Restoring a butterfly hot spot by large ungulates refaunation: The case of the Milovice military training range, Czech Republic

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Abstract

Background: Refaunation by large ungulates represents a cost-efficient approach to managing natural biotopes. We studied butterflies inhabiting grasslands of the former Milovice military training range in the Czech Republic, grazed since 2015 by a combination of Exmoor pony (“wild” horse), Tauros cattle (“aurochs”) and European wisent.

Methods: We compared presence-absence patterns from two surveys preceding the refaunation, one immediately after military use termination (early 1990s), the other prior to the refaunation (2009), with the current (2016–19) results of monitoring 8 grazed and 8 ungrazed plots. We analysed the life history, climatic and conservation-related traits of recorded butterflies to gain insights into the ungulates’ impacts.

Results: Following the termination of military use, several poorly mobile species displaying broad oceanity niches were lost. Newly gained species are more mobile and prefer warmer continental conditions. Currently, numbers of butterfly species do not differ between refaunated and neglected plots, but the former hosted higher abundances. Butterflies developing on coarse grasses and shrubs inclined towards neglected plots, whereas refaunated plots supported species developing on small forbs.

Conclusion: The changes in species composition following the cessation of military use are attributable to successional change, coupled with climate-driven expansion of thermophilous and mobile species. By blocking succession, large ungulates support butterflies depending on competitively poor plants. Preserving butterflies of late-successional stages is ensured by the diverse setting of the reserves and by the relatively low ungulate densities.

Introduction

In most continents, late-Pleistocene and early Holocene human pressure extirpated, or drastically reduced, the populations of large ungulate herbivores, which reshaped the ecological dynamics of entire biomes (Corlett 2004; Johnson et al. 2009; Galetti et al. 2018). This affected nutrient cycling (e.g., Zimov et al. 1995), fire regimes (e.g., Galetti 2004; Navarro and Pereira 2012), seed dispersal and germination (Vera 2000; Doughty et al. 2012) and overall vegetation physiognomy (Feurdean et al. 2018). Subsequent activities of preindustrial agriculturists and pastoralists resumed the role of large wild ungulates, maintaining the disturbance-succession dynamics exploited by numerous species, including invertebrates. Many enigmas and paradoxes encountered in European insects conservation – such as the affiliation of many taxa to purportedly “cultural” grasslands (Thomas 1993; van Swaay et al. 2012), ancient ways of forests use (Warren 1991; Dolek et al. 2018), frequently disturbed habitats (Rehoukova et al. 2016; Tropek et al. 2017) or finely-grained landscapes (Slanoverova et al. 2012, Slancarova et al. 2014) – are resolved, once the large ungulates activity is factored in. The current biodiversity has evolved in a megafaunal world (Bartanova et al. 2018; Todisco et al. 2010). Modern ecosystems are functionally incomplete, with entire trophic levels impoverished or missing, and if not actively managed, they fail to provide habitats for a sizeable portion of associated biota (Dirzo et al. 2014; Sebek et al. 2015).

The current rewilding/ refaunation /naturalistic grazing movements (Seddon et al. 2014; Merckx and Pereira 2015; Svenning et al. 2016; Cromsigt et al. 2018; Garrido et al. 2019) strive to reverse the transformation of ecosystems that started in the late Pleistocene and culminated with recent land use intensification (Dirzo et al. 2014; Jirku and Dostal 2015; Henning et al. 2017; Pedersen et al. 2019). Although insect conservationists have long advocated habitat management by ungulate grazing (e.g., Öckinger et al. 2006; Cizek et al. 2013; Slancarova et al. 2014), relatively few megafauna refaunation projects have focused primarily on insect conservation or are systematically monitored for impacts on insects (van Klink et al. 2018). Each refaunation project develops within specific sociocultural constraints, rarely allowing for proper replications (cf. Root-Bernstein et al. 2018; but see Garrido et al. 2019). There is an urgent need to study refaunation effects on invertebrates, both as encouragement for others and as feedback for the wider conservation community (Jepson 2016; van Klink et al. 2018).

The initial refaunation plans for the Czech Republic (cf. Jirku and Dostal 2015) aimed at large protected areas, or actively used military training ranges. It was believed that in these large and biotically rich areas, restoring populations of large ungulates would be most feasible. The first refaunation project, however, has materialised on a relatively small scale of two grazing reserves within a disused military training range, in otherwise densely populated Central Bohemia (Figure 1). Since 2015, three once near-extirpated components of native European megafauna roam on grasslands formerly used for army training: the European bison or wisent, a species rescued from near-certain extinction (Ölech 2008); the back-bred “aurochs” in its restored Tauros form derived from several taurine breeds of domestic cattle (Bos taurus) (Goderie et al. 2015); and the horse (Equus caballus), in an ancient feral Exmoor pony breed (Baker 1993; Hovens and Rijkers 2013).

Coincidently, the area was surveyed for butterflies immediately after the cessation of its military use (Matouš 1994), and again in the following decade, in the context of comparing fauna of abandoned military ranges (Cizek et al. 2013). The setting thus offers a unique opportunity to study butterfly assemblages’ responses to the abandonment of a military-used landscape, and to monitor effects of large ungulates refaunation on such assemblages.

A promising approach to generalisation from single-site results focuses on life history (= functional) traits of constituent species. It links habitat properties and the species composition of assemblages via species-specific traits (Legendre et al. 1997; Carmona et al. 2016). By linking species traits to results of habitat manipulation, it may disclose the mechanisms of species responses to habitat change (Slancarova et al. 2014; Gamez-Virues et al. 2015).
The development of insect assemblages over two decades is naturally subject to other influences than locally operating disturbance-succession dynamics. An obvious factor is changing climate, restructuring faunas on the continental scale (Warren et al. 2001, Essens et al. 2017), but non-climatic drivers of species distribution shifts, such as large-scale land use changes, were also detected (Aguirre-Gutierrez et al. 2016).

Here, we first compare butterfly records from the three subsequent surveys: shortly after the military use termination, shortly before the refaunation by large ungulates, and under the large ungulates’ impact. For this past-present comparison, we hypothesised that cessation of military use was followed by losses of specialists of early-successional disturbed grounds (H1), whereas subsequent species gains should reflect current faunal changes, explicable by the recently warming climate (H2) and such conservation-related attributes as ongoing changes in distribution extent (H3). We then analyse the current monitoring of butterfly assemblages on refaunated versus neglected plots. We hypothesise that the whole-year grazing by wild ungulates should not decrease species richness and abundances compared to neglected plots (H4), but the assemblage should shift from species associated with competitively dominant host plants, typical for successionally advanced and little disturbed vegetation, towards species associated with competitively inferior host plants typical for frequently disturbed conditions (H5).

**Materials And Methods**

**Study area, refaunation and earlier butterfly surveys**

The Milovice military training range (50.26N, 14.89E, attitude 200–250 m a.s.l., mean annual temperature 8–9 °C, annual precipitation 500–600 mm) (Figure 1) was established in 1904, originally on 34.6 km². It was subsequently used by all armies that operated on Czech territory, gradually expanding its area to 40 km². The last users were the Soviets, who operated an air force base and headquarters here for the former Czechoslovakia until 1991. The natural setting is the gently rolling Středočeská Tabule Plain formed by Mesozoic carbonate-rich sandstones, siltstones and claystones, and covered by brown soils, rendzinas and carbonate rich sands. Woodlands dominated by Quercus petrea, Pinus sylvestris and Betula pendula are interspersed by finely grained mosaics of shrublands, grasslands and early successional vegetation that developed on former farmlands and were utilised for training troops for over 80 years (Cížek et al. 2013).

Following the cessation of military use, parts of the open training fields were developed (golf course, amusement park, industrial zone), while three large areas were proclaimed a Site of European Community Importance (SCI) Milovice-Mladá. The Central site (local toponym: Pozorovatelna, hereinafter “C”, 50.254N, 14.881E) has been partly managed by conventional sheep grazing, while the Northern (Traviny, “N”, 50.278N, 14.883E) and Southern (Pod Benáteckým vrchem, “S”, 50.241N, 14.886E) sites remained unmanaged, except for occasional disturbance of S by armoured vehicles practiced by military history enthusiasts and for domestic cattle grazing in a corner of N in 2014–2016. Much of all three sites had suffered succession-driven homogenisation of the once diverse vegetation mosaic by competitively dominant grasses (mainly Calamagrostis epigejos and Arhenatherum elatius), ruderal forbs and shrubs (mainly Crataegus, Prunus and Rosa).

The site S (2015–2017, 40 ha; 106 ha since 2018) has been grazed since spring 2015 by ≈35 Exmoor ponies (hereinafter “horse”) and ≈20 Tauros cattle (hereinafter “aurochs”). Since spring 2016, ≈35 horses and ≈20 wisents graze the site N (125 ha) (Figure 1). Both S and N are thus year-round cross-grazed by horses and big bovids ( wisent or aurochs) living in naturally structured social units, i.e. mixed sex/age harems/herds. To provide variable management regimes, both temporally and permanently ungrazed plots of various sizes (units to tens of hectares) are present both within and outside the grazing reserves at any given time. The animals receive no supplementary feeding and no medication, except for strictly determined individual cases, and predators enter the sites freely (Jíru et al. 2018). To control grazing intensity, facilitate gene-flow and avoid social stress, two to three year-old surplus animals are transferred to similar projects in the Czech Republic and abroad.

The first modern butterfly survey of the area was conducted immediately after the cessation of military use. Matouš (1994) published a commented list of species, treating the entire military range as a single locality. Fifteen years later, in 2009, the training fields S, C and N were surveyed in a semiquantitative manner (Čížek et al., 2013). The current monitoring of the refaunation impact, launched in spring 2016, thus represents the third survey.

**Current butterfly monitoring**

We set 16 rectangular plots (50 x 200 m) at both refaunated (n = 7) and non-refaunated (n = 9) sections of N and S sites (n = 8 each) (Figure 1). From spring 2016 onwards, the plots were visited five times each year (May, early June, late June, July, August) to cover seasonal aspects of butterfly assemblages. The recording followed the timed survey protocol by Kadlec et al. (2010), appropriate for heterogeneous environments with temporally changing locations of butterfly resources, such as flower patches. Each visit to a plot lasted 30 min, all butterfly species present were recorded using a net when necessary and taking vouchers of species not recognisable in the field. Abundances were recorded, using rounded numbers for species seen in large abundances. We also recorded the closest hour, cloudiness (3-points ordinal scale, from clear sky = 1 to overcast = 3), wind (Beaufort scale 1 – 4, i.e., calm to gentle breeze), and nectar supply (0 – no flowers within the plot, 1 – flowers scarce but present, 2 – flowers moderately abundant, 3 – flowers abundant). We restricted the visits to the highest butterfly activity period (10 AM – 4 PM) and to weather suitable for butterflies, randomising their sequence with respect to time of day. A single round of visits took 2–3 consecutive days.
Statistical analyses

For the past-present comparison, we visualised the patterns defined by species presences/absences recorded by Matouš (1994), Cizek et al. (2013) and the current monitoring, the latter collated across the four years, using the correspondence analysis (CA), an unconstrained ordination appropriate for 1/0 data, in CANOCO, v. 5.0 (Ter Braak and Smilauer 2013). We computed four variants of CAs: 1) based on three “samples” defined by the three consecutive surveys; 2) differentiating records from the locations N, C and S (possible using Cizek et al. (2013) and the current data), thus obtaining six “samples”; and 3+4) as in the previous two cases, but after exclusion of migrant and arboreal species.

We interpreted the CA results by three sets of the constituent species traits (Table 1, Appendix 1): (a) life history traits, mostly associated with feeding modes, dispersal and population structure, as compiled for Central Europe by Bartonova et al. (2014); (b) climatic niche traits, compiled by Schweiger et al. (2014) on the basis of species ranges in Europe and known to contribute to population trends (Essens et al. 2017); and (c) conservation attributes describing the distribution and Red-list status in the Czech Republic. We used the CANOCO option “explanation of species scores for functional traits”. This analysis, a multivariate version of the fourth-corner approach (Legendre et al. 1997; Dray et al. 2014), relates the species ordination scores from the CA ordination to trait values of the species, testing for strengths of the relationship using redundancy analysis (RDA), a multivariate version of linear regression (Ter Braak and Smilauer 2013). We analysed the three sets of traits separately, using the forward selection process to attain best-fitting traits combinations.

To compare numbers of butterfly species and individuals recorded during the current monitoring, we used 2-way analysis of variance with factors year (4 levels) and management (2 levels, refaunation vs. neglect, the latter including the plots grazed by cattle in 2016–17). Cumulative numbers of species and summed numbers of individuals across the five yearly visits were the dependent variables.

To study the current composition of butterfly assemblages, we used canonical correspondence analysis (CCA), a constrained ordination method relating the species composition of samples to external predictors and testing the relationships of species composition to predictors using the Monte Carlo test with 999 permutations, again in CANOCO. We reflected the temporal dependency in our data using a hierarchical split-plot permutation design, permuting the individual plots randomly, and the 20 subsequent visits per plot as mutually dependent cyclic shifts. We first ran separate tests for all possible nuisance covariables, i.e. year (both as 4-levels factor and as a linear value), site (N vs. S), hour (as factor and 2nd-degree polynomial), weather (a combination of cloudiness and wind), nectar and plots position (forward-selected from latitude, longitude, their polynomials and interaction).

For the pivotal effect of refaunation itself, we used two different codings, aiming to answer two slightly different questions. Refaunation (3-levels factor: refaunation, cattle and neglect) aimed to disclose the effect of wild ungulates, while ungulates (5-levels: horse, auroch, wisent, cattle and neglect) aimed to decipher effects of the three ungulate animals. We also tested for military vehicle effect (2-levels factor tanks). We developed the models by systematically adding the covariates that had significant effects in the single-term CCA into the refaunation and ungulates models, until we reached models that significantly explained the distribution of monitored butterflies while being stringently controlled for nuisance effects.

Analogously to the past-present comparison, we interpreted the final CCA current monitoring models by species traits, relating the CCA scores to the three sets of traits and using forward selection to select the best-fitting traits combinations.

Results

Past-present comparison

The early 1990s survey of the entire military range (Matouš et al. 1994) detected 72 butterfly species (14 currently red-listed); the interim survey of the sites N, C, S (Čížek et al. 2013) detected 51 (6 red-listed) species and the 2016–19 monitoring at the sites N and S detected 58 (7 red-listed) species (Table 2). The numbers are comparable only with caution. The earliest survey covered all biotopes in the area, including wooded parts outside the grasslands. It recorded a higher representation of arboreal species (n = 9) than the two latter surveys (4 and 5). The two latter surveys focused on grasslands, but while the interim survey consisted of five visits in a single year, the current monitoring consisted of 20 visits in four years. Still, even after exclusion of arboreal species and migrants whose abundances vary greatly among years, the earliest survey detected more species than the latter two surveys pooled.

The indirect CA analyses (Figure 2) revealed differences among the three surveys in butterfly species composition (total variation = 0.22, Axis 1 separating the earliest and the two subsequent surveys: 59.5%, Axis 2 distinguishing the interim and the current survey: 40.5%). The pattern held if the localities N, C, S were treated separately (variation = 0.40; % subsequent axes: 45.2, 29.0). Removing 11 arboreal and vagrant species (cf. Table 2) decreased the explained variation (three samples variant: 0.25, six samples variant: 0.35) without changing the overall pattern (% successive axes: 59.9, 40.1, vs. 47.9, 31.0).

Interpreting the CA ordinations by life history traits gave results consistent across the four variants (Table 3). Species present shortly after the cessation of military use and lost subsequently tended to be less mobile. Their ranges were characterised by a broader oceanity niche, narrower continentality niche and lower mean annual temperatures (Figure 2). Species inclining towards the current survey require higher mean annual
temperatures and higher numbers of growing degree days. The species lost since the earliest survey are declining in the Czech Republic, while those gained recently display rather restricted distributions in the country (Table 3).

**Current monitoring of refaunation effects**

The 61 species currently recorded (Table 2) were observed in 25,322 individuals. The mean(±SD) /median/ range per plot and year, summed across the five yearly visits, were 24.6(±4.88)/ 24 /15–38 species, and 395.7(±214.55)/ 343/ 99–1,057 individuals. The numbers of species per plot differed significantly among years and refaunation performed marginally better than neglect, except for 2018 with an opposite pattern, resulting into marginally significant year x year interaction (ANOVA, year F(356,df) = 3.72, P < 0.05; management F(156,df) = 2.88, P = 0.095; interaction F(356,df) = 2.36, P = 0.082). The numbers of individuals also differed among years and were consistently higher at refaunated plots (ANOVA, year F(356,df) = 3.11, P < 0.05; management F(156,df) = 4.93, P < 0.05; interaction F(356,df) = 1.45, P = 0.237) (Figure 3).

In the CCA analyses (Table 4), the potential nuisance covariates nectar, hour or weather did not affect the composition of assemblages, implying that nectar was available rather evenly across the plots and visits, and visits were carried out under suitable weather. The strong effect of factorially coded year explained the highest variation of all (co)variables. It was followed by plots position, specifically latitude, colinear with the effect of site. Tanks as a separate predictor had no effect. For ungulates, the first axis, which distinguished auroch and horse from neglect, cattle and wisent, was not significant. The significant second and (still canonical) third axes distinguished neglect and cattle, respectively, from other situations. Refaunation alone had no significant effect.

Adding tanks into either ungulates or refaunation models increased the models' statistical significance, suggesting that some butterflies responded to thus created intensive disturbance. The models also dramatically improved after inclusion of factorial year + latitude as covariates (Table 4, Figure 4). Apparently, the originally weak effects of the focal predictors were due to variation among years and collinearity between grazing regimes and positions of plots.

For refaunation, the ordination now clearly separated plots grazed by large ungulates from neglected plots at the first axis, and cattle plus tanks from all the other factors at the second axis. The butterflies closely associated with large ungulates were narrowly specialised herb-feeders, such as the obligatorily myrmecophilous *Phengaris alcon*, multiple other Lycaenidae (*Plebejus argus, Polyommatus coridon*), but also some more specialised Pieridae (*Colias alfacariensis* and Hesperiidae (*Erynnis tages, Pyrgus malvae*). Species associated with neglect were those preferring coarse grasslands (the fritillary *Boloria dia*; the satyriinae *Melanargia galathea, Maniola jurtina*, the hesperids *Ochloides venatus, Hesperia comma*) and shrubs (*Iphiclides podalirius, Coenonympha arcania*). Cattle pasture was associated with common generalists (*Pieris brassicae, Vanessa cardui, Thymelicus lineola*), but also with the declining hesperid *Spialia sertorius*, which was also closely associated with Tanks. Almost identical patterns arose in analysis with ungulates. The first axis distinguished neglect from the three megafaunal species, the second axis distinguished cattle plus tanks, and the (still canonical) third and fourth axes separated auroch from wisent and cattle from other management types, respectively (see Appendix 3 for detailed positions of all butterfly species).

Interpreting results of the CCA model ~refaunation+tanks | factorial year+latitude by species traits returned a single significant pattern (Figure 4), associating the first and second ordination axes with the life history trait host plant form. For a similarly structured model ungulates+tanks | factorial year+locality, the result was identical, although only marginally significant, and the same held for analogous models not containing covariates (Appendix 4). Species inclining towards neglect (first axis) but also towards tanks and cattle (second axis), tended to develop on woody plants or coarse grasses, while those inclining towards refaunation tended to develop on small forbs. No climatic niche or conservation-related trait performed significantly in these analyses.

**Discussion**

The former Milovice military training area harbours rich butterfly assemblages, arguably preserved there owing to exclusion of intensive agriculture and forestry, combined with the past finely-grained disturbance-succession dynamics typical for military areas (Reif et al. 2011; Cizek et al. 2013; Busek and Reif 2017). Following the cessation of military use, several species were lost, while others were gained subsequently. The presence of aurochs, horses and wisents increases local butterfly abundances and contributes to maintaining their diversity, providing for multiple species of conservation concern, including the critically endangered obligatorily myrmecophilous *Phengaris alcon* (cf. Thomas and Settele 2014).

**Changes since termination of military use**

The termination of military activities was followed by the successional overgrowth of the disturbed sparsely vegetated surfaces, and gradual dominance of coarse grasses and tall forbs. We therefore expected (hypothesis H1) decrease of specialists associated with small competitively inferior forbs, which our analyses of life history traits did not support. The only life history trait responding to the past-present ordination was mobility. Poorly mobile species were associated with the past military use. Among European butterflies, high mobility is a generalist trait associated with broad
trophic ranges, long flight period and other features facilitating survival in human-dominated landscapes (Dapporto and Dennis 2013, Bartonova et al. 2014), whereas poor mobility increases extinction risks (Birkhofer et al. 2017; Essens et al. 2017). Because mobility relates inversely to local population density (Bartonova et al. 2016), some poorly mobile species may need large habitat areas to sustain viable populations. The changes after cessation of military use probably led to shrinking habitats supply for poorly mobile specialists.

Associations of lost and gained species with climatic niche traits (H2) were more straightforward. In agreement with the warmer and drier climate in Central Europe during the last few decades (Stuhlbrecher and Fortmann 2018), the lost species shared broad oceanity or precipitation niches, whereas the newly gained species require higher temperatures. Also, in agreement with H3, the locally lost species display decreasing distribution trends in the Czech Republic and elsewhere in Central and Western Europe (cf. van Swaay et al. 2010).

A combination of restricted mobility and broad oceanity or precipitation niches applies to several locally lost and nationally threatened species (cf. Benes et al. 2002; Hejda et al. 2017): the hesperid Pyrgus amoricanus and Thymelicus acteon and the satyrides Hipparchia semele, Hyponephele lycaon and Erebia aethiops. The latter is a sparse woodland species (Slamova et al., 2012) only loosely associated with grasslands, but its current occurrence in the area was safely excluded by concurrent targeted searches. The remaining four, all strongly declining in Central Europe (van Swaay et al. 2010), require sparsely vegetated pastures and often colonise such landforms as disused quarries and post-industrial barrens (Bourn and Thomas 2002; Benes et al. 2003; Tropek et al. 2010; Tropek et al. 2017). Broad oceanity tolerance certainly applies to Hipparchia semele, distributed from Eastern Europe to Atlantic coastal dunes (Schirmel and Fortmann 2014), but also to Hyponephele lycaon and Pyrgus amoricanus, whose distribution follows maritime climates far north to southern Fennoscandia (Fourcade et al. 2017; Mikkola 1979). The species newly gained during the last two decades include Iphiclides podalirius, Satyrium accaciae, S. spini, Lycaena dispar and Polyommatus bellargus, all currently (re)expanding in Central Europe. The first three are associated with shrubs (Benes et al. 2002), the fourth with tall ruderal forbs (Strausz et al. 2012), and only the fifth, gained as late as 2018, with sparsely vegetated surfaces (Benes et al. 2003). The gains and losses thus reflect the interacting forces of climate and land use change (Reif et al. 2008; Thomas et al. 2015).

**Refaunation by large ungulates**

Large ungulates did not demonstrably change per plot species richness while increasing butterfly abundances, only partly supporting our hypothesis H4. In the same time, refaunation affected the local assemblages’ composition. It favoured species developing on small forbs over those developing on large forbs, grasses or shrubs, supporting our hypothesis H5.

As in other studies (cf. Hennig et al. 2017, Cromsigt et al. 2018b, Zielke et al. 2019), the immediate effects of year-round wild ungulates presence included reduction of tall coarse grasses, slowing down scrub growth due to browsing and bark peeling, reduction of grass blooming by feeding on grass inflorescences, and exposing barren ground around tracks and walls. As in experiments with feral horses (Garrido et al. 2019), some richly blooming forbs, including species that rarely bloomed in the years preceding the refaunation (unpublished data), became notably more abundant. The differences in butterfly assemblages composition between refaunated and neglected plots became apparent only after statistical control for the effect of years and to monitored plots position. Still, species benefitting from refaunation included the iconic Phengaris alcon f. rebeli, whose host plant, the poorly competitive perennial Gentiana cruciata (cf. Petanidou et al. 1995), boomed shortly after the establishment of grazing. This obligatorily myrmecophilous butterfly is likely host plant limited, because its females prefer oviposition on plants overtopping surrounding vegetation (Meyer-Hozak 2000; Vilbas et al. 2016).

The simplest explanation of the higher butterfly numbers at refaunated plots, attraction to increased nectar, is unlikely, as covariable nectar had no effect in ordinations. A tempting explanation is the smaller size of the forbs-feeding specialists, related to higher local population densities and lower mobility. In any case, it is intriguing that many plant groups avoided by horses (e.g., Rosaceae, Fabaceae, Polygonaceae, Scrophulariaceae: Chodkiewicz 2020), once the dominant grazers of West-Palaearctic grasslands, are frequent in the larval diet of European butterflies. Possible coevolutionary relationships between mammalian megafauna and herbivorous insects, and their conservation implications, deserve further investigation.

The patterns revealed by ordinations relating species composition to refaunation were admittedly less convincing than in studies comparing starkly contrasting habitats, such as close woodlands vs. clearings (cf. Benes et al. 2006; Sebek et al. 2015). It appears that the refaunated and neglected plots were interconnected by individual movements. The distances among study plots were within the routine movement abilities of most of butterflies (Fric et al. 2010; Stevens and Baguette 2010; Vodickova et al. 2019), although this may not apply for the least mobile species (Korosi et al. 2008). Possibly, individual butterflies located some of their vital resources at both grazed and ungrazed sections of the area, in line with the resource-based understanding of (animal) habitats (Dennis et al. 2006; Turlure et al. 2019).

The setting of our study did not allow distinguishing between the effects of horses and big bovids, as both pastures contained combinations of these ungulates. The literature on refaunation/rewilding in temperate (e.g., Vera 2000, Zielke et al. 2020) and northern boreal (Macias-Fauria et al. 2020) regions agree that these two ungulate groups supplement each other in effects on vegetation, as well as seasonal and diurnal habitat use. Additionally, both horses and bovids acted as dominant grazers in late Quaternary European ecosystems, and both were present in traditional rural landscapes.
The mechanical disturbance by armoured vehicles (factor tanks), had no significant separate effect, seemingly countering the claims (e.g. Heneberg et al. 2016) that it provides disturbed conditions beneficial for some insects. Inclusion of tanks, however, increased the explanatory power of models containing ungulates or refaunation effects (Table 4), suggesting a complementarity with large grazers for some butterfly species. This might be the case of Sphialia sertorius, a skipper closely associated with tanks in ordination diagrams and developing on Sanguisorba minor; a competitively inferior forb preferring sparsely vegetated surfaces (cf. Gros 2002). It is tempting to postulate that on military lands, and in the current Milovice reserves, the heavy vehicles supplement yet another lost component of the megaherbivore fauna of interglacial Europe, proboscideans (van Kolfschoten 2000).

The effect of domestic cattle, grazed at two plots for two years of the project, was orthogonal to the ordination gradient distinguishing refaunation and neglect. The cattle were grazed with high stocking and supplementary feeding during the vegetation season and were not grazed in winter. Such grazing style suppresses forbs and fails to suppress coarse grasses. Grazing by domestic breeds in more biodiversity-friendly ways is possible (Enri et al. 2017; Henning et al. 2017, Hall and Bunce 2019), but this was not the case in our system.

While being demonstrably positive for butterflies associated with poorly competitive forbs, the refaunation did not detectably imperil species associated with coarse grasses or shrubs. In this respect, the Milovice situation differs from some projects with documented negative outcomes for insect assemblages (cf. Lorimer and Driessen 2014; van Klink and WallisDeVries 2018). It seems beneficial that contrary to some refaunation sites amidst urbanised landscapes (Lorimer and Driessen 2014), our study system is situated in a diverse rural setting, including ungrazed/neglected plots, which provide conditions contrasting with the grazed sites. This habitat diversity likely allows for resource compensation/supplementation by the butterflies (Ouin et al. 2004), enabling coexistence of species requiring different disturbance levels (Bergman et al. 2018). The current grazing pressure \( \approx 0.5 \text{ grazers} \cdot \text{ha}^{-1} \) does not deplete the sites of larval host plants or nectar. There is a potential long-term risk, as the whole operation is funded from the EU Agri-environmental scheme “grazing”, which requires maintaining stable grazing intensity. More flexibility may be necessary, as grazing levels appropriate for restoring overgrown sites may become too high once species-rich dry grasslands develop.

**Conclusion**

We demonstrated that abandonment and successional changes of a former military area restructured the rich local butterfly fauna, and that refaunation of parts of the area by megafaunal grazers contributes to maintaining high butterfly abundances and species richness. Analysing traits of the constituent butterfly species revealed that the post-abandonment changes, spanning across two decades, affected butterfly assemblages via different mechanisms than does the current megaherbivores activity. The post-abandonment changes led to losses of some poorly mobile species and gains of some regionally expanding species, presumably rather good dispersers. The changes also had an important climatic component, consistent with current climate change. The megaherbivores affect butterfly assemblages by transforming vegetation, and hence supporting species developing on small forbs on the expense of large forbs, coarse grasses and woody plants feeders. Local heterogeneity of conditions, and existence of ungrazed sections in the vicinity of the grazed ones, ensure that species from the other group are not locally imperilled. Given that many of the species lost since abandonment of the area by the military were poor dispersers, reintroductions of some of the lost species, whose habitats the ungulates have restored, is a logical next step.

Unresolved questions include differences among ungulate species in affecting butterfly resources, possible legacies of coevolution between temperate butterflies and ungulates, and future development of the butterfly assemblages. The latter question is tractable by sustained monitoring, whereas the former two can be approached by expansion of studies similar to ours to sites varying in composition of both buttery assemblages and ungulate species. This ambitious programme is increasingly feasible, as the rewilding/refaunation movement expands and the number of potential study systems rapidly increases. In the Czech Republic alone, progeny of the Milovice ungulate herds currently roam at additional seven sites, offering rich opportunities for future research.

**Declarations**

**Ethics approval and consent to participate:** Any handling of the butterflies fully conformed to valid legal provisions of the Czech Republic. Welfare of vertebrate animals was not affected by this study.

**Consent to publish:** Not applicable.

**Availability of data and materials:** All the primary data are included to this manuscript as electronic appendices.

**Competing interests:** We declare no competing interests, financial or otherwise.

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**Authors’ contributions:** M. Konvicka (MK), V. Vodickova (VV) and M. Jirku (MJ) planned and designed the study. J. Benes (JB) analysed the historical records and contributed current knowledge on butterfly species distribution. David Ricl (DR) and JB performed the current butterfly monitoring. MJ contributed background details about the refaunation project. MK, VV and MJ analysed the current monitoring data and jointly wrote the manuscript.
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Description Of Appendices

Appendix 1 – primary data on presence/absence of butterflies during three subsequent surveys of the (former) Milovice military training area, used for past-present comparison, together with life history traits, climatic niche traits and conservation-related attributes of the species.

Appendix 2 – primary data on butterfly abundances obtained during current monitoring, plus environmental characteristics of monitoring plots, and life history, climatic niche and conservation related-attributes of the species.

Appendix 3 – CCA scores of individual butterfly species, plus species weights, from the current monitoring ordination analyses, final covariate models for refaunation + Tanks and ungulates + Tanks effects.

Appendix 4 – detailed results of the RDA analyses, interpreting the results of CCA analyses of refaunation / ungulates effects on butterfly assemblages by the constituent butterflies’ life history traits.

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**Tables**

Table 1. List of life history traits, climatic niche traits and conservation-related traits used for analyses of butterfly assemblages inhabiting the former Milovice military range, currently grazed by large ungulates.
| Life history traits (Bartonova et al., 2014) | Description                                                                                                                                                                                                 | Character                  |
|--------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------|
| Wing span                                  |                                                                                                                                             | Numeric (mm)              |
| Host plant form                            | Ephemerals – 1, larger forbs – 2, grasses and sedges – 3, trees and shrubs – 4                                                                | Ranked                    |
| Voltinism                                  | Average number of generations, C. Europe                                                                                                          | Numeric                   |
| Fertility                                  | Average number of eggs per female at eclosion                                                                                                    | Categories 1–9             |
| Mobility                                   | Ranked tendency to disperse                                                                                                                    | Categories 1–9             |
| Density                                    | Ranked average density per area of habitat                                                                                                      | Ranked 1–9                 |
| Diet breadth                               | Number of plant families fed on by larvae in the Czech Republic                                                                                   | Numeric                   |
| Flight period length                       | Number of adult occurrence months (hibernation months excluded)                                                                                   | Numeric                   |
| Overwintering stage                        | Ranked, larva – 1, adult – 5                                                                                                                    |                           |

| Climatic niche traits (Schweiger et al., 2014) | Description                                                                                                                                 | Character                  |
|-----------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------|---------------------------|
| Annual temperature                           | Mean annual temperature                                                                                                                   | °C                        |
| Annual temperature niche breadth [NB]        | SD of the above                                                                                                                           |                           |
| Continentality                               | Annual range in monthly temperatures                                                                                                    | °C                        |
| Contintality NB                              | SD of the above                                                                                                                           |                           |
| Precipitation                               | Annual precipitation sum                                                                                                                  | mm                        |
| Precipitation NB                             | SD of the above                                                                                                                           |                           |
| Oceanity                                     | Annual range in monthly precipitation sum                                                                                                 | mm                        |
| Oceanity NB                                  | SD of the above                                                                                                                           |                           |
| Winter GDD                                   | Accumulated growing degree days [GDD] (> 5°C), January – February                                                                        | °C                        |
| Winter GDD NB                                | SD of the above                                                                                                                           |                           |
| Spring GDD                                   | Accumulated GDD, January – April                                                                                                          | °C                        |
| Spring GDD NB                                | SD of the above                                                                                                                           |                           |
| Early summer GDD                             | Accumulated GDD, January – June                                                                                                           | °C                        |
| Early summer GDD NB                          | SD of the above                                                                                                                           |                           |
| Summer GDD                                   | Accumulated GDD, January – August                                                                                                         | °C                        |
| Summer GDD NB                                | SD of the above                                                                                                                           |                           |
| Water availability                           | Soil water content of the upper horizon (0.5 m)                                                                                           | No unit (0–1)             |
| Water availability NB                        | SD of the above                                                                                                                           |                           |

| Conservation related traits                   |                                                                                                                                            | Character                  |
|----------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------|---------------------------|
| Red list status                              | Czech Republic (1– no status, 2– near threatened, 3– vulnerable, 4– endangered, 5– critically endangered)                                 | Ranked 1–5                 |
| Present CZ distribution                      | Occupied Czech Republic 10x10 km grid squares 2002–2014                                                                                   | Numeric                   |
| Past CZ distribution                         | Occupied Czech Republic grid squares 1951–2001                                                                                             | Numeric                   |
| CZ distribution trend                        | [1– (Present distribution/Past distribution)], details in Bartonova et al. (2014)                                                           | Numeric                   |
| Global range size                            | Categorized, (1 – smaller than Europe, 5 – larger than the Palaearctic)                                                                   | Numeric                   |
| European range size                          | Number of occupied (ca 70 x 70 km) grid squares in Europe (Schweiger et al. 2014)                                                          | Numeric                   |
Table 2. List of butterfly species (nomenclature and system: Wiemers et al., 2018) recorded from the former Milovice military training range during the three consecutive surveys, split into respective localities if possible, with their Czech Republic red-list [RL] status (NT – near threatened, VU – vulnerable, EN – endangered, CR – critically endangered) following Hejda et al. (2017). The study of Matouš (1994) refers to the early 1990s, Čížek et al. (2013) to 2009, and this study to the 2016–2019 monitoring. Abbreviations are used in the ordination diagram at Figure 4.

$^{\$}$Arboreal and $^{\$}$migrant species excluded from some analyses.

*The “rebel” ecological form, developing on Gentiana cruciata.
| Species                  | Abbreviation | RL status | Matouš (1994) | Čížek et al. (2013) | This study |
|--------------------------|--------------|-----------|---------------|---------------------|------------|
| *Hesperidae*             |              |           | Entire area   | Site S | Site C | Site N | Pooled | Site S | Site N | Pooled |
| Carcharodus alceae       | Calc         | NT        | +             | -      | -      | -      | +      | +      | +      | +      |
| *Erynnis tages*          | Etag         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Carteroccephalus palaemon* | Cpal     | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Pyrgus malvae*          | Pmal         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Pyrgus armoricanus*     | EN           | +         | -             | -      | -      | -      | -      | -      | -      | -      |
| *Ochlodes sylvanus*      | Osyl         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Thymelicus lineola*     | Tiin         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Thymelicus sylvestris*  | Tsyl         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Thymelicus acteon*      | EN           | +         | -             | -      | -      | -      | -      | -      | -      | -      |
| *Spialia sertorius*      | Sser         | VU        | +             | +      | +      | +      | +      | +      | +      | +      |
| *Hesperia comma*         | Hcom         | VU        | +             | -      | -      | -      | -      | +      | +      | +      |
| *Papilionidae*           |              |           |               |        |        |        |        |        |        |        |
| *Papilio machaon*        | Pmac         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Iphiclides podalirius*  | Ipod         | NT        | -             | -      | -      | -      | -      | +      | +      | +      |
| *Pieridae*               |              |           |               |        |        |        |        |        |        |        |
| *Pieris brassicae*       | Pbra         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Pieris napi*            | Pnap         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Pieris rapae*           | Prap         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Leptidea juvernica*     | Ljuv         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Gonepteryx rhamni*      | Grha         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Colias alfacariensis*   | Calf         | VU        | +             | +      | +      | +      | +      | +      | +      | +      |
| *Colias hyalo*           | Chya         | -         | -             | -      | -      | -      | -      | -      | -      | -      |
| *Colias croce*           | Ccro         | -         | +             | -      | -      | -      | -      | +      | +      | +      |
| *Anthocaris cardamines*  | Acar         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Pontia edusa*           | Pedu         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Nymphalidae*            |              |           |               |        |        |        |        |        |        |        |
| *Apatura iris*           | Airi         | -         | +             | -      | -      | -      | +      | -      | -      | +      |
| *Apatura ilia*           |              | -         | +             | -      | -      | -      | -      | -      | -      | -      |
| *Aglais urticae*         | Aurt         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Nymphalis antiopa*      |              | -         | +             | +      | +      | +      | -      | -      | -      | -      |
| *Nymphalis polychloros*   |              | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Vanessa atalanta*       | Vata         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Vanessa cardui*         | Vcar         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Inachis io*             | Iio          | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Araschnia levana*       | Alev         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Polygonia c-album*      | Pc-a         | -         | +             | +      | +      | +      | +      | +      | +      | +      |
| *Malitaea athalia*       | NT           | +         | -             | +      | +      | +      | -      | -      | -      | -      |
| *Malitaea cinxia*        | VU           | +         | -             | -      | -      | -      | -      | -      | -      | -      |
| Taxon | Scientific Name | Status | Count 1 | Count 2 | Count 3 | Count 4 | Count 5 | Count 6 | Count 7 | Count 8 | Count 9 | Count 10 |
|-------|----------------|--------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| Issoria lathonia | Ilat | - | + | + | + | + | + | + | + | + | + | + |
| Argynnis aglaja | Aagl | - | + | + | + | + | - | + | + | - | + | + |
| Argynnis adippe | Aadi | - | + | + | - | - | + | - | - | - | + | - |
| Argynnis paphia | Apap | - | + | + | + | + | + | + | + | + | + | + |
| Boloria dia | Bdia | - | + | + | + | + | + | + | + | + | + | + |
| Boloria selene | NT | + | - | - | - | - | - | - | - | - | - | - |
| Coenonympha arcania | Cacr | NT | + | + | + | + | + | + | + | + | + | + |
| Coenonympha glycerion | Cgly | - | + | + | - | - | + | + | + | + | + | + |
| Coenonympha pamphilus | Cpam | - | + | + | + | + | + | + | + | + | + | + |
| Erebia medusa | Emed | NT | + | - | - | - | + | + | + | + | + | + |
| Erebia aethiops | EN | + | - | - | - | - | - | - | - | - | - | - |
| Melanargia galathea | Mgal | - | + | + | + | + | + | + | + | + | + | + |
| Lasiommata megera | Lmeg | - | + | + | + | + | + | + | + | + | + | + |
| Lasiommata maera | NT | + | - | - | - | - | - | - | - | - | - | - |
| Pararge aegeria | Paeg | - | + | + | - | - | - | - | + | + | + | + |
| Maniola jurtina | Mjur | - | + | + | + | + | + | + | + | + | + | + |
| Hyponephele lycaon | CR | + | - | - | - | - | - | - | - | - | - | - |
| Aphantopus hyperanthus | Ahyp | - | + | + | + | + | + | + | + | + | + | + |
| Hipparchia semele | CR | + | - | - | - | - | - | - | - | - | - | - |
| **Lycaenidae** | | | | | | | | | | | | |
| Thecla betulae | - | + | - | - | - | - | - | - | - | - | - | - |
| Neozephyrus quercus | NT | + | - | - | - | - | - | - | - | - | - | - |
| Callophrys rubi | Crub | NT | + | + | + | + | + | + | + | - | + | + |
| Satyrium acaciae | Saca | - | - | - | - | - | - | - | + | - | - | - |
| Satyrium pruni | Spru | NT | + | - | - | - | - | + | - | - | - | - |
| Satyrium w-album | NT | - | + | - | + | - | - | - | - | - | - | - |
| Satyrium spini | Sspi | VU | - | - | - | - | - | + | + | + | + | + |
| Lycaena alciphron | VU | - | - | - | + | + | - | - | - | - | - | - |
| Lycaena dispar | Ldis | - | - | - | - | - | - | + | + | + | + | + |
| Lycaena phlaeas | Lphl | - | + | + | + | + | + | + | + | + | + | + |
| Lycaena tityrus | Ltit | - | + | + | + | + | + | + | + | + | + | + |
| Lycaena virgaureae | Lvir | NT | + | + | + | + | + | + | + | + | + | + |
| Celastrina argiolus | Carg | - | + | + | + | + | + | + | + | + | + | + |
| Cupido minimus | Cmin | VU | + | + | - | - | + | + | + | + | + | + |
| Aricia agestis | Aage | - | + | + | + | + | + | + | + | + | + | + |
| Aricia eumedon | NT | + | - | - | - | - | - | - | - | - | - | - |
| Plebejus argyrognomon | Pargy | - | + | - | - | - | - | + | + | + | + | + |
| Plebejus argus | Parg | NT | + | + | + | + | + | + | + | + | + | + |
| Cyaniris semiargus | VU | - | - | - | + | + | - | - | - | - | - | - |
| Phengaris alcon* | Palc | EN | + | + | + | + | + | + | + | + | + | + |
| Polyommatus bellargus | Pbcall | VU | - | - | - | - | - | + | + | + | + | + |
Table 3. Results of explaining species scores obtained from the correspondence analyses (CA) of three successive butterfly assemblages surveys (early 1990s, 2009, 2016–19) in the (former) Milovice military training range, by life history traits, climatic niche traits and conservation attributes of constituent species.

Three-levels analyses pooled individual sites surveyed, while six-levels analyses treated the grasslands sites S, C and N separately, if allowed by the data. -/+ signs preceding the Ax1-Ax4 values indicate the direction of the correlation with respective CA axes. %var, F and P values refer to Monte Carlo tests for the significance of the relationships between trait values and CA ordination scores.
| Analysis | Ordination axes: traits correlations | % var. | Axis 1 | All axes | Ordination axes: traits correlations | % var. | Axis 1 | All axes | Ordination axes: traits correlations | % var. | Axis 1 | All axes |
|----------|------------------------------------|--------|--------|---------|------------------------------------|--------|--------|---------|------------------------------------|--------|--------|---------|
| 3-levels | Ordination axes: traits correlations | 17.0   | 14.5***| 5.0***  | -Ax1: Distribution trend CZ; Current range CZ | 17.3   | 19.9***| 9.2***  |
| 3-levels | Mobility                           | 2.2    | 2.5+   | -       | +Ax1: Oceanity NB;                 |        |        |         | -Ax1: Distribution trend CZ; Current range CZ | 20.8   | 13.4***| 9.8***  |
| 6-levels | Mobility                           | 2.0    | 2.6*   | -       | -Ax1: Precipitation NB             | 2.5    | 3.0*   | -       | -Ax1: Distribution trend CZ; Current range CZ | 18.6   | 15.9***| 12.5***|
| 6-levels | Mobility                           | 3.5    | 3.4*   | -       | +Ax1: Early summer GDD, Summer GDD | 2.4    | 2.6*   | -       | +Ax1: Distribution trend, CZ current range; -Ax2: CZ current range | 20.2   | 14.7***| 9.5***  |

Table 4. Results of CCA analyses, comparing the 2016–19 current monitoring results from plots refaunated by large ungulates versus neglected plots. E1–E4 are eigenvalues of respective canonical axes, F and P values refer to results of Monte Carlo tests for the first canonical axis and all canonical axes.
| Model                                      | E1    | E2   | E3  | E4   | % variation | Axis1 F, P   | All axes F, P |
|-------------------------------------------|-------|------|-----|------|-------------|--------------|---------------|
| ~nectar                                   | 0.039 |      |     |      | 1.2         | 4.9, ns      |               |
| ~weather                                  | 0.034 | 0.027|      |      | 1.7         | 4.2, ns      | 3.8, ns       |
| ~factorial hour                           | 0.021 | 0.016| 0.012| 0.008| 0.6         | 2.5, ns      | 1.3+          |
| ~polynomial hour                          | 0.017 | 0.006|      |      | 0.3         | 2.1, ns      | 1.4*          |
| ~factorial year                           | 0.114 | 0.049| 0.019|      | 6.1         | 14.5*        | 7.9**         |
| ~linear year                              | 0.045 |      |      |      | 1.4         | 5.6, ns      |               |
| ~position (forward selected: latitude)    | 0.026 |      |      |      | 0.7         | 3.2**        |               |
| ~site                                     | 0.026 |      |      |      | 0.7         | 3.2**        |               |
| Covariate model\(^1\) (~factorial year +latitude) | 0.114 | 0.054| 0.020| 0.019| 8.0         | 14.5*        | 6.8***        |
| ~tanks                                    | 0.015 |      |      |      | 0.3         | 1.9, ns      |               |
| ~tanks | latitude +factorial year                  | 0.007 |      |      |      | 0.1         | 1.0, ns      |               |
| ~refaunation\(^2\)                        | 0.019 | 0.010|      |      | 0.5         | 2.3, ns      | 1.8, ns       |
| ~refaunation | latitude +factorial year                  | 0.019 | 0.006|      |      | 0.4         | 2.5**, 1.7**  |               |
| ~refaunation +tanks                       | 0.029 | 0.014| 0.010|      | 1.1         | 3.6, ns      | 2.2*          |
| ~refaunation +tanks | latitude +factorial year                  | 0.022 | 0.009| 0.004|      | 0.5         | 2.9**, 1.6**  |               |
| ~ungulates\(^3\)                          | 0.023 | 0.015| 0.010| 0.005| 0.8         | 2.8, ns      | 1.6*          |
| ~ungulates | latitude +factorial year                  | 0.020 | 0.009| 0.006| 0.003| 0.3         | 2.5*, 1.2+    |               |
| ~ungulates +tanks                         | 0.035 | 0.015| 0.015| 0.010| 1.4         | 4.2*, 1.9*   |               |
| ~ungulates +tanks | latitude +factorial year                  | 0.023 | 0.010| 0.009| 0.004| 0.4         | 3.0*, 1.3*    |               |

\(^+\): P < 0.1, \(^*\): P < 0.05, \(^**\): P < 0.01, \(^***\): P < 0.001

\(^1\)Obtained by forward selection from all significant terms above. \(^2\)3-levels factor (refaunation, cattle, and neglect); \(^3\)5-levels factor (horse, auroch, wisent, cattle, and neglect).

**Figures**
Figure 1

Former Milovice military training range. (a) The position of the range within the Czech Republic. (b) Aerial view of the area with the military range borders (green line), positions of the grasslands N, C, S, and borders of the two grazing reserves (red lines). (c, d) More detailed view at the 16 monitored plots within grasslands S and N. Original work; the aerial photo taken from Český Ústav Zeměměřický a Kartografický (www.mapy.cz); freely available for non-commercial research.
Figure 2

Interpreting three subsequent butterfly surveys of the Milovice military range by species traits. Results of CA analysis of presence/absence data obtained from the Milovice military training range (analysis six levels, arboREAL and migrant species excluded, cf. Table 3) interpreted by life history traits (top left), climatic niche traits (top right) and conservation attributes (bottom left) of constituent species. Positions of the three surveys, based on CA ordination of recorded butterflies, in bottom right.

Figure 3

Butterfly species richness and abundance of neglected and refaunated plots. Numbers of butterfly species (black bars) and individuals (grey bars) recorded in the former Milovice military training range, with respect to management of the plots. Means ±SD recorded per the plot and year are shown.
Figure 4

Ordination analysis of large ungulates refaunation effects on butterfly assemblages. Left: CCA biplot relating the current (2016–19) butterfly species composition at monitored plots in the Milovice former military training area to refaunation; model after inclusion of covariables (~refaunation +tanks | factorial year +latitude; see Table 4 for the CCA model parameters). Right: RDA biplot relating ordination scores of the model to the butterfly species life history traits. In this model, Host plant form was the only trait selected by a forward selection (eigenvalue 0.049, adjusted explained variation 3.3%, Monte Carlo test F = 3.0, P = 0.039).

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- Appendix1.xlsx
- Appendix2.xlsx
- Appendix3.xlsx
- Appendix4.xlsx