their seeds represent an optimal gradient of size/mass, from larger to smaller: *Schinus patagonicus* ($18.81 \text{ mm}^2 \pm 0.21 \text{ mm}^2$; $0.607 \text{ g} \pm 0.019 \text{ g}$); *Maytenus boaria* ($4.95 \text{ mm}^2 \pm 0.03 \text{ mm}^2$; $0.368 \text{ g} \pm 0.029 \text{ g}$); and *Aristotelia chilensis* ($3.25 \text{ mm}^2 \pm 0.02 \text{ mm}^2$; $0.185 \text{ g} \pm 0.017 \text{ g}$) (Supplementary Material, Figure S1). These species are representative of the understory (as pioneers of clearing colonization), unaffected by masting behavior (enabling us to extrapolate to the medium-term) and endozoochorous (thus homogenizing the functional group and its implications in expected patterns of spatial

distribution of deposition). Seeds were obtained from fruits randomly collected on plants at the study site, in order to estimate specific individual seed mass/size and to prepare a seed pool for experimental depots.

In the experiment, we offered seeds to predators in the field by attaching them to wooden popsicle sticks holding three seeds of each species (nine seeds total per stick, Fig. 1c). Seeds were fastened to the sticks in a random order, using non-toxic glue, wearing gloves to prevent human scent to impregnate them (García et al. 2011). At each forest and gap sites, we randomly placed seed depots (= set of three wooden popsicle sticks; Fig. 1b, c) at a minimum distance of 25 cm each, nailed to the ground with a wire staple over each stick center. Because understory cover is an important factor influencing seed predation rate (Caccia et al. 2009; Royo and Carson 2008), we placed seed depots under parental species shrubs. This also controls for possible differences in real seed rain densities, usually expected to be stronger under bush, than far from bush (especially in clearings; García et al. 2011). Initially, 10 seed depots were placed separated at least 30 m from each other, and sites were more than 200 m apart (Fig. 1b). We evaluated seed predation after 48 h of installing the experiment, a period comparable with previous studies in several environments (Hulme 1994; Kollman et al. 1998; Hulme & Borelli 1999; Orrock 2015). Both the seeds removed from the popsicle sticks and those damaged (with obvious bite marks) but remaining in place were considered as predated.

Statistical analyses

To determine if seed predation (response variable) differed between “habitats” (“forest interior,” “forest gaps”), we used generalized linear mixed models (GLMM) (Fig. 1). Seed predation was calculated as the proportion of predated seeds after 48 h. To evaluate if there was a preference for seed “species,” we included it as a predictive variable, with factors “Small” (*A. chilensis*), “Medium” (*M. boaria*), and “Large” (*S. patagonicus*). We also considered the interaction between factors, in order to test if potential differences between species depended on the habitat type. We assumed a binomial distribution, using a GLMM based on Laplace approximation and a logit link function (lme4 package, *glmer* function, Bates et al. 2015). Since our experimental design had

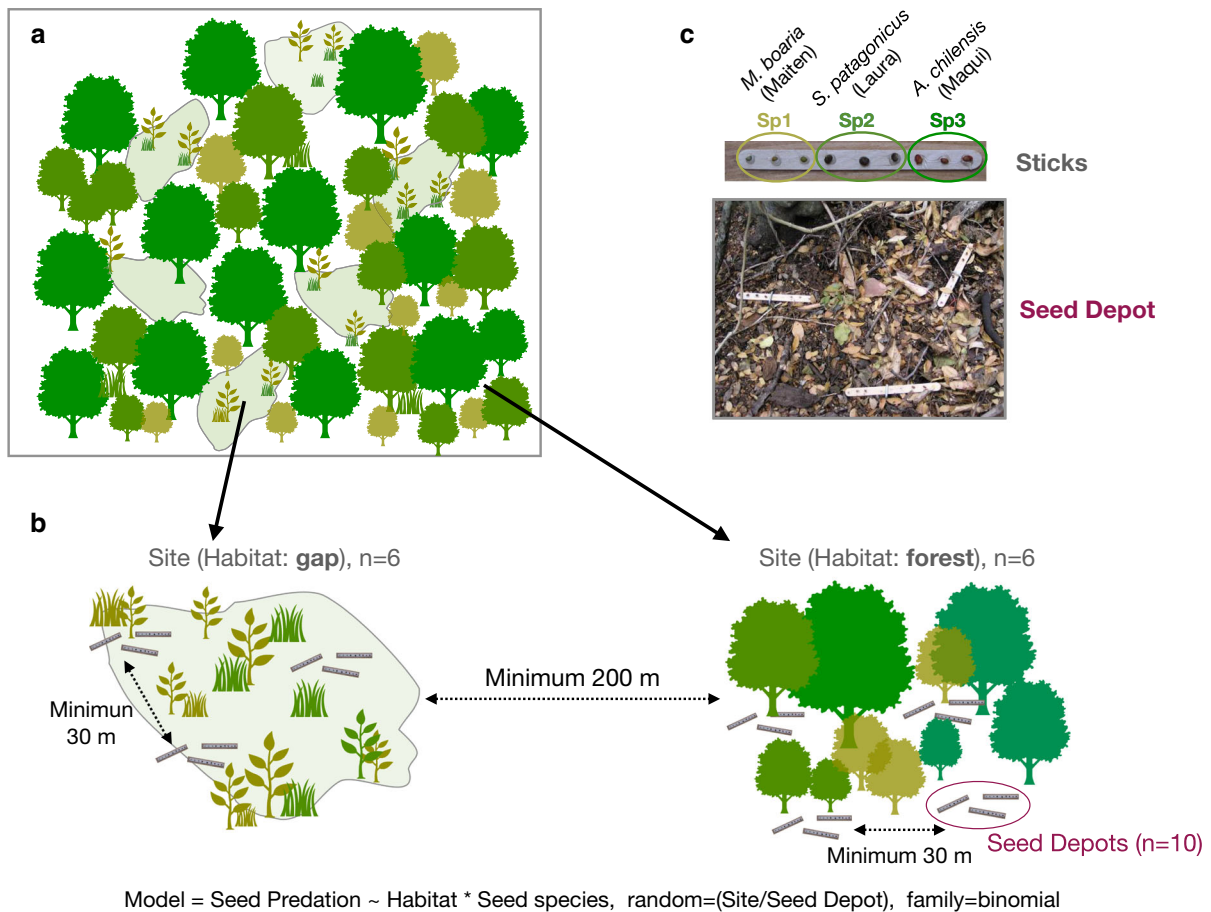


Fig. 1 Schematic representation of experimental design, and model construction (please see “Methods” section): **a**, **b** Forest gap array, number of sites, replicates, and model description; **c** popsicle sticks with seeds attached and depot arrangement

different gaps immersed in a large native forest, we tested and corroborated there were no differences among gap sites using a factorial analysis (Table S1). Finally, since sticks within each depot are pseudoreplicates, we used “seed depot” nested in “sites” as a random variable (Fig. 1). During monitoring, we found variable numbers of seed depots (minimum $n = 3$, maximum $n = 10$; blown, broken, or lost), but GLMMs contemplate uneven number of pseudoreplicates. To study the amount of total variation explained by each model, we used analysis of deviance (*pseudo* r^2 , BaylorEdPsych package; Beaujean 2012). Additionally, we performed a False Discovery Rate (FDR) post hoc test (Benjamini and Hochberg 1995) to compare the proportion of predation among seed species. All analyses were performed using R 3.5.0 (R Development Core Team 2018).

Results and discussion

We found that seed predation by rodents differed between habitats and, independently, between seed species, as shown by the non-significant interaction between factors (Table 1; Fig. 3b; Table S2). Seed predation was more than $2 \times$ higher in forest gaps than in forest interior ($P < 0.001$, Table 1; Fig. 2).

Table 1 Anova of global factors’ effects and GLMM results

Global fixed effects	Chisq	Df	<i>P</i> value	(pseudo) r^2
Habitat	9.945	1	0.001	0.40
Seed species	11.355	1	0.003	
Habitat*seed species	0.742	2	0.689	

Statistically significant values are in bold

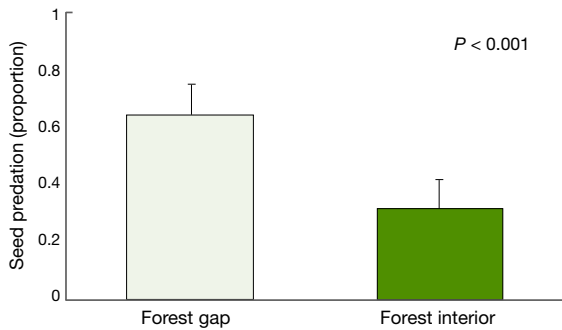


Fig. 2 Seed predation was higher in forest gaps. Proportion of seed predation in forest gap and forest interior after 48 h of field exposure. Letters mean significant difference between treatments ($p < 0.05$); bars represent means $\pm$ standard error

This result may seem counterintuitive, given that higher seed consumption in areas such as gaps would contradict “predation fear” behavior (Bleicher 2017). Several studies show evidence that rodents prefer to forage in sheltered habitats providing refuge from predators (Kollman and Buschor 2003; Yang et al. 2016; Zhang et al. 2016). For example, Germain et al. (2013) showed that seed predation varied spatially as seed predation decreased with decreasing vegetation cover. Yet, this behavior might vary among individuals (McArthur et al. 2014) and also can be influenced by the context (Steele et al. 2015) and the spatial scale considered (García et al. 2011). For instance, the ability of some species to accurately perceive changes in predation risk (Sundell et al. 2004) and the presence of other factors constraining foraging behavior (e.g., strong intra e interspecific competition; Yunker et al. 2002; Dupuch et al. 2014) might lead rodents to forage in riskier habitats. Maybe the fact that the ‘seeds are there (in the sticks) make them more visible and available for the rodents. Because of the short period that seeds were exposed (Díaz et al. 1999), we assume that consumers were efficient in finding the seeds offered. Why rodents make the tradeoff of searching for good food in risky places is probably related to the fact that good food in safe places is harder to find (McArthur et al. 2014). On the other hand, although forest gaps from our study almost lacked tree canopy cover, they did present understory vegetation (see *Study area* section), which has shown to enhance seed predation rates (Kollman and Buschor 2003), as rodents suffer higher predation risk in areas with reduced vegetation cover of low height (Booman et al. 2009; Pons and Pausas 2007). Such a positive effect on

seed predation has been in fact, previously reported for bamboo patches in forest gaps of the temperate Patagonian region (Caccia et al. 2006). Complementarily, habitat differences may emerge from a higher availability of fruits and seeds in forest gaps compared to forest interior, leading to positive responses among seed predators (García et al. 2011). Thus, although the present study does not enable us to discern a specific mechanism, we assume that both perception of risk and resource availability are underpinning the present habitat effects on seed predation.

Besides higher predation in gaps than in forests, we also found that the proportion of predated seeds depended on seed identity rather than on seed size (Table 1, Fig. 3a, b). The biggest seed species (*S. patagonicus*) was 41% and 17% more predated than medium-sized seed species (*M. boaria*) ($P < 0.05$) and the smaller ones, *A. chilensis* ($P = 0.305$),

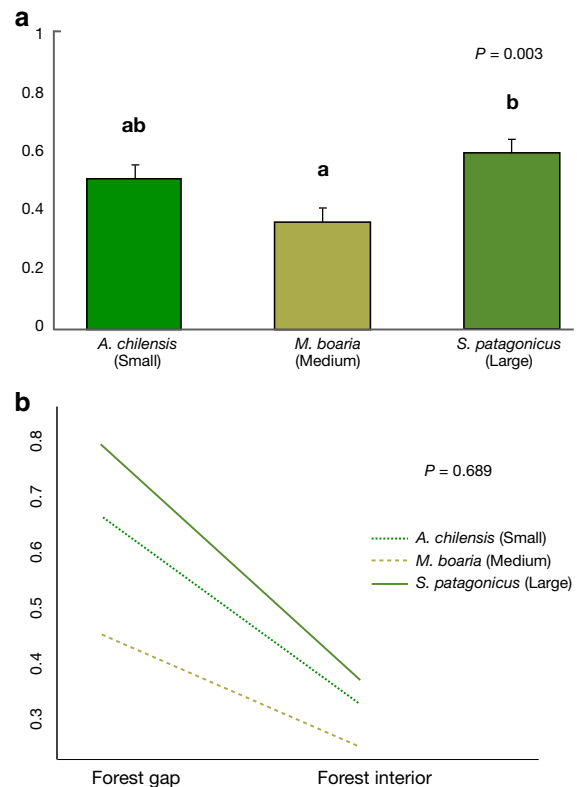


Fig. 3 *S. patagonicus* and *A. chilensis* were more predated regardless of habitat. **a** Average predation (proportion) of each seed species during the experiment. Letters mean significant difference among treatments ($p < 0.05$); bars represent means $\pm$ standard error; **b** Non-significant interaction between seed species and habitat

respectively (Fig. 3; Supp. Mat., Figure S1, Table S3). In addition, we found no differences on the proportion of predated seeds between *M. boaria* and *A. chilensis* ($P = 0.101$, Fig. 3; Figure S1, Table S3). Although seed traits promoting foraging behavior of rodents are controversial (Dylewski et al. 2020), several authors found that rodents prefer larger and heavier seeds (Nuñez et al. 2008; Carrillo-Gavilán et al. 2010; Chen et al. 2017; Wang and Ives 2017). However, consumption differences in our experiment mostly emerged between *S. patagonicus* and the intermediate-sized *M. boaria*, suggesting that size is not the only seed trait determining rodent choices. In any case, by making such “choices,” rodents can generate inter-specific differences in recruitment potential and influence forest regeneration dynamics (García et al. 2005; Larios et al. 2017 and references therein; Hegstad and Maron 2019; Moyano et al. 2019). Whether seed predation by rodents will finally leave an imprint in the composition of forest gaps will ultimately depend on the specific responses of seed species to other post-dispersal forces (drought, frost, and light tolerances, e.g., Manríquez et al. 2016; Promis and Allen 2017).

Our results on habitat and seed species effects on seed survival are based on a short-term, single estimation of seed predation, precluding somehow our ability to infer long-term and large-scale predictable patterns (see also Caccia et al. 2006). This is especially true in the case of interspecific differences in seed predation, which may be affected by the occurrence of seed masting events, especially from the highly erratic and low-frequency masting tree (e.g., *Nothofagus dombeyi*) or understory species (e.g., *Chusquea culeou*; Kitzberger et al. 2007). In spite of this, none of these plant species was masting in the year of our study, suggesting that our results may be at least extrapolated to the non-masting years. Regarding seasonal variability, it is also known that differential seed predation may change according to the variable proportion of different species in the seed rain or to increasing rodent densities (e.g., Díaz et al. 1999; but see Kollmann et al. 1998). In our case, we set up our experiment in the co-occurring peak of the fruiting season of the three fleshy-fruited plants under study, and thus our findings relate to the maximum potential densities of these seed species in the field. Concerning the spatial extent of our findings, we consider it to represent one of the main environmental conditions in forest ecosystems: forest vs. gaps. In fact, our

additional factorial analysis revealed that predation rates were similar across gaps (Table S1), suggesting that the strong inter-habitat differences found here are generalized across the forest landscape.

Understanding how foraging activity of post-dispersal seed predator changes according to habitat patchiness and seed species identity is essential given their influence on forest composition and its regeneration process (Côté et al. 2003; Caccia et al. 2006; Bricker et al. 2010; Hegstad and Maron 2019). Yu et al. (2014) tested whether rodent seed predation or dispersal was beneficial for gap regeneration, and found that scatter-hoarding rodents rarely retrieved seeds from forest gaps, suggesting that rodent seed predation patterns contributed to the regeneration of the dominant species in gaps. In our case, the higher seed predation found in forest gaps might negatively impact on the recruitment of seedlings and slow down the forest regeneration of certain species. Our study then remarks the importance of considering species identity, given the fact that our results cannot be explained based on seed mass/size, and reinforce the idea that factor-associated habitat use by rodents at multiple spatial scales are important in mediating composition and regeneration of temperate southern forest communities.

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Availability of data and material See Supplementary Material.

Declarations

Conflict of interest The authors declare that they have no conflict of interest.

Consent to participate All persons entitled to authorship have been so named.

Consent for publication All authors have approved its submission for publication in Plant Ecology.

References

- Abe M, Miguchi H, Nakashizuka T (2001) An interactive effect of simultaneous death of dwarf bamboo, canopy gap, and predatory rodents on beech regeneration. *Oecologia* 127:281–286
- Amico GC, García D, Rodríguez-Cabal MA (2008) Spatial structure and scale-dependent microhabitat use of endemic “tapaculos” (Rhinocryptidae) in a temperate forest of southern South America. *Ecol Austral* 18:169–180
- Armesto JJ, Díaz I, Papic C, Willson MF (2001) Seed rain of fleshy and dry propagules in different habitats in the temperate rainforests of Chiloé Island, Chile. *Ecol Austral* 26:311–320
- Bates D, Mächler M, Bolker B, Walker S (2015) Fitting linear mixed-effects models using lme4. 67: 48. <https://doi.org/10.18637/jss.v067.i01>
- Beaujean A (2012) BaylorEdPsych: R package for Baylor University educational psychology quantitative courses. R package version 0.5. R Foundation for Statistical Computing, Vienna, Austria. <http://CRAN.R-project.org/package=BaylorEdPsych>
- Benjamini Y, Hochberg Y (1995) Controlling the false discovery rate: a practical and powerful approach to multiple testing. *J R Stat Soc Series B* 57:289–300
- Bleicher SS (2017) The landscape of fear conceptual framework: definition and review of current applications and misuses. *Peer J* 5:e3772
- Booman GC, Laterra P, Comparatore V, Murillo N (2009) Post-dispersal predation of weed seeds by small vertebrates: interactive influences of neighbor land use and local environment. *Agric Ecosyst Environ* 129:277–285
- Bricker M, Pearson D, Maron J (2010) Small-mammal seed predation limits the recruitment and abundance of two perennial grassland forbs. *Ecology* 91:85–92
- Bustamante R, Armesto JJ (1995) Regeneration dynamics in canopy gaps of a montane forest of Chiloé Island, Chile. *Rev Chil Hist Nat* 68:391–398
- Cabrera, AL (1976) Regiones fitogeográficas argentinas. In: Kugler WF (ed) *Enciclopedia argentina de agricultura y jardinería*. 2nd edn. Acme, Buenos Aires, p 85
- Caccia FD, Chaneton EJ, Kitzberger T (2006) Trophic and non-trophic pathways mediate apparent competition through post-dispersal seed predation in a Patagonian mixed forest. *Oikos* 113:469–480
- Caccia FD, Chaneton EJ, Kitzberger T (2009) Direct and indirect effects of understorey bamboo shape tree regeneration niches in a mixed temperate forest. *Oecologia* 161:771–780
- Carrillo-Gavilán MA, Lalagüe H, Vilà M (2010) Comparing seed removal of 16 pine species differing in invasiveness. *Bio Inv* 12:2233–2242
- Chen Q, Tomlinson KW, Cao L, Wang B (2017) Effects of fragmentation on the seed predation and dispersal by rodents differ among species with different seed size. *Integr Zool* 12:468–476
- Côté M, Ferron J, Gagnon R (2003) Impact of seed and seedling predation by small rodents on early regeneration establishment of black spruce. *Can J for Res* 33:2362–2371
- Díaz I, Papic C, Armesto JJ (1999) An assessment of post-dispersal seed predation in temperate rain forest fragments in Chiloé Island, Chile. *Oikos* 87:228–238
- Dupuch A, Morris DW, Ale SB, Wilson DJ, Moore DE (2014) Landscapes of fear or competition? Predation did not alter habitat choice by Arctic rodents. *Oecologia* 174:403–412
- Dylewski Ł, Ortega YK, Bogdziewicz M, Pearson DE (2020) Seed size predicts global effects of small mammal seed predation on plant recruitment. *Ecol Lett*. <https://doi.org/10.1111/ele.13499>
- Echeverría ME, Sottile GD, Mancini MV, Fontana SL (2014) Nothofagus forest dynamics and palaeoenvironmental variations during the mid and late Holocene, in southwest Patagonia. *Holocene* 24:957–969
- Fedriani JM (2005) Do frugivorous mice choose where or what to feed on? *J Mamm* 86:576–586
- García D, Chacoff NP (2007) Scale-dependent effects of habitat fragmentation on hawthorn pollination, frugivory, and seed predation. *Conserv Biol* 21:400–411
- García D, Obeso JR, Martínez I (2005) Rodent seed predation promotes differential recruitment among bird-dispersed trees in temperate secondary forests. *Oecologia* 144:435–446
- García D, Zamora R, Amico GC (2011) The spatial scale of plant-animal interactions: effects of resource availability and habitat structure. *Ecol Monogr* 81:123–139
- Germain RM, Johnson L, Schneider S, Cottenie K, Gillis EA, MacDougall AS (2013) Spatial variability in plant predation determines the strength of stochastic community assembly. *Am Nat* 182:169–179
- Guidetti BY, Amico GC, Dardanelli S, Rodríguez-Cabal MA (2016) Artificial perches promote vegetation restoration. *Plant Ecol* 217:935–942
- Gutiérrez AG, Armesto JJ, Aravena JC, (2004) Disturbance and regeneration dynamics of an old-growth North Patagonian rain forest in Chiloé Island, Chile. *J Ecol* 92:598–608
- Hegstad RJ, Maron JL (2019) Productivity and related soil properties mediate the population-level consequences of rodent seed predation on Blanketflower, *Gaillardia aristata*. *J Ecol* 107:34–44
- Heinemann K, Kitzberger T (2006) Effects of position, understorey vegetation and coarse woody debris on tree regeneration in two environmentally contrasting forests of north-western Patagonia: a manipulative approach. *J Biogeogr* 33:1357–1367
- Hulme PE (1994) Post-dispersal seed predation in grassland: its magnitude and sources of variation. *J Ecol* 82:645–652
- Hulme PE (1998) Post-dispersal seed predation: consequences for plant demography and evolution. *Perspect Plant Ecol Evol Syst* 1:32–46
- Hulme PE, Borelli T (1999) Variability in post-dispersal seed predation in deciduous woodland: relative importance of location, seed species, burial and density. *Plant Ecol* 145:149–156
- Jackson ST, Wong A (1994) Using forest patchiness to determine pollen source areas of closed-canopy pollen assemblages. *J Ecol* 82:89–99
- Jansen PA, Hemerik L, Bongers F (2004) Seed mass and mast seeding enhance dispersal by a neotropical scatter-hoarding rodent. *Ecol Monogr* 74:369–389

- Kitzberger T, Chaneton EJ, Caccia F (2007) Indirect effects of prey swamping: differential seed predation during a bamboo masting event. *Ecology* 88:254–2554
- Kollmann J, Buschor M (2003) Edges effects on seed predation by rodents in deciduous forests of northern Switzerland. *Plant Ecol* 164:249–261
- Kollmann J, Coomes DA, White SM (1998) Consistencies in post-dispersal seed predation of temperate fleshy-fruited species among seasons, years and sites. *Func Ecol* 12:683–690
- Kotler BP, Brown J, Mukherjee S, Berger-Tal O, Bouskila A (2010) Moonlight avoidance in gerbils reveals a sophisticated interplay among time allocation, vigilance and state-dependent foraging. *Proc R Soc B* 277:1469–1474
- Larios L, Pearson DE, Maron JL (2017) Incorporating the effects of generalist seed predators into plant community theory. *Funct Ecol* 31:1856–1867
- Lobo N, Duong M, Millar JS (2009) Conifer-seed preferences of small mammals. *Can J Zool* 87:773–780
- MacArthur RH, Pianka ER (1966) On optimal use of a patchy environment. *Am Nat* 100:603–609. <https://doi.org/10.1086/282454>
- Manríquez MT, Mestre L, Lencinas MV, Promis Á, Pastur GM, Soler R (2016) Flowering and seeding patterns in pure and mixed Nothofagus forests in Southern Patagonia. *Ecol Process* 5:21
- Maron JL, Pearson DE, Potter T, Ortega YK (2012) Seed size and provenance mediate the joint effects of disturbance and seed predation on community assembly. *J Ecol* 100:1492–1500
- McArthur C, Banks PB, Boonstra R, Forbey JS (2014) The dilemma of foraging herbivores: dealing with food and fear. *Oecologia* 176:677–689
- Mermoz M, Martín C (1986) Mapa de vegetación del Parque y la Reserva Nacional Nahuel Huapi. Delegación Regional Patagonia, Bariloche, Argentina.
- Morris DW, Davidson DL (2000) Optimal foraging mice match patch use with habitat differences in fitness. *Ecology* 8:2061–2066
- Moyano J, Chiuffo MC, Nuñez MA, Rodríguez-Cabal MA (2019) Seed predation does not explain pine invasion success. *Oecologia* 189:981–991
- Nuñez MA, Simberloff D, Relva MA (2008) Seed predation as a barrier to alien conifer invasions. *Bio Inv* 10:1389–1398
- Orrock JL, Holt RD, Baskett ML (2010) Refuge-mediated apparent competition in plant–consumer interactions. *Ecol Lett* 13:11–20
- Orrock JL, Borer ET et al (2015) A continent-wide study reveals clear relationships between regional abiotic conditions and post-dispersal seed predation. *J Biogeogr* 42:662–670
- Parkes D, Newell G, Cheal D (2003) Assessing the quality of native vegetation: the ‘habitat hectares’ approach. *Ecol Manag Restor* 4:S29–S38
- Pearson DE, Hierro JL, Chiuffo M, Villarreal D (2014) Rodent seed predation as a biotic filter influencing exotic plant abundance and distribution. *Bio Inv* 16:1185–1196
- Pollmann W (2003) Stand structure and dendroecology of an old-growth Nothofagus forest in Conguillio National Park, south Chile. *For Ecol Manag* 176:87–103
- Pons J, Pausas JG (2007) Rodent acorn selection in a Mediterranean oak landscape. *Ecol Res* 22:535–541
- Promis A, Allen RB (2017) Tree seedlings respond to both light and soil nutrients in a Patagonian evergreen-deciduous forest. *PLoS ONE* 12:e0188686
- R Core Team (2018) R: A language and environment for statistical computing. Foundation for Statistical Computing. <http://www.Rproject.org/>
- Rodríguez-Cabal MA, Aizen MA, Novaro AJ (2007) Habitat fragmentation disrupts a plant-disperser mutualism in the temperate forest of South America. *Biol Conserv* 139:195–202
- Royo AA, Carson WP (2008) Direct and indirect effects of a dense understory on tree seedling recruitment in temperate forests: habitat-mediated predation versus competition. *Can J For Res* 38:1634–1645
- Schnurr JL, Canham CD, Ostfeld RS, Inouye RS (2004) Neighborhood analyses of small-mammal dynamics: impacts on seed predation and seedling establishment. *Ecology* 85:741–755
- Schreiner M, Bauer EM, Kollmann J (2000) Reducing predation of conifer seeds by clear-cutting *Rubus fruticosus* agg. in two montane forest stands. *For Ecol Manag* 126:81–290
- Sivy KJ, Ostoja SM, Schupp EW, Durham S (2011) Effects of rodent species, seed species, and predator cues on seed fate. *Acta Oecol* 37:321–328
- Steele MA, Rompre G, Stratford JA, Zhang H, Suchocki M, Marino S (2015) Scatterhoarding rodents favor higher predation risks for cache sites: the potential for predators to influence the seed dispersal process. *Integr Zool* 10:257–266
- Sundell J, Dudek D, Klemme I, Koivisto E, Pusenius J, Ylönen H (2004) Variation in predation risk and vole feeding behaviour: a field test of the risk allocation hypothesis. *Oecologia* 139:157–162
- Ushio M, Kitayama K, Balser TC (2010) Tree species-mediated spatial patchiness of the composition of microbial community and physicochemical properties in the topsoils of a tropical montane forest. *Soil Biol Biochem* 42:1588–1595
- Veblen T (1985) Forest development in tree-fall gaps in the temperate rain forests of Chile. *NGS* 1:162–183
- Wang B, Ives AR (2017) Tree-to-tree variation in seed size and its consequences for seed dispersal versus predation by rodents. *Oecologia* 183:751–762. <https://doi.org/10.1007/s00442-016-3793-0>
- Wilson SD, Keddy PA (1986) Species competitive ability and position along a natural stress/disturbance gradient. *Ecology* 67:236–242
- Yang Y, Zhang M, Yi X (2016) Small rodents trading off forest gaps for scatter-hoarding differs between seed species. *Forest Ecol Manag* 379:226–231
- Yu F, Shi X, Wang D, Wang T, Yi X, Lou Y (2014) Seed predation patterns favor the regeneration of dominant

species in forest gaps compared with the understory in an oak-pine mixed forest. *Acta Theriol* 59:495–502

Yunger JA, Meserve PL, Gutiérrez JR (2002) Small-mammal foraging behavior: mechanisms for coexistence and implication for population dynamics. *Ecol Monogr* 72:561–577

Zhang Y, Yu J, Sichilima AM, Wang W, Lu J (2016) Effects of thinning on scatter-hoarding by rodents in temperate forest. *Integr Zool* 11:182–190

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