

# Trophic plasticity of the endemic Japanese weasel in a lowland agricultural landscape

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## Abstract:

The conversion of natural ecosystems into human-modified landscapes (HMLs) is the main driver of biodiversity loss in terrestrial ecosystems, particularly the loss of large predators. Their demise can alter food webs substantially, sometimes releasing smaller carnivores, such as members of the Mustelidae. Nevertheless, even small carnivores must adapt to anthropogenic impacts on food availability, altering their resource use. In this context, the crops grown in agrarian habitats can profoundly affect community assembly. Here, we conducted dietary analysis on 75 Japanese weasel (*Mustela itatsi*) scats, collected between July 2017 and August 2018, to determine their seasonal food habits in a landscape dominated by rice paddy fields in Saitama prefecture, eastern Japan. From spring to autumn, Japanese weasels consumed predominantly (semi-)aquatic and terrestrial animal taxa, specifically invasive crayfish (*Procambarus clarkii*), insects (e.g., Coleoptera and Odonata) as well as adult anurans, which are all readily available prey species. In winter, Japanese weasels consumed predominantly fruit (e.g., figs, *Ficus carica*), with a relative decrease in combined animal content in scats, due to the scarcity of animal prey in dried-out paddy fields and irrigation ditches. Although frugivory is unusual in *Mustela* species diets, our findings demonstrate that Japanese weasels are capable of adaptive trophic plasticity, enabling them to survive atypical resource conditions in paddy field habitats. To enhance broad efforts to conserve *Mustela itatsi* in Japan, we recommend the diversification of rice paddy monocultures and encourage winter flooding to increase the availability of aquatic and semi-aquatic animal prey.

**Keywords:** agricultural landscape, paddy fields, Mustelidae, frugivory, human modified landscapes (HMLs), trophic plasticity.

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## Short title

Diets of the Japanese weasel in an agricultural lowland

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26 **Key words:** agricultural landscape, frugivory, paddy field, human modified landscapes  
27 (HMLs), Mustelidae, trophic plasticity.

## 28 **Introduction**

29 The conversion of natural ecosystems into human-modified landscapes (HMLs) is the  
30 main driver of biodiversity loss in terrestrial ecosystems (Foley et al., 2005; Tucker et  
31 al., 2018). However, the consequences of habitat degradation arising from agricultural  
32 land conversion remain poorly understood (Magioli et al., 2019). Apex predators are  
33 often the first to be lost from HMLs because their extensive home ranges are sensitive  
34 to habitat loss and fragmentation (Dirzo et al., 2014; Tucker et al., 2018). When apex  
35 predators are lost, the ecological release (Ritchie and Johnson, 2009) of subordinate  
36 predators results in 'winner and loser' replacements (Filgueiras et al., 2021) causing the  
37 alteration or loss of trophic interactions that disrupt local ecological networks (Hanski,  
38 2005; Valiente-Banuet, 2015; Galiana et al., 2022). Those predators that can persist in or  
39 colonize HMLs are usually more generalist and often omnivorous, opportunistic species  
40 with broader ecological niches (Fleming and Bateman, 2018; Magioli et al., 2019). In  
41 this context, the crops grown in agrarian habitats can profoundly determine community  
42 assembly according to a trade-off between productivity and disturbance (Gorzynski et  
43 al., 2021).

44 In the absence of large mammalian predators (Order Carnivora) from HMLs  
45 (Ordiz et al., 2021), small-medium sized carnivores (generally defined as  $\leq 21.5$  kg; Do  
46 Linh San et al., 2022) are elevated to the highest remaining trophic level in depauperate  
47 food webs (Gehrt et al., 2010; Marneweck et al., 2021; 2022). Nevertheless, their  
48 population dynamics and life history traits remain fundamentally different to those of

50 larger carnivores (Wallach et al., 2015). Due to energy lost in trophic level transitions  
51 (Shurin and Seabloom, 2005), small carnivores predate toward the broad base of trophic  
52 pyramids (Norrdhal and Kopimaki, 2000; Barbier and Loreau, 2019), and typically  
53 affect primary consumers, such as rodents (Lambin, 2017) and invertebrates. In turn,  
54 this can affect primary producer communities (e.g., plant biomass, regeneration,  
55 pollination, and seed dispersal) through trophic cascade effects (Hamback et al., 2004).  
56 Ultimately, however, pyramids topped by smaller carnivores are less steep and diverse  
57 than unperturbed systems (Duffy, 2003; Marneweck et al., 2022), with lower ecosystem  
58 functionality (Duffy et al., 2007). Furthermore, even smaller carnivores must adapt their  
59 resource use (i.e., food choice and habitat use), spatial organization, and life-histories to  
60 continue to persist in habitats altered substantially by human activities (Gehrt et al.,  
61 2010; Fischer et al., 2012; Rosalino et al., 2014). A better understanding of small  
62 carnivore adaptability, especially their trophic plasticity, is therefore vital for  
63 maintaining residual ecosystem functionality in HMLs and for planning future  
64 conservation management (Macdonald et al., 2017; Marneweck et al., 2021; 2022).

65 Rice (*Oryza sativa*) crops are grown extensively across Asia (Bandumula,  
66 2018), often as a monoculture that structurally alters former natural habitats and  
67 ecological communities (Katayama et al., 2015). Rice cultivation in Japan dates back  
68 over 2000 years (Verschuer and Cobcroft, 2016), with rice paddy fields now covering c.  
69 2,335,000 hectares (Ministry of Agriculture, Forestry and Fisheries, 2024a), producing  
70 approximately 7,165,000 tons of rice per year (Ministry of Agriculture, Forestry and  
71 Fisheries, 2024b). Across Japan, these extensively modified farmland areas lack large  
72 predators, but do support a guild of medium-sized generalist carnivores, including  
73 raccoon dogs (*Nyctereutes procyonoides*), invasive raccoons (*Procyon lotor*) and  
74 introduced masked palm civets (*Paguma larvata*), although they are less suitable for

76 Japanese martens (*Martes melampus*) and Japanese badgers (*Meles anakuma*) (Ohdachi  
77 et al., 2015).

78 The much smaller (110–600 g, with male-biased sexual size dimorphism;  
79 Masuda and Watanabe, 2015) Japanese weasel, *Mustela itatsi*, an endemic mustelid, is  
80 also distributed broadly over agricultural lowlands and urbanized areas across Honshu,  
81 Shikoku, and Kyusyu Islands of Japan (Masuda and Watanabe, 2015). The Japanese  
82 weasel population has gradually declined across Japan, especially in lowlands (Sasaki et  
83 al., 2014), resulting in their ‘Near Threatened’ IUCN Red List status (under A2, A3 and  
84 A4), equivalent to a decline of c. 25% over the last three generations (Kaneko et al.,  
85 2016). Furthermore, populations in western Japan face competition with the invasive  
86 Siberian weasel (*M. sibirica*) (Sasaki et al., 2014). The Japanese weasel is thus a species  
87 of urgent conservation concern in lowland HMLs (Kaneko et al., 2016; Suzuki, 2018b).

88 Typically, weasel species, such as *Mustela nivalis*, *M. erminea*, and *M. frenata*  
89 (now *Neogale frenata*), rely mostly on predating rodent prey in wooded habitat  
90 (McDonald et al., 2000; Zub et al., 2008; Vaca-León et al., 2019). In its natural hilly or  
91 mountain habitats, however, the Japanese weasel exhibits wider trophic plasticity than  
92 other weasel species, hunting in terrestrial and riverine habitats and consuming not only  
93 small mammals (i.e., mice, voles, shrews, and moles) but also aquatic or semi-aquatic  
94 prey (e.g., herptiles, fish, insects, crustaceans and anurans: Fujii et al., 1998; Suda et al.,  
95 2014). Furthermore, they occasionally supplement their diet with seeds, fruits, and  
96 berries (Kaneko et al., 2009; 2013; Okawara et al., 2014). This diet is similar to that of  
97 the larger (500–1500g) western polecat (*M. putorius*) that also predaes both terrestrial  
98 and aquatic prey (e.g., Hammershoj et al., 2004; Lode, 1997; Sainsbury et al., 2020), as  
99 well as occasionally eating fruits (Santos et al., 2009).

100 In this study, we conducted faecal analysis to investigate how trophic plasticity

enables Japanese weasels to survive in rice paddy monoculture farmland in Saitama prefecture, eastern Japan. Here, intense agricultural activities, habitat modifications, water pollution, and the introduction of invasive species (Fujioka and Lane, 1997; Takeuchi, 2010; Tsunoda et al., 2010) affect food web structure and integrity, and rodents are uncommon due to being controlled (Lorica et al., 2020; Singleton et al., 2021) and limited by summer flooding (Aplin et al., 2006). We hypothesize that Japanese weasels may (i) consume substantial quantities of (semi-)aquatic prey (e.g., anurans and fish as well as introduced red-swamp crayfish, *Procambarus clarkii*), and (ii) sustain themselves by consuming eclectic food sources in winter, when other prey are less available due to the drainage of rice paddy fields. Finally, we use our findings to make recommendations for improving the conservation value of rice paddy fields in Japan for weasels and broader biodiversity.

## Materials and methods

### Study area

The study area, located in Kazo-shi, Saitama, Japan, was approximately 1.2 km<sup>2</sup> of agricultural land entirely covered by paddy fields, with a few small buildings but no forest cover (36°04'N, 139°32'E; Fig. 1 and Fig. S1 in Supporting information, SI). In addition, a few fig trees (*Ficus carica*) were cultivated along the side of a local floriculture farm (Figs. S1 and S2 in SI).

There was one paved road (Prefectural route 38) through this area, with a traffic volume of av. 12,466 cars per day (Saitama prefecture, 2017). This region has a humid-temperate climate (Köppen climate classification: Cfa) with hot, humid summers and cold, dry winters. Average monthly temperature and total precipitation during the

126 study period (July 2017 to August 2018) ranged from 2.5°C in January to 28.2°C in  
127 August and from 11.0 mm in February to 453.0 mm in October (Automated  
128 Meteorological Data Acquisition System at Kuki station; data obtained from Japan  
129 Meteorological Agency, <http://www.jma.go.jp/jma/menu/menureport.html>, accessed 27  
130 May 2020). Rice was planted in mid-May, with harvesting completed by mid-November  
131 (see Figs. S3 and S4 in SI). There were many small irrigation ditches between paddy  
132 fields, which dried up during the fallow season (i.e., between December to April),  
133 making aquatic prey scarce (see Fig. S4 in SI). In this area, raccoon dogs and introduced  
134 masked palm civets, as well as invasive raccoons also occurred at low densities  
135 (author's unpublished camera-trapping data) and potentially competed with Japanese  
136 weasels in the predatory guild; however, the Japanese marten and Japanese badger were  
137 absent (Saitama prefecture, 2018).

### 138 **Field survey protocol**

139 To collect Japanese weasel scats, we used all farmland roads and tracks (i.e., c. 60%  
140 paved and 40% gravel roads, c. 5 m wide) between paddy fields, as well as an adjacent  
141 margin of c. 2 m, as our survey transect, following the general scat detection protocol  
142 (Martinoli et al. 2001; Zhou et al., 2013; Lei et al., 2023). This amounted to a total  
143 tortuous transect distance of ca. 5.4 km (represented as thin lines in the polygon  
144 depicted in Fig. 1). We repeated these surveys three to four times a month from July  
145 2017 to August 2018. Japanese weasel scats were easy to differentiate from other  
146 candidate sympatric carnivore species (mentioned above) based on their diminutive size  
147 (mean diameter of scats sampled = 5.6 (S.D. 1.2) mm: standardized size  $\leq$  8.7 mm from  
148 Tsuji et al., 2011) and appearance. Japanese weasel scats were collected and sealed in  
149 plastic bags, then taken to the laboratory and frozen at  $-20$  °C.

## Faecal analysis

Scat samples were rinsed through a sieve (0.5 mm mesh) with water and sorted under a  $\times 10$  magnifying lens. All food items were dried in an incubator at  $80^{\circ}\text{C}$  and then weighed (to 0.01 g). We also collected the rinse water into a glass flask, which we left undisturbed for ca. 15 min to settle. We then collected 10 ml of bottom sediment with a pipette, which we plated on a petri dish and examined at  $\times 20$  magnification for earthworm chaetae, according to methodology used by Kaneko et al. (2009). We divided scat contents into ten categories, after excluding non-food materials (e.g., sand and gravel): rodents; herptiles (i.e., reptiles and amphibians); fish; insects; crayfish (the only crayfish species observed in our study area); earthworms; other animals (e.g., myriapods, land and aquatic snails); seeds (implying they consumed fruits); other plant parts (i.e., leaves and stems); and unidentified items.

For dietary analyses, we estimated the frequency of occurrence (FO; %) for all food categories and average proportion of dry weight (PDW; %) values for the nine categories (excluding earthworms) using the following equations (see Kaneko et al., 2009; Hisano et al., 2016):

$$\text{FO (\%)} = \left( \frac{\text{the number of occurrences of a food category present in sampled scats}}{\text{total number of sampled scats}} \right) \times 100$$
$$\text{PDW (\%)} = \left( \frac{\text{dry weight of a food category in a scat}}{\text{total dry weight of all food items in that scat}} \right) \times 100.$$

To assess seasonal dietary changes empirically, we compared PDW of all food

173 categories within each season using Kruskal-Wallis tests. When statistical significances  
174 were identified using Kruskal-Wallis tests, we performed post hoc *t*-test with Bonferroni  
175 corrections for multiple comparisons. Finally, to determine seasonal dietary breadth, we  
176 estimated Shannon-Wiener indices using the PDWs of all food categories in each  
177 season. All analyses were performed using the R ver. 4.1.3 (R Core Team, 2022).

## 178 Results

179 In total, we sampled 75 scats, including 11 from spring, 20 from summer, 23 from  
180 autumn and 21 from winter. Overall, crayfish were the dominant food category (FO =  
181 44.0%; PDW = 26.1%), followed by the insect (mainly Coleoptera and Odonata larvae,  
182 FO = 41.3%; PDW = 16.4%), herptile (FO = 36.0%; PDW = 15.8%), and seed (FO =  
183 26.7%; PDW = 15.8%) categories (Table 1 and Table S1 in SI). The FO and PDW of the  
184 rodent and earthworm categories comprised a relatively small proportion (< 10%) of the  
185 overall diet (Table 1 and Table S1 in SI).

186 In spring, the insect and herptile categories predominated, and were found in  
187 90.9% and 72.7% of the weasel scats sampled, respectively. Crayfish dominated weasel  
188 diet through summer (FO = 45.0%) and autumn (FO = 73.9%), followed by the insect,  
189 herptile, and other animal categories (Table 1). In winter, vegetable matter, i.e., seed  
190 (FO = 66.7%) and other plant part categories (FO = 61.9%), occurred more frequently  
191 than combined animal remains (FO = 4.8–23.8%; Table 1).

192 The PDWs of food categories differed significantly across all four seasons  
193 (Kruskal-Wallis tests,  $P < 0.01$ ; Fig. 2). In spring, the PDW of the insect category was  
194 predominant, occurring at significantly higher proportions ( $P < 0.01$ ) than five of the  
195 other food categories (Fig. 2-a). In summer and autumn, the PDW of the crayfish



197 category was predominant, followed by the insect and herptile categories (Fig. 2-b and  
198 c). Multiple pairwise comparison tests between all food categories revealed that the  
199 PDW of crayfish was significantly higher than that of either the rodent or fish category  
200 in summer ( $P < 0.05$ ) and significantly higher than seven of the other food categories  
201 (excluding comparison with herptiles,  $P = 0.08$ ) in autumn ( $P < 0.05$ ). Furthermore, in  
202 autumn, the PDWs of the herptile and insect categories were significantly higher than  
203 that of the fish category ( $P < 0.05$ ). In winter, the PDW of the seed category (indicating  
204 Rosales fruits, e.g., fig; Fig. S2 in SI) predominated, followed by the other plant part  
205 category, while combined animal remains comprised a relatively small percentage dry  
206 weight of each scat (Fig. 2-d and Table S1 in SI). Multiple comparison tests indicated  
207 the PDW of the seed category was significantly higher than for any single animal  
208 category ( $P < 0.01$ ; Fig. 2), while there was no statistical significance between the seed  
209 category and the other plant part category ( $P = 0.30$ ).

210 Shannon-Wiener indices for seasonal dietary breadth of the Japanese weasel  
211 (estimated using PDW) were 2.5 in spring, 2.6 in summer, 2.3 in autumn and 2.4 in  
212 winter.

## 213 Discussion

214 From spring to autumn, Japanese weasels ate prey species that were readily available in  
215 our study area. In support of our first hypothesis, weasels predominantly consumed  
216 (semi-) aquatic and terrestrial animal taxa (specifically crayfish, insects, and herptiles),  
217 but with only a modest FO for the rodent category in spring (18.2%), with no rodent  
218 consumption in summer. Among prey species, the consumption of the insect category  
219 decreased gradually from spring to autumn, while the consumption of crayfish increased

(Table 1 and Fig. 2). The consumption of the herptile category decreased from spring to summer and then increased to autumn. These seasonal patterns reflect the phenological cycles of prey taxa as they reproduce and proliferate in paddy fields from spring to autumn, benefitting from warm temperatures and water supplied by irrigation for rice cultivation (Washitani, 2007; Takeuchi, 2010). In spring when rice cultivation and irrigation commences, aquatic insect larvae (e.g., Coleoptera and Odonata) and a diverse abundance of adult anurans (e.g., *Pelophylax porosus porosus* and *Fejervarya kawamurai*) are available, due to immigration, emergence and/or reproduction (specifically for anurans) (Ban and Kiritani, 1980; Fujioka and Lane, 1997); similarly, around mid-May, red swamp crayfish emerge from underground hibernation in paddy field irrigation ditches, where they over-winter. From summer to autumn, the availability of insects gradually decreases as temperatures cool (e.g., Nezu et al., 2011), while juvenile anurans and crayfish mature. Importantly, water depth in paddy fields is around 5 cm when rice crops start to grow in spring (typically in mid-May; Fig. S3 in SI), but levels drop and irrigation ditches become shallower from summer to autumn (Fig. S4 in SI), allowing weasels greater predatory access to aquatic animals.

In winter, paddy fields and irrigation ditches dry up entirely, causing (semi-)aquatic animal prey to become scarce. In support of our second hypothesis, this resulted in Japanese weasels switching to a more frugivorous diet composed of plant food categories (i.e., seeds, especially fig fruits, as well as other plant material).

Although rodents are common in the winter diets of Japanese weasels in other parts of their range (Otsu, 1971; Fujii et al., 1998; our Table 2), in our study, the rodent category FO was just 4.8% in winter (Table 1). This is likely because the homogeneous paddy field monoculture in our study area did not include any scrub or wooded habitats that support terrestrial rodent populations over winter. We did, however, find that Japanese

247 weasels continued to consume red swamp crayfish in winter (FO = 14.3%; Table 1).  
248 These were likely either dug out of dry irrigation ditch beds where they over-winter  
249 (Gherardi et al., 2002; Sato et al., 2023) or taken from small pools (typically less than  
250 20 cm water depth) that persist at intersections between ditches in during winter (Fig.  
251 S5 in SI) and likely provide an important feeding site for Japanese weasels through the  
252 winter months.

253 Although our study area was relatively small (i.e., 1.2 km<sup>2</sup>), home range sizes  
254 of male weasels typically range from 0.1 km<sup>2</sup> (in an agricultural landscape) to 0.35 km<sup>2</sup>  
255 (in suburban riverine habitats) with range overlap between several individuals (Masuda  
256 and Watanabe, 2015; Watanabe, 2005). This suggests that our study area likely  
257 supported several individuals. Carnivore home range sizes are determined by the  
258 dispersion of food resources (Macdonald and Johnson, 2015), and thus it is possible that  
259 not all resident weasels had access to fig trees in their territories or chose to eat fruit.  
260 Whether only a few weasels eat fruit could be determined by future studies that track  
261 which foraging sites individuals use.

262 Consistent with our findings, previous studies on Japanese weasel diet in  
263 HMLs have also reported a high FO value for red swamp crayfish (>20 %) and seeds  
264 (19.0–63.1 %) (Fujii et al. 1998) with one report of the high rates of frugivory in a  
265 mountainous habitat (Furuya et al., 1979; see Table 2). In contrast to our study site,  
266 weasels in natural habitats typically consume terrestrial insects (e.g., Coleoptera and  
267 Orthoptera) and/or reptiles (e.g., Furuya et al., 1979; Sekiguchi et al., 2002), as well as  
268 rodents in winter (Otsu, 1971). In HMLs, specifically in urban and suburban areas  
269 where the land is covered by buildings and paved roads, riverine habitats are important  
270 for weasels, providing both foraging and resting sites (Fujii et al., 1998; Suda et al.,  
271 2014; Suzuki, 2018a; Watanabe, 2005). This can also cause weasels to rely more heavily

273 on aquatic prey, such as red swamp crayfish (Fujii et al. 1998). Furthermore, urbanized  
274 HMLs may have fruit available from cultivated trees and bushes planted in parks,  
275 gardens, and orchards (Fujii et al. 1998; Kaneko et al. 2009).

276 In conclusion, our findings broadly corroborate previous reports that the  
277 Japanese weasel is far less specialized on mammalian (especially rodent) prey than  
278 other small *Mustela* species (e.g., McDonald et al., 2000; Zub et al., 2008; Vaca-León et  
279 al., 2019), due to its adaptable and plastic trophic niche (Table 2). The consumption of  
280 red swamp crayfish and other (semi-)aquatic animals by Japanese weasels reveals the  
281 species' capacity for opportunism and efficient dietary switching to exploit  
282 environmental prey availability; a trait also seen in the western polecat (Lode, 1997)  
283 that can adapt to various environments (Lode, 1994) as well as in larger mustelids (e.g.,  
284 Zhou et al., 2011; 2015). Simultaneously, this predation effect may provide some degree  
285 of biological control on invasive red swamp crayfish, which are a pest species in Japan  
286 (Nakata et al., 2005), although likely not sufficient to substantially reduce numbers. The  
287 high frequency of seed category (FO = 67.7%; Table 1) in Japanese weasel scats further  
288 exemplifies their trophic plasticity, where, among related mustelids, switching to  
289 seasonal fruits is rare among *Mustela* species (McDonald et al., 2000; Martinoli et al.,  
290 2001; Zub et al., 2008; Vaca-León et al., 2019) and a trait more typically seen in  
291 martens (*Martes* spp., Zhou et al., 2011). This winter frugivory is important because  
292 Japanese weasels are too small and slender to hibernate (Newman et al., 2011;  
293 Wereszczuk and Zalewski, 2015; Macdonald and Newman, 2017).

294 A broader guild of mesocarnivores were present in this study area, especially  
295 raccoons and raccoon dogs (author's unpublished camera trapping data and footprint  
296 track observations in mud), and these may exert intra-guild competitive pressures on  
297 smaller Japanese weasels, also affecting their access to food resources. Certainly,

299 raccoons are known to predate crayfish (Boncompagni et al., 2021) and to consume  
300 seasonal fruit, with raccoon dogs also eating a significant proportion of available fruit  
301 and crustaceans (Xu et al., 2023), including crayfish (Takatsuki and Kobayashi, 2023).  
302 Masked palm civets are similarly adaptive opportunists that consume various fruits and  
303 invertebrates in Japan (Iwama et al., 2017).

304 While the versatile feeding habits of Japanese weasels enable them to exploit  
305 alternative food resources, and thus to survive in HMLs, their population has been in  
306 steady decline across Japan, especially in lowland areas (Sasaki et al., 2014; Kaneko et  
307 al., 2016). Consequently, the Japanese government has banned the hunting of female  
308 Japanese weasels (Kaneko et al., 2016). In addition to the difficult resource conditions  
309 that agricultural and urbanized habitats present (Sasaki et al., 2014; Masuda and  
310 Watanabe, 2015), this decline is exacerbated by competitive exclusion resulting from  
311 the spread of the invasive alien Siberian weasel through western Honshu, Shikoku and  
312 Kyushu Islands of Japan (Sasaki et al., 2014)—although not detected in our study site.

313 To further conserve weasel populations in agricultural lowlands and especially  
314 in paddy field monocultures, we recommend that habitat management strategies aimed  
315 specifically at enhancing winter food availability are implemented (Korpimäki et al.,  
316 2004). For example, at a microhabitat scale, allowing winter flooding of paddy fields  
317 would result in the year-round availability of more aquatic and semi-aquatic animal prey  
318 (Washitani, 2007) and waterfowl (Katayama et al., 2020), especially in the south of the  
319 Japanese archipelago. At landscape scale, breaking rice monoculture with a mosaic of  
320 habitats through reforestation and managing abandoned fields as a biotope or ecological  
321 park could also be advantageous. This could result in greater year-round prey diversity,  
322 particularly enhancing the availability of rodents (Alain et al., 2006; Panzacchi et al.,  
323 2010; Rey Benayas and Bullock, 2015), invertebrates, and fruits (Haggard et al., 2019;

Katayama et al., 2020). Furthermore, Japanese weasels use small burrows on river banks or fallow paddy fields as reproductive nests (Watanabe, 2005). Therefore, natural embankments of rivers and irrigation ditches are important to maintain their viable populations. However, any intervention must be monitored carefully to ensure this does not disadvantage Japanese weasels by favouring other small carnivore species to a greater extent, and thus promoting intra-guild competition (Linnell and Strand, 2000).

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**Figure legends:**

**Fig. 1.** Location of the study area, depicting the use of roads and adjacent road margins (shown as thin lines in the dashed-line polygon) as the transect route.

**Fig. 2.** Seasonal scat content for Japanese weasels in a paddy area in Saitama, Japan, divided into food categories and then expressed as mean proportions of dry weight (PDW; %) with 95% confidence intervals (represented by vertical bars). Kruskal-Wallis tests comparing PDWs among food categories in each season are shown at the top of each panel. Horizontal bars with star(s) represent statistically significant pairwise comparisons of food categories, using post hoc multi-comparison t tests with Bonferroni correction.

**Table 1.** Seasonal frequency of occurrence (FO; %) of 10 food categories (including unidentified items) in the diet of Japanese weasels in a paddy area in Saitama, Japan (*N*, number of items occurred).

Season	Spring		Summer		Autumn		Winter		Whole period	
Number of scats	11		20		23		21		75	
Food category	<i>N</i>	FO (%)	<i>N</i>	FO (%)	<i>N</i>	FO (%)	<i>N</i>	FO (%)	<i>N</i>	FO (%)
Rodent	2	18.2	0	0.0	1	4.3	1	4.8	4	5.3
Herptile	8	72.7	5	25.0	11	47.8	3	14.3	27	36.0
Fish	0	0.0	0	0.0	0	0.0	2	9.5	2	2.7
Insect	10	90.9	7	35.0	12	52.2	2	9.5	31	41.3
Crayfish	4	36.4	9	45.0	17	73.9	3	14.3	33	44.0
Earthworm	0	0.0	0	0.0	2	8.7	2	9.5	4	5.3
Other animal	2	18.2	7	35.0	6	26.1	5	23.8	20	26.7
Seed	1	9.1	2	10.0	3	13.0	14	66.7	20	26.7
Other plant part	5	45.5	3	15.0	10	43.5	13	61.9	31	41.3
Unidentified item	5	45.5	4	20.0	5	21.7	3	14.3	17	22.7

**Table 2.** Frequency of occurrence (%) of nine food categories in the diet of Japanese weasels observed in previous studies from different habitat types across Japan in comparison to our study (*N*, number of faeces or stomachs analyzed). Seasons are abbreviated as ‘Su’, summer; ‘Au’, autumn; ‘Wi’, winter; and ‘Whole’, all year round.

Site	Zamami Island, Okinawa	Nishikuma valley, Kochi	Hiwa, Hiroshima	Asahi & Iide Mts, Yamagata	Hamura, Tokyo	Tachikawa, Tokyo	Hamura, Tokyo	Mito, Ibaraki	Kazo, Saitama
Habitat type	Hilly forest	Montane forests	Montane forests	Montane foothill	Riverine, suburban	Riverine, suburban	Riverine, suburban	Suburban	Paddy fields
Season	Su–Au	Whole	Au	Wi	Whole	Whole	Whole	Wi	Whole
<i>N</i>	141	317	46	75	285	65	33	237	75
Food category									
Mammal	17.0	21.8	13.0	50.7	26.5	32.3	6.1	21.5	5.3
Bird	1.4	8.8	6.5	13.3	8.1	4.6	3.0	2.5	0.0
Herptile	34.0	10.4	54.3	5.3	6.7	1.5	9.1	0.0	36.0
Fish	0.0	0.0	17.4	6.7	20.5	7.7	21.2	18.1	2.7
Insect	85.8	63.1	8.3	0.0	32.3	44.6	75.8	39.2	41.3
Crustacean	14.9	2.2	8.7	5.3	43.9	27.7	21.2	20.3	44.0
Earthworm	0.0	0.0	15.2	0.0	0.0	0.0	0.0	8.3	5.3
Other animal	3.5	0.0	2.2	0.0	6.7	7.7	12.1	12.5	26.7
Seed (fruits)	7.1	83.9	0.0	0.0	63.1	60.0	33.3	19.0	26.7
Reference	Sekiguchi et al., (2002)	Furuya et al., (1979)	Yukawa, (1968)*	Otsu, (1971)	Fujii et al., (1998)	Fujii et al., (1998)	Okawara et al., (2014)	Kaneko et al., (2009)	This study

\*Data cited from the Table 3 in Sekiguchi et al., (2002).

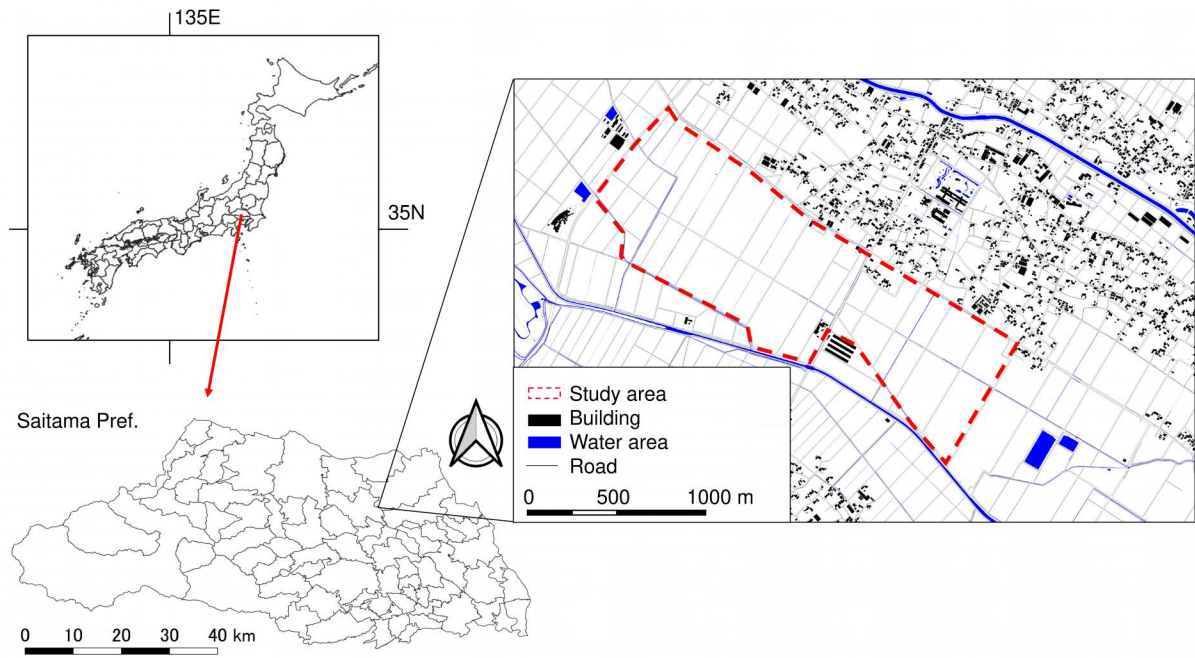


Fig. 1

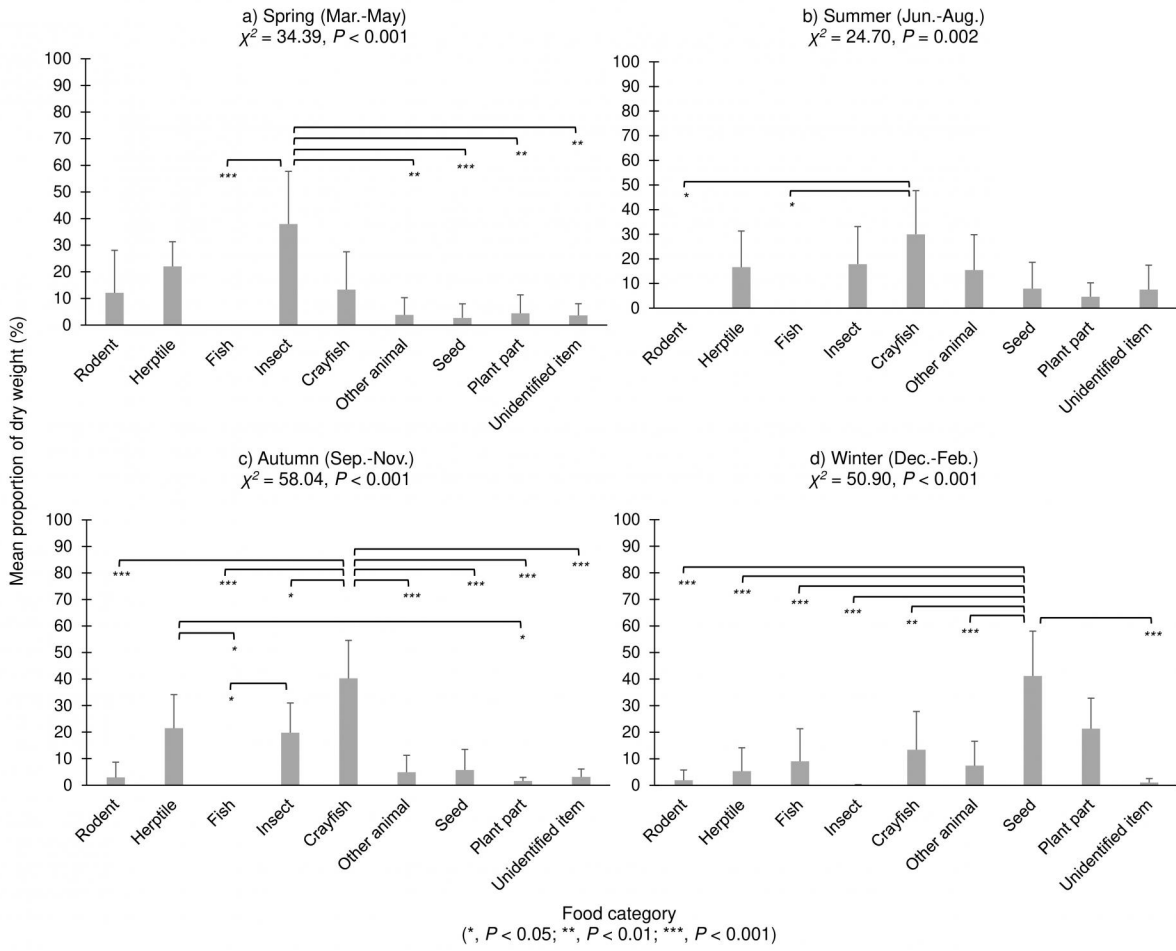


Fig.2

**Supplementary Materials:**

**Title:** Trophic plasticity of the endemic Japanese weasel in a lowland agricultural landscape

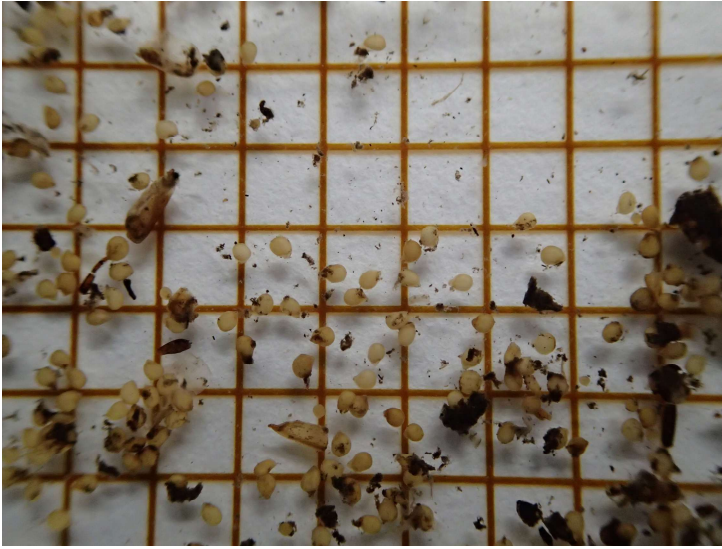
This supplementary material includes Figs. S1–S5 and Table S1.





**Fig. S1.** Aerial photograph of the study area taken on non-cropping season of 2015. The brown areas are (dry) paddy fields. The red dashed polygon represents the area sampled and grey thin lines in the polygon represent farm roads (i.e., representing sampled transects). The yellow square (identified by an arrow) indicates the location of a small-floriculture farm, where ca. two fig trees were planted (see also Fig. S4). Aerial photographed sourced from the Geospatial Information Authority of Japan (taken in 2015, <https://maps.gsi.go.jp/development/ichiran.html>).

(a)



(b)



**Fig. S2.** (a) seeds of fig fruits (*Ficus carica*) found in a Japanese weasel scat, supplementarily sampled in Jan. 2022 (grid size: 5 mm); and (b) a cultivated fig tree planted along the roadside in a small floricultural-farm (indicated by yellow square in Fig. S1) located in the study site (taken by author in Nov. 2020).

(a) May. 2018



(b) Jun. 2018



**Fig. S3.** View of paddy fields in late spring (a) and mid summer (b). Photos taken by author.



(a) Nov. 2020



(b) Jan. 2022



(c) Nov. 2020



(d) Jan. 2022



**Fig. S4.** View of paddy fields (a, b), ditch (c) and pool (d) dried up during fallow (winter) season. Photos taken by author.

(a) Nov. 2020



(b) Jan. 2022



**Fig. S5.** Examples of temporary pools at an intersection of irrigation ditches. Yellow arrow in photograph (a) indicates the remains of a claw from a red-swamp crayfish (*Procambarus clarkii*). Photo taken by author.

**Table S1.** Seasonal mean proportions of dry weight (PDW; %) of 10 food categories (including unidentified item) in the diet of Japanese weasels in a paddy area in Saitama, Japan.

Season	Spring (Mar.-May)		Summer (Jun.-Aug.)		Autumn (Sep.-Nov.)		Winter (Dec.-Feb.)		Whole priod	
Number of scats	11		20		23		21		75	
Food category	<i>N</i>	PDW (%)	<i>N</i>	PDW (%)	<i>N</i>	PDW (%)	<i>N</i>	PDW (%)	<i>N</i>	PDW (%)
Rodent	2	12.1	0	0.0	1	2.9	1	1.6	4	3.2
Herptile	8	22.0	5	16.7	11	24.5	3	5.3	27	15.8
Fish	0	0.0	0	0.0	0	0.0	2	9.1	2	2.5
Insect	10	38.0	7	17.9	12	19.8	2	0.1	31	16.4
Crayfish	4	13.3	9	30.0	17	37.3	3	13.4	33	26.1
Earthworm*	0	-	0	-	2	-	2	-	4	-
Other animal	2	3.8	7	15.4	6	4.9	5	7.1	20	8.3
Seed	1	2.7	2	7.9	3	5.7	14	41.2	20	15.8
Other plant part	5	4.4	3	4.6	10	1.6	13	21.2	31	8.4
Unidentified item	5	3.6	4	7.5	5	3.3	3	1.1	17	3.8

\*dry weights of earthworm chaetae were not measured.