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9	INFLUENCE SMALL MAMMAL COMMUNITIES IN GRASSLANDS
10	
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31 ABSTRACT

Ecosystem/habitat restoration has become a major goal of international biodiversity policy. 32 However, restorations are often limited in space or time, and we know little on whether and how 33 34 restoration and management affect vertebrates. Here we assessed the local and landscape-scale effects of habitat restoration and management on small mammal communities in the Egyek-35 Pusztakócs marsh system (Hortobágy National Park, Hungary), site of the largest active restoration 36 of grasslands on former croplands in Europe. We live-trapped mice, voles and shrews in spring and 37 autumn in 2 years (four sampling periods) at two sites in six habitat types: croplands, grasslands 38 restored three to six years earlier, and natural grasslands. Data on 421 individuals of 12 species 39 showed that restored grasslands were similar to croplands and natural grasslands in species richness, 40 abundance and composition. At the local scale, management influenced abundance because there 41 were more small mammals in unmanaged and early-mown grasslands with taller vegetation than in 42 late-mown or grazed grasslands with lower vegetation, or in ploughed croplands. Elevation was also 43 important because sites at higher elevation provided refuges during spring floods or summer 44 droughts. At the landscape scale, the proportion of restored and natural grasslands positively 45 affected the abundance of small mammals, whereas the proportion of linear habitats (roads, canals) 46 had a negative effect on abundance. Our results show that management is more important than 47 48 restoration per se at the local scale, which is expected for habitat generalists such as small mammals in contrast to specialists such as plant-feeding invertebrates. However, restoration provides 49 landscape-scale benefits by increasing the area of grasslands which can serve as refuges for small 50 mammals in unfavourable periods. We thus conclude that a mosaic of restored and appropriately 51 managed grasslands with tall vegetation will provide the best chances for the persistence of small 52 53 mammal communities in dynamic landscapes.

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Key-words: community assembly, ecosystem services, grassland restoration, local extinction,
 population dynamics, recolonisation, rodent, shrew

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58 INTRODUCTION

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Agriculture and other human land uses have transformed the surface of Earth, resulting in the 60 61 decline, degradation and fragmentation of natural habitats (Lubchenco et al., 1991). The restoration of ecosystems/habitats can theoretically counter these processes (Young, 2000) but is rarely 62 implemented at spatial and temporal scales that are adequate to make a difference (Lengvel *et al.*, 63 2014; Woodcock et al., 2010). Restoration has recently been incorporated as explicit targets into 64 relevant policy instruments (e.g. Aichi Biodiversity Targets, EU Biodiversity Strategy for 2020), yet 65 our knowledge on the design, implementation and impact of restoration remains limited in several 66 aspects. Habitat restoration is typically followed up by monitoring vegetation development, and 67 studies of trophic groups other than plants are scarce (Brudvig, 2011; Mortimer et al., 1998; 68 Woodcock et al., 2008). This is surprising because many invertebrate and vertebrate animals are 69 important for ecosystem functioning and a full evaluation of restoration success needs to integrate 70 higher trophic levels (animals) (Longcore, 2003; Young, 2000). Despite these calls, the number of 71 72 studies of restoration effects on vertebrates is still low and most of them focus on birds in forests and shrubs (e.g. Brawn, 1998; Germaine and Germaine, 2002; Hoover, 2008; Machmer, 2002). 73 74

Small mammals (voles, mice and shrews) play important roles in ecosystems. Most voles and mice 75 are herbivores or granivores and several species hoard plant seeds for storage. Small mammals may 76 thus alter plant species composition (Howe and Brown, 2000; Torre et al., 2007) and provide 77 ecosystem services such as seed dispersal (Mohammadi, 2010; Schnurr et al., 2004; Williams et al., 78 2001) and pollination (Vieira et al., 2009). Subterranean small mammals play a role in maintaining 79 soil structure (Medin and Clary, 1989). Finally, small mammals are important in grassland food 80 webs, both as consumers or top-down regulators of vegetation and arthropods (e.g. most shrews are 81 insectivores), and as a food resource for reptiles, birds and mammals (Castién and Gonsálbez, 1999; 82 Mohammadi, 2010; Torre et al., 2007; Vieira et al., 2009). 83

The proportion of grasslands decreased considerably due to agriculture, and grasslands have 85 become the focus of many conservation actions in Europe (Hedberg and Kotowski, 2010; Kiehl et 86 87 al., 2010) and North America (Gerla et al., 2012). The impact of grassland restoration on small mammals has been addressed in a handful of studies in North America. Stone (2007) found that 88 restoration led to a short-term decline in the number of captures, total biomass and species richness 89 of small mammals, followed by a partial recovery 3-5 years after restoration. A comparison of a 90 91 successional gradient from prairie to forest (Moro and Gadal, 2007) showed that time since restoration (abandonment) did not directly affect small mammals and that their abundance was 92 93 highest in middle, rather than early or late, successional stages, where the structural diversity of vegetation was highest. Finally, in the most comprehensive study of grassland restoration and small 94 mammals to date, Mulligan (2012) found that the colonization of restored fields occurred rapidly 95 and was positively related to the connectivity of habitats, and that restored grasslands served as 96 refuges during regional declines. Only the latter study involved a landscape context, which should 97 98 be addressed and considered in practice (Brudvig, 2011; Lengyel et al., 2014) because local studies 99 may overlook larger-scale factors such as refuges and dispersal corridors that may influence restoration outcome. 100

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The aim of this study was to evaluate the local and landscape-scale effects of habitat restoration and 102 management on small mammal communities after a large-scale restoration of grasslands on former 103 croplands. We used a chronosequence design (space-for-time substitution) to sample small 104 mammals on natural grasslands (restoration target), restored grasslands (restoration process) and 105 106 croplands (restoration start). We then analysed the data to evaluate the effects of local factors (management, elevation) and landscape configuration (proportion of natural grasslands, restored 107 grasslands, croplands, woodlands, and linear habitats such as roads and canals) on the species 108 richness, abundance, and composition of small mammal communities. We further tested whether 109

110 small mammal communities change with time since restoration, and also evaluated the links

between the dynamics of small mammal communities and the spatial and temporal changes in

112 habitats due to weather, restoration and management.

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114 METHODS

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116 The Egyek-Pusztakócs marsh system and the sampling design

Our study site was the Egyek-Pusztakócs marsh and grassland complex (4073 ha; N 47.54-47.62°, E 20.86-20.99°) in Hortobágy National Park (eastern Hungary). Grassland restoration was carried out in 2005-2008 on a total of 760 ha of former cropland by sowing of two seed mixtures after ploughing, followed by mowing or grazing as post-restoration management. To our knowledge, this is currently the largest non-spontaneous grassland restoration in Europe. More details on restoration and vegetation development are given in Lengyel et al. (2012) and on changes in arthropod communities in Déri *et al.* (2011) and Rácz *et al.* (2013).

124

We sampled small mammals (mice, voles and shrews) at 12 sites in three habitat types (Table 1): 125 126 two on croplands (wheat fields, starting point of restoration), eight on restored grasslands (process of restoration) and two on natural grasslands (restoration targets). Within restored grasslands, four 127 levels of restoration age (3- to 6-year-old restorations) were replicated at two sites each (n = 8128 sites). Sampling was conducted in the spring and autumn of 2011 and 2012 (four sampling periods 129 total). Spring 2011 was characterized by floods after snowmelt, whereas the spring and summer 130 were relatively dry (total precipitation: 350 mm). Significant precipitation in winter led to intense 131 132 floods in spring 2012 followed by a long drought (270 mm) between May and September (Fig. S1). Sampling sites were at least 800 m away from each other to minimise spatial non-independence. 133 The sampled habitat patches ranged from 16 to 300 hectares in area [mean 66.6 ha \pm standard 134 deviation (SD) 81.68]. None of the response variables (see below) showed significant correlations 135

136	with patch area (Spearman rank correlations, n.s.), therefore, we did not control for it in statistical
137	analyses. Sites were chosen to represent the management practices characteristic to the general area.
138	Restored grasslands chosen were managed either by mowing early in the season (June, $n = 4$ sites),
139	by mowing late in the season (August, $n = 2$) or by grazing by sheep (from April to November, $n =$
140	2) (Lengyel et al., 2012). The two natural grasslands sampled were not managed, whereas the two
141	croplands were regularly ploughed. We also recorded elevation and vegetation height at each site.
142	Although the differences in elevation were small (89-91 m a.s.l.), lower-lying areas were more often
143	flooded and less suitable for small mammals than were higher plateaus. Elevation of the sites was
144	determined from a detailed topographic map (accuracy: 0.5 m). We also measured the height of
145	vegetation (accuracy 5 cm) 0.5 m from each trap in a random direction.
146	
147	To characterise landscape configuration, we calculated the proportion of five habitat types
148	(croplands, restored grasslands, natural grasslands, woodlands, and linear habitats such as roads and
149	canals) at each sampling site. We used a habitat map prepared in 2010 of the entire Egyek-
150	Pusztakócs system as part of other studies and used ESRI ArcMap GIS 10.0 to calculate the
151	proportion of each habitat type in non-overlapping buffers of 400-m radius around each sampling
152	site.
153	
154	Small mammal sampling
155	We live-trapped small mammals using Sherman live traps (H. B. Sherman Traps, Tallahassee,
156	Florida, U.S.). Sherman traps have been repeatedly found to perform as well or, more frequently,
157	better than other live traps in multi-species studies of small mammals (e.g. Anthony et al., 2005;
158	Belant and Windels, 2007; Hayes et al., 1996; Kaufman and Kaufman, 2007; Morris, 1968; Santos-
159	Filho et al., 2006; Torre et al., 2010), and were found to cause less mortality than other live traps
160	(Dizney et al., 2008). Because Sherman traps may be less effective at capturing small shrews than

other live traps (Sibbald *et al.*, 2006), we performed all analyses with or without data on shrews

(i.e., mice and voles only). However, as the inclusion or exclusion of shrews led to qualitatively 162 similar results, we presented results based on the full dataset. Although Sherman traps are 163 particularly effective in combination with other types of traps (e.g. pitfall traps, Longworth traps) to 164 165 attain a full complement of small mammals found in a habitat (Sibbald et al., 2006), our study did not aim to find all species, rather, it aimed to characterise differences among habitat and 166 management types by an adequately high, appropriately replicated constant sampling effort 167 designed to achieve a reasonable compromise between the requirements of spatiotemporal 168 replication and on-site effort intensity. 169

170

At each site, we installed 36 traps in a 6×6 grid covering 1 hectare (distance between traps: 20 m). 171 We baited the traps with seed mixture (grains, sunflower) and bacon following the 172 recommendations of the trap manufacturer and our previous experience. Traps were operated for 173 five nights to ensure compatibility with the National Biodiversity Monitoring System of Hungary. 174 Five nights appeared appropriate because captures peaked on day 2-4 and decreased by day 5 (Fig. 175 176 S2). Total sampling effort amounted to 8640 trap-nights (36 traps×5 nights×12 sites×4 sampling periods). The traps were checked twice a day (7:00-9:00, 18:00-20:00 hours). Before each check, 177 we counted potential avian predators in and 200 m around the sampling grids for 5 minutes from a 178 distance. Small mammals captured in the traps were identified to species and weighed. We marked 179 individuals by a permanent felt pen at the base of the tail to enable the detection of recaptures. Our 180 previous experience showed that this marking could be detected for at least a week. 181

182

183 Statistical analysis

Response variables were the number of species (species richness), and the number of individuals (abundance) of small mammals (rodents, i.e., mice and voles, and shrews). For both response variables, data from traps at one site (n = 36 traps) during one sampling period (n = 5 nights) were pooled to obtain one datapoint per site and per period (total n = 48). Exploratory analyses revealed a

highly significant interactive effect of year and season on both species richness and abundance 188 (richness, $F_{1,33} = 43.918$, p < 0.0001; abundance, $F_{1,33} = 36.242$, p < 0.0001) because richness and 189 abundance increased substantially from spring to autumn in 2011 but not in 2012, while the effect 190 of location was not significant (p > 0.32). To evaluate the effects of local and landscape-scale 191 variables on the species richness and abundance of small mammals while also considering this 192 temporal fluctuation, we constructed generalized linear mixed-effects models (GLMMs) with 193 194 season (spring/autumn) nested within year as a random effect. GLMMs allow the incorporation of 195 nested random effects even when variances are unequal and within-group errors are correlated, and are robust to unbalanced designs (Crawley, 2007). We fitted GLMMs using the 'glmer' function of 196 197 R with Poisson error distribution and log link function, as recommended for count data (R Core Team, 2014). Local predictor variables were elevation (in m) and management (early mowing, late 198 mowing, grazing on restored grasslands, no management on natural grasslands, ploughing on 199 croplands). Landscape-scale predictor variables were the proportion of five habitat types in 400-m 200 buffers around the sampling sites. The five landscape-scale variables were not correlated (Pearson 201 correlation, n.s.). To allow for testing whether predictor variables had similar or different effects in 202 the four sampling periods, we also used sampling period as a fixed effect with four levels. 203

204

To test the effect of restoration age on small mammals, we repeated the GLMMs using data only 205 from restored grasslands. In these models, both the local and landscape-scale predictors described 206 above were included, and we also included the time since restoration (in years) to test whether and 207 how small mammal communities change after restoration. In all GLMMs, we started with the full 208 model including all main effects and their interactions with sampling period and removed non-209 210 significant terms in a backward stepwise algorithm (removal if $p \ge 0.1$) to obtain minimum adequate models. We then used the reduced model to estimate coefficients for predictors. For post-211 hoc comparison of management, a categorical variable, we specified contrasts to compare each 212 management type to regularly ploughed croplands (restoration start) in the analysis of all sites, and 213

to grazing in the analyses restricted to restored sites. In additional statistical tests, data were log-

transformed (log x + 1) for parametric tests (one-way ANOVA), if necessary, to meet the

assumption of homogeneous variances. All statistical tests were implemented in R (version 3.1.1.).

217

218 RESULTS

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In total, we captured 421 individuals of 12 species (full list of species and number of individuals captured are given in **Table S1**). We detected 20 individuals of three species in the spring and 359 individuals of 12 species in the autumn of 2011. In 2012, we caught 24 individuals of five species in the spring and 18 individuals of eight species in the autumn. Recapture rate was on average 9% (n = 465 captures total) and ranged between 0 and 30% per site (n = 12). Recaptured individuals were counted only once in all statistical analyses.

226

227 Effects on species richness of small mammals

GLMMs showed that species richness was little influenced by the local and landscape factors 228 studied. There was a weak interaction between sampling period and management (GLMM, z =229 2.178, p = 0.029), likely because the number of species decreased less from autumn 2011 to spring 230 2012 in late-mown restorations than in other habitat types (Fig. 1A). Furthermore, there were 231 marginally non-significant effects of sampling period (z = -1.907, p = 0.057) and the interaction 232 between sampling period and elevation (z = 1.915, p = 0.056). The latter was mostly because small 233 mammals disappeared from autumn 2011 to spring 2012 in three of four sites under 90 m a.s.l. and 234 only one individual was found in the fourth site. In areas at or above 90 m, at least one individual 235 was caught in five of eight sites (total n = 23 individuals), suggesting higher chances for survival 236 during the harsh late winter and spring flooding in early 2012. Neither the remaining local and 237 landscape variables nor their interactions with sampling period influenced species richness 238 significantly. 239

241 Factors influencing small mammal abundance

At the local scale, the abundance of small mammals was significantly influenced by management, 242 243 elevation, and interactions between sampling period and management and between sampling period and elevation (Table 2). Significant coefficient estimates showed that early-mown restorations had 244 more individuals, whereas late-mown and grazed restorations had fewer individuals than ploughed 245 croplands (Table 2), particularly when abundance was generally high (autumn 2011, Fig. 1B). The 246 interaction between sampling period and management was because abundance decreased more from 247 2011 to 2012 in early-mown restorations than in late-mown restorations (Fig. 1). The interaction 248 between sampling period and elevation was because the relationship between elevation and 249 abundance was negative in autumn 2011 and slightly positive or constant in the three periods of 250 251 lower abundance (Fig. 2).

252

In autumn 2011, when abundance was generally high, non-managed natural grasslands had taller 253 vegetation (mean $5.8 \pm S.D. 2.98$ cm, n = 72 traps) than either early-mown (3.6 ± 2.18 , n = 144), 254 late-mown $(3.5 \pm 1.96, n = 72)$ or grazed restored grasslands $(3.2 \pm 1.99, n = 72)$ (one-way ANOVA 255 on log-transformed data, $F_{3,356} = 12.060$, p < 0.0001; Tukey HSD-test, p < 0.001 for each of the 256 three comparisons), and thus probably provided better chances to avoid predators than did restored 257 grasslands. This was supported by a negative correlation between average vegetation height and the 258 number of predators in autumn 2011 (Pearson r = -0.695, n = 10 grasslands, p = 0.026). Predators 259 most often recorded at the sites included Marsh Harrier Circus aeruginosus, Common Buzzard 260 Buteo buteo, Kestrel Falco tinnunculus and Great Egret Egretta alba, and other potential predators 261 262 observed in the vicinity were Red Fox Vulpes vulpes, Least Weasel Mustela nivalis, European polecat Mustela putorius and Steppe Polecat M. eversmanii. 263

At the landscape scale, the proportion of natural and restored grasslands influenced abundance 265 positively, whereas the proportion of linear habitats had a negative effect on abundance (Table 2). 266 None of the interaction terms between landscape variables and either sampling period or 267 268 management influenced abundance significantly. 269 **Differences by restoration age** 270 In the smaller sample of restored fields (n = 8), restoration age strongly influenced species richness 271 (GLMM, z = 29.020, p < 0.0001) because sites restored in 2005 had fewer species (1.3 ± SE 0.74) 272 than sites restored in later years (2006: 2.3 ± 0.67 , 2007: 2.3 ± 0.80 , 2008: 1.9 ± 0.99). Restoration 273 age also influenced the abundance of small mammals (Table 2), however, no consistent temporal 274 trend could be observed (2005: 5.4 ± 4.54 individuals, 2006: 13.3 ± 5.54 , 2007: 6.3 ± 2.82 , 2008: 275 8.4 ± 6.18). 276

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Although the spatial distribution of captures was rather consistent across periods for common species, species composition varied greatly in the four sampling periods due to rare species and there was no sign of a clear separation in species composition either by habitat type, management or restoration age (**Fig. S3**).

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283 DISCUSSION
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At the local scale, management had the largest influence on small mammals. The management effect was mediated by vegetation height because sites with taller vegetation had more species and individuals than sites with lower vegetation. Elevation was also important because higher areas provided better chances for survival during unfavourable periods such as floods. At the landscape scale, small mammals were influenced positively by the proportion of natural and restored grasslands and negatively by the proportion of linear habitats. These results suggest that restored

grasslands can be important landscape-level refuges, although when conditions are favourable, e.g. in autumn 2011, local management and not restoration or time since restoration *per se* appears primarily important in shaping small mammal communities in grasslands. Our study is the first to provide evidence of both local and landscape-scale factors influencing small mammal assemblages after grassland restoration. The patterns found support the view that the effects of population fluctuations and restoration are difficult to disentangle and that restorations play an unexpected but important role as refuges during regional declines of small mammals (Mulligan, 2012).

298

Several results showed that restoration provided habitats that were comparable or better than croplands for small mammals. Only three restorations and one cropland had small mammals after the spring 2011 bottleneck, and only restorations and natural grasslands had small mammals after the extreme summer drought of 2012. Moreover, the proportion of restored grasslands positively affected small mammal abundance after the spring flood in 2011 and the summer drought of 2012. These results show that in only three to six years, restoration can lead to grasslands that serve as refuges for small mammals during unfavourable periods.

306

Our results also suggest that the effect of management was mediated by vegetation height. 307 Vegetation cover was highest in unmanaged areas, and was low in mown or grazed sites in autumn 308 2011, and predators were more numerous in sites with lower vegetation. These patterns may explain 309 the differences in small mammal abundance because vegetation provides safe microsites for the 310 movement, hiding, feeding or food gathering activities of small mammals (Moro and Gadal, 2007). 311 The risks of predation on small mammals both by birds and mammals are probably higher when 312 vegetation cover is low because small mammals can be more easily spotted and captured by 313 predators at sites with poor vegetation cover (Sutherland and Dickman, 1999). In extremely dry 314 years such as 2012, areas with higher vegetation cover probably also provided more abundant food 315 than areas which are mowed or grazed. A study in Switzerland found the highest density of small 316

mammals in non-mown grasslands (Aschwanden *et al.*, 2007), whereas an African study found that
intensive grazing negatively affects the abundance and diversity of small mammals (Yarnell *et al.*,
2007). Consequently, the differences in abundance in relation to management found here are likely
to exist due to the mediating effect of vegetation cover.

321

322 Our results showed that the large temporal, weather-driven fluctuations typical in small mammal communities (Butet et al., 2006; Merritt et al., 2001) may be modified by both local effects 323 (elevation, management) and landscape-scale effects (proportion of restored/natural grasslands). 324 Almost all species disappeared from lower-lying areas (< 90 m a.s.l.) during the winter, whereas 325 populations of some species survived in higher areas. Higher areas may thus be particularly 326 important refuges for several small mammal species during harsh winters and springs with 327 extensive floods (Wijnhoven et al., 2005). During the winter, small mammal populations typically 328 decrease to a minimum due to the combined effects of predation, cold weather, food shortage or 329 competition for resources (Aars and Ims, 2002; Hansen et al., 1999). The chances of successful 330 survival during the winter are highest in refuge patches where vegetation cover is high enough and 331 food supply is adequate (van Deventer and Nel, 2006). Our observations followed these patterns 332 because species richness and abundance increased with the proportion of restored or natural 333 grasslands around the sites in each spring period, but they also provided further insight. For 334 example, the connectivity of patches, as estimated by the proportion of linear habitats, negatively 335 affected small mammal richness in spring 2012, likely due to the higher exposure of individuals to 336 predators. These patterns suggest that homogeneous patches that are rich in natural and restored 337 grasslands with no management or early mowing and that are poor in croplands, roads or canals 338 339 thus can serve as refuges and can mitigate the fluctuations of small mammals.

340

The lack of a direct, local effect of restoration on small mammals is likely to be related to their
broad food spectrum and life history traits (fast reproduction, good dispersal ability, quick

colonisation after local extinction etc.). The studied restoration led to the acceleration of secondary 343 succession in vegetation development towards the target natural grasslands, with grassland-344 dominated vegetation forming as early as Year 2 after restoration (Lengyel et al., 2012). The 345 346 species composition of arthropods (orthopterans, true bugs, spiders, carabid beetles) also changed considerably from Year 1 to Year 2 because species characteristic to target grasslands appeared in 347 greater numbers in Year 2 and after (Déri et al., 2011). In only four years, the species richness of 348 orthopterans (crickets and grasshoppers) has doubled and their abundance increased nearly ten-fold 349 on restored grasslands compared to croplands (Rácz et al., 2013). The restoration of phytophagous 350 insects, most of which specialize on certain plants, positively correlated with the restoration success 351 of the vegetation in restored calcareous and mesotrophic grasslands (Woodcock et al., 2010), 352 indicating that more specialized animal groups may more closely follow the vegetation restoration 353 process. The generally broad food spectrum and the life history characteristics of small mammals, 354 in combination with their weather-driven population dynamics, make it likely that the restoration of 355 small mammal communities does not correlate well with the restoration process of vegetation or 356 other animal taxa. 357

358

In conclusion, habitat management was primarily important in shaping small mammal communities 359 at the local scale and restoration had little effect locally. Management by mowing and grazing 360 decreased vegetation height, which increased predation risks on small mammals. However, 361 grassland restoration provided benefits to small mammals at the landscape scale by increasing the 362 total area of grasslands, which provide better conditions for the persistence of small mammals 363 during unfavourable periods. Grassland restoration and management should thus re-create a mosaic 364 of habitat types that combines restored and natural vegetation and no management in at least a 365 portion of the landscape to maximise the chances of persistence for small mammals. The policy 366 implications of our study are that both the local effects and landscape configuration should be 367 considered in the design of restoration and management, and in the monitoring of biodiversity and 368

- 369 ecosystem services. The restoration of small mammals as important providers of ecosystem services
- 370 should also consider the inherent spatiotemporal dynamics of populations that operates at the
- 371 landscape scale and that can be managed locally.
- 372

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497 TABLES

	Habitat Year		Area	Local factors		Landscape factors: Proportion of				
Sampling site	type	restored	(ha)	Elevation (m)	Management	Cropland	Grassland	Restoration	Linear	Woodland
Sóút É szántó	cropland	_	60	90.5	ploughing	0.99	0.00	0.00	0.00	0.00
Kilátó szántó	cropland	_	50	90.0	ploughing	0.56	0.29	0.00	0.02	0.00
27 ha-os gyep	restoration	2005	27	89.5	mowing early	0.22	0.21	0.32	0.01	0.01
Hagymás-hát	restoration	2005	26	91.0	sheep-grazing	0.00	0.08	0.36	0.00	0.00
Górés észak	restoration	2006	48	90.0	mowing early	0.00	0.03	0.86	0.02	0.04
Nyírházi halom	restoration	2006	28	91.0	mowing early	0.00	0.24	0.67	0.02	0.04
Csepregi szántó	restoration	2007	46	89.0	mowing late	0.00	0.06	0.60	0.00	0.00
K Csepregi	restoration	2007	21	90.0	mowing late	0.00	0.36	0.41	0.01	0.01
Csattag ÉK	restoration	2008	28	89.5	mowing early	0.00	0.00	0.59	0.00	0.00
Görbeszék-halom	restoration	2008	16	90.0	sheep-grazing	0.06	0.47	0.35	0.00	0.02
Csattag-hát	grassland	_	300	89.0	none	0.00	0.77	0.00	0.00	0.00
Fekete-rét DK	grassland	_	150	90.0	none	0.00	0.99	0.00	0.01	0.00

Table 1. General characteristics, local and landscape factors of sampling sites. Grassland: natural grasslands, restoration: restored grasslands, linear
 habitats: roads, canals.

Table 2. Results of generalized linear mixed-effects models testing the effects of management,

- elevation and landscape variables on abundance of small mammals (random effect: season within
- year) in all sites and in restored grasslands only. Significant effects are highlighted in Bold.

Response	Species richness	Estimate ± S.E.	z value	Pr(> z)
Abundance	(Intercept)	79.52 ± 37.444	2.124	0.034
(all sites)	Sampling period	-44.31 ± 16.942	-2.616	0.009
	Management _{None}	-0.08 ± 0.560	-0.140	0.888
	Management _{Early-mown}	1.99 ± 0.417	4.762	< 0.0001
	Management _{Late-mown}	-2.20 ± 0.514	-4.275	< 0.0001
	Management _{Grazed}	-1.80 ± 0.677	-2.659	0.008
	Elevation	-0.89 ± 0.417	-2.106	0.035
	Proportion of grassland	1.85 ± 0.817	2.269	0.023
	Proportion of restoration	1.59 ± 0.703	2.262	0.023
	Proportion of linear habitats	-35.78 ± 11.824	-3.026	0.002
	$Period \times Mgmt_{None}$	0.07 ± 0.204	0.344	0.731
	Period × Mgmt _{Early-mown}	-0.64 ± 0.186	-3.452	0.001
	Period × Mgmt _{Late-mown}	$\textbf{0.88} \pm \textbf{0.208}$	4.238	< 0.0001
	$Period \times Mgmt_{Grazed}$	0.24 ± 0.285	0.830	0.407
	Period × Elevation	$\boldsymbol{0.49 \pm 0.188}$	2.609	0.009
Abundance	Sampling period	-34.76 ± 17.135	-2.028	0.043
(restored sites)	Management _{Early-mown}	2.11 ± 0.395	5.343	< 0.0001
	Management _{Late-mown}	-0.97 ± 0.460	-2.100	0.036
	Elevation	-0.36 ± 0.430	-0.831	0.406
	Restoration age	-0.47 ± 0.146	-3.204	0.001
	Proportion of cropland	3.56 ± 1.670	2.132	0.033
	Proportion of linear habitat	29.24 ± 14.529	2.012	0.044
	Period × Mgmt _{Early-mown}	-0.68 ± 0.173	-3.914	< 0.0001
	Period × Mgmt _{Late-mown}	$\boldsymbol{0.57 \pm 0.188}$	3.006	0.003
	Period × Elevation	$\textbf{0.39} \pm \textbf{0.190}$	2.027	0.043

507 FIGURE LEGENDS

508

Figure 1. Mean \pm S.E. species richness (A) and abundance (B) in unmanaged natural grasslands, early-mown, late-mown and grazed grassland restorations and ploughed croplands in four sampling periods. Datapoints were jittered along the X axis for clarity.

512

Figure 2. Relationships between abundance of small mammals and elevation. Spring datapoints were jittered along the X axis for clarity and lines are for visual guidance only (please see statistics in Table 2).









523	SUPPORTING INFORMATION
524	Additional Supporting Information may be found in the online version of this article at the
525	publisher's web-site.
526	(see below)
527	
528	
529	
530	SUPPORTING INFORMATION
531	
532	to accompany "Local habitat management and landscape-scale restoration influence small mammal
533	communities in grasslands" by Mérő et al.
534	
535	Contents:
536	WEATHER INFORMATION: Fig. S1
537	SUPPORTING INFORMATION RESULTS
538	Number of daily captures: Fig. S2
539	Species and individuals captured: Table SI
540	Changes in species composition: Fig. S3
541	
542	
543	WEATHER INFORMATION
544	The study site is characterized by a continental climate with large annual fluctuations in weather
545	The study site is characterised by a continental chinate with large annual fuctuations in weather. The mean annual temperature is 0.5 °C and the mean total annual precipitation is 550 mm. We
540	sampled small mammals in the spring (March April) and autumn (Sontember October) of both 2011
547	and 2012 (four periods total) Spring 2011 was characterised by high water and large flooded areas
540	due to extremely high precipitation in 2010 (Fig. S1: total 2010 precipitation in Hortohágy region:
550	950 mm source: Hungarian Meteorological Service, http://www.met.hu). Summer 2011 was dry
551	(total precipitation: 350 mm) and water receded gradually from all grasslands. Winter 2011/2012
552	again brought significant precipitation (Fig. S1) and intense flooding at snowmelt. Summer 2012
553	had a long drought (total precipitation 270 mm), with hardly any rain in August and September
554	(Fig. S1).



Figure S1. The number of hours per day with precipitation assembled from hourly present weather reports (days/months on the horizontal and hours of the day on the vertical axis) from the closest weather station with detailed measurement (Debrecen airport, c. 50 km E from the study site). Colour coding: green - rain; blue - snow, orange - thunderstorms, light blue shaded area - climate normals (average probability of precipitation per day). Bars at the top of the graphs indicate days with any precipitation (green) or no precipitation (white). Dark red horizontal bars in 2011 and 2012 indicate the sampling periods, data for 2010 and 2013 are shown for comparison only. Source: WeatherSpark (http://www.weatherspark.com).

568 SUPPORTING INFORMATION RESULTS

569

570 Number of daily captures571

The number of individuals captured per day was highest on day 4 (croplands and restored
grasslands) or on day 3 (natural grasslands) and decreased by day 5 in each of the three habitat
types (Fig. S2).



596 597

Figure S2. Mean \pm S.E. number of individuals captured on day 1 (white columns) through day 5 (black column) in three habitat types based on n = 421 individuals, with recaptures (n = 44) excluded (A), and proportion of all captures falling on day 1 through 5 in the four sampling periods (B).

602

603 Species and individuals captured

604

Table S1. Number of males, females and juveniles by species caught in the four sampling periods.

Species	Year	Season	Males	Females (gravid/lactating)	Juveniles (males/females)		
Source analysis	2011	spring autumn	3	4			
sorex araneus	2012	spring autumn	1	1			
S minutus	2011	spring autumn	1	3 (1)			
S. minutus	2012	spring autumn		2			
Crocidura leucodon	2011	spring autumn	12	1 (1) 47 (7)	1 (0/1)		
	2012	spring autumn			1 (0/1)		
C suaveolens	2011	spring autumn	2	5 (2)			
	2012	spring autumn		1			
Apodemus agrarius	2011	spring autumn	15	16	3 (1/2)		
	2012	spring autumn	3				
A. flavicollis	2011	spring autumn	1				
	2012	spring autumn		1			
A. sylvaticus	2011	autumn	7	6(1)	5 (3/2)		
	2012	autumn		1			
A. uraliensis	2011	autumn	2	2			
	2012	autumn		2 1			
Mus spicilegus	2011	autumn	2 6	6			
	2012	spring autumn	2	1			
Micromys minutus	2011	autumn		3			
	2012	autumn	0	0.(2)			
Microtus arvalis	2011	autumn	8 35 7	9 (3) 100 (47) 8 (1)	58 (12/46)		
	2012	autumn	/	8 (1) 3 (1)	6 (2/4)		
M. subterraneus	2011	autumn		3	1 (1/0)		
	2012	autumn	10	10 (4)	0		
Total	2011	autumn	10 84 12	10 (4) 193 (58) 12 (1)	0 68 (17/51)		
	2012	autumn	2	12 (1) 9 (1)	7 (2/5)		

607 Changes in species composition

We compared the small mammal species composition of restored and natural grasslands and
croplands by non-metric multidimensional scaling (NMDS) using Sørensen similarity as
implemented in the 'metaMDS' function of the R package 'vegan' (Oksanen et al. 2011).

611 612

Ordination based on NMDS showed that species composition varied greatly through the four sampling periods (**Fig. S3**). There was no sign of a clear separation of species composition of the sites either by habitat type or management. The extent of the scatter of centroids confirmed that species composition was most diverse in autumn 2011, in the period of highest abundance, followed by 2012 autumn and the two spring periods (**Fig. S3**).

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619

Figure A3. Changes in species composition of small mammals on restored grasslands (indicated by year of restoration and management), croplands and natural grasslands in four sampling periods from spring 2011 to autumn 2012. Symbols are omitted for clarity and arrows indicate temporal trends (trajectories) at the 12 sampling sites. Ordination was conducted by non-metric multidimensional scaling using Sørensen similarity based on the presence/absence of species (stress: 0.130).