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Large regional variation in global impacts of agriculture on terrestrial insects and other arthropods

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ABSTRACT

Many insects and other arthropods are reported to be in rapid decline worldwide, mainly driven by changes in land use and climate. At the same time, arthropods provide many important services that benefit agriculture, and thus their losses may pose risks to food security. Although biodiversity responses vary between global realms, this spatial heterogeneity is not well-understood and is rarely addressed in global analyses. To understand the geographical variation of biodiversity change better, we investigated the responses of arthropod diversity to agricultural land use and intensity for six geopolitical regions, using a global dataset of biodiversity records. We show that agricultural intensification generally leads to average biodiversity declines, but we find large variation in responses per region. Notably, responses in Europe are minimal or even significantly positive. Further investigation suggests that responses in Europe may be attributable to a combination of a primary vegetation baseline that is subject to comparatively high levels of disturbance and fragmentation, and significant compositional changes that have favoured widespread generalists and non-native species. Conversely, narrow-ranging habitat specialists are associated with significant declines. These findings have important consequences for global biodiversity assessments and associated predictions of ecosystem service delivery, which are likely underestimating land use impacts on biodiversity by relying on statistical model coefficients that are heavily biased towards Europe and may fail to report important compositional changes by focussing analyses solely on species richness and total abundance.

1 Introduction

Insects and other arthropods are reported to be in rapid decline worldwide, driven principally by the expansion and intensification of agriculture (Raven and Wagner, 2021; Wagner et al., 2021). While there is ongoing speculation as to whether insects are in decline globally, given considerable variation in trends between species, orders and regions, and a significant monitoring bias towards the northern hemisphere (Dirzo et al., 2014; Klink et al., 2020; Outhwaite et al., 2020), the data that is available suggests that terrestrial insect abundance is decreasing globally at ca. 10% per decade, while freshwater insect abundance is increasing at approximately the same rate (Klink et al., 2020). It is, moreover, estimated that 5-10% of insect species may have gone extinct since the start of the industrial era, numbering between 250,000 and 500,000 species (Cardoso et al., 2020). Many arthropod species provide valuable ecosystem services for agricultural systems, such as crop pollination and pest regulation (Costanza et al., 1997; Dainese et al., 2019; Potts et al., 2016b; Yang and Gratton, 2014), and thus declines in arthropod abundance and diversity may pose substantial risks to food production and human well-being (Millard et al., 2023; Pereira et al., 2024; Potts et al., 2016a), particularly in regions that rely heavily on animal pollination and natural pest control.

While biodiversity is changing worldwide, the magnitude and direction of change has been shown to vary widely between biogeographic realms (Almond et al., 2022; Olson et al., 2001), with the strongest declines observed in the neotropics. Although large-scale analyses have highlighted stark differences in biodiversity responses to land use between the tropical and temperate zones (Millard et al., 2021; Newbold et al., 2020; Outhwaite et al., 2022), it remains common to report on global statistics, which, in turn, are frequently used in combination with land-use projections of integrated assessment models (IAMs) to estimate future impacts on biodiversity and ecosystem service delivery (e.g., Kim et al., 2018; Kozicka et al., 2023; Leclère et al., 2020; Pereira et al., 2024; Schipper et al., 2020). Such approaches may be missing important regional context (Bevan et al., 2025), and since global ecological data availability tends to be highly biased towards Europe and North America (Proença et al., 2017; Trimble and Aarde, 2012), where contemporary biodiversity change appears to be less pronounced (Almond et al., 2022), global statistics are likely underestimating impacts on biodiversity for large parts of the planet.

In order to estimate the impacts of land use on biodiversity better across the globe, an

32 improved understanding of the regional heterogeneity of responses is needed. In this study,
33 we investigated differences in the responses of arthropod diversity to agricultural land use and
34 land-use intensity across six world regions, with a particular focus on Europe, where preliminary
35 results show changes in diversity from primary vegetation to be either minimal or even signifi-
36 cantly positive (Fig. 1). Although this is not an entirely unexpected result, as it is known that
37 impacts of land use on biodiversity are less pronounced in temperate regions (Newbold et al.,
38 2020), it is important to understand better what drives these responses in Europe, particularly
39 since biodiversity data collection tends to be highly biased towards the European continent. In-
40 deed, the fact that Europe is a data-rich region means that further investigation is statistically
41 feasible.

42 Impacts of anthropogenic stressors such as agricultural expansion and intensification on
43 species assemblages can be estimated by comparing biodiversity in anthropogenic land-use types
44 against an undisturbed baseline that is assumed to represent species assemblages in their natural
45 state, typically termed 'primary vegetation' (Hudson et al., 2017). However, it is difficult to
46 determine a 'virgin' state of nature, and it is known that anthropogenic stressors in Europe
47 have been high (and accelerating) since the 18th and 19th centuries, long before widespread
48 biodiversity monitoring has taken place (Mihoub et al., 2017), impacting biodiversity within
49 the natural habitats that are used as baseline condition. Among these stressors, deforestation
50 to clear land for pasture and cropland has likely been the most influential (Darby, 1955),
51 rapidly transforming the mixed deciduous and boreal forests that had come to dominate Europe
52 following the end of the last glacial period (Williams, 2003). This deforestation has been
53 persistent, and reconstructions suggest that by 1850, deforestation had reached 60-70% in central
54 Europe and in excess of 90% in Britain (Kaplan et al., 2017). Although the industrial revolution
55 helped to alleviate pressure on European forests, and helped intensify agriculture to put aside
56 space for nature, the associated reliance on agro-chemicals has come to be identified as one of the
57 main contemporary drivers of insect declines in Europe (Benton et al., 2002; Fox, 2013). As a
58 result of these persistent pressures on the natural environment and the domination of landscapes
59 by anthropogenic habitats, the baseline condition of the remainder of natural habitat in Europe
60 has thus likely been gradually degraded (Soga and Gaston, 2018). The result of using natural
61 habitat of this contemporary nature as a baseline in statistical models of the impact of land use
62 on biodiversity is that impacts are likely to be less negative than if they had been compared
63 against truly undisturbed primary vegetation.

64 In terms of species compositions, this long history of land-use change on the European
65 subcontinent has likely produced both winners and losers (Newbold et al., 2018), as forest
66 clearings and the loss of primary vegetation would have benefited open-habitat generalists at
67 the expense of closed-canopy, forest specialists. Additionally, winners of land-use change tend to
68 be non-native and invasive (Chytrý et al., 2008; Liu et al., 2023; Pyšek et al., 2020), which may
69 mask declines in native biodiversity (Van Kleunen et al., 2015). This appears to be particularly
70 relevant in Europe, North America and Australasia, where the number of introductions of
71 non-native species dominate compared to other regions (Seebens et al., 2025). The baseline
72 natural habitat in regions like Europe may thus be relatively more dominated by non-native,
73 widespread and generalist species than in regions without a long history of human impact on
74 the environment. Depending on the extent of change in species compositions within natural
75 habitat, differences in arthropod (alpha) diversity between primary vegetation and agricultural
76 land use may thus be considerably less pronounced in Europe than in other regions.

77 We explore three main hypotheses that may explain differences in biodiversity–land-use re-
78 sponses of arthropods between Europe and other regions: We hypothesise that as a result of
79 Europe’s long history of land-use change (1) structurally, natural habitat in Europe is more
80 fragmented, and thus less surrounded by other natural habitat, than in other regions, with the
81 result that many of the species that would have been present within large tracts of natural
82 habitat will have disappeared; (2) homogenization in European ecosystems over time has led
83 to compositional similarities between natural vegetation and agricultural sites, and thus species
84 turnover between natural habitat and agricultural sites is lower than in other regions; and (3)
85 winners, i.e. species that respond positively in agricultural land use, tend to be widespread
86 habitat generalists and non-native species, which have had more opportunity to establish them-
87 selves in Europe than in other regions, with the result that overall responses in Europe are less
88 negative than in other regions. In testing the third hypothesis we compare the effect of range
89 extent, habitat specificity and non-nativeness on arthropod diversity and specifically compare
90 Europe against North America. These regions span a similar latitudinal range and are thus
91 subject to similar climatic conditions, and are comparable in terms of economic development
92 and agricultural land use. Crucially, ecosystems in Europe and North America have been sub-
93 ject to different land-use histories and we thus expect to see differences in the degree to which
94 widespread generalists and non-native species mask declines of native specialists.

2 Methods

2.1 Biodiversity data

The arthropod records used in this study stem from the PREDICTS database, which contains worldwide, site-level species occurrence and abundance records sourced from published studies that sampled from land-use types along a gradient of intensity (Hudson et al., 2017). These data are nested at four levels: 'source' (a unique publication), 'study' (a collection of data within a source that has a consistent sampling methodology), 'block' (a collection of sites in a distinct spatial cluster) and 'site' (a geographic location at which records were sampled). We used the updated version of the PREDICTS database (Hudson et al., 2023, version 1.1), and an additional data release (Contu et al., 2022), to obtain 1,079,780 records of 10,200 arthropod species, across 7978 sites worldwide (Table 1 and Fig. 1). The sampled species belong to the orders Coleoptera (26.1% of records), Hymenoptera (17.8%), Lepidoptera (17.7%), Sarcop-
tiformes (13.6%), Mesostigmata (7.6%), Araneae (4.5%), Hemiptera (4.4%), Diptera (2.7%), Entomobryomorpha (2.0%) , Poduromorpha (1.4%) and other orders (5.3%), with slight variation between regions (Table S2). We investigated biodiversity responses for agricultural sites (cropland, pasture and tree plantation) at three use-intensities (minimal, light and intense) and compared these against responses in primary vegetation (native vegetation not known or inferred to have ever been completely destroyed) and secondary vegetation (vegetation that is in various stages of recovery after the original primary vegetation was completely destroyed, e.g., for a period of agricultural use). In the latter category, we did not include 'young' (early successional stage) secondary vegetation, which may be very similar to disturbed agricultural land. Exact definitions of these land-use types and their use-intensities, as used in the PREDICTS database, can be found in the supplementary information A (Table S1).

We compared the responses of arthropod diversity to land use across six world regions: North America, Europe, South and Southeast Asia, South and Central America, sub-Saharan Africa and Australasia (Australia and New Zealand), largely similarly to Almond et al. (2022). Although these regions broadly mirror the world's biogeographic realms (Olson et al., 2001), they were selected primarily as geopolitical regions with largely shared land-use histories. Although Europe and East Asia share the same biogeographic realm (the Palearctic) their land-use histories differ in important ways, which we expect to have led to differences in the contemporary responses of biodiversity to land use and land-use change. These regions were, furthermore,

126 selected on the basis of data availability, which meant that Oceania, North Africa, Central Asia
127 and East Asia were omitted from the analysis. Following a preliminary sensitivity analysis (see
128 supplementary information A; Section B.7) in which we found a disproportionately strong influ-
129 ence of a single study from the Azores islands on European biodiversity responses, we decided
130 to remove the Azores islands from the dataset, given its geographical isolation from the rest
131 of Europe. As such, records from Europe are restricted to continental Europe, Great Britain
132 and Ireland. For consistency, we removed records from similarly sized island groups from other
133 regions, which led to the removal of Comoros from sub-Saharan Africa and Puerto Rico from
134 South and Central America (as a result this region does not contain data from the Caribbean).

135 For the majority of this regional analysis, we modelled arthropod diversity responses to an
136 aggregate agricultural land-use category (combining cropland, pasture and plantations), but
137 for Europe we further investigated responses to individual land-use types (where sample sizes
138 were sufficient) and compared these against global responses. In order to determine whether the
139 diversity responses of arthropods are indicative of a general European pattern, we also modelled
140 biodiversity responses for vertebrate species (in Europe and globally). Preprocessed and filtered
141 to natural and agricultural land use, vertebrate records in PREDICTS totalled 559,074 globally
142 (5032 species), distributed across 7842 sites and 224 studies. Europe comprised 51,665 records
143 (183 species), 1157 sites and 15 studies. The majority of vertebrate records stemmed from
144 bird species, with Passeriformes (song birds) as the largest order (46.2% of global records).
145 Vertebrate records in Europe consisted entirely of bird species, with Passeriformes comprising
146 60.3% of records.

147 For the further comparative analysis between Europe and North America, we extracted
148 species data on range extent, habitat specificity and non-native status from additional data
149 sources (see Sections 2.5, 2.6 and 2.7). To maximise the number of species for which we could
150 extract these data, we manually checked species' Latin binomial names in the PREDICTS
151 database (for Europe and North America) for errors such as abbreviations and spelling mis-
152 takes or inconsistencies, and compared names with the Global Biodiversity Information Facility
153 (GBIF) taxonomic backbone to ensure that, where possible, synonyms were harmonised to a
154 single species.

Table 1: Counts of unique species, studies, sites and total records for arthropods in the PREDICTS database, filtered for natural vegetation and agricultural land-use types. Europe comprises the most data-rich region in the database.

| Region | Unique species | Studies | Sites | Total records | Records % |
|---------------------------|----------------|---------|-------|---------------|-----------|
| North America | 2141 | 43 | 988 | 141,634 | 14.5 |
| Europe | 4098 | 118 | 3412 | 396,728 | 40.5 |
| South and Southeast Asia | 898 | 30 | 525 | 77,555 | 7.9 |
| South and Central America | 1494 | 85 | 1125 | 79,986 | 8.2 |
| Sub-Saharan Africa | 659 | 21 | 426 | 44,263 | 4.5 |
| Australasia | 708 | 26 | 712 | 238,825 | 24.4 |
| Tropical Zone | 2732 | 122 | 1858 | 192,389 | 19.7 |
| Temperate Zone | 7127 | 202 | 5330 | 786,602 | 80.3 |
| Total | 9800 | 323 | 7188 | 978,991 | 100.0 |

2.2 Main modelling framework

As biodiversity metrics, we primarily modelled species richness, total abundance and Simpson’s diversity index, which is a measure of evenness (Simpson, 1949). Rather than modelling Simpson’s diversity index directly, we modelled its inverse to convert the index to a diversity measure that reflects the effective number of species, allowing responses to be comparable with species richness (Jost, 2006). We modelled these three metrics with mixed effects models using R Statistical Software (R Core Team, 2024, version 4.5.1) with the Bayesian modelling package `brms` (Bürkner, 2021, version 2.22.0). We modelled species richness with a zero-inflated negative binomial distribution, in order to account for overdispersion and zero-inflation. Total abundance and Simpson’s diversity index were modelled with a log-normal distribution, which we found fit the data well. In cases where sampling effort varied between sites within a study, we corrected abundance measurements by dividing by relative sampling effort (relative to the maximum sampling effort in the study), assuming that recorded abundance scales linearly with sampling effort (Newbold et al., 2015). This applied to 9.1% of studies in our dataset and 17.6% of abundance measurements. Due to these corrections, total abundance was not modelled as a count variable but rather as a continuous variable with a log-normal distribution (Newbold et al., 2015).

We did not rescale abundance relative to the maximum recorded abundance within studies, as in De Palma et al. (2021), for the reason that model diagnostics of log-transformed total

174 abundance indicated better model fit than either models of log-transformed rescaled abundance
175 or models of rescaled abundance using a zero-and-one-inflated beta distribution (see Section E
176 of Supplementary Information B). We suspect that the model diagnostics of log-transformed
177 total abundance showed better fit because we modelled diversity records within a single phylum
178 (Arthropoda), which means that problematic between-study sampling and reporting differences
179 that abundance rescaling is meant to address are possibly less pronounced than in the case
180 of modelling records across various phyla and kingdoms (De Palma et al., 2021). We did
181 investigate differences in modelled abundance responses to land use between these models,
182 including a zero-inflated negative binomial model of total abundance, and note that while there
183 is some variation between models and the significance of certain responses (predominantly in
184 data-scarce regions), model choice does not meaningfully change general regional patterns and
185 conclusions drawn from these in this study (Fig. S20).

186 For all three diversity metrics, we estimated regional differences in biodiversity responses to
187 land use by modelling land use and region as interactive fixed effects, using study-level random
188 intercepts to account for differences in sampling methods and study-design as well as block-
189 level random intercepts to account for any spatial structuring in the sites sampled within each
190 study (Eq. 1). We present model predictions as the median and 95% credible interval from a
191 sample of 1,000 posterior draws of the expected response conditional on the fixed effects. Where
192 predictions are presented as changes from primary vegetation, we divided the posterior response
193 samples of all land-use types by primary vegetation and computed the median and 95% credible
194 intervals from these relative responses.

$$\text{Diversity} \sim \text{Land use} \times \text{Region} + (1|\text{Study/Block}) \quad (1)$$

195

196 **2.3 Natural habitat**

197 In order to investigate the landscape composition of natural habitat surrounding PREDICTS
198 sites, we computed the percentage of natural habitat within a 5km radius of the PREDICTS
199 sites (square buffer). We used the land-use layers from Matej et al. (2025), which have a 30
200 arcsecond resolution (approximately 1km² at the equator) and span the years 1992–2020. These
201 land-use classes are derived from the land cover product of the European Space Agency Climate
202 Change Initiative (ESA CCI LC) and use statistics from the Food and Agriculture Organization
203 (FAO) to discern land cover into 32 land-use classes. From these classes, we aggregated the

204 wilderness and forestry classes, as well as grazing land characterized by open wooded lands, to
205 constitute natural habitat.

206 We also used the dataset’s land-use intensity layers to investigate differences in the use-
207 intensity of natural habitat between regions. Land-use intensity is estimated as human appro-
208 priation of net primary production (HANPP), which is an indicator that quantifies the effects of
209 human-induced changes in productivity and harvest on ecological biomass flows (Haberl et al.,
210 2014). The total HANPP of the natural habitat in a buffer is comprised of the HANPP due to
211 land use change (HANPP_{luc}, which equals the difference between the actual NPP of the habitat
212 and the potential NPP in the hypothetical absence of land use) and the HANPP due to harvest
213 (HANPP_{harv}, which equals the difference between the actual NPP and the NPP remaining in
214 ecosystems after harvest, NPP_{eco}). By definition, the HANPP of wilderness areas, and the
215 HANPP_{luc} of forestry areas, are zero. We converted HANPP to a percentage of potential NPP
216 ($NPP_{pot} = NPP_{eco} + HANPP_{harv} + HANPP_{luc}$) to obtain a metric that is comparable across
217 regions and land use types (%HANPP) (Erb et al., 2013). We computed %HANPP and the
218 percentage of natural habitat area within the buffers surrounding sites from the year in which
219 the midpoint of the sampling duration of the site occurred (i.e., the date half-way between the
220 first and last sampling dates) and for each site computed their mean values in the buffer. Fi-
221 nally, we estimated the connectivity of the natural habitat surrounding PREDICTS sites within
222 a larger buffer with 10km radius with three metrics – mean patch size, patch density and the
223 aggregation index (He et al., 2000) – using the R package `landscapemetrics` (Hesselbarth et
224 al., 2019, version 2.2.1). To achieve this, the layers of fractional natural habitat were converted
225 into binary layers using three different natural habitat (NH) thresholds: $NH \geq 0.3$, $NH \geq 0.5$ and
226 $NH \geq 0.7$.

227 2.4 Compositional dissimilarity

228 We compared compositional dissimilarity between species in natural vegetation (primary and
229 secondary vegetation) and in agricultural land use with Sørensen’s index (Mueller-Dombois,
230 1974). Sørensen’s dissimilarity $S = \frac{b+c}{2a+b+c} \in [0, 1]$ is a popular dissimilarity measure in the
231 study of ecological communities (McCune and Grace, 2002), where a designates the number of
232 species observed in both sites, and b and c designate the number of unique species sampled in
233 each site individually. We further partitioned Sørensen’s index into nestedness and turnover
234 (Baselga, 2010), using the `betapart` R package (Baselga and Orme, 2012, version 1.6.1), to

235 determine the extent to which dissimilarity is the result of species replacement (turnover) or
236 species loss (nestedness; i.e., the extent to which composition in one land use is a subset of the
237 other). We only compared sites within the same study and computed dissimilarity exclusively
238 for site combinations in the off-diagonal of the pairwise dissimilarity matrix to avoid problems
239 of non-independence from using all possible pairwise comparisons (Longacre et al., 2005).

240 Species compositions in agricultural sites were compared against primary and secondary
241 vegetation, rather than just primary vegetation, because within-study comparisons between
242 primary vegetation and agriculture were found to be scarce in the PREDICTS database, limit-
243 ing the land-use coefficients that we were able to estimate. This was found to be particularly
244 the case for light- and intense-use agriculture in Europe, where within-study comparisons with
245 secondary vegetation are much more abundant than with primary vegetation. In addition,
246 dissimilarity between secondary vegetation and primary vegetation was not found to be signifi-
247 cantly different from dissimilarity between primary vegetation and other primary vegetation in
248 any region (Fig. S8), which suggests that the comparison of dissimilarity between agricultural
249 sites and all natural vegetation sites may be sufficiently representative of a comparison with
250 just primary vegetation. We used Sørensen’s index, which is an occurrence-based index, rather
251 than indices that incorporate individual species abundances, to further maximise the number
252 of possible land-use comparisons by being able to make use of both occurrence and abundance
253 records (which were coerced into presence/absence).

254 We estimated compositional dissimilarity for each region and between each land use and
255 natural vegetation pair (including natural vegetation with other natural vegetation sites) using
256 a generalised linear mixed effects model (Eq. 2). Differently to our models of diversity, we mod-
257 elled dissimilarity with a zero-and-one-inflated beta distribution as the variable is bounded by
258 zero and one and was found to be inflated at these boundaries. Since between-study differences
259 in sampling methods and taxa studied may lead to different levels of zero- and one-inflation,
260 we modelled the zero- and one-inflation parameters of the model with study-level random inter-
261 cepts (Equation 2). Furthermore, we included the geographical distance between site pairs as
262 an explanatory variable to account for the fact that nearby assemblages are expected to share
263 more species than distant assemblages. Similarly to Newbold et al. (2016), we randomised
264 the order of pairwise comparisons of sites 100 times, which obtained 100 instantiations of our
265 dataset, and thus 100 individual model fits. For each model, we sampled 100 predictions from
266 the posterior distribution, keeping geographical distance fixed at the mean. Median values and

267 95% credible intervals were then computed over the full range of the 10,000 (100 models \times 100
268 posterior samples) predictions.

$$\begin{aligned} & \text{Dissimilarity} \sim \text{Land use pair} \times \text{Region} + \text{GeoDistance} + (1|\text{Study}) \\ & \text{beta}(\text{zero inflation}) \sim (1|\text{Study}) \\ & \text{beta}(\text{one inflation}) \sim (1|\text{Study}) \end{aligned} \tag{2}$$

269

270 **2.5 Range extent**

271 We computed range extent estimates for arthropods in Europe and North America from ob-
272 servation data from the Global Biodiversity Information Facility (GBIF, 2024a,b), which we
273 filtered to include only presences and exclude fossil and living specimens (e.g., from zoos). This
274 was done by computing a convex hull around the observation points using the R package `con-`
275 `caveman` (Gombin et al., 2020, version 1.1.0), and then summing the total area of the hull using
276 `terra` (Hijmans, 2024, version 1.8.60). Range extent examples for four selected species are
277 provided in Fig. S16. We obtained range extents for 3579 out of 4098 arthropods from the
278 PREDICTS database for Europe (297,672 records), and 1859 out of 2141 arthropods for North
279 America (93,644 records). We divided these data into two categories, based on the percentiles
280 within each region: Narrow range (percentiles 0–50%; 1790 species and 124,953 records in Eu-
281 rope; 930 species and 46,139 records in North America) and wide range (percentiles 50–100%;
282 1789 species and 172,719 records in Europe; 929 species and 47,505 records in North America).
283 Diversity metrics (richness, abundance and Simpson’s index) were then modelled for each region
284 individually with linear mixed effects models of land use and range extent as interactive fixed
285 effects (Eq. 3).

$$\text{Diversity} \sim \text{Land use} \times \text{Range extent} + (1|\text{Study}/\text{Block}) \tag{3}$$

286

287 **2.6 Habitat specificity**

288 We obtained data on habitat specificity for European arthropods from the European habitat as-
289 sessments of the International Union for Conservation of Nature (IUCN) Red List of Threatened
290 Species (IUCN (International Union for Conservation of Nature), 2024). Habitat requirements
291 are unknown for the majority of arthropod species, even in well-studied regions like Europe and

Table 2: Weighting table used to estimate habitat specificity from IUCN habitat types. The habitat specificity score HSS is computed as the sum of all weights of habitat types. A high score implies low habitat specificity, and vice versa.

| Suitability | Major importance | Weight |
|---------------------|-------------------|--------|
| Suitable | Yes | 1.0 |
| Suitable | No, Unknown or NA | 0.5 |
| Marginal or Unknown | - | 0.3 |

292 North America, which meant that we were able to obtain habitat assessments for only 744 out
 293 of 4098 arthropods from the PREDICTS database for Europe (82,696 records). Analogously
 294 to Sykes et al. (2020), we then computed a habitat specificity score HSS as a weighted sum of
 295 the number of habitat types recorded as being used by a species (Table 2). Due to a scarcity of
 296 IUCN habitat assessments for North America, we obtained North American assessments from
 297 the NatureServe database instead (NatureServe, 2024). This resulted in assessments for 377 out
 298 of 2141 arthropods (14,024 records). In this case, we computed HSS by attributing unit weight
 299 to each habitat type as the NatureServe assessments do not include measures of 'suitability' or
 300 'major importance'. We divided habitat specificity into two categories: High specificity (HSS
 301 percentiles 0–50%) and low specificity (HSS percentiles 50–100%). Although percentiles for
 302 Europe and North America are computed from different data sources, we observe high simi-
 303 larity between HSS density plots (Fig. S18) and thus expect a comparison of distributional
 304 percentiles to be sensible. Finally, diversity metrics were modelled for each region individually
 305 with linear mixed models of land use and habitat specificity as interactive fixed effects (Eq. 4).

$$\text{Diversity} \sim \text{Land use} \times \text{Habitat specificity} + (1|\text{Study/Block}) \quad (4)$$

306

307 2.7 Non-native species

308 Data on non-native species in Europe were obtained from the European Alien Species Informa-
 309 tion Network (EASIN; European Commission - Joint Research Centre, 2024), which includes
 310 all species that are non-native to Europe as determined by the European Commission. These
 311 species are not necessarily invasive (i.e., causing harm by outcompeting native species, altering
 312 habitats, or spreading disease in a new environment), but can become invasive and thus detri-
 313 mental to local ecosystems once established. In total, 3389 arthropods species were extracted

314 from the database. Following the extraction from EASIN, we cross-referenced the species names
315 with GBIF to obtain a list of known synonyms for each species. Comparing the full list of
316 species names and synonyms with species records in the PREDICTS database for Europe, we
317 then determined which species in the database are non-native to Europe. In this way, 209 out
318 of the 4098 European arthropods in our subset of the PREDICTS database were identified as
319 non-native (14,000 records). We note that some of the species sampled in PREDICTS and not
320 identified as non-native may actually be non-native, but not recorded in EASIN. Moreover, since
321 the EASIN classifications treat species as either native or non-native to Europe as a whole, we
322 were not able to address cases where species are native in some parts of Europe but non-native
323 in other parts.

324 For North America, data on non-native species were obtained from the Global Register of
325 Introduced and Invasive Species (GRIIS; Pagad et al., 2018). Extracting data from the contigu-
326 ous United States (Simpson et al., 2023b), Alaska (Simpson et al., 2023a) and Canada (Pagad,
327 2022) obtained 3902 arthropod species. After cross-referencing with GBIF and obtaining syn-
328 onyms, 73 out of the 2141 North American arthropods our subset of the PREDICTS database
329 were identified as non-native (4659 records).

330 We computed the proportion of non-native species in a site 1) as the proportion of non-
331 native species to the total number of species and 2) as the proportion of the total abundance of
332 non-native species to total abundance of all species in a site. We modelled these variables as a
333 function of land use using a generalised linear mixed effects model with a zero-and-one-inflated
334 beta distribution (Eq. 5). Similarly to our model of compositional similarity, we modelled the
335 zero- and one-inflation parameters of the model with study-level random intercepts.

$$\begin{aligned} \text{Proportion} &\sim \text{Land use} + (1|\text{Study}/\text{Block}) \\ \text{beta}(\text{zero inflation}) &\sim (1|\text{Study}) \\ \text{beta}(\text{one inflation}) &\sim (1|\text{Study}) \end{aligned} \tag{5}$$

336

337 2.8 Model testing

338 We inspected model fit by generating observed versus fitted plots and posterior predictive checks,
339 and used Moran's I test to check for residual spatial autocorrelation. We ensured the converge
340 of `brms` models by inspecting R-hat values and visually ensuring that Markov chains were well-
341 mixed. Additionally, we inspected the empirical cumulative distribution function (ECDF) of

342 probability integral transform (PIT) values, which can be interpreted as the Bayesian counter-
343 part to the QQ-plot, to check whether predictive distributions were well-calibrated (deviations
344 from the diagonal can indicate misfit, bias and over- or underdispersion). Information on good-
345 ness of fit are summarised for all main models in Table S2 of Supplementary Information A,
346 while individual model diagnostic plots and estimated parameters are available in Supplemen-
347 tary Information B for complete transparency.

348 **3 Results**

349 **3.1 Regional responses to land use**

350 Changes in arthropod diversity from primary to secondary vegetation and agricultural land use
351 are significantly negative globally and for most individual regions, with the notable exception
352 of Europe, where diversity responses are not significantly different from primary vegetation, or
353 even significantly positive in the case total abundance change in minimal-use agriculture (Fig.
354 1). Although diversity in Europe decreases marginally with increasing agricultural use-intensity,
355 changes remain statistically indistinguishable from the primary vegetation baseline (zero) and
356 are substantially elevated over global responses. The fact that a large number of PREDICTS
357 sites stem from agricultural sites in Europe (Fig. 1; samples size), while the samples size for
358 primary vegetation in Europe is comparable to that of other regions (Tables S3 to S5), lends
359 confidence to the accuracy of these responses.

360 We observe the most consistently negative responses of diversity with agricultural use-
361 intensity for North America, South and Central America, sub-Saharan Africa and South and
362 Southeast Asia. Responses of species richness and total abundance are especially negative
363 in intense-use agriculture in sub-Saharan Africa. Moreover, responses in sub-Saharan Africa
364 are significantly more negative than global responses, which implies that estimating land use
365 impacts of agriculture on biodiversity in the region with global PREDICTS responses would
366 severely underestimate impacts. In North America, declines in evenness (Simpson's diversity in-
367 dex) tend to be more extreme than declines in species richness, which suggests that in addition
368 to substantial changes in species numbers, assemblages in agriculture are disproportionately
369 dominated by few species. To a lesser extent, the same can be observed for Europe. For all
370 other regions, declines in Simpson's diversity tend to be less extreme than declines in species
371 richness. Diversity declines with agricultural use-intensity are less consistent in Australasia,

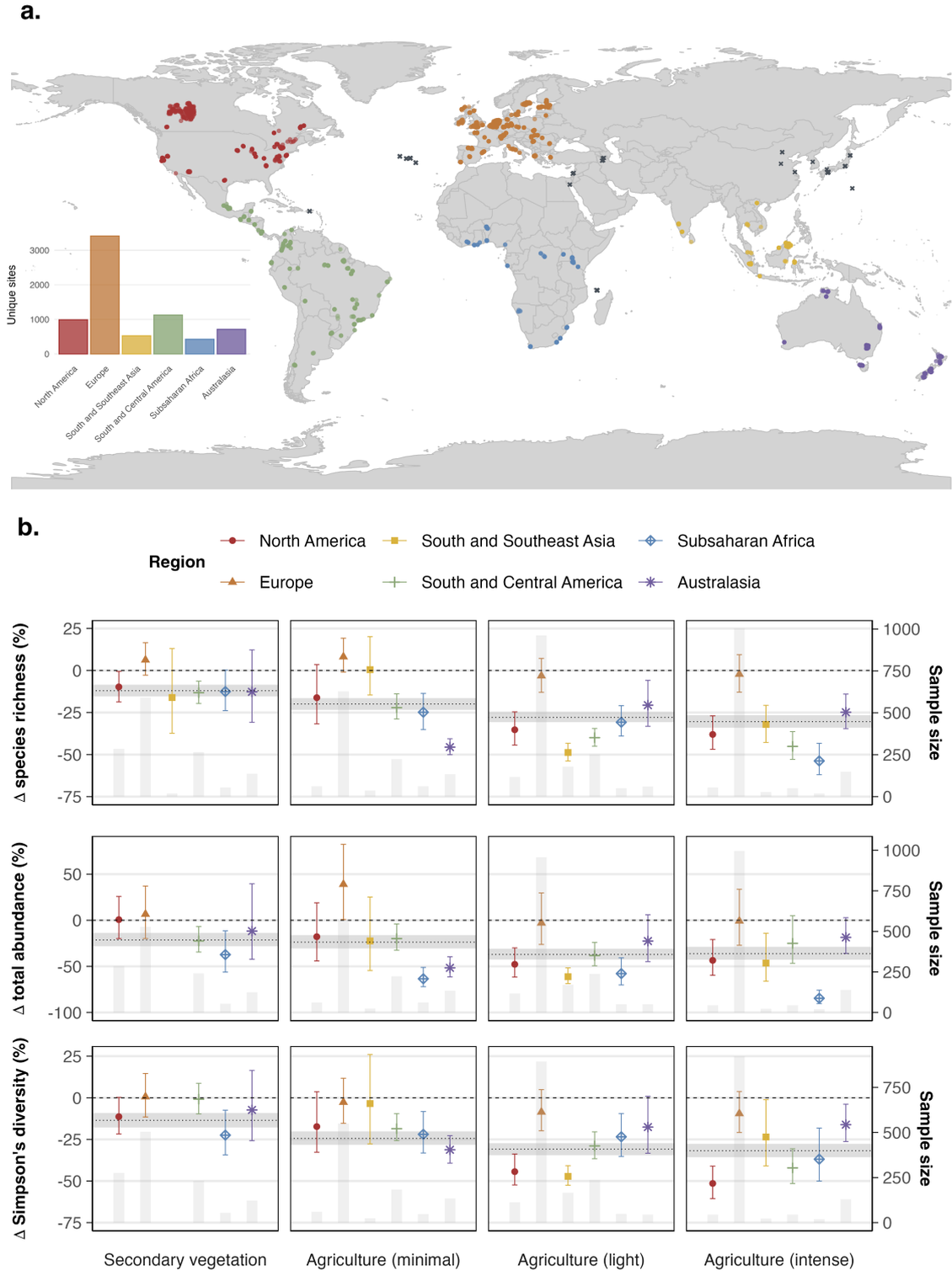


Figure 1: (a) Sites with arthropod records from the PREDICTS database used in this study. Sites discarded from the regional analysis (stemming from regions that lacked enough data and from small islands that were removed following a sensitivity analysis) are displayed as crosses. (b) Responses of species richness, total abundance and Simpson's diversity index per land use and region, relative to primary vegetation. Points indicate median modelled responses and error bars represent 95% credible intervals. Dotted lines indicate global responses, which include sites discarded from the regional analysis (a), with 95% credible intervals shaded. The right y-axis shows sample sizes (number of sites), plotted as grey bars. Only model coefficients computed from at least 10 sites are shown. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

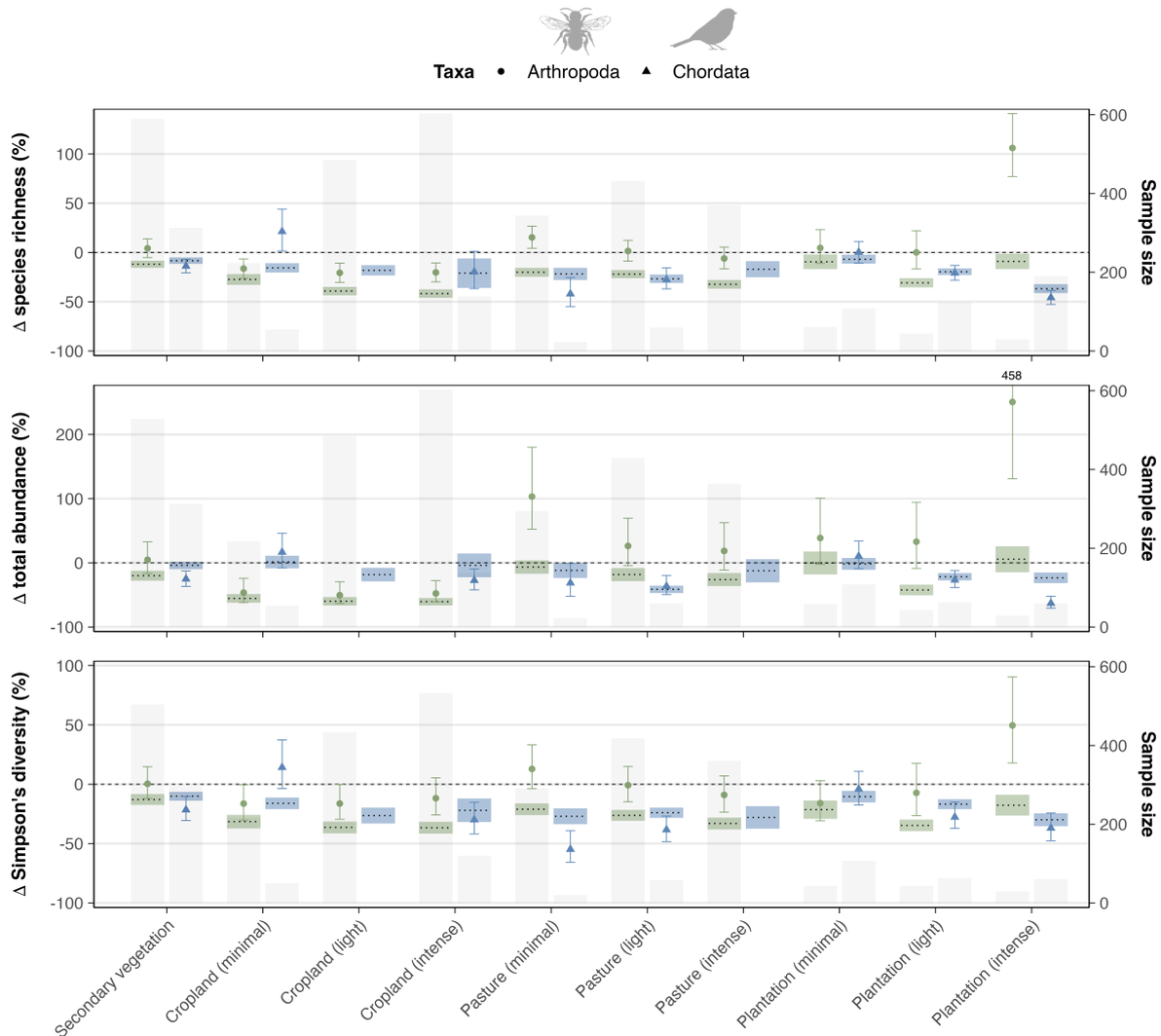


Figure 2: Responses of species richness, total abundance and Simpson's diversity index, relative to primary vegetation, per agricultural land-use type & intensity in Europe, for arthropods (circles), vertebrates (triangles) and plants (squares). Dotted lines indicate global responses, with 95% credible intervals shaded. Points indicate median modelled responses and error bars represent 95% credible intervals. The right y-axis shows sample sizes (number of sites), plotted as grey bars. Only model coefficients computed from at least 10 sites are shown.

372 where declines in species richness are significantly negative but changes in total abundance tend
 373 to be marginal and non-significant (with the exception of minimal-use agriculture).

374 Although biodiversity responses to agricultural intensification in Europe show little overall
 375 change from primary vegetation (Fig. 1), species richness and total abundance are nevertheless
 376 significantly reduced in European cropland (Fig. 2). Positive responses in Europe are, instead,
 377 limited to minimal-use pasture and intense-use tree plantations. Nevertheless, responses in all
 378 individual agricultural land use types (cropland, pasture and tree plantation) and intensities
 379 are consistently elevated over global responses (Fig. 2). Interestingly, responses of arthropod
 380 diversity in Europe are not mirrored by vertebrates (phylum Chordata; Fig. 2). In fact,

381 responses of vertebrates in Europe tend to match global responses very closely (with the notable
382 exception of minimal-use cropland).

383 **3.2 Natural habitat availability**

384 The availability of natural habitat surrounding PREDICTS primary vegetation sites in Europe is
385 substantially lower compared to South and Southeast Asia and South and Central America, but
386 is comparable to North America, sub-Saharan Africa and Australasia (Fig. 3b). Natural habitat
387 availability tends to decrease with increasing agricultural use-intensity in all regions. At the
388 same time, the %HANPP of the natural habitat surrounding primary vegetation sites is notably
389 higher in Europe than in most other regions except sub-Saharan Africa. In fact, the %HANPP of
390 natural habitat surrounding European primary vegetation is comparable to that of secondary
391 vegetation and agricultural sites. Furthermore, natural habitat tends to be less aggregated
392 (more fragmented) in Europe (Fig. 3c). Habitat fragmentation is particularly evident when
393 computed for higher natural habitat thresholds, which suggests that it is particularly dense
394 patches of natural vegetation (high coverage of natural habitat) that are fragmented. Results
395 for other habitat connectivity metrics such as mean patch size and patch density are similar
396 although differences between Europe and other regions are smaller (Fig. S4).

397 It is clear, however, that the sampling of primary vegetation sites in PREDICTS is subject
398 to considerable bias (Fig. 3a). While primary vegetation sites in Europe and the Americas
399 are sampled relatively representatively across the region, South and Southeast Asia and sub-
400 Saharan Africa lack samples from important biodiversity hotspots such as New Guinea and the
401 Congo Basin, and large tracts of wilderness in central (outback) Australia are data-deficient
402 (Fig. S5). Compared to regional averages, natural habitat surrounding PREDICTS primary
403 vegetation sites tends to considerably under-represent wilderness areas while over-representing
404 forestry areas (Fig. S6). In Europe, it is particularly evident that the proportion of wilderness
405 making up natural habitat in the region is lower than in other regions (restricted primarily to
406 patches in the Alps, Iceland and northern Scandinavia; Fig. S5), and that PREDICTS primary
407 vegetation sites severely undersample this wilderness (0.11% of natural habitat surrounding
408 primary vegetation sites within a 5km radius is wilderness versus a regional average of 8.36%;
409 Fig. S6).

410 Regional averages of the %HANPP of natural habitat are highest in Europe and South and
411 Southeast Asia, but whereas the %HANPP of natural habitat surrounding PREDICTS primary

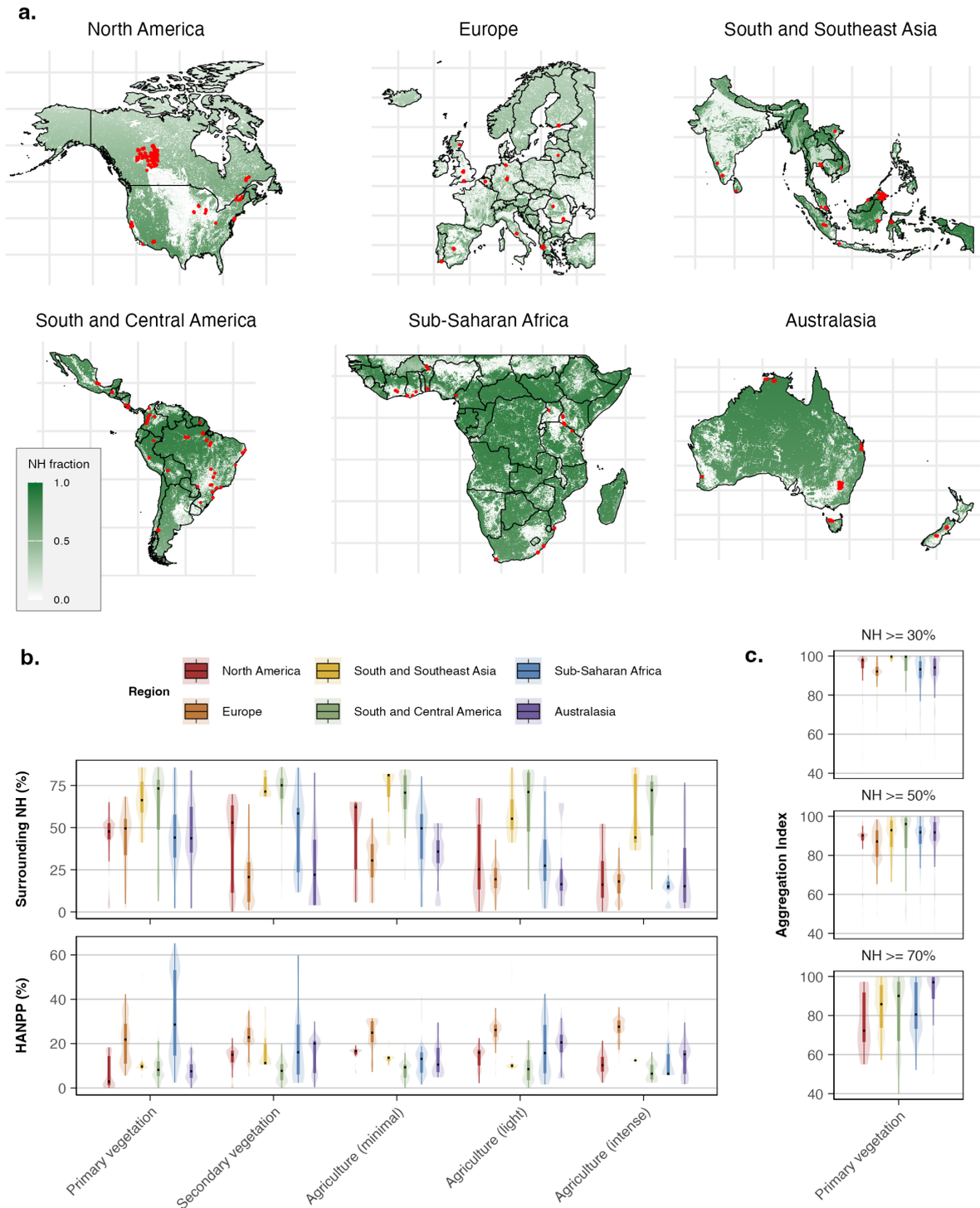


Figure 3: (a): Natural habitat (wilderness, forestry areas and open woodland grazing land) availability in each region, from Matej et al. (2025) for the year 2005, with the primary vegetation sites from PREDICTS that were used in this study plotted as red points. (b): The percentage of surrounding natural habitat and HANPP as a percentage of the potential NPP of natural habitat, within a 5km-radius of PREDICTS sites, separated by land use in each region. (c): Degree of connectivity of natural habitat surrounding primary vegetation sites within a larger 10km-radius buffer, estimated with the aggregation index (He et al., 2000) and shown for three different thresholds of natural habitat. Europe is missing from the 70% threshold as there were no PREDICTS primary vegetation sites in Europe with natural habitat exceeding 70% anywhere within a 5-km buffer. (b) and (c) show violin plots of the data distributions, overlaid on box and whisker plots with the median values plotted as black lines. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

412 vegetation sites in Europe is considerably elevated above the regional average, the opposite is
413 true in South and Southeast Asia (Fig. S7), suggesting that PREDICTS primary vegetation
414 sites are biased towards higher use-intensity in Europe and towards lower use-intensity in South
415 and Southeast Asia. Moreover, we observe a strong bias of PREDICTS primary vegetation
416 sites in sub-Saharan Africa towards higher %HANPP and thus use-intensity (particularly for
417 forestry), such that %HANPP surrounding primary vegetation sites is comparable with Europe,
418 although regional averages are considerably lower. These sampling biases help explain why the
419 average %HANPP surrounding a site tends to be higher in primary vegetation sites in sub-
420 Saharan Africa than in Europe (Fig. 3b). In the Americas, %HANPP values of natural habitat
421 surrounding PREDICTS primary vegetation samples mirrors regional averages fairly well, but
422 wilderness areas are nevertheless undersampled and forestry areas oversampled (Fig. S6).

423 3.3 Species Composition

424 In Europe, compositional dissimilarity between minimal-use agriculture and natural vegeta-
425 tion (primary and secondary vegetation) is 41.9% (95% credible interval: [29.8, 55.7]) higher
426 than the baseline dissimilarity between natural vegetation and other natural vegetation sites
427 (Fig. 4). This dissimilarity is overwhelmingly the result of species turnover, which suggests
428 that arthropod assemblages in minimal-use agriculture tend to consist of novel, possibly more
429 disturbance-tolerant species not found in natural vegetation. Fitting the models against indi-
430 vidual land-use types in Europe shows that this turnover stems predominantly from cropland
431 and pasture sites, while higher compositional similarity is observed between natural vegetation
432 and tree plantations (Fig. S10). At increasing agricultural use-intensities in Europe, we ob-
433 serve more homogeneity between agriculture and natural vegetation as total dissimilarity and
434 turnover drop to comparable levels as the natural vegetation baseline. However, sample sizes of
435 dissimilarity between natural vegetation and intense-use agriculture tend to be very small (in
436 all regions), which means that these responses are uncertain and unlikely to be representative.

437 In comparison, increasing agricultural intensification in North America is not associated
438 with significant differences in overall compositional dissimilarity (although this may partly re-
439 flect limited sample sizes). Nevertheless, median nestedness in both light- and intense-use
440 agriculture is considerably elevated above the baseline, indicating that compositional differ-
441 ences primarily reflect species loss rather than turnover. This mirrors significant declines in
442 richness observed with agricultural intensification (Fig. 1) and thus suggests that assemblages

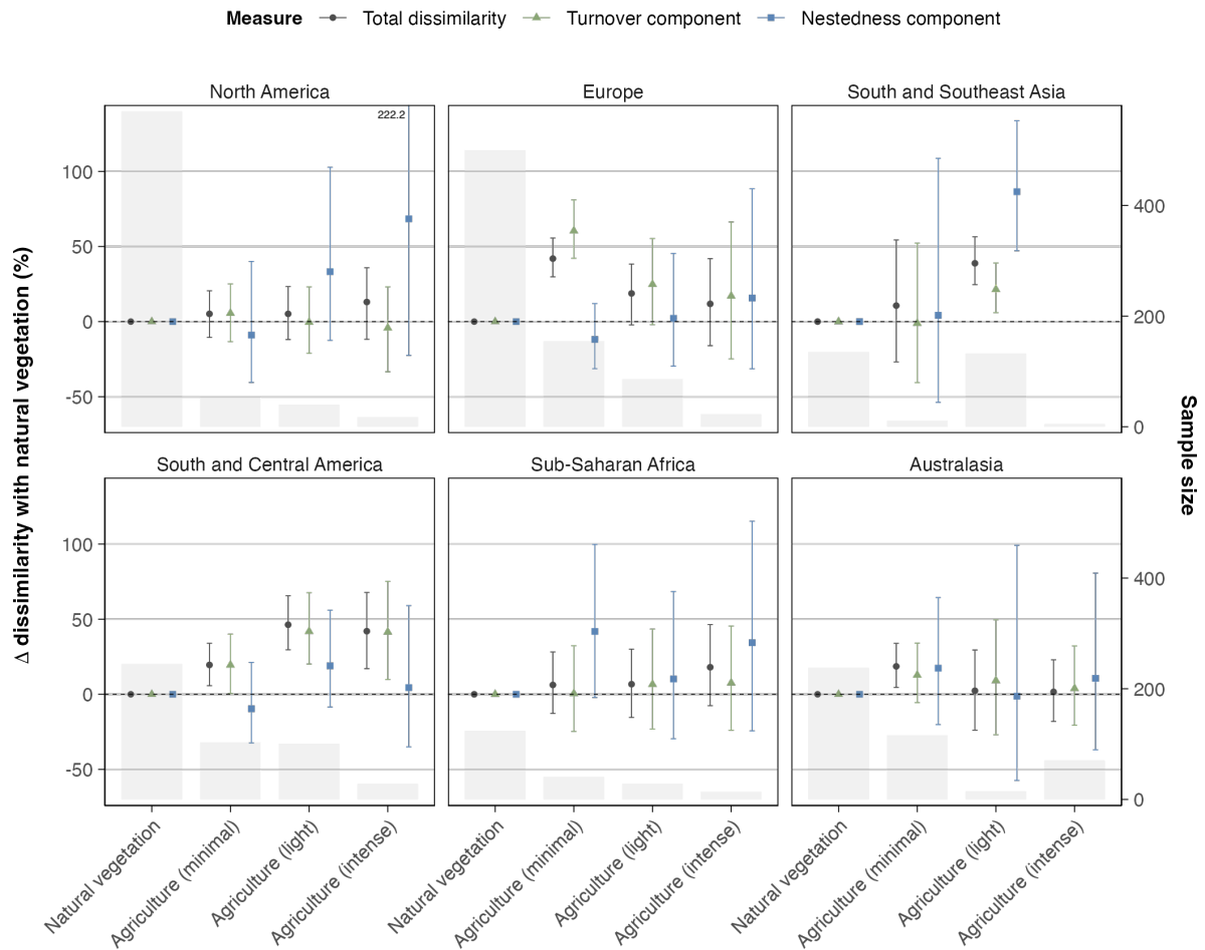


Figure 4: Compositional dissimilarity (Sørensen's Index) of each land use with natural vegetation (primary and secondary vegetation) per region, shown relative to the dissimilarity of natural vegetation with other natural vegetation sites (%). A more positive relative response of turnover indicates that dissimilarity is driven more by species replacement, whereas a more positive relative response of nestedness indicates that dissimilarity is driven more by species loss (i.e., the extent to which composition in one land use is a subset of the other). Points indicate median modelled responses and error bars represent 95% credible intervals. The right y-axis shows sample sizes (number of sites), plotted as grey bars. Only model coefficients computed from at least 10 sites are shown.

443 in light- and intense-use agriculture tend to represent species-poor subsets of those found in nat-
444 ural vegetation. Increased compositional dissimilarity comparable to that found in European
445 minimal-use agriculture can also be observed in South and Southeast Asia (light-use agricul-
446 ture) and in South and Central America (light- and intense-use agriculture), each coinciding
447 with substantial biodiversity declines (Fig. 1). The dominant components of this dissimilarity
448 differ, however: in South and Southeast Asia, dissimilarity is driven primarily by nestedness,
449 whereas in South and Central America it is dominated by turnover. Nevertheless, where sample
450 sizes are larger, relative responses of species richness and total abundance to agricultural land
451 use tend to be positively associated with relative responses of species turnover and negatively
452 with nestedness (Fig. S11 and Fig. S12).

453 **3.4 Effects of range extent, habitat specificity and non-native species**

454 In Europe, we observe highly significant differences between the responses of species richness
455 and total abundance of narrow- and wide-ranging species, and species with high- and low habitat
456 specificity (Fig. 5): Wide-ranging species and habitat generalists (low habitat specificity) are
457 associated with substantial increases in agricultural sites relative to primary vegetation, while
458 narrow-ranging species and habitat specialists (high habitat specificity) are associated with
459 considerable declines. In North America, on the other hand, differences in responses between
460 these groups of species is negligible: Both narrow- and wide-ranging species, and both habitat
461 generalists and specialists are associated with declines in species richness and total abundance
462 in agriculture. Further analysis per individual land-use types in Europe, which were estimable
463 for cropland and pasture, highlights that differences in responses tend to be considerably larger
464 in cropland than in pasture (Fig. S14), which suggests that it is particularly in cropland that
465 wide-ranging species and habitat generalists are able to outcompete narrow-ranging species and
466 habitat specialists.

467 In both regions, the proportion of non-native species as a fraction of species richness is
468 slightly elevated in light- and intense-use agriculture over primary vegetation, but this is only
469 significant in North America (Fig. 5, right and top). Slight increases over primary vegetation
470 can also be observed for non-native proportions as a fraction of total abundance, but these are
471 not significant in either region (Fig. 5, right and bottom). Indeed, for North America, it should
472 be noted that sample sizes are small and thus 95% credible intervals can be large. In absolute
473 terms, however, the proportion of non-native species is consistently higher in Europe than in

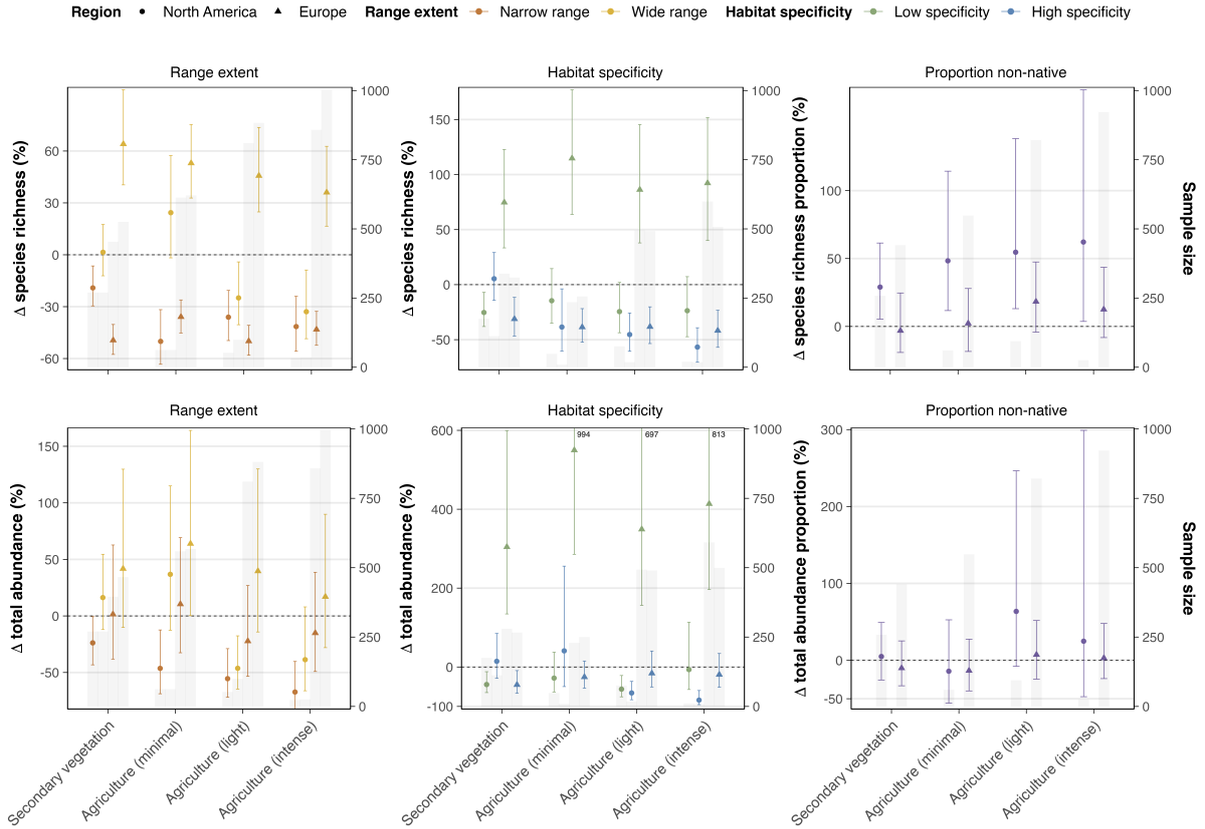


Figure 5: Comparison of European and North American arthropod diversity responses (species richness and total abundance) to agricultural land use, relative to primary vegetation. Shown are the responses of species with narrow and wide range extents (left; range extents below or above the median, respectively), species with low or high habitat specificity (centre; habitat specificity scores above or below the median, respectively), and the relative proportion of non-native species in a given site, either as a fraction of species richness (top) or of total abundance (bottom). Points indicate median modelled responses and error bars represent 95% credible intervals. The right y-axis shows sample sizes (number of sites). Only model coefficients computed from at least 10 sites are shown.

474 North America, particularly in primary vegetation, although credible intervals largely overlap
475 (Fig. S13).

476 **4 Discussion**

477 This analysis has revealed important differences in the biodiversity responses of arthropods to
478 agricultural land use and intensity between six major world regions. Our results show that in
479 Europe, overall arthropod diversity in agricultural sites is comparable to primary vegetation,
480 and tends to be higher in minimal-use agriculture, while other regions, and global averages,
481 tend to show significant declines (Fig. 1). Responses in Europe are consistently more positive
482 than global averages, but non-negative responses are limited to pasture and plantation sites,
483 while diversity is nevertheless significantly reduced in cropland (Fig. 2). Similar variation
484 between land use types is likely to exist in other regions, but a lack of data currently prohibits
485 such comparisons. The effect sizes of the responses in intense-use plantation in Europe are
486 surprisingly large, although it should be noted that the data underpinning these responses stem
487 from a single study (on micro-arthropods sampled from soil cores in the Netherlands), which
488 suggests it is unlikely to be representative. It is notable, however, that almost half (42.3%) of
489 PREDICTS data on arthropods in Europe were sampled from soil cores, but none of these data
490 were sampled in primary vegetation, which may draw into question the extent to which responses
491 between primary vegetation and other land-use types can be reliably compared. Refitting our
492 models on PREDICTS data without soil core samples did not meaningfully change biodiversity–
493 land use responses, however (Fig. S23 and Fig. S24). Moreover, it is interesting that European
494 biodiversity responses of vertebrates differ substantially from those of arthropods (Fig. 2). We
495 did not investigate the responses of vertebrates further in this work, and thus can only speculate
496 on possible reasons for this disparity, but possible explanations may be linked to the ability of
497 arthropods to successfully exploit microhabitats in agricultural landscapes (García-Tejero and
498 Taboada, 2016) or to characteristically short lifespans and rapid reproduction rates, which allow
499 populations to rebound quickly after declines.

500 The result that biodiversity change in Europe is more negative in cropland than in pasture
501 or plantation forest mirrors previous findings of land-use impacts on insects in Germany, where
502 changes in biomass and richness were found to be most negative in arable fields, followed by
503 meadow and forest, and, overall, agricultural landscapes were found to be higher in biomass

504 (although not richness) than semi-natural ones (Uhler et al., 2021). Our finding of biodiversity
505 gains in minimal-use agriculture in Europe is thus not entirely unexpected. Indeed, it has long
506 been understood that low-intensity environmental disturbance can benefit ecological diversity,
507 as stipulated by the intermediate disturbance hypothesis (Connell, 1978). Indeed, mixed natu-
508 ral and agricultural landscapes typically boast a higher diversity of habitats than the original
509 vegetation – habitat diversity being one of strongest predictors of ecological diversity (Thomas,
510 2013). Nevertheless, it is unexpected that neither light nor intense-use agriculture in Europe
511 is associated with significant biodiversity declines. Moreover, it is note-worthy that responses
512 in North America are more similar to the predominantly tropical regions of South and Cen-
513 tral America, sub-Saharan Africa and South and Southeast Asia than to Europe, which draws
514 into question to what extent previous findings that land-use impacts on biodiversity are lower in
515 temperate than in tropical zones are the result of actual differences in climate rather than differ-
516 ences between Europe and other regions (Millard et al., 2021; Newbold et al., 2020; Outhwaite
517 et al., 2022).

518 Our regional analysis of natural habitat and %HANPP suggest that the representativeness
519 of primary vegetation sites sampled for arthropods in the PREDICTS database is relatively
520 poor in all investigated regions, showing bias towards low proportions of wilderness and high
521 %HANPP (use-intensity). This selection bias is likely the result of accessibility issues, given that
522 intact primary vegetation in wilderness areas is neither easy to sample, nor in close proximity
523 to other land-use types of interest (e.g., agricultural and urban areas). This suggests that, in
524 addition to real shifts in baseline biodiversity that may reduce differences in biodiversity between
525 natural and anthropogenic land use types, the compounding effect of sampling bias towards
526 more accessible, and thus often more degraded, primary vegetation suggests that biodiversity
527 estimates in primary vegetation are likely to be underestimates. This implies that estimates
528 of biodiversity change between primary vegetation and agricultural sites are also likely to be
529 underestimates (the only exception to this is South and Southeast Asia, where %HANPP of
530 natural habitat surrounding PREDICTS primary vegetation sites is considerably lower than
531 the regional average, and thus biodiversity in primary vegetation may be overestimated for the
532 region). An underestimation of biodiversity change for many of the world’s regions is concerning,
533 given that our estimates of richness and abundance change between primary vegetation and
534 intense-use agriculture are already high (e.g., median reductions of 53.5% in species richness and
535 84.6% in total abundance in sub-Saharan Africa; Fig. 1). This may have important implications

536 for previous studies of land-use impacts on biodiversity using the PREDICTS database, which
537 are typically measured relative to primary vegetation sites.

538 Our analysis of compositional dissimilarity shows that biodiversity gains in minimal-use agri-
539 culture in Europe coincide with significant species turnover compared with natural vegetation,
540 whereas dissimilarity between natural habitat and light- or intense-use agriculture is compara-
541 ble to that among natural habitats, suggesting that increasing agricultural use-intensity is not
542 necessarily associated with greater compositional divergence (Fig. 4). This demonstrates the
543 limitations of commonly used diversity metrics such as species richness and total abundance,
544 which may show little overall change despite important compositional changes (Fletcher Jr.
545 et al., 2025), but it also highlights that on average, assemblages found in natural vegetation
546 in Europe are no more compositionally dissimilar than intensively farmed agricultural land is
547 to natural vegetation, which suggests that baseline diversity in European primary vegetation
548 may be considerably degraded. Indeed, compared to North America, primary vegetation in Eu-
549 rope is more fragmented and subject to considerably higher %HANPP (Fig. 3). Furthermore,
550 proportions of non-native species, which tend to establish more easily in disturbed habitats
551 (Liu et al., 2023), are consistently (though not significantly) higher in primary vegetation in
552 Europe than in North America (Fig. S13). The investigation into European arthropod char-
553 acteristics shows that biodiversity gains in agriculture are predominantly associated with wide-
554 ranging habitat-generalists, while narrow-ranged species and habitat specialists are marked by
555 significant declines (Fig. 5). Taken together, this paints a picture of winners and losers, and
556 widespread ecosystem transformation, despite little overall change in biodiversity metrics such
557 as richness, abundance and evenness between primary vegetation and agricultural sites (Fig.
558 1). Indeed, as a result of widespread ecosystem transformation, primary vegetation may cease
559 to function as an ecologically informative baseline, and important declines in the diversity of
560 native habitat specialists may become masked by increases of (possibly non-native) widespread
561 habitat generalists.

562 As opposed to Europe, we observe large declines in arthropod diversity in light- and intense-
563 use agriculture in North America (Fig. 1), find no evidence of turnover between natural veg-
564 etation and agricultural sites (Fig. 4), and only a marginal effect of range extent and habitat
565 specificity in the region (Fig. 5). This suggests that North American ecosystems, natural
566 or anthropogenic, have possibly undergone less transformation, perhaps as a result of a more
567 recent history of land-use change. Interestingly, we did find significant increases in the pro-

568 portion of non-native species with increasing agricultural land-use intensity, which may drive
569 changes in species composition in the future. We do note, however, that the PREDICTS data
570 for North America contained records for only a very small number of the total number of non-
571 native arthropod species obtained for North America from the GRIIS database (73 out of 3902
572 species), which means that absolute model predictions of non-native proportions are almost
573 certainly underestimates. Similarly, we were only able to obtain data on habitat specificity for
574 very few species in North America, and these data were heavily biased towards Hymenopteran
575 species in both regions, which suggests that the presented biodiversity responses may not be
576 representative estimates for other arthropod orders (Fig. S19). Range extents were obtained
577 for a larger group of species, whose distributions of taxonomic orders better reflect the complete
578 distributions for each region (Fig. S2 and Fig. S17). It should also be noted that within-study
579 comparisons between natural vegetation and agricultural sites, from which compositional dis-
580 similarity was estimated, were scarce in all regions (Fig. 4). This is particularly evident for
581 Europe, where agricultural sites are otherwise very well sampled (Fig. 1) which highlights an
582 important data deficiency that will need to be addressed by future studies in order to allow for
583 more reliable estimates of compositional change between natural vegetation and anthropogenic
584 land-use types.

585 **5 Conclusion**

586 In this study, we have found considerable differences in the responses of arthropod diversity
587 to agricultural land use and intensity between six major world regions, highlighting that Eu-
588 rope exhibits minimal declines with agricultural intensification. As the best-sampled region in
589 our dataset with major influence over global biodiversity statistics, which go on to be used in
590 integrated assessments to estimate the impacts of land-use change on biodiversity and ecosys-
591 tem provisions, it is important to understand these findings better. We have found evidence
592 consistent with the hypothesis that a long history of land-use change in Europe and associated
593 deforestation of wild forests has resulted in natural vegetation that is composed of very little
594 true wilderness, and is more fragmented and subject to higher use-intensity than in other re-
595 gions. Our results support the hypothesis that this history of land-use change and agricultural
596 expansion has primarily benefited wide-ranging habitat generalists and non-native species, while
597 specialists of the forested natural habitat that was once widely available on the landmass have

598 been largely filtered out. At the same time, we have found that primary vegetation sites sampled
599 for arthropod records in the PREDICTS database tend to be biased towards natural habitat
600 that is subject to considerably higher use-intensity than regional averages and is composed of
601 disproportionately low proportions of wilderness area. With the exception of South and South-
602 east Asia, these biases suggest that biodiversity estimates in primary vegetation are likely to be
603 underestimates, and thus changes in biodiversity relative to primary vegetation are also likely
604 to be underestimates. Despite suspected shifts in baselines and sampling biases, our results do
605 highlight the possible value that minimal-use agriculture in Europe, such as organic, regenera-
606 tive and mixed-cropping systems may hold to arthropod diversity, where large species turnover
607 compared with natural vegetation points to distinct assemblages that may have adapted well
608 to low-intensity disturbance.

609 Our results underscore the complexity and nuance of modelling threats to local biodiver-
610 sity. In addition, the heterogeneity in responses between regions and species groups echoes
611 the findings of mixed biodiversity trends worldwide (Johnson et al., 2024; Klink et al., 2020;
612 Martins et al., 2022) and reinforces the picture that ecosystem change in the Anthropocene
613 can boost local biodiversity (Thomas, 2013), at the same time as species are being lost glob-
614 ally at an unprecedented rate (Dirzo et al., 2014; Wagner et al., 2021). In Europe, changes in
615 species composition appear to have resulted in novel arthropod assemblages that may prove to
616 be resilient to low-intensity agricultural disturbance and well-adapted to open and mixed-use
617 landscapes. For global analyses of impacts on biodiversity, which tend to be biased towards
618 Europe, our results imply, however, that global estimates of biodiversity change likely underes-
619 timate agriculture-driven declines in most of the world’s regions, while overestimating declines
620 for the European continent. As such, region-specific analyses appear necessary to avoid mask-
621 ing important regional variation. This aim is conditional, however, on a considerable increase
622 in systematic biodiversity sampling in many of the world’s regions, since most are currently
623 severely undersampled, hindering robust statistical analysis.

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Supplementary Information A

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A Tables

A.1 PREDICTS definitions of land use types and intensities

Table S1: Definitions of land use types and levels of intensity used by PREDICTS database (Hudson et al., 2017).

| Land use | Definition | Minimal use | Light use | Intense use |
|--------------------|--|--|--|--|
| Primary vegetation | <p>PRIMARY vegetation (forest or non-forest) is native vegetation that is not known or inferred to have ever been completely destroyed, before the year in which the biodiversity was sampled, by human actions or by extreme natural events that do not normally play a role in ecosystem dynamics. Sites where primary vegetation has been destroyed by natural events that are part of the normal ecosystem dynamic (e.g. fire in Mediterranean ecosystems) remain as primary vegetation provided that colonization from adjacent habitat and regeneration is possible. Sites in urban and suburban settings where the vegetation has never been completely destroyed should also be classed as primary vegetation. Synonyms include “ancient woodland”, “old-growth forest” and “natural grassland” (unless any indication is given of a previous land use class). Primary vegetation can be used by people (e.g., fruit harvesting, selective logging). Primary vegetation includes sites where people have tried to restore degraded habitat, so long as the native vegetation has never been destroyed. Sites where the original vegetation is known to have been completely destroyed should not be classed as primary vegetation.</p> | <p>Any disturbances identified are very minor (e.g., a trail or path) or very limited in the scope of their effect (e.g., hunting of a particular species of limited ecological importance).</p> | <p>One or more disturbances of moderate intensity (e.g., selective logging) or breadth of impact (e.g., bushmeat extraction), which are not severe enough to markedly change the nature of the ecosystem. Primary sites in suburban settings are at least Light use.</p> | <p>One or more disturbances that is severe enough to markedly change the nature of the ecosystem; this includes clear-felling of part of the site too recently for much recovery to have occurred. Primary sites in fully urban settings should be classed as Intense use.</p> |

Continued on next page

| Land use | Definition | Minimal use | Light use | Intense use |
|----------------------|---|---|--|---|
| Secondary vegetation | SECONDARY vegetation is where the original primary vegetation was completely destroyed. This could be by human actions (including fire), and includes where sites are recovering to a natural state following a period of human-dominated land use (cropland, plantation forest, pasture or urban). Also counted as secondary are places where natural events (fires, storms etc.) have destroyed the vegetation, but not where the vegetation is naturally maintained by fire (such as climatically Mediterranean systems), which would be primary. Secondary vegetation includes areas where humans have made an active attempt (through planting etc.) to return an area where the natural vegetation was previously destroyed to a more natural state. Synonyms include “old-field”, “abandoned” and “fallow”. Although not managed as intensively as the human-dominated classes, such sites can be used by people in much the same way as primary vegetation sites. | As for Primary Vegetation-Minimal use | As for Primary Vegetation-Light use | As for Primary Vegetation-Intense use |
| Cropland | CROPLAND is land that people have planted with herbaceous crops, even if these crops will be fed to livestock once harvested. Sites described as “fields”, “arable”, “ploughed” or “tilled” all qualify as cropland. If cropland is abandoned, including temporary abandonment (i.e. fallow), it becomes Secondary vegetation. | Low-intensity farms, typically with small fields, mixed crops, crop rotation, little or no inorganic fertiliser use, little or no pesticide use, little or no ploughing, little or no irrigation, little or no mechanisation. | Medium intensity farming, typically showing some but not many of the following: large fields, annual ploughing, inorganic fertiliser application, pesticide application, irrigation, no crop rotation, mechanisation, monoculture crop. Organic farms in developed countries often fall within this category, as may high-intensity farming in developing countries. | High-intensity monoculture farming, typically showing many of the following features: large fields, annual ploughing, inorganic fertiliser application, pesticide application, irrigation, mechanisation, no crop rotation. |

Continued on next page

| Land use | Definition | Minimal use | Light use | Intense use |
|-------------------|---|---|--|---|
| Pasture | PASTURE is land where livestock is known to be grazed regularly or permanently. The plant species may be predominantly native (as in rangelands) or strongly associated with humans (as in European-style pastures). Land that is planted with a crop for harvesting and subsequently feeding to livestock is Cropland, not Pasture. However, if vegetation is planted and livestock are grazed directly on this vegetation, then this does count as pasture. | Pasture with minimal input of fertiliser and pesticide, and with low stock density (not high enough to cause significant disturbance or to stop regeneration of vegetation). | Pasture either with significant input of fertiliser or pesticide, or with high stock density (high enough to cause significant disturbance or to stop regeneration of vegetation). | Pasture with significant input of fertiliser or pesticide, and with high stock density (high enough to cause significant disturbance or to stop regeneration of vegetation). |
| Plantation forest | PLANTATION FOREST applies to previously cleared areas that people have planted with crop trees or crop shrubs for commercial or subsistence harvesting of wood and/or fruit. The species planted may or may not be native. Planting an area with native woody plants for habitat restoration rather than for goods does not constitute plantation forest; rather this would be secondary vegetation with the stage dependent on the architectural complexity (see definitions under secondary vegetation). Likewise, natural regrowth is not a plantation forest, even if the regrowth will be harvested. If plantation forest is abandoned, it becomes Secondary vegetation – and may have a greater architectural complexity than the time since abandonment might suggest. | Extensively managed or mixed timber, fruit/coffee, oil-palm or rubber plantations in which native understorey and/or other native tree species are tolerated, which are not treated with pesticide or fertiliser, and which have not been recently (< 20 years) clear-felled. | Monoculture fruit/coffee/rubber plantations with limited pesticide input, or mixed species plantations with significant inputs. Monoculture timber plantations of mixed age with no recent (< 20 years) clear-felling. Monoculture oil-palm plantations with no recent (< 20 years) clear-felling. | Monoculture fruit/coffee/rubber plantations with significant pesticide input. Monoculture timber plantations with similarly aged trees or timber/oil-palm plantations with extensive recent (< 20 years) clear-felling. |

A.2 Model diagnostic summaries

Table S2: Summaries of model fit (R^2) and convergence (R-hat values) for the main models fitted in this study.

| Model | Distribution family | Model equation | Conditional R^2 | Marginal R^2 | R-hats | nSites |
|---|----------------------------|---|-------------------------------|-------------------------------|-----------|--------|
| Global model of species richness | zero_inflated_-negbinomial | Species_richness \sim UI2 + (1 SS/SSB), $z_i \sim 1$ | 0.905 (95% CI [0.893, 0.912]) | 0.005 (95% CI [0.003, 0.007]) | 1.00-1.01 | 7604 |
| Regional model of species richness | zero_inflated_-negbinomial | Species_richness \sim UI2 * Region + (1 SS/SSB), $z_i \sim 1$ | 0.918 (95% CI [0.908, 0.925]) | 0.025 (95% CI [0.006, 0.059]) | 1 | 7188 |
| Global model of total abundance | gaussian | LogAbund \sim UI2 + (1 SS/SSB) | 0.905 (95% CI [0.903, 0.907]) | 0.005 (95% CI [0.003, 0.006]) | 1.00-1.01 | 7122 |
| Regional model of total abundance | gaussian | LogAbund \sim UI2 * Region + (1 SS/SSB) | 0.910 (95% CI [0.908, 0.912]) | 0.053 (95% CI [0.023, 0.086]) | 1.00-1.01 | 6751 |
| Global model of Simpson's diversity | gaussian | LogSimp \sim UI2 + (1 SS/SSB) | 0.694 (95% CI [0.685, 0.702]) | 0.035 (95% CI [0.027, 0.043]) | 1 | 6726 |
| Regional model of Simpson's diversity | gaussian | LogSimp \sim UI2 * Region + (1 SS/SSB) | 0.707 (95% CI [0.699, 0.716]) | 0.056 (95% CI [0.036, 0.084]) | 1 | 6355 |
| Global model of species richness (individual land use types, phylum: Arthropoda) | zero_inflated_-negbinomial | Species_richness \sim LU + (1 SS/SSB), $z_i \sim 1$ | 0.903 (95% CI [0.892, 0.911]) | 0.008 (95% CI [0.005, 0.011]) | 1 | 7604 |
| Europe model of species richness (individual land use types, phylum: Arthropoda) | zero_inflated_-negbinomial | Species_richness \sim LU + (1 SS/SSB), $z_i \sim 1$ | 0.917 (95% CI [0.908, 0.924]) | 0.007 (95% CI [0.003, 0.012]) | 1 | 3412 |
| Global model of total abundance (individual land use types, phylum: Arthropoda) | gaussian | LogAbund \sim LU + (1 SS/SSB) | 0.907 (95% CI [0.905, 0.909]) | 0.018 (95% CI [0.014, 0.022]) | 1.00-1.01 | 7122 |
| Europe model of total abundance (individual land use types, phylum: Arthropoda) | gaussian | LogAbund \sim LU + (1 SS/SSB) | 0.938 (95% CI [0.937, 0.940]) | 0.025 (95% CI [0.019, 0.032]) | 1.00-1.01 | 3211 |
| Global model of Simpson's diversity (individual land use types, phylum: Arthropoda) | gaussian | LogSimp \sim LU + (1 SS/SSB) | 0.694 (95% CI [0.686, 0.702]) | 0.044 (95% CI [0.034, 0.055]) | 1.00-1.01 | 6726 |
| Europe model of Simpson's diversity (individual land use types, phylum: Arthropoda) | gaussian | LogSimp \sim LU + (1 SS/SSB) | 0.690 (95% CI [0.677, 0.703]) | 0.020 (95% CI [0.010, 0.032]) | 1 | 3032 |
| Global model of species richness (individual land use types, phylum: Chordata) | zero_inflated_-negbinomial | Species_richness \sim LU + (1 SS/SSB), $z_i \sim 1$ | 0.855 (95% CI [0.848, 0.862]) | 0.011 (95% CI [0.007, 0.015]) | 1 | 7842 |
| Europe model of species richness (individual land use types, phylum: Chordata) | zero_inflated_-negbinomial | Species_richness \sim LU + (1 SS/SSB), $z_i \sim 1$ | 0.850 (95% CI [0.837, 0.861]) | 0.084 (95% CI [0.031, 0.160]) | 1 | 1157 |
| Global model of total abundance (individual land use types, phylum: Chordata) | gaussian | LogAbund \sim LU + (1 SS/SSB) | 0.838 (95% CI [0.834, 0.841]) | 0.008 (95% CI [0.006, 0.010]) | 1.00-1.01 | 7007 |
| Europe model of total abundance (individual land use types, phylum: Chordata) | gaussian | LogAbund \sim LU + (1 SS/SSB) | 0.790 (95% CI [0.778, 0.801]) | 0.052 (95% CI [0.035, 0.070]) | 1 | 966 |
| Global model of Simpson's diversity (individual land use types, phylum: Chordata) | gaussian | LogSimp \sim LU + (1 SS/SSB) | 0.749 (95% CI [0.743, 0.755]) | 0.019 (95% CI [0.014, 0.024]) | 1 | 6708 |
| Europe model of Simpson's diversity (individual land use types, phylum: Chordata) | gaussian | LogSimp \sim LU + (1 SS/SSB) | 0.769 (95% CI [0.756, 0.782]) | 0.057 (95% CI [0.036, 0.082]) | 1 | 923 |

Continued on next page

| Model | Distribution family | Model equation | Conditional R^2 | Marginal R^2 | R-hats | nSites |
|---|---------------------------|--|-------------------------------|-----------------------------------|-----------|--------|
| Regional model of total dissimilarity | zero_one.inflated_beta | Dissimilarity \sim LU * Region + Dist_geo + (1 SS), zoi \sim (1 SS), coi \sim (1 SS) | 0.434 (95% CI [0.412, 0.457]) | 0.102 (95% CI [0.071, 0.136]) | 1 | 2858 |
| Regional model of turnover | zero_one.inflated_beta | Turnover \sim LU * Region + Dist_geo + (1 SS), zoi \sim (1 SS), coi \sim (1 SS) | 0.368 (95% CI [0.344, 0.391]) | 0.074 (95% CI [0.049, 0.104]) | 1 | 2858 |
| Regional model of nestedness | zero_one.inflated_beta | Nestedness \sim LU * Region + Dist_geo + (1 SS), zoi \sim (1 SS), coi \sim (1 SS) | 0.155 (95% CI [0.134, 0.177]) | 0.036 (95% CI [0.019, 0.061]) | 1 | 2858 |
| Europe model of species richness and range extent | zero.inflated_negbinomial | Species_richness \sim UI2 * Quants + (1 SS/SSB), zi \sim 1 | 0.800 (95% CI [0.780, 0.820]) | 0.037 (95% CI [0.021, 0.056]) | 1.00-1.01 | 6114 |
| North America model of species richness and range extent | zero.inflated_negbinomial | Species_richness \sim UI2 * Quants + (1 SS/SSB), zi \sim 1 | 0.848 (95% CI [0.831, 0.861]) | 0.005 (95% CI [0.001, 0.012]) | 1 | 1697 |
| Europe model of total abundance and range extent | gaussian | LogAbund \sim UI2 * Quants + (1 SS/SSB) | 0.825 (95% CI [0.820, 0.830]) | 0.030 (95% CI [0.026, 0.034]) | 1 | 5741 |
| North America model of total abundance and range extent | gaussian | LogAbund \sim UI2 * Quants + (1 SS/SSB) | 0.581 (95% CI [0.559, 0.600]) | 0.023 (95% CI [0.011, 0.037]) | 1 | 1677 |
| Europe model of species richness and habitat specificity | zero.inflated_negbinomial | Species_richness \sim UI2 * Quants + (1 SS/SSB), zi \sim 1 | 0.757 (95% CI [0.747, 0.767]) | 0.044 (95% CI [0.023, 0.073]) | 1 | 3389 |
| North America model of species richness and habitat specificity | zero.inflated_negbinomial | Species_richness \sim UI2 * Quants + (1 SS/SSB), zi \sim 1 | 0.646 (95% CI [0.610, 0.677]) | 0.040 (95% CI [0.007, 0.102]) | 1 | 605 |
| Europe model of total abundance and habitat specificity | gaussian | LogAbund \sim UI2 * Quants + (1 SS/SSB) | 0.693 (95% CI [0.683, 0.704]) | 0.126 (95% CI [0.114, 0.138]) | 1 | 3233 |
| North America model of total abundance and habitat specificity | gaussian | LogAbund \sim UI2 * Quants + (1 SS/SSB) | 0.630 (95% CI [0.595, 0.663]) | 0.061 (95% CI [0.032, 0.101]) | 1 | 585 |
| Europe model of non-native proportions of richness | zero_one.inflated_beta | SiteSpeciesProps \sim UI2 + (1 SS/SSB), zoi \sim (1 SS), coi \sim (1 SS) | 0.353 (95% CI [0.309, 0.402]) | 0.003 (95% CI [2.961e-04, 0.008]) | 1 | 2837 |
| North America model of non-native proportions of richness | zero_one.inflated_beta | SiteSpeciesProps \sim UI2 + (1 SS/SSB), zoi \sim (1 SS), coi \sim (1 SS) | 0.658 (95% CI [0.552, 0.715]) | 0.002 (95% CI [7.614e-08, 0.012]) | 1 | 832 |
| Europe model of non-native proportions of abundance | zero_one.inflated_beta | SiteTotAbunProps \sim UI2 + (1 SS/SSB), zoi \sim (1 SS), coi \sim (1 SS) | 0.364 (95% CI [0.313, 0.412]) | 0.004 (95% CI [1.334e-04, 0.010]) | 1 | 2837 |
| North America model of non-native proportions of abundance | zero_one.inflated_beta | SiteTotAbunProps \sim UI2 + (1 SS/SSB), zoi \sim (1 SS), coi \sim (1 SS) | 0.576 (95% CI [0.440, 0.648]) | 0.003 (95% CI [2.082e-06, 0.027]) | 1 | 832 |

A.3 Regional responses

Table S3: Regional responses of species richness to land use, relative to primary vegetation (median, 2.5% and 97.5% percentiles of model responses), and number of studies and sites per coefficient.

| Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|---------------------------|-----------------------|--------|-------|-------|----------|--------|
| Global | Primary vegetation | 0.0 | NA | NA | 166 | 1908 |
| Global | Secondary vegetation | -12.0 | -15.7 | -8.3 | 174 | 1488 |
| Global | Agriculture (minimal) | -19.9 | -23.2 | -16.1 | 108 | 1201 |
| Global | Agriculture (light) | -27.8 | -31.0 | -24.9 | 145 | 1691 |
| Global | Agriculture (intense) | -30.4 | -33.8 | -26.7 | 94 | 1316 |
| Europe | Primary vegetation | 0.0 | NA | NA | 22 | 230 |
| Europe | Secondary vegetation | 5.5 | -3.4 | 14.6 | 61 | 590 |
| Europe | Agriculture (minimal) | 7.3 | -1.5 | 17.0 | 40 | 628 |
| Europe | Agriculture (light) | -4.0 | -12.4 | 5.8 | 51 | 961 |
| Europe | Agriculture (intense) | -3.2 | -12.5 | 7.2 | 46 | 1003 |
| North America | Primary vegetation | 0.0 | NA | NA | 23 | 469 |
| North America | Secondary vegetation | -9.9 | -18.7 | 0.3 | 19 | 285 |
| North America | Agriculture (minimal) | -16.3 | -33.3 | 4.2 | 6 | 62 |
| North America | Agriculture (light) | -35.1 | -45.1 | -23.7 | 16 | 117 |
| North America | Agriculture (intense) | -37.9 | -47.8 | -26.6 | 13 | 55 |
| South and Southeast Asia | Primary vegetation | 0.0 | NA | NA | 30 | 264 |
| South and Southeast Asia | Secondary vegetation | -16.1 | -35.5 | 11.1 | 7 | 18 |
| South and Southeast Asia | Agriculture (minimal) | 0.5 | -15.0 | 18.9 | 8 | 36 |
| South and Southeast Asia | Agriculture (light) | -48.9 | -54.2 | -43.3 | 12 | 179 |
| South and Southeast Asia | Agriculture (intense) | -32.1 | -42.5 | -20.0 | 6 | 28 |
| South and Central America | Primary vegetation | 0.0 | NA | NA | 56 | 335 |
| South and Central America | Secondary vegetation | -13.0 | -19.9 | -6.5 | 57 | 265 |
| South and Central America | Agriculture (minimal) | -22.1 | -28.6 | -14.3 | 32 | 223 |
| South and Central America | Agriculture (light) | -39.7 | -44.8 | -34.0 | 40 | 252 |
| South and Central America | Agriculture (intense) | -45.1 | -52.9 | -36.2 | 11 | 50 |
| Subsaharan Africa | Primary vegetation | 0.0 | NA | NA | 17 | 241 |
| Subsaharan Africa | Secondary vegetation | -11.6 | -23.2 | 1.2 | 8 | 54 |
| Subsaharan Africa | Agriculture (minimal) | -24.8 | -34.8 | -14.5 | 11 | 62 |
| Subsaharan Africa | Agriculture (light) | -30.2 | -38.1 | -21.3 | 9 | 50 |
| Subsaharan Africa | Agriculture (intense) | -53.5 | -61.9 | -43.0 | 6 | 19 |

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| Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|-------------|-----------------------|--------|-------|-------|----------|--------|
| Australasia | Primary vegetation | 0.0 | NA | NA | 10 | 232 |
| Australasia | Secondary vegetation | -13.0 | -31.7 | 12.2 | 10 | 136 |
| Australasia | Agriculture (minimal) | -45.4 | -50.0 | -41.0 | 5 | 134 |
| Australasia | Agriculture (light) | -20.1 | -32.2 | -5.8 | 10 | 61 |
| Australasia | Agriculture (intense) | -24.7 | -34.9 | -14.0 | 11 | 149 |

Table S4: Regional responses of total abundance to land use, relative to primary vegetation (median, 2.5% and 97.5% percentiles of model responses), and number of studies and sites per coefficient.

| Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|---------------------------|-----------------------|--------|-------|-------|----------|--------|
| Global | Primary vegetation | 0.0 | NA | NA | 146 | 1711 |
| Global | Secondary vegetation | -21.3 | -28.1 | -13.8 | 162 | 1363 |
| Global | Agriculture (minimal) | -23.7 | -30.6 | -16.1 | 103 | 1131 |
| Global | Agriculture (light) | -37.1 | -42.5 | -30.9 | 136 | 1640 |
| Global | Agriculture (intense) | -36.2 | -42.7 | -28.8 | 88 | 1277 |
| Europe | Primary vegetation | 0.0 | NA | NA | 19 | 161 |
| Europe | Secondary vegetation | 6.6 | -19.9 | 37.1 | 56 | 528 |
| Europe | Agriculture (minimal) | 39.0 | 0.9 | 82.3 | 37 | 570 |
| Europe | Agriculture (light) | -3.0 | -26.2 | 29.6 | 49 | 957 |
| Europe | Agriculture (intense) | -0.8 | -27.2 | 33.6 | 45 | 995 |
| North America | Primary vegetation | 0.0 | NA | NA | 22 | 443 |
| North America | Secondary vegetation | 0.7 | -19.9 | 25.7 | 19 | 285 |
| North America | Agriculture (minimal) | -17.8 | -44.1 | 19.0 | 6 | 62 |
| North America | Agriculture (light) | -47.8 | -61.6 | -30.0 | 16 | 117 |
| North America | Agriculture (intense) | -43.5 | -59.7 | -20.9 | 12 | 45 |
| South and Southeast Asia | Primary vegetation | 0.0 | NA | NA | 22 | 230 |
| South and Southeast Asia | Secondary vegetation | -39.6 | -73.8 | 34.3 | 5 | 5 |
| South and Southeast Asia | Agriculture (minimal) | -22.4 | -54.5 | 25.2 | 6 | 24 |
| South and Southeast Asia | Agriculture (light) | -61.2 | -68.6 | -51.6 | 10 | 171 |
| South and Southeast Asia | Agriculture (intense) | -46.5 | -66.1 | -14.2 | 4 | 23 |
| South and Central America | Primary vegetation | 0.0 | NA | NA | 52 | 308 |
| South and Central America | Secondary vegetation | -22.4 | -34.5 | -6.8 | 55 | 241 |
| South and Central America | Agriculture (minimal) | -19.8 | -32.5 | -4.0 | 32 | 223 |
| South and Central America | Agriculture (light) | -38.2 | -49.3 | -24.0 | 37 | 238 |

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| Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|---------------------------|-----------------------|--------|-------|-------|----------|--------|
| South and Central America | Agriculture (intense) | -25.0 | -46.7 | 4.9 | 10 | 44 |
| Subsaharan Africa | Primary vegetation | 0.0 | NA | NA | 16 | 235 |
| Subsaharan Africa | Secondary vegetation | -37.3 | -56.2 | -11.5 | 8 | 54 |
| Subsaharan Africa | Agriculture (minimal) | -63.5 | -72.0 | -51.1 | 11 | 62 |
| Subsaharan Africa | Agriculture (light) | -57.8 | -70.1 | -40.7 | 9 | 50 |
| Subsaharan Africa | Agriculture (intense) | -84.6 | -90.3 | -75.8 | 6 | 19 |
| Australasia | Primary vegetation | 0.0 | NA | NA | 8 | 212 |
| Australasia | Secondary vegetation | -11.7 | -42.3 | 39.5 | 8 | 125 |
| Australasia | Agriculture (minimal) | -51.6 | -61.4 | -39.6 | 5 | 134 |
| Australasia | Agriculture (light) | -22.6 | -44.8 | 5.9 | 9 | 51 |
| Australasia | Agriculture (intense) | -18.6 | -36.0 | 2.7 | 10 | 139 |

Table S5: Regional responses of Simpson’s diversity index to land use, relative to primary vegetation (median, 2.5% and 97.5% percentiles of model responses), and number of studies and sites per coefficient.

| Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|--------------------------|-----------------------|--------|-------|-------|----------|--------|
| Global | Primary vegetation | 0.0 | NA | NA | 146 | 1597 |
| Global | Secondary vegetation | -13.6 | -17.8 | -9.1 | 161 | 1320 |
| Global | Agriculture (minimal) | -24.4 | -28.2 | -20.2 | 101 | 1056 |
| Global | Agriculture (light) | -30.9 | -34.5 | -27.3 | 136 | 1558 |
| Global | Agriculture (intense) | -31.8 | -35.7 | -27.7 | 88 | 1195 |
| Europe | Primary vegetation | 0.0 | NA | NA | 19 | 161 |
| Europe | Secondary vegetation | 0.6 | -11.5 | 14.6 | 56 | 504 |
| Europe | Agriculture (minimal) | -2.7 | -15.3 | 11.7 | 37 | 550 |
| Europe | Agriculture (light) | -8.5 | -19.8 | 4.9 | 49 | 894 |
| Europe | Agriculture (intense) | -9.5 | -20.8 | 3.8 | 45 | 923 |
| North America | Primary vegetation | 0.0 | NA | NA | 22 | 436 |
| North America | Secondary vegetation | -11.3 | -21.7 | 0.3 | 19 | 276 |
| North America | Agriculture (minimal) | -17.3 | -32.7 | 3.6 | 5 | 60 |
| North America | Agriculture (light) | -44.4 | -52.4 | -33.8 | 16 | 113 |
| North America | Agriculture (intense) | -51.5 | -60.5 | -41.1 | 12 | 45 |
| South and Southeast Asia | Primary vegetation | 0.0 | NA | NA | 22 | 230 |
| South and Southeast Asia | Secondary vegetation | -13.7 | -44.8 | 35.7 | 5 | 5 |
| South and Southeast Asia | Agriculture (minimal) | -3.5 | -27.7 | 26.0 | 6 | 24 |

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| Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|---------------------------|-----------------------|--------|-------|-------|----------|--------|
| South and Southeast Asia | Agriculture (light) | -47.2 | -52.6 | -40.8 | 10 | 166 |
| South and Southeast Asia | Agriculture (intense) | -23.5 | -40.9 | -1.1 | 4 | 23 |
| South and Central America | Primary vegetation | 0.0 | NA | NA | 52 | 286 |
| South and Central America | Secondary vegetation | -0.8 | -9.7 | 8.7 | 54 | 233 |
| South and Central America | Agriculture (minimal) | -18.6 | -25.7 | -9.5 | 32 | 184 |
| South and Central America | Agriculture (light) | -28.9 | -36.6 | -20.5 | 37 | 236 |
| South and Central America | Agriculture (intense) | -42.1 | -51.5 | -30.6 | 10 | 44 |
| Subsaharan Africa | Primary vegetation | 0.0 | NA | NA | 16 | 154 |
| Subsaharan Africa | Secondary vegetation | -22.4 | -34.3 | -7.4 | 8 | 54 |
| Subsaharan Africa | Agriculture (minimal) | -21.9 | -33.2 | -8.2 | 10 | 48 |
| Subsaharan Africa | Agriculture (light) | -23.4 | -35.2 | -9.5 | 9 | 49 |
| Subsaharan Africa | Agriculture (intense) | -36.9 | -50.1 | -18.2 | 6 | 19 |
| Australasia | Primary vegetation | 0.0 | NA | NA | 8 | 208 |
| Australasia | Secondary vegetation | -7.3 | -25.7 | 16.4 | 8 | 123 |
| Australasia | Agriculture (minimal) | -31.2 | -39.2 | -22.8 | 5 | 134 |
| Australasia | Agriculture (light) | -17.6 | -33.3 | 0.9 | 9 | 44 |
| Australasia | Agriculture (intense) | -16.1 | -26.3 | -3.8 | 10 | 129 |

A.4 Regional responses per land use type and intensity

Table S6: Global and European responses of species richness to land use for arthropods and vertebrates, relative to primary vegetation (median, 2.5% and 97.5% percentiles of model responses), and number of studies and sites per coefficient.

| Taxa | Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|--------|----------------------|--------|-------|-------|----------|--------|
| Arthropoda | Global | Primary vegetation | 0.0 | NA | NA | 166 | 1908 |
| Arthropoda | Global | Secondary vegetation | -11.9 | -15.6 | -8.2 | 174 | 1488 |
| Arthropoda | Global | Cropland (minimal) | -27.4 | -32.8 | -21.8 | 43 | 364 |
| Arthropoda | Global | Cropland (light) | -39.1 | -43.3 | -34.7 | 58 | 697 |
| Arthropoda | Global | Cropland (intense) | -41.8 | -45.9 | -37.5 | 52 | 716 |
| Arthropoda | Global | Pasture (minimal) | -20.0 | -24.4 | -15.3 | 44 | 664 |
| Arthropoda | Global | Pasture (light) | -22.0 | -25.9 | -17.8 | 63 | 641 |
| Arthropoda | Global | Pasture (intense) | -32.3 | -36.6 | -27.8 | 29 | 461 |
| Arthropoda | Global | Plantation (minimal) | -9.5 | -16.9 | -2.2 | 43 | 173 |
| Arthropoda | Global | Plantation (light) | -30.8 | -35.2 | -26.1 | 45 | 353 |
| Arthropoda | Global | Plantation (intense) | -9.1 | -16.6 | -1.2 | 22 | 139 |
| Arthropoda | Europe | Primary vegetation | 0.0 | NA | NA | 22 | 230 |
| Arthropoda | Europe | Secondary vegetation | 4.2 | -5.3 | 13.8 | 61 | 590 |
| Arthropoda | Europe | Cropland (minimal) | -16.4 | -26.2 | -6.5 | 23 | 223 |
| Arthropoda | Europe | Cropland (light) | -20.7 | -30.4 | -10.9 | 26 | 485 |
| Arthropoda | Europe | Cropland (intense) | -20.2 | -29.9 | -10.6 | 33 | 603 |
| Arthropoda | Europe | Pasture (minimal) | 15.3 | 4.3 | 26.7 | 21 | 344 |
| Arthropoda | Europe | Pasture (light) | 1.5 | -8.7 | 12.3 | 25 | 432 |
| Arthropoda | Europe | Pasture (intense) | -6.1 | -16.5 | 5.4 | 18 | 371 |
| Arthropoda | Europe | Plantation (minimal) | 4.6 | -11.1 | 23.2 | 11 | 61 |
| Arthropoda | Europe | Plantation (light) | 0.1 | -16.5 | 22.0 | 8 | 44 |
| Arthropoda | Europe | Plantation (intense) | 106.0 | 77.0 | 141.0 | 1 | 29 |
| Chordata | Global | Primary vegetation | 0.0 | NA | NA | 173 | 3473 |
| Chordata | Global | Secondary vegetation | -8.4 | -11.6 | -5.1 | 91 | 1348 |
| Chordata | Global | Cropland (minimal) | -15.6 | -19.9 | -10.5 | 20 | 319 |
| Chordata | Global | Cropland (light) | -18.1 | -23.4 | -12.9 | 17 | 135 |
| Chordata | Global | Cropland (intense) | -20.9 | -34.5 | -4.7 | 5 | 146 |
| Chordata | Global | Pasture (minimal) | -21.7 | -27.6 | -15.3 | 19 | 142 |
| Chordata | Global | Pasture (light) | -26.7 | -31.0 | -22.4 | 24 | 308 |
| Chordata | Global | Pasture (intense) | -17.0 | -24.9 | -8.7 | 10 | 109 |

Continued on next page

| Taxa | Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|----------|--------|----------------------|--------|-------|-------|----------|--------|
| Chordata | Global | Plantation (minimal) | -6.9 | -11.2 | -2.2 | 33 | 490 |
| Chordata | Global | Plantation (light) | -19.6 | -23.1 | -16.2 | 46 | 985 |
| Chordata | Global | Plantation (intense) | -36.8 | -41.2 | -32.1 | 22 | 387 |
| Chordata | Europe | Primary vegetation | 0.0 | NA | NA | 7 | 147 |
| Chordata | Europe | Secondary vegetation | -13.9 | -20.6 | -7.0 | 6 | 313 |
| Chordata | Europe | Cropland (minimal) | 21.3 | 1.5 | 44.0 | 1 | 54 |
| Chordata | Europe | Cropland (intense) | -19.7 | -36.7 | 1.3 | 1 | 138 |
| Chordata | Europe | Pasture (minimal) | -42.0 | -54.8 | -25.3 | 2 | 22 |
| Chordata | Europe | Pasture (light) | -27.5 | -36.9 | -15.5 | 4 | 60 |
| Chordata | Europe | Pasture (intense) | 0.7 | -45.5 | 69.4 | 1 | 1 |
| Chordata | Europe | Plantation (minimal) | 0.3 | -10.8 | 11.2 | 4 | 108 |
| Chordata | Europe | Plantation (light) | -20.9 | -28.1 | -13.2 | 6 | 124 |
| Chordata | Europe | Plantation (intense) | -45.8 | -52.7 | -38.6 | 5 | 190 |

Table S7: Global and European responses of total abundance to land use for arthropods and vertebrates, relative to primary vegetation (median, 2.5% and 97.5% percentiles of model responses), and number of studies and sites per coefficient.

| Taxa | Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|--------|----------------------|--------|-------|-------|----------|--------|
| Arthropoda | Global | Primary vegetation | 0.0 | NA | NA | 146 | 1711 |
| Arthropoda | Global | Secondary vegetation | -20.1 | -27.2 | -11.6 | 162 | 1363 |
| Arthropoda | Global | Cropland (minimal) | -55.5 | -61.7 | -48.4 | 41 | 358 |
| Arthropoda | Global | Cropland (light) | -59.8 | -65.6 | -52.6 | 58 | 697 |
| Arthropoda | Global | Cropland (intense) | -60.6 | -66.4 | -55.1 | 50 | 705 |
| Arthropoda | Global | Pasture (minimal) | -6.7 | -17.2 | 3.8 | 41 | 614 |
| Arthropoda | Global | Pasture (light) | -18.3 | -27.8 | -7.1 | 58 | 624 |
| Arthropoda | Global | Pasture (intense) | -26.2 | -36.2 | -15.9 | 27 | 448 |
| Arthropoda | Global | Plantation (minimal) | -0.3 | -17.5 | 18.6 | 41 | 159 |
| Arthropoda | Global | Plantation (light) | -42.3 | -49.5 | -33.1 | 40 | 319 |
| Arthropoda | Global | Plantation (intense) | 5.6 | -13.0 | 27.6 | 19 | 124 |
| Arthropoda | Europe | Primary vegetation | 0.0 | NA | NA | 19 | 161 |
| Arthropoda | Europe | Secondary vegetation | 4.7 | -16.6 | 32.7 | 56 | 528 |
| Arthropoda | Europe | Cropland (minimal) | -46.5 | -61.9 | -24.1 | 22 | 218 |
| Arthropoda | Europe | Cropland (light) | -50.4 | -63.4 | -29.7 | 26 | 485 |
| Arthropoda | Europe | Cropland (intense) | -47.7 | -60.9 | -27.6 | 32 | 602 |

Continued on next page

| Taxa | Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|--------|----------------------|--------|-------|-------|----------|--------|
| Arthropoda | Europe | Pasture (minimal) | 103.2 | 52.4 | 180.1 | 18 | 294 |
| Arthropoda | Europe | Pasture (light) | 26.2 | -4.4 | 69.5 | 23 | 429 |
| Arthropoda | Europe | Pasture (intense) | 18.5 | -11.4 | 62.5 | 17 | 364 |
| Arthropoda | Europe | Plantation (minimal) | 38.6 | -2.4 | 100.6 | 10 | 58 |
| Arthropoda | Europe | Plantation (light) | 32.9 | -8.8 | 94.2 | 7 | 43 |
| Arthropoda | Europe | Plantation (intense) | 250.6 | 130.9 | 458.1 | 1 | 29 |
| Chordata | Global | Primary vegetation | 0.0 | NA | NA | 156 | 3045 |
| Chordata | Global | Secondary vegetation | -3.8 | -9.5 | 2.2 | 81 | 1209 |
| Chordata | Global | Cropland (minimal) | 1.3 | -8.1 | 11.6 | 20 | 319 |
| Chordata | Global | Cropland (light) | -18.4 | -28.2 | -6.9 | 14 | 131 |
| Chordata | Global | Cropland (intense) | -3.9 | -21.1 | 16.2 | 5 | 146 |
| Chordata | Global | Pasture (minimal) | -11.8 | -22.8 | 0.6 | 18 | 139 |
| Chordata | Global | Pasture (light) | -41.3 | -46.8 | -35.2 | 24 | 308 |
| Chordata | Global | Pasture (intense) | -12.4 | -27.2 | 8.8 | 10 | 109 |
| Chordata | Global | Plantation (minimal) | -1.7 | -10.3 | 8.0 | 30 | 477 |
| Chordata | Global | Plantation (light) | -21.7 | -27.7 | -15.8 | 38 | 875 |
| Chordata | Global | Plantation (intense) | -23.4 | -31.2 | -14.5 | 19 | 249 |
| Chordata | Europe | Primary vegetation | 0.0 | NA | NA | 7 | 147 |
| Chordata | Europe | Secondary vegetation | -25.1 | -36.8 | -12.6 | 6 | 313 |
| Chordata | Europe | Cropland (minimal) | 16.6 | -7.9 | 46.0 | 1 | 54 |
| Chordata | Europe | Cropland (intense) | -27.4 | -42.1 | -9.8 | 1 | 138 |
| Chordata | Europe | Pasture (minimal) | -31.3 | -52.3 | -1.0 | 2 | 22 |
| Chordata | Europe | Pasture (light) | -36.5 | -49.5 | -19.7 | 4 | 60 |
| Chordata | Europe | Pasture (intense) | 6.7 | -65.3 | 228.9 | 1 | 1 |
| Chordata | Europe | Plantation (minimal) | 10.2 | -9.2 | 33.9 | 4 | 108 |
| Chordata | Europe | Plantation (light) | -26.8 | -38.6 | -12.0 | 5 | 63 |
| Chordata | Europe | Plantation (intense) | -63.0 | -70.7 | -52.2 | 4 | 60 |

Table S8: Global and European responses of Simpson’s diversity index to land use for arthropods and vertebrates, relative to primary vegetation (median, 2.5% and 97.5% percentiles of model responses), and number of studies and sites per coefficient.

| Taxa | Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|--------|----------------------|--------|-------|-------|----------|--------|
| Arthropoda | Global | Primary vegetation | 0.0 | NA | NA | 146 | 1597 |
| Arthropoda | Global | Secondary vegetation | -12.9 | -17.4 | -8.2 | 161 | 1320 |

Continued on next page

| Taxa | Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|--------|----------------------|--------|-------|-------|----------|--------|
| Arthropoda | Global | Cropland (minimal) | -31.5 | -37.0 | -25.5 | 38 | 331 |
| Arthropoda | Global | Cropland (light) | -36.3 | -41.3 | -31.2 | 58 | 645 |
| Arthropoda | Global | Cropland (intense) | -36.6 | -41.3 | -31.2 | 50 | 634 |
| Arthropoda | Global | Pasture (minimal) | -21.1 | -26.0 | -16.1 | 41 | 584 |
| Arthropoda | Global | Pasture (light) | -26.2 | -30.6 | -21.6 | 57 | 605 |
| Arthropoda | Global | Pasture (intense) | -33.1 | -37.8 | -27.7 | 27 | 442 |
| Arthropoda | Global | Plantation (minimal) | -21.4 | -28.2 | -12.7 | 40 | 141 |
| Arthropoda | Global | Plantation (light) | -34.7 | -39.5 | -29.7 | 40 | 308 |
| Arthropoda | Global | Plantation (intense) | -17.7 | -26.8 | -9.2 | 19 | 119 |
| Arthropoda | Europe | Primary vegetation | 0.0 | NA | NA | 19 | 161 |
| Arthropoda | Europe | Secondary vegetation | 0.4 | -12.1 | 14.8 | 56 | 504 |
| Arthropoda | Europe | Cropland (minimal) | -16.3 | -30.3 | -0.5 | 21 | 216 |
| Arthropoda | Europe | Cropland (light) | -16.3 | -29.5 | -0.2 | 26 | 433 |
| Arthropoda | Europe | Cropland (intense) | -11.8 | -25.9 | 5.4 | 32 | 533 |
| Arthropoda | Europe | Pasture (minimal) | 12.8 | -3.8 | 33.1 | 18 | 291 |
| Arthropoda | Europe | Pasture (light) | -0.8 | -14.8 | 14.9 | 23 | 418 |
| Arthropoda | Europe | Pasture (intense) | -9.0 | -23.5 | 7.0 | 17 | 361 |
| Arthropoda | Europe | Plantation (minimal) | -16.0 | -30.8 | 3.0 | 9 | 43 |
| Arthropoda | Europe | Plantation (light) | -7.3 | -26.4 | 17.5 | 7 | 43 |
| Arthropoda | Europe | Plantation (intense) | 49.6 | 17.9 | 90.4 | 1 | 29 |
| Chordata | Global | Primary vegetation | 0.0 | NA | NA | 156 | 2953 |
| Chordata | Global | Secondary vegetation | -10.1 | -13.9 | -6.4 | 81 | 1093 |
| Chordata | Global | Cropland (minimal) | -16.1 | -20.9 | -11.0 | 20 | 313 |
| Chordata | Global | Cropland (light) | -26.4 | -32.6 | -19.1 | 14 | 131 |
| Chordata | Global | Cropland (intense) | -22.0 | -31.1 | -11.2 | 5 | 128 |
| Chordata | Global | Pasture (minimal) | -27.0 | -33.6 | -20.3 | 18 | 134 |
| Chordata | Global | Pasture (light) | -24.0 | -28.1 | -19.6 | 24 | 299 |
| Chordata | Global | Pasture (intense) | -28.0 | -37.2 | -18.2 | 10 | 109 |
| Chordata | Global | Plantation (minimal) | -10.4 | -15.3 | -5.4 | 30 | 476 |
| Chordata | Global | Plantation (light) | -16.8 | -20.8 | -12.3 | 38 | 833 |
| Chordata | Global | Plantation (intense) | -29.9 | -34.8 | -24.0 | 18 | 239 |
| Chordata | Europe | Primary vegetation | 0.0 | NA | NA | 7 | 147 |
| Chordata | Europe | Secondary vegetation | -21.6 | -30.5 | -10.4 | 6 | 297 |
| Chordata | Europe | Cropland (minimal) | 14.2 | -3.6 | 37.3 | 1 | 50 |
| Chordata | Europe | Cropland (intense) | -30.0 | -41.8 | -15.3 | 1 | 120 |

Continued on next page

| Taxa | Region | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|----------|--------|----------------------|--------|-------|-------|----------|--------|
| Chordata | Europe | Pasture (minimal) | -54.8 | -65.7 | -39.2 | 2 | 20 |
| Chordata | Europe | Pasture (light) | -38.3 | -48.4 | -26.9 | 4 | 58 |
| Chordata | Europe | Pasture (intense) | -29.9 | -70.1 | 67.0 | 1 | 1 |
| Chordata | Europe | Plantation (minimal) | -4.1 | -17.5 | 10.8 | 4 | 107 |
| Chordata | Europe | Plantation (light) | -27.7 | -37.1 | -15.2 | 5 | 63 |
| Chordata | Europe | Plantation (intense) | -36.9 | -47.7 | -24.3 | 4 | 60 |

A.5 Compositional dissimilarity

Table S9: Compositional dissimilarity (total, turnover and nestedness) for each region and land use compared with natural vegetation, relative to the dissimilarity between natural vegetation and other natural vegetation sites within the same region (median, 2.5% and 97.5% percentiles of model responses), and number of sites per coefficient.

| Measure | Region | Land use | median | lower | upper | nSites |
|----------------------|---------------------------|-----------------------|--------|-------|-------|--------|
| Total dissimilarity | Europe | Natural vegetation | 0.0 | 0.0 | 0.0 | 500 |
| Total dissimilarity | Europe | Agriculture (minimal) | 41.9 | 29.8 | 55.7 | 155 |
| Total dissimilarity | Europe | Agriculture (light) | 18.8 | -2.2 | 38.3 | 86 |
| Total dissimilarity | Europe | Agriculture (intense) | 11.8 | -16.0 | 41.9 | 23 |
| Total dissimilarity | North America | Natural vegetation | 0.0 | 0.0 | 0.0 | 570 |
| Total dissimilarity | North America | Agriculture (minimal) | 5.2 | -10.5 | 20.5 | 54 |
| Total dissimilarity | North America | Agriculture (light) | 5.2 | -11.9 | 23.3 | 40 |
| Total dissimilarity | North America | Agriculture (intense) | 13.1 | -11.8 | 35.8 | 18 |
| Total dissimilarity | South and Southeast Asia | Natural vegetation | 0.0 | 0.0 | 0.0 | 136 |
| Total dissimilarity | South and Southeast Asia | Agriculture (minimal) | 10.6 | -26.8 | 54.4 | 11 |
| Total dissimilarity | South and Southeast Asia | Agriculture (light) | 38.8 | 24.7 | 56.5 | 132 |
| Total dissimilarity | South and Southeast Asia | Agriculture (intense) | NA | NA | NA | 5 |
| Total dissimilarity | South and Central America | Natural vegetation | 0.0 | 0.0 | 0.0 | 245 |
| Total dissimilarity | South and Central America | Agriculture (minimal) | 19.5 | 5.7 | 33.9 | 103 |
| Total dissimilarity | South and Central America | Agriculture (light) | 46.3 | 29.6 | 65.6 | 100 |
| Total dissimilarity | South and Central America | Agriculture (intense) | 42.0 | 17.1 | 67.8 | 28 |
| Total dissimilarity | Sub-Saharan Africa | Natural vegetation | 0.0 | 0.0 | 0.0 | 124 |
| Total dissimilarity | Sub-Saharan Africa | Agriculture (minimal) | 6.2 | -12.7 | 28.2 | 41 |
| Total dissimilarity | Sub-Saharan Africa | Agriculture (light) | 6.7 | -15.4 | 30.0 | 28 |
| Total dissimilarity | Sub-Saharan Africa | Agriculture (intense) | 18.0 | -7.6 | 46.4 | 14 |
| Total dissimilarity | Australasia | Natural vegetation | 0.0 | 0.0 | 0.0 | 238 |
| Total dissimilarity | Australasia | Agriculture (minimal) | 18.6 | 4.5 | 33.9 | 116 |
| Total dissimilarity | Australasia | Agriculture (light) | 2.3 | -23.9 | 29.5 | 15 |
| Total dissimilarity | Australasia | Agriculture (intense) | 1.6 | -18.1 | 23.0 | 70 |
| Nestedness component | Europe | Natural vegetation | 0.0 | 0.0 | 0.0 | 500 |
| Nestedness component | Europe | Agriculture (minimal) | -11.9 | -31.3 | 12.0 | 155 |
| Nestedness component | Europe | Agriculture (light) | 2.2 | -29.6 | 45.4 | 86 |
| Nestedness component | Europe | Agriculture (intense) | 15.7 | -31.4 | 88.4 | 23 |
| Nestedness component | North America | Natural vegetation | 0.0 | 0.0 | 0.0 | 570 |

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| Measure | Region | Land use | median | lower | upper | nSites |
|----------------------|---------------------------|-----------------------|--------|-------|-------|--------|
| Nestedness component | North America | Agriculture (minimal) | -9.0 | -40.5 | 40.1 | 54 |
| Nestedness component | North America | Agriculture (light) | 33.3 | -12.5 | 102.7 | 40 |
| Nestedness component | North America | Agriculture (intense) | 68.4 | -22.5 | 222.2 | 18 |
| Nestedness component | South and Southeast Asia | Natural vegetation | 0.0 | 0.0 | 0.0 | 136 |
| Nestedness component | South and Southeast Asia | Agriculture (minimal) | 4.2 | -53.8 | 108.7 | 11 |
| Nestedness component | South and Southeast Asia | Agriculture (light) | 86.5 | 47.2 | 133.7 | 132 |
| Nestedness component | South and Southeast Asia | Agriculture (intense) | NA | NA | NA | 5 |
| Nestedness component | South and Central America | Natural vegetation | 0.0 | 0.0 | 0.0 | 245 |
| Nestedness component | South and Central America | Agriculture (minimal) | -9.6 | -32.4 | 21.2 | 103 |
| Nestedness component | South and Central America | Agriculture (light) | 18.9 | -8.5 | 56.0 | 100 |
| Nestedness component | South and Central America | Agriculture (intense) | 4.4 | -35.0 | 59.0 | 28 |
| Nestedness component | Sub-Saharan Africa | Natural vegetation | 0.0 | 0.0 | 0.0 | 124 |
| Nestedness component | Sub-Saharan Africa | Agriculture (minimal) | 41.8 | -2.1 | 99.8 | 41 |
| Nestedness component | Sub-Saharan Africa | Agriculture (light) | 10.2 | -29.6 | 68.4 | 28 |
| Nestedness component | Sub-Saharan Africa | Agriculture (intense) | 34.3 | -24.4 | 115.2 | 14 |
| Nestedness component | Australasia | Natural vegetation | 0.0 | 0.0 | 0.0 | 238 |
| Nestedness component | Australasia | Agriculture (minimal) | 17.4 | -20.2 | 64.4 | 116 |
| Nestedness component | Australasia | Agriculture (light) | -1.3 | -57.3 | 99.2 | 15 |
| Nestedness component | Australasia | Agriculture (intense) | 10.6 | -37.0 | 80.7 | 70 |
| Turnover component | Europe | Natural vegetation | 0.0 | 0.0 | 0.0 | 500 |
| Turnover component | Europe | Agriculture (minimal) | 60.4 | 42.1 | 81.0 | 155 |
| Turnover component | Europe | Agriculture (light) | 24.8 | -2.0 | 55.3 | 86 |
| Turnover component | Europe | Agriculture (intense) | 17.0 | -24.9 | 66.4 | 23 |
| Turnover component | North America | Natural vegetation | 0.0 | 0.0 | 0.0 | 570 |
| Turnover component | North America | Agriculture (minimal) | 5.6 | -13.3 | 25.1 | 54 |
| Turnover component | North America | Agriculture (light) | -0.4 | -21.1 | 23.2 | 40 |
| Turnover component | North America | Agriculture (intense) | -4.2 | -33.4 | 23.2 | 18 |
| Turnover component | South and Southeast Asia | Natural vegetation | 0.0 | 0.0 | 0.0 | 136 |
| Turnover component | South and Southeast Asia | Agriculture (minimal) | -1.1 | -40.7 | 52.2 | 11 |
| Turnover component | South and Southeast Asia | Agriculture (light) | 21.4 | 5.9 | 39.0 | 132 |
| Turnover component | South and Southeast Asia | Agriculture (intense) | NA | NA | NA | 5 |
| Turnover component | South and Central America | Natural vegetation | 0.0 | 0.0 | 0.0 | 245 |
| Turnover component | South and Central America | Agriculture (minimal) | 19.5 | 0.6 | 40.0 | 103 |
| Turnover component | South and Central America | Agriculture (light) | 41.9 | 20.1 | 67.7 | 100 |
| Turnover component | South and Central America | Agriculture (intense) | 41.5 | 9.8 | 75.2 | 28 |

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| Measure | Region | Land use | median | lower | upper | nSites |
|--------------------|--------------------|-----------------------|--------|-------|-------|--------|
| Turnover component | Sub-Saharan Africa | Natural vegetation | 0.0 | 0.0 | 0.0 | 124 |
| Turnover component | Sub-Saharan Africa | Agriculture (minimal) | 0.5 | -24.8 | 32.3 | 41 |
| Turnover component | Sub-Saharan Africa | Agriculture (light) | 6.6 | -23.1 | 43.4 | 28 |
| Turnover component | Sub-Saharan Africa | Agriculture (intense) | 7.6 | -23.9 | 45.4 | 14 |
| Turnover component | Australasia | Natural vegetation | 0.0 | 0.0 | 0.0 | 238 |
| Turnover component | Australasia | Agriculture (minimal) | 12.8 | -5.6 | 34.1 | 116 |
| Turnover component | Australasia | Agriculture (light) | 9.1 | -26.9 | 49.6 | 15 |
| Turnover component | Australasia | Agriculture (intense) | 3.9 | -20.6 | 32.2 | 70 |

A.6 Effects of range extent, habitat specificity and non-native proportions

Table S10: Responses of species richness to land use for Europe and North America, relative to primary vegetation (median, 2.5% and 97.5% percentiles of model responses), and number of studies and sites per coefficient. Prop. non-native: The proportion of non-native species as a fraction of species richness at a site.

| Region | Category | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|------------------|-----------------------|--------|-------|-------|----------|--------|
| Europe | Narrow range | Primary vegetation | 0.0 | NA | NA | 16 | 174 |
| Europe | Narrow range | Secondary vegetation | -49.4 | -57.6 | -40.3 | 42 | 453 |
| Europe | Narrow range | Agriculture (minimal) | -35.9 | -45.2 | -26.1 | 37 | 613 |
| Europe | Narrow range | Agriculture (light) | -49.9 | -58.1 | -40.8 | 37 | 810 |
| Europe | Narrow range | Agriculture (intense) | -43.2 | -52.3 | -32.7 | 38 | 858 |
| Europe | Wide range | Primary vegetation | 0.0 | NA | NA | 16 | 174 |
| Europe | Wide range | Secondary vegetation | 64.0 | 40.4 | 95.4 | 49 | 525 |
| Europe | Wide range | Agriculture (minimal) | 53.0 | 32.8 | 75.4 | 38 | 621 |
| Europe | Wide range | Agriculture (light) | 45.7 | 24.8 | 73.6 | 43 | 883 |
| Europe | Wide range | Agriculture (intense) | 36.1 | 16.5 | 62.7 | 46 | 1003 |
| Europe | High specificity | Primary vegetation | 0.0 | NA | NA | 7 | 77 |
| Europe | High specificity | Secondary vegetation | -31.2 | -46.7 | -11.5 | 20 | 324 |
| Europe | High specificity | Agriculture (minimal) | -38.6 | -52.3 | -21.8 | 18 | 256 |
| Europe | High specificity | Agriculture (light) | -38.2 | -53.5 | -20.2 | 26 | 492 |
| Europe | High specificity | Agriculture (intense) | -41.7 | -56.9 | -23.3 | 28 | 507 |
| Europe | Low specificity | Primary vegetation | 0.0 | NA | NA | 6 | 66 |
| Europe | Low specificity | Secondary vegetation | 74.7 | 33.4 | 122.7 | 21 | 338 |
| Europe | Low specificity | Agriculture (minimal) | 114.8 | 63.8 | 177.0 | 16 | 235 |
| Europe | Low specificity | Agriculture (light) | 86.3 | 37.9 | 145.4 | 26 | 495 |
| Europe | Low specificity | Agriculture (intense) | 92.3 | 40.1 | 151.7 | 29 | 599 |
| Europe | Prop. non-native | Primary vegetation | 0.0 | NA | NA | 13 | 105 |
| Europe | Prop. non-native | Secondary vegetation | -3.1 | -19.1 | 24.6 | 46 | 441 |
| Europe | Prop. non-native | Agriculture (minimal) | 2.3 | -18.4 | 28.0 | 36 | 548 |
| Europe | Prop. non-native | Agriculture (light) | 18.3 | -4.2 | 47.5 | 42 | 821 |
| Europe | Prop. non-native | Agriculture (intense) | 12.5 | -8.2 | 43.6 | 45 | 922 |
| N. America | Narrow range | Primary vegetation | 0.0 | NA | NA | 19 | 405 |
| N. America | Narrow range | Secondary vegetation | -19.1 | -29.6 | -6.5 | 18 | 270 |
| N. America | Narrow range | Agriculture (minimal) | -50.1 | -63.3 | -31.7 | 6 | 62 |

Continued on next page

| Region | Category | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|------------------|-----------------------|--------|-------|-------|----------|--------|
| N. America | Narrow range | Agriculture (light) | -36.1 | -49.6 | -20.6 | 10 | 52 |
| N. America | Narrow range | Agriculture (intense) | -41.5 | -55.7 | -24.0 | 8 | 33 |
| N. America | Wide range | Primary vegetation | 0.0 | NA | NA | 20 | 409 |
| N. America | Wide range | Secondary vegetation | 1.5 | -12.1 | 17.5 | 18 | 270 |
| N. America | Wide range | Agriculture (minimal) | 24.4 | -1.8 | 57.4 | 6 | 62 |
| N. America | Wide range | Agriculture (light) | -24.9 | -40.4 | -4.1 | 13 | 99 |
| N. America | Wide range | Agriculture (intense) | -32.9 | -48.5 | -8.8 | 9 | 35 |
| N. America | High specificity | Primary vegetation | 0.0 | NA | NA | 7 | 57 |
| N. America | High specificity | Secondary vegetation | 5.4 | -14.2 | 29.2 | 7 | 112 |
| N. America | High specificity | Agriculture (minimal) | -38.5 | -60.4 | -3.9 | 1 | 9 |
| N. America | High specificity | Agriculture (light) | -45.3 | -60.5 | -25.8 | 4 | 18 |
| N. America | High specificity | Agriculture (intense) | -56.7 | -70.5 | -39.2 | 3 | 18 |
| N. America | Low specificity | Primary vegetation | 0.0 | NA | NA | 8 | 73 |
| N. America | Low specificity | Secondary vegetation | -25.3 | -38.0 | -7.0 | 8 | 175 |
| N. America | Low specificity | Agriculture (minimal) | -14.6 | -35.0 | 14.6 | 2 | 48 |
| N. America | Low specificity | Agriculture (light) | -24.5 | -43.7 | 2.1 | 6 | 75 |
| N. America | Low specificity | Agriculture (intense) | -23.7 | -47.3 | 7.3 | 4 | 20 |
| N. America | Prop. non-native | Primary vegetation | 0.0 | NA | NA | 20 | 395 |
| N. America | Prop. non-native | Secondary vegetation | 29.0 | 5.4 | 61.4 | 18 | 258 |
| N. America | Prop. non-native | Agriculture (minimal) | 48.3 | 11.7 | 114.3 | 5 | 60 |
| N. America | Prop. non-native | Agriculture (light) | 54.6 | 13.1 | 138.3 | 13 | 94 |
| N. America | Prop. non-native | Agriculture (intense) | 62.1 | 3.7 | 174.2 | 8 | 25 |

Table S11: Responses of total abundance to land use for Europe and North America, relative to primary vegetation (median, 2.5% and 97.5% percentiles of model responses), and number of studies and sites per coefficient. Prop. non-native: The proportion of non-native species as a fraction of total abundance at a site.

| Region | Category | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|--------|--------------|-----------------------|--------|-------|-------|----------|--------|
| Europe | Narrow range | Primary vegetation | 0.0 | NA | NA | 13 | 105 |
| Europe | Narrow range | Secondary vegetation | 1.5 | -38.2 | 62.7 | 39 | 395 |
| Europe | Narrow range | Agriculture (minimal) | 10.5 | -32.6 | 69.2 | 35 | 559 |
| Europe | Narrow range | Agriculture (light) | -22.3 | -53.4 | 26.9 | 37 | 810 |
| Europe | Narrow range | Agriculture (intense) | -15.1 | -49.1 | 38.6 | 38 | 858 |
| Europe | Wide range | Primary vegetation | 0.0 | NA | NA | 13 | 105 |

Continued on next page

| Region | Category | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|------------------|-----------------------|--------|-------|-------|----------|--------|
| Europe | Wide range | Secondary vegetation | 41.8 | -10.1 | 129.7 | 45 | 466 |
| Europe | Wide range | Agriculture (minimal) | 63.9 | 0.3 | 164.1 | 36 | 567 |
| Europe | Wide range | Agriculture (light) | 39.6 | -14.2 | 129.9 | 42 | 881 |
| Europe | Wide range | Agriculture (intense) | 17.0 | -28.0 | 89.9 | 45 | 995 |
| Europe | High specificity | Primary vegetation | 0.0 | NA | NA | 6 | 73 |
| Europe | High specificity | Secondary vegetation | -45.2 | -66.4 | -9.2 | 17 | 266 |
| Europe | High specificity | Agriculture (minimal) | -25.4 | -53.7 | 15.1 | 17 | 250 |
| Europe | High specificity | Agriculture (light) | -16.1 | -50.9 | 40.4 | 25 | 490 |
| Europe | High specificity | Agriculture (intense) | -19.4 | -51.3 | 34.3 | 27 | 499 |
| Europe | Low specificity | Primary vegetation | 0.0 | NA | NA | 5 | 62 |
| Europe | Low specificity | Secondary vegetation | 304.4 | 134.5 | 598.7 | 18 | 280 |
| Europe | Low specificity | Agriculture (minimal) | 549.8 | 285.6 | 993.7 | 15 | 229 |
| Europe | Low specificity | Agriculture (light) | 349.2 | 156.3 | 697.0 | 25 | 493 |
| Europe | Low specificity | Agriculture (intense) | 413.7 | 196.4 | 813.3 | 28 | 591 |
| Europe | Prop. non-native | Primary vegetation | 0.0 | NA | NA | 13 | 105 |
| Europe | Prop. non-native | Secondary vegetation | -10.1 | -33.2 | 25.2 | 46 | 441 |
| Europe | Prop. non-native | Agriculture (minimal) | -13.5 | -39.8 | 27.4 | 36 | 548 |
| Europe | Prop. non-native | Agriculture (light) | 7.3 | -24.5 | 51.7 | 42 | 821 |
| Europe | Prop. non-native | Agriculture (intense) | 2.9 | -23.8 | 47.9 | 45 | 922 |
| N. America | Narrow range | Primary vegetation | 0.0 | NA | NA | 19 | 405 |
| N. America | Narrow range | Secondary vegetation | -23.8 | -43.3 | -0.6 | 18 | 270 |
| N. America | Narrow range | Agriculture (minimal) | -46.4 | -68.8 | -12.4 | 6 | 62 |
| N. America | Narrow range | Agriculture (light) | -55.5 | -72.0 | -28.9 | 10 | 52 |
| N. America | Narrow range | Agriculture (intense) | -67.3 | -82.5 | -40.2 | 7 | 23 |
| N. America | Wide range | Primary vegetation | 0.0 | NA | NA | 20 | 409 |
| N. America | Wide range | Secondary vegetation | 16.2 | -12.0 | 54.5 | 18 | 270 |
| N. America | Wide range | Agriculture (minimal) | 36.8 | -12.7 | 115.0 | 6 | 62 |
| N. America | Wide range | Agriculture (light) | -46.3 | -64.7 | -17.6 | 13 | 99 |
| N. America | Wide range | Agriculture (intense) | -38.7 | -66.2 | 8.0 | 8 | 25 |
| N. America | High specificity | Primary vegetation | 0.0 | NA | NA | 7 | 57 |
| N. America | High specificity | Secondary vegetation | 14.4 | -28.6 | 85.0 | 7 | 112 |
| N. America | High specificity | Agriculture (minimal) | 40.9 | -49.6 | 255.6 | 1 | 9 |
| N. America | High specificity | Agriculture (light) | -65.9 | -83.1 | -36.5 | 4 | 18 |
| N. America | High specificity | Agriculture (intense) | -84.0 | -95.4 | -59.6 | 2 | 8 |
| N. America | Low specificity | Primary vegetation | 0.0 | NA | NA | 8 | 73 |

Continued on next page

| Region | Category | Land use | Median | 2.5% | 97.5% | nStudies | nSites |
|------------|------------------|-----------------------|--------|-------|-------|----------|--------|
| N. America | Low specificity | Secondary vegetation | -44.5 | -64.8 | -12.0 | 8 | 175 |
| N. America | Low specificity | Agriculture (minimal) | -28.2 | -63.3 | 37.5 | 2 | 48 |
| N. America | Low specificity | Agriculture (light) | -56.3 | -75.7 | -21.6 | 6 | 75 |
| N. America | Low specificity | Agriculture (intense) | -6.7 | -57.0 | 113.3 | 3 | 10 |
| N. America | Prop. non-native | Primary vegetation | 0.0 | NA | NA | 20 | 395 |
| N. America | Prop. non-native | Secondary vegetation | 4.9 | -25.6 | 49.2 | 18 | 258 |
| N. America | Prop. non-native | Agriculture (minimal) | -14.2 | -55.7 | 52.6 | 5 | 60 |
| N. America | Prop. non-native | Agriculture (light) | 63.6 | -7.8 | 246.4 | 13 | 94 |
| N. America | Prop. non-native | Agriculture (intense) | 24.8 | -47.4 | 299.0 | 8 | 25 |

B Figures

B.1 Taxonomic representation

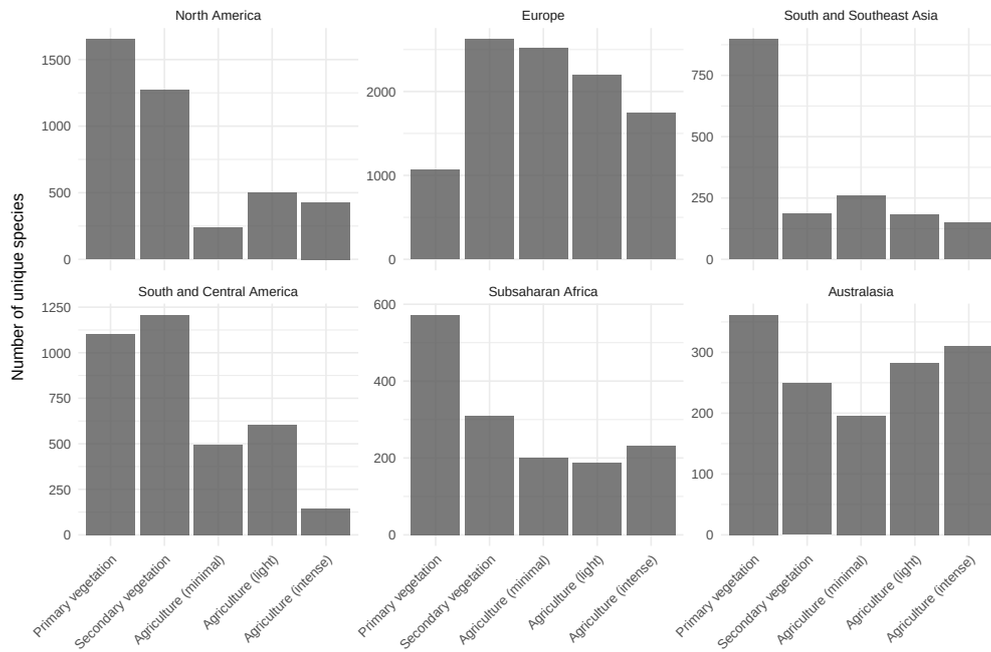


Figure S1: Number of unique arthropod species per land use and region in our dataset of the PREDICTS database.

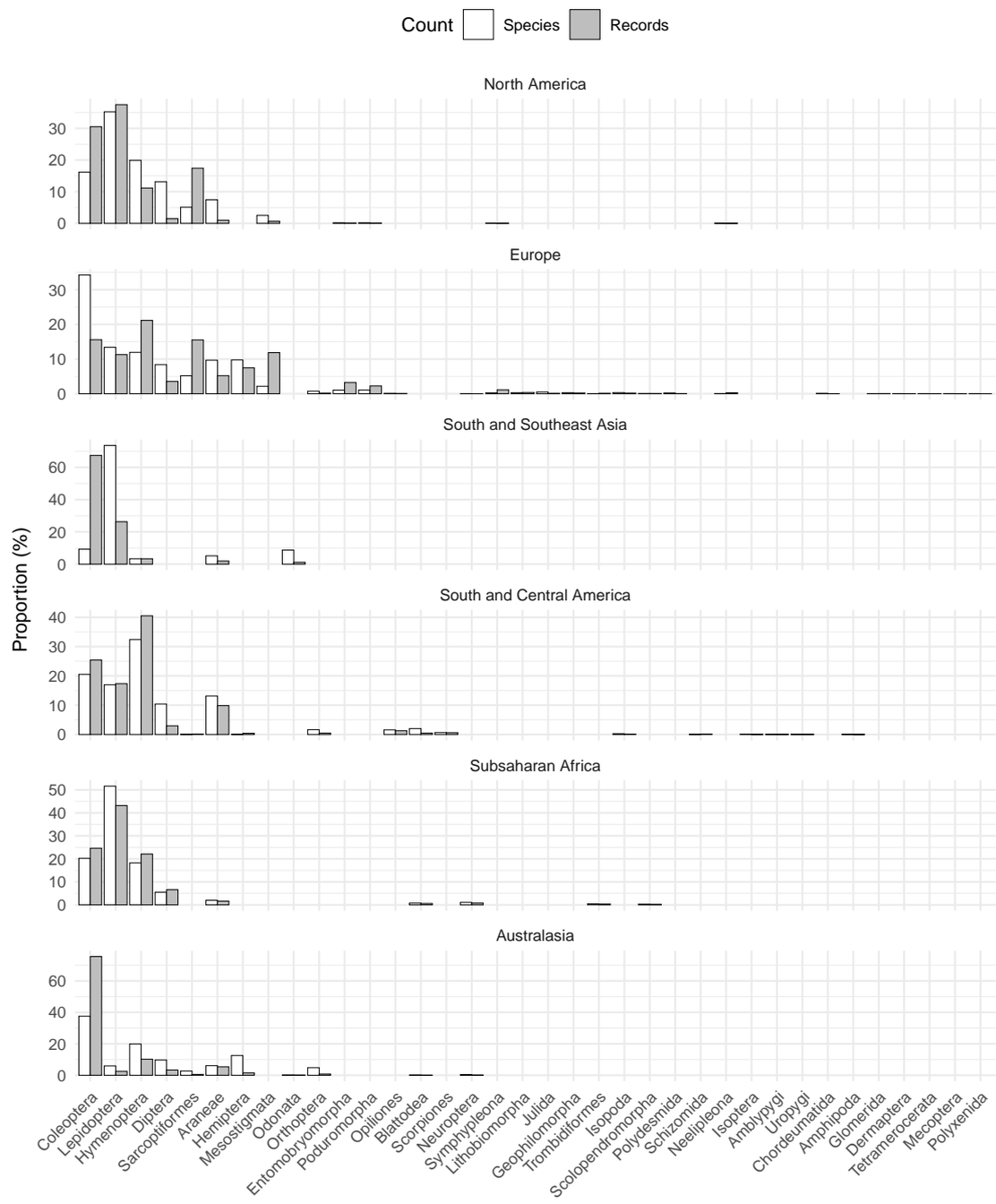


Figure S2: Percentages of unique arthropod species and records per taxonomic order and region, relative to the total per region, in our dataset of the PREDICTS database.

B.2 Vertebrate records

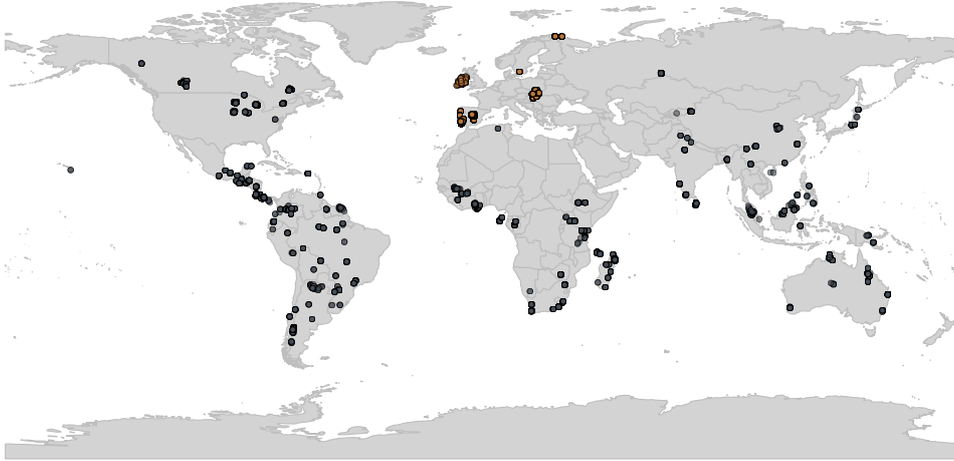


Figure S3: Sites with vertebrate records (phylum Chordata) from the PREDICTS database. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

B.3 Natural habitat availability

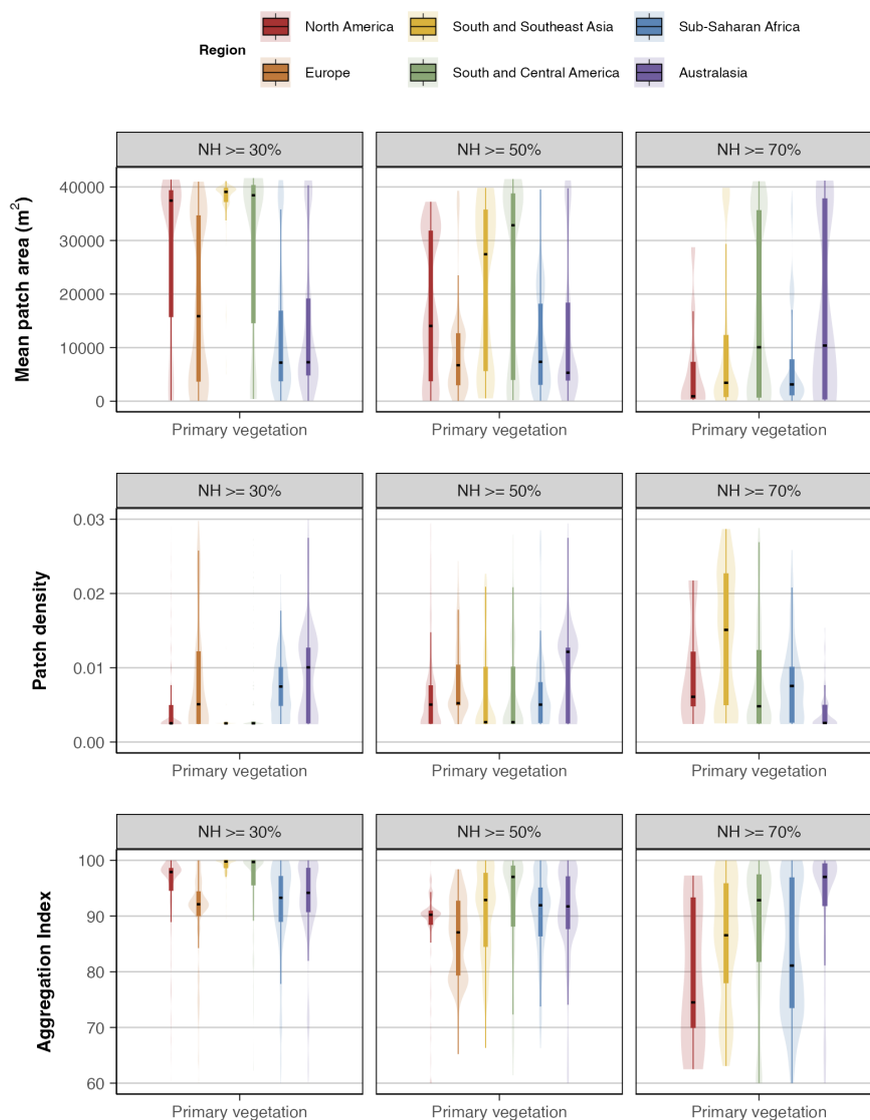


Figure S4: Connectivity of natural habitat (wilderness, forestry areas and open woodland grazing land) within 10km-radius buffers surrounding the primary vegetation sites from PREDICTS that were used in this study. Plotted are three standard habitat connectivity measures of categorical habitat layers: Mean patch area, patch density and aggregation index (He et al., 2000). Binary layers were obtained by converting continuous fractional layers to binary layers using a threshold of 0.5. In Europe, mean patch area tends to be lower, patch density higher and aggregation index lower, indicating more fragmentation. Differences are larger where the mean natural habitat in the buffer is higher, suggesting that it is particularly dense natural habitat that is more fragmented in Europe.

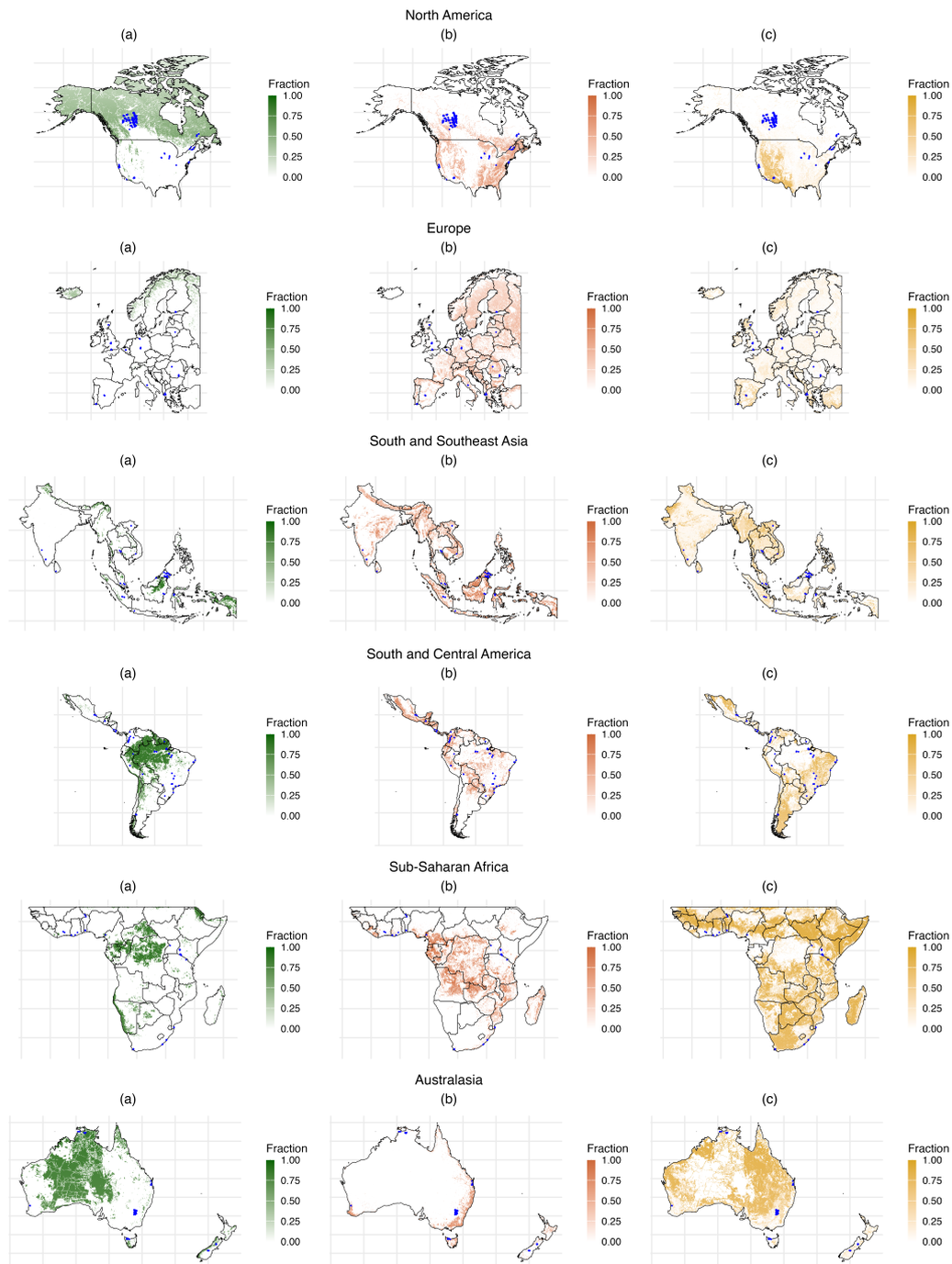


Figure S5: Maps of wilderness (a), forestry areas (b) and open woodland grazing land (c) in each region, plotted for the year 2005, and PREDICTS primary vegetation sites plotted in blue. Regions were cropped to exclude overseas territories like Hawaii from North America, or parts of a region in which no records were sampled in PREDICTS, such as Madagascar in sub-Saharan Africa or Iceland in Europe. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

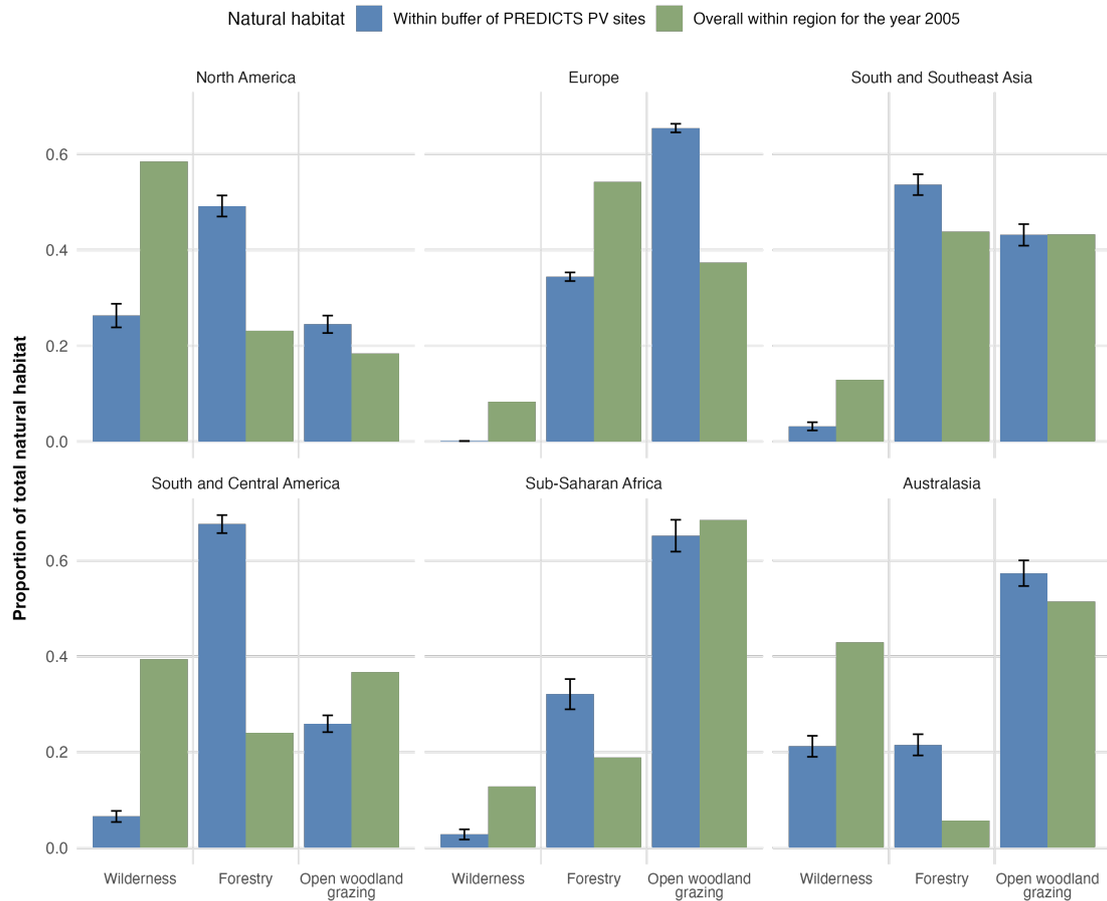


Figure S6: Proportions of wilderness, forestry areas and open woodland grazing land, of total natural habitat within 5-km buffers surrounding PREDICTS sites of primary vegetation (mean and 95% confidence interval), and overall within each region for the year 2005. The latter were computed directly from the natural habitat. The figure reveals a clear selection biases towards areas with lower proportions of wilderness. Regions were cropped to exclude overseas territories like Hawaii from North America, or parts of a region in which no records were sampled in PREDICTS, such as Madagascar in sub-Saharan Africa or Iceland in Europe. Corresponding maps are plotted in Fig. S5.

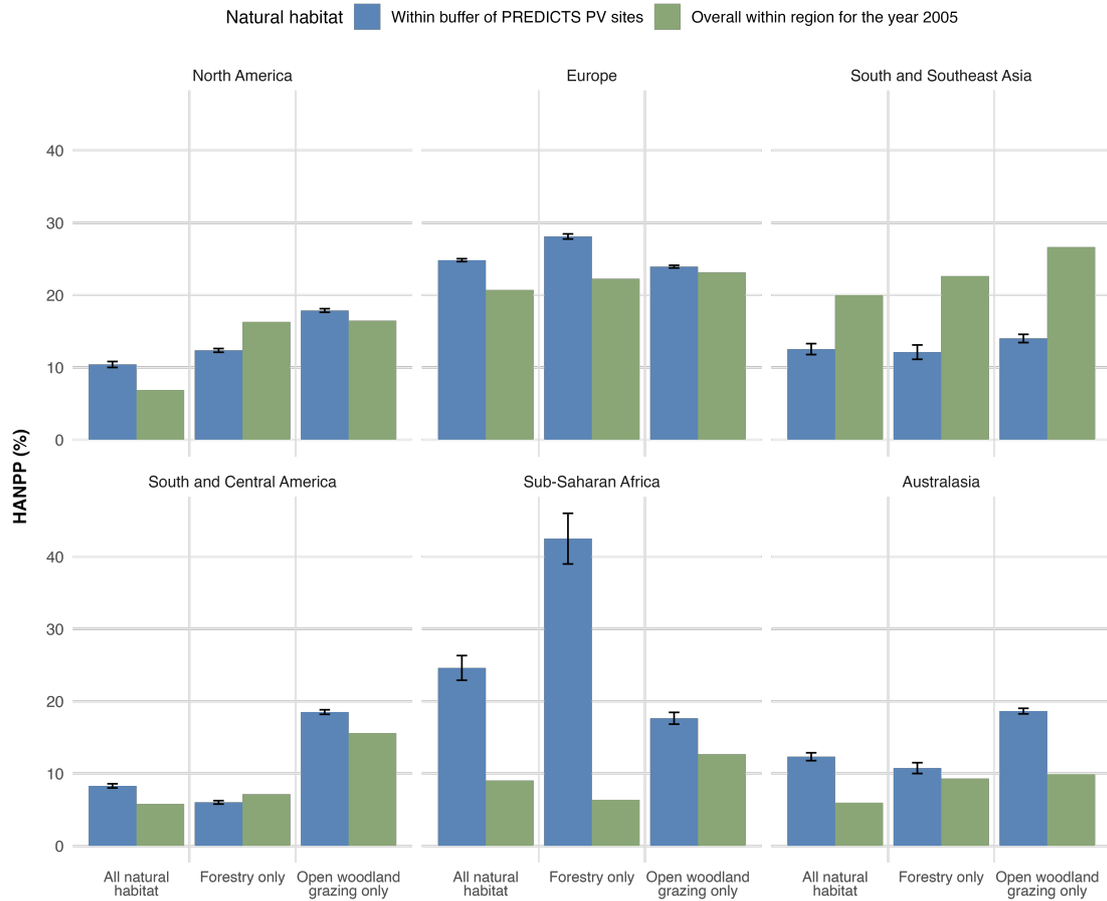


Figure S7: %HANPP per region within 5-km buffers surrounding PREDICTS sites of primary vegetation (mean and 95% confidence interval), and overall within each region for the year 2005. 'All natural habitat': Total HANPP as a percentage of the total potential NPP of the natural habitat. 'Forestry only': Forestry HANPP as a percentage of the potential NPP of forestry areas only. 'Open woodland grazing only': Open woodland grazing HANPP as a percentage of the potential NPP of open woodland grazing land only. The figure reveals a clear selection bias towards higher %HANPP in all region except South and Southeast Asia. This bias is particularly evident for forestry in sub-Saharan Africa, where sampled primary vegetation sites in the PREDICTS database (for arthropods) are sampled in forestry that is subject to almost 6.38 times the %HANPP of forestry areas overall in the region. Regions were cropped to exclude overseas territories like Hawaii from North America, or parts of a region in which no records were sampled in PREDICTS, such as Madagascar in sub-Saharan Africa or Iceland in Europe. Corresponding maps can be found in Fig. S5.

B.4 Compositional dissimilarity

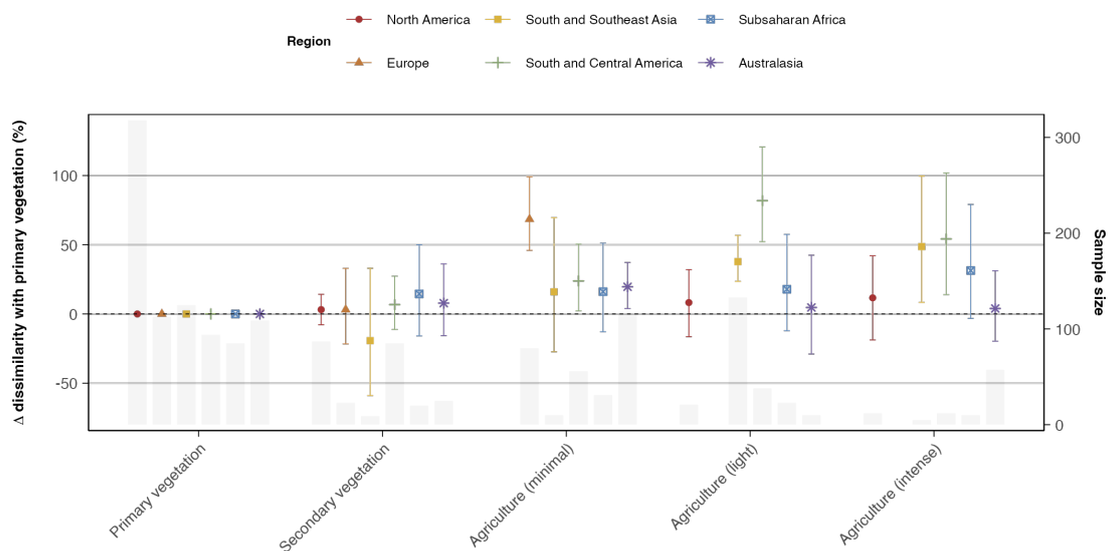


Figure S8: Compositional dissimilarity (Sørensen's Index) of each land use with primary vegetation per region, shown relative to the dissimilarity of primary vegetation with other primary vegetation sites (%). Points indicate median modelled responses and error bars represent 95% credible intervals. The right y-axis shows sample sizes (number of sites), plotted as grey bars. Only model coefficients computed from at least 10 sites are shown.

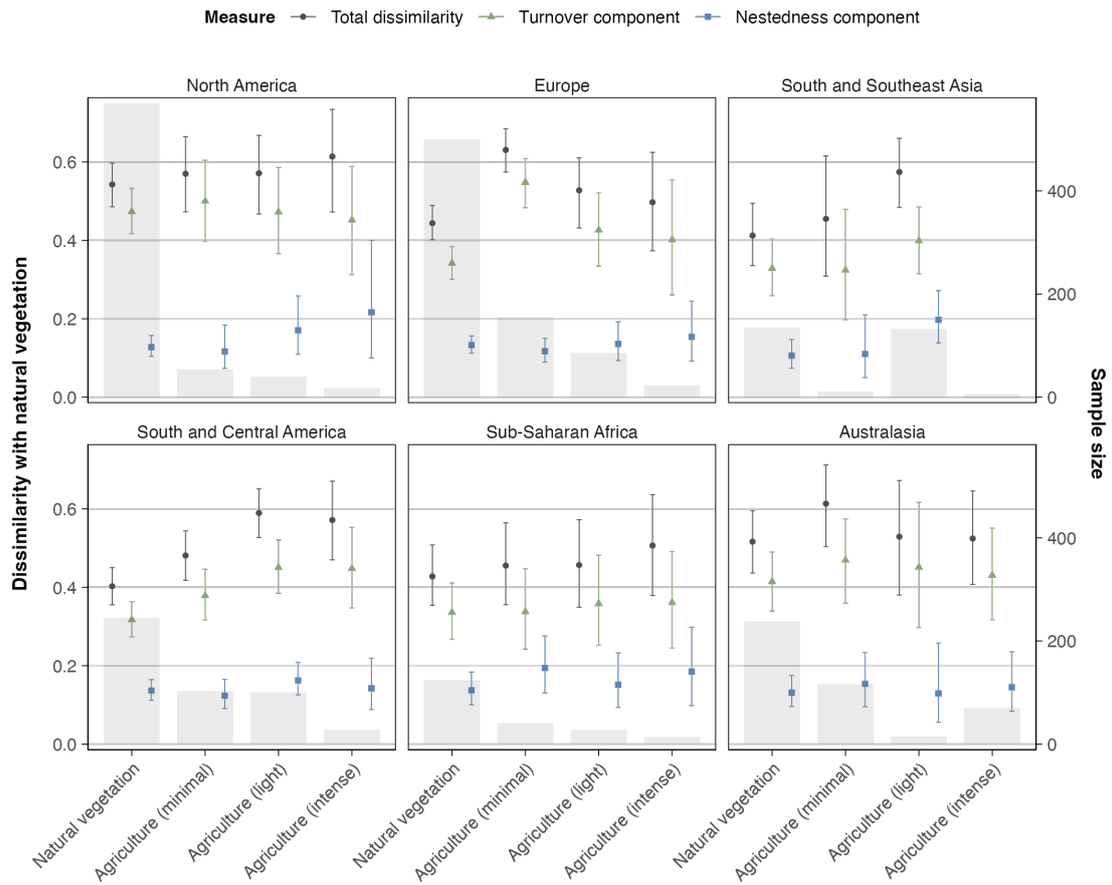


Figure S9: Absolute predictions of compositional dissimilarity (Sørensen's Index) of each land use with natural vegetation per region (%). Points indicate median modelled responses and error bars represent 95% credible intervals. The right y-axis shows sample sizes (number of sites), plotted as grey bars. Only model coefficients computed from at least 10 sites are shown.

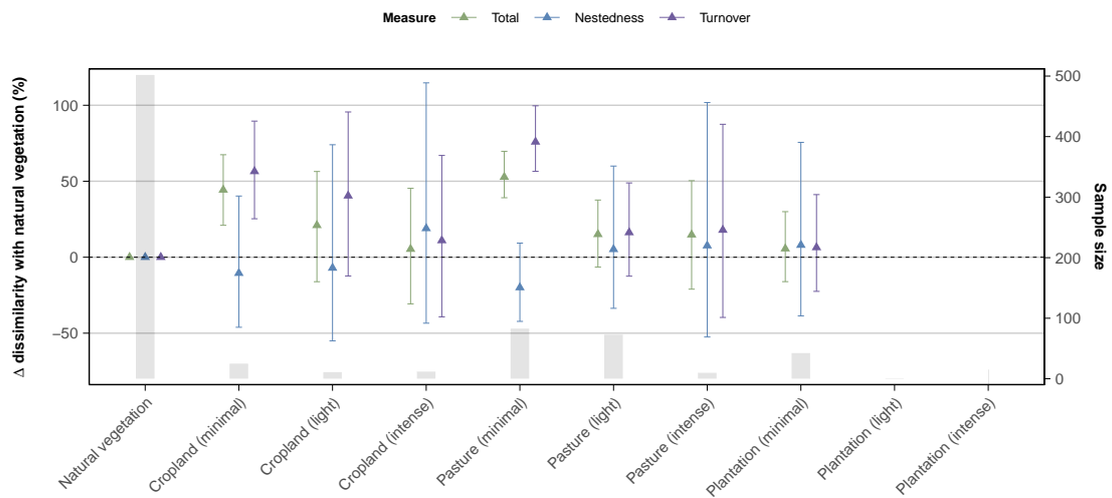


Figure S10: Compositional dissimilarity (Sørensen's Index) of each land use with natural vegetation in Europe, shown relative to the dissimilarity of natural vegetation with other natural vegetation sites (%). Points indicate median modelled responses and error bars represent 95% credible intervals. The right y-axis shows sample sizes (number of sites), plotted as grey bars. Only model coefficients computed from at least 10 sites are shown. The figure shows that species turnover in minimal-use agriculture stems from cropland and pasture sites. It also shows that direct, within-study comparisons of agricultural sites with primary or secondary vegetation are very low, despite the large number of PREDICTS records in agricultural sites in Europe, highlighting an important data deficiency.

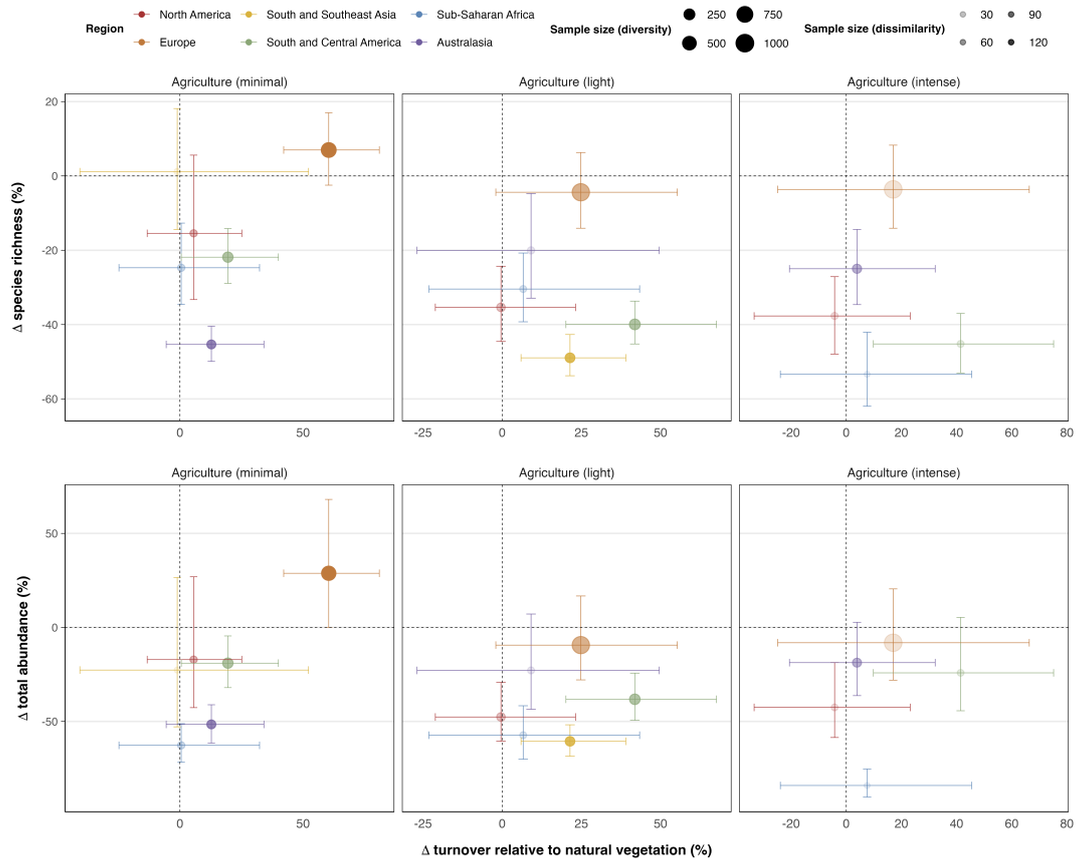


Figure S11: Turnover relative to natural vegetation (primary and secondary vegetation) of main text Fig. 4 plotted against the responses of species richness and total abundance (relative to primary vegetation) of main text Fig. 1. Points indicate median modelled responses and error bars represent 95% credible intervals. The sample sizes (number of sites) of diversity coefficients are indicated by the size of the points while the sample sizes of turnover coefficients are indicated by the transparency of the points.

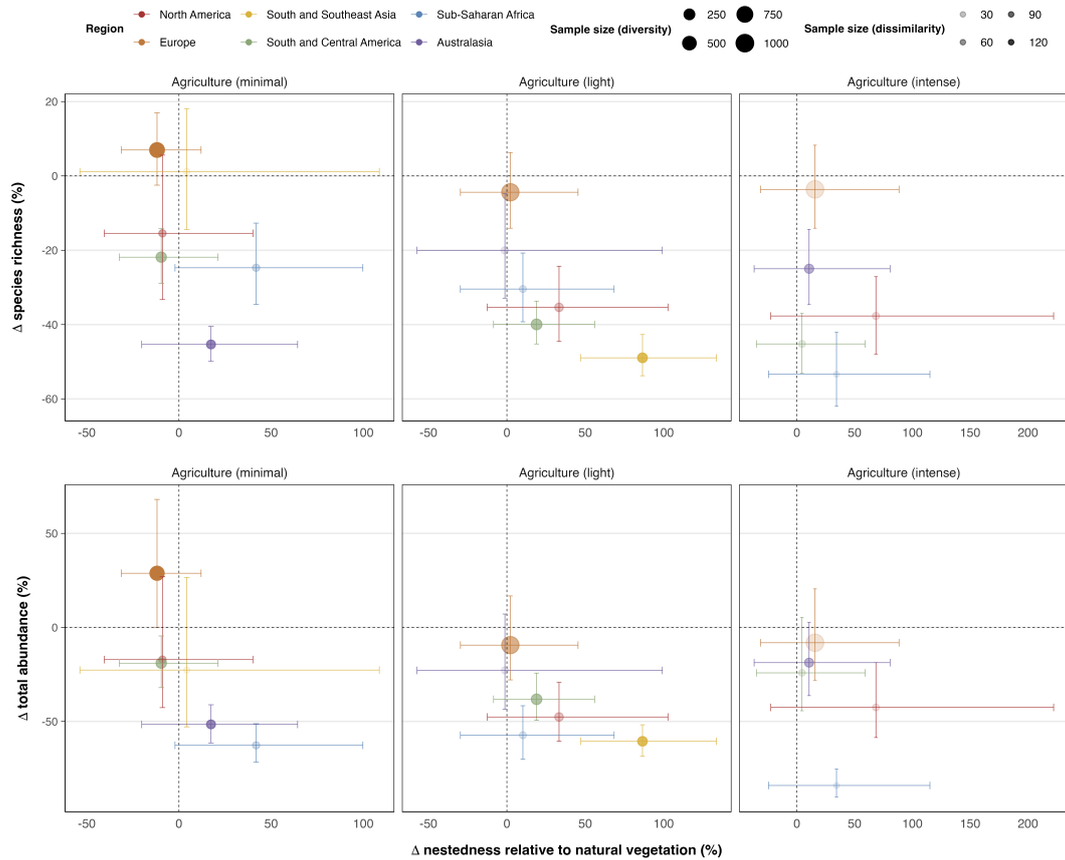


Figure S12: Nestedness relative to natural vegetation (primary and secondary vegetation) of main text Fig. 4 plotted against the responses of species richness and total abundance (relative to primary vegetation) of main text Fig. 1. Points indicate median modelled responses and error bars represent 95% credible intervals. The sample sizes (number of sites) of diversity coefficients are indicated by the size of the points while the sample sizes of nestedness coefficients are indicated by the transparency of the points.

B.5 Effects of range extent, habitat specificity and non-native species

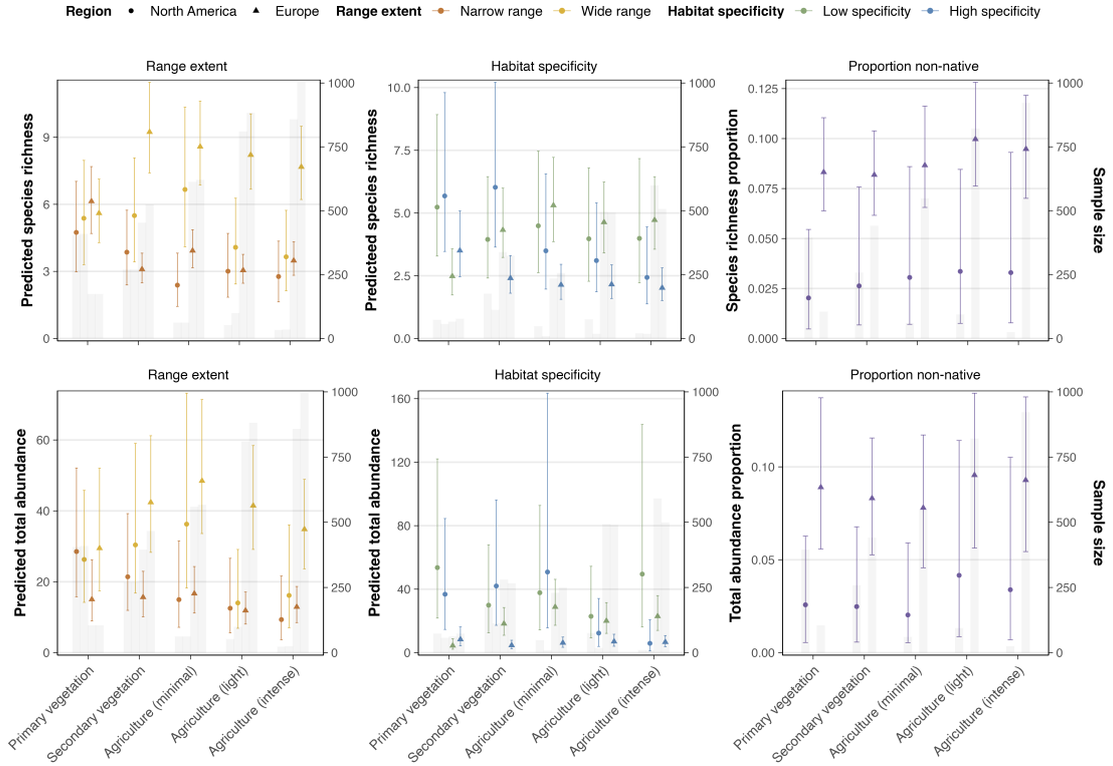


Figure S13: Absolute predictions of species richness and total abundance per land use in Europe and North America, comparing responses of species with narrow and wide range extent, and high and low habitat specificity. Non-native proportion: The proportion of non-native species in a site, as a fraction of species richness or total abundance. Points indicate median modelled responses and error bars represent 95% credible intervals. The right y-axis shows sample sizes (number of sites). Only model coefficients computed from at least 10 sites are shown.

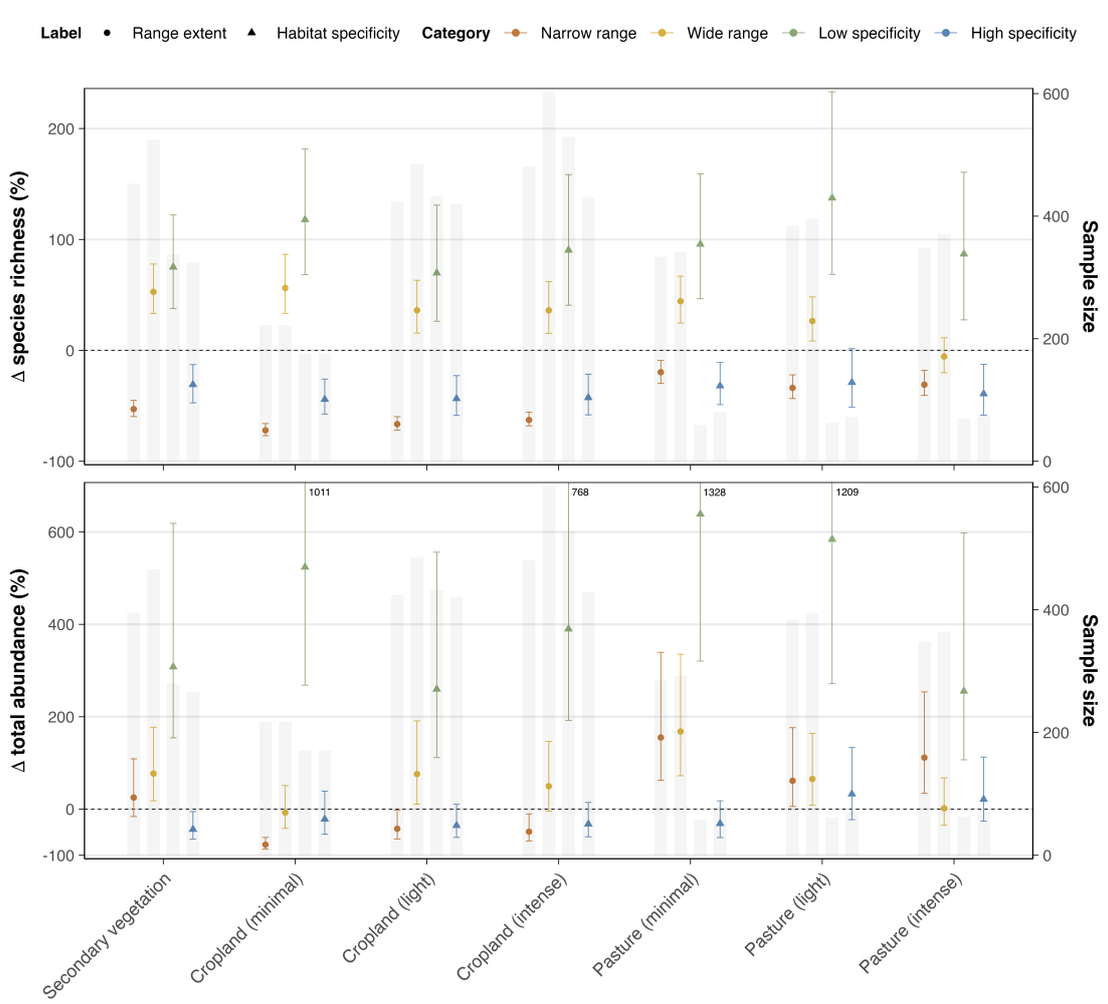


Figure S14: Arthropod diversity responses (species richness and total abundance) to individual agricultural land use types in Europe, relative to primary vegetation, comparing species with narrow and wide range extent, and high and low habitat specificity. Points indicate median modelled responses and error bars represent 95% credible intervals. The right y-axis shows sample sizes (number of sites). Only model coefficients computed from at least 10 sites are shown.

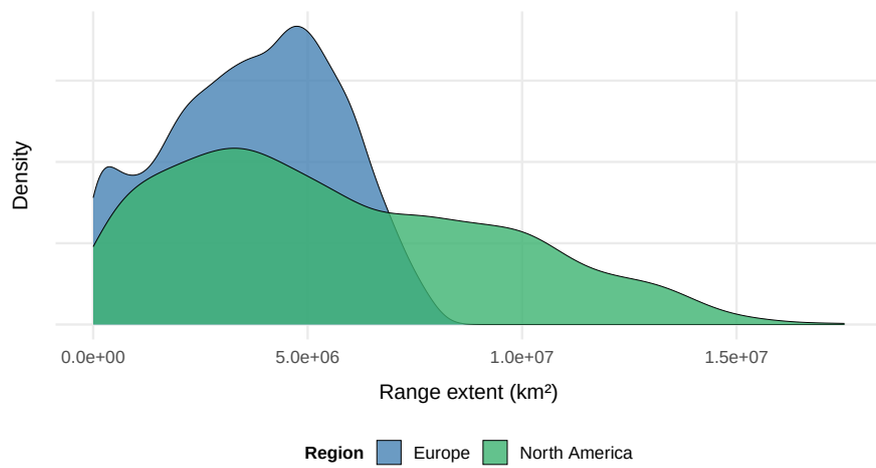


Figure S15: Densities of range extents for Europe and North America.

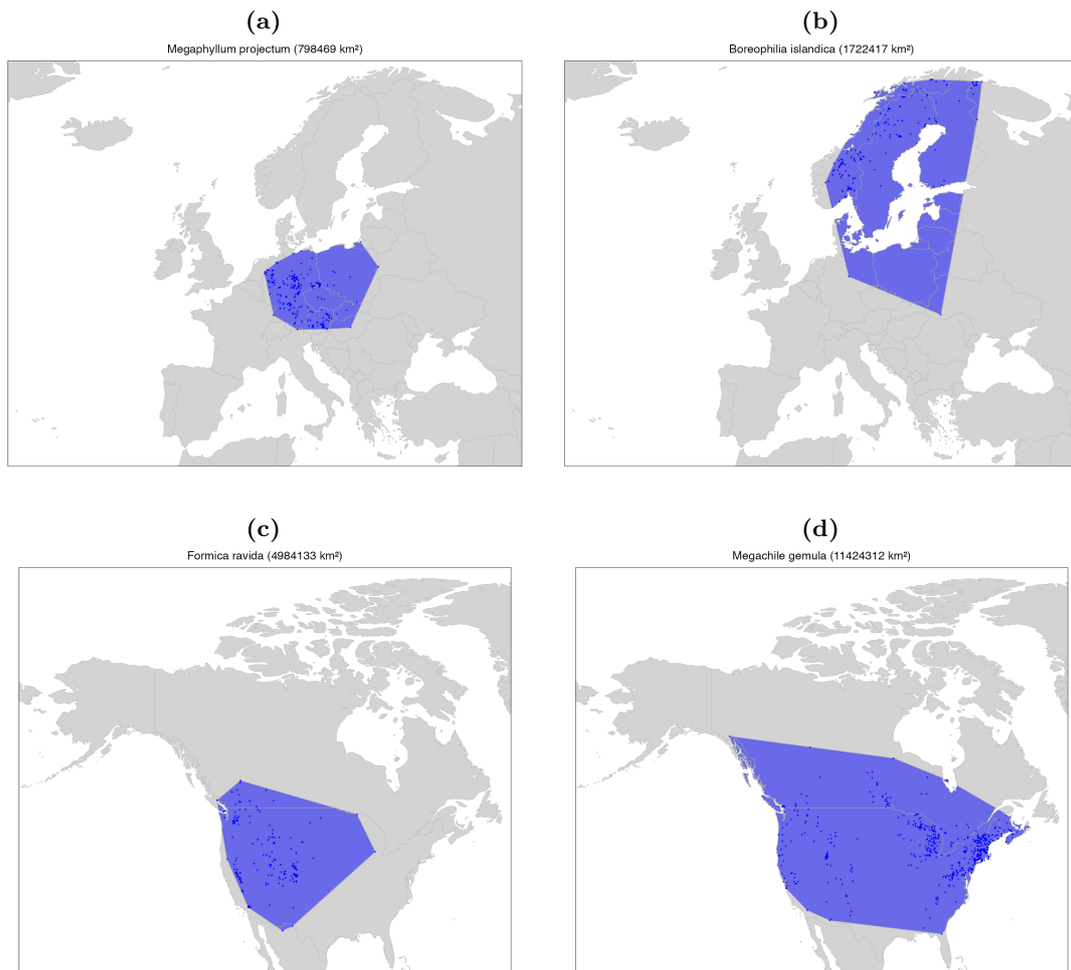


Figure S16: Examples of range extents for Europe (a, b) and North America (c, d). Map lines delineate study areas and do not necessarily depict accepted national boundaries.

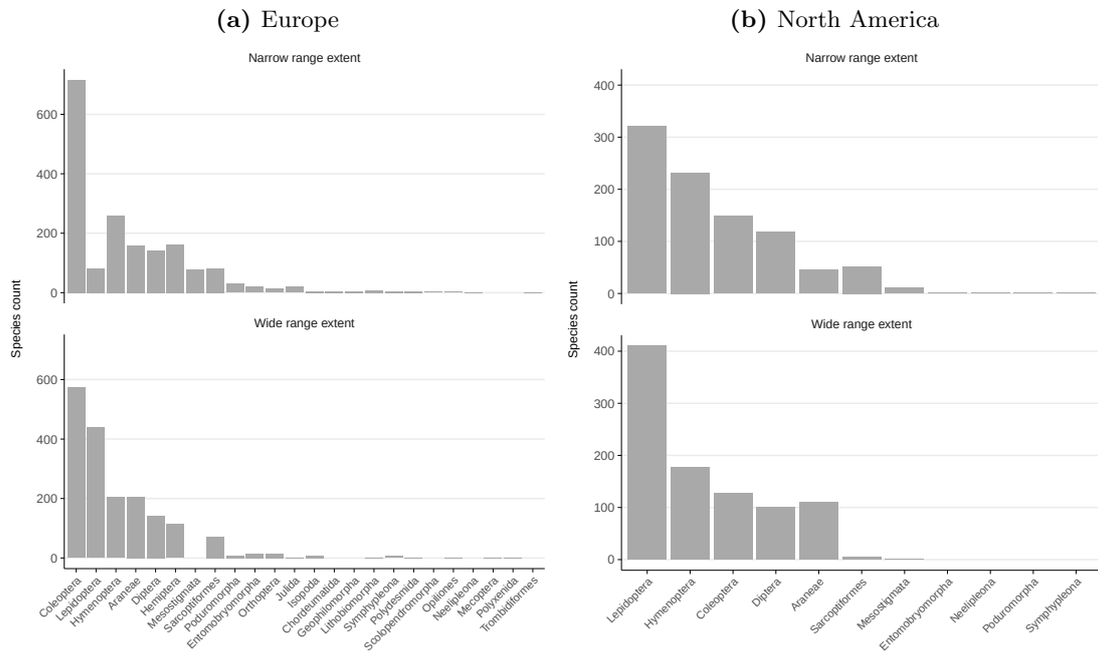


Figure S17: Species counts for taxonomic orders per range extent category in Europe (a) and North America (b).

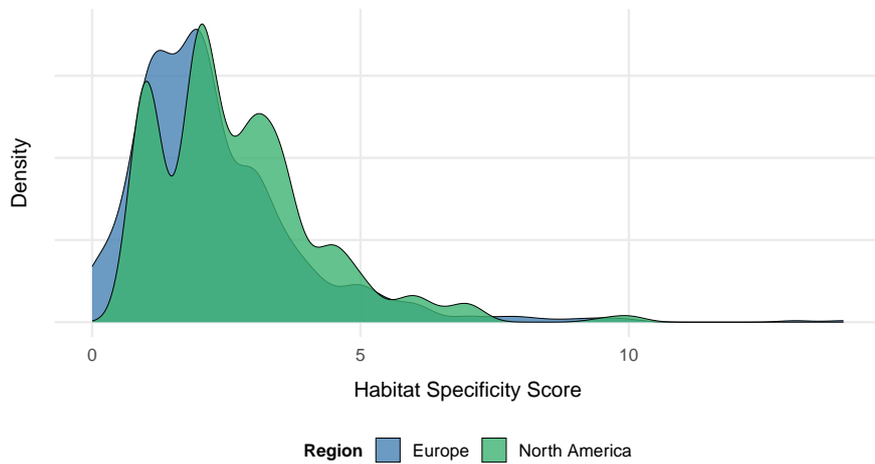


Figure S18: Densities of habitat specificity score for Europe and North America. A higher score implies less habitat specificity.

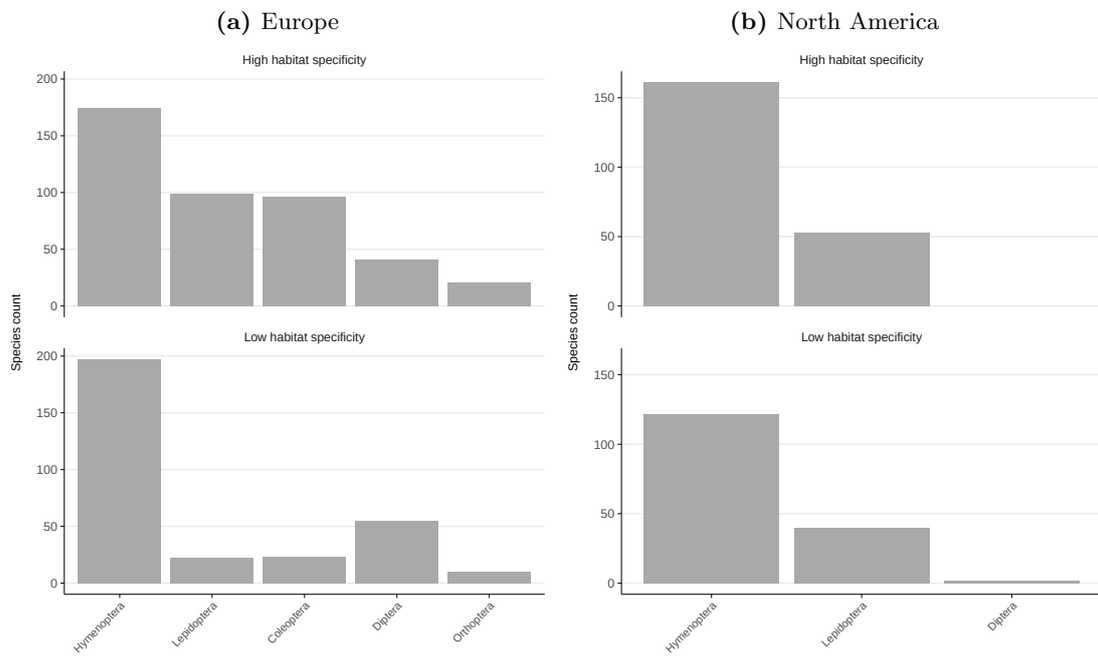


Figure S19: Species counts for taxonomic orders per habitat specificity category in Europe (a) and North America (b).

B.6 Abundance model comparison

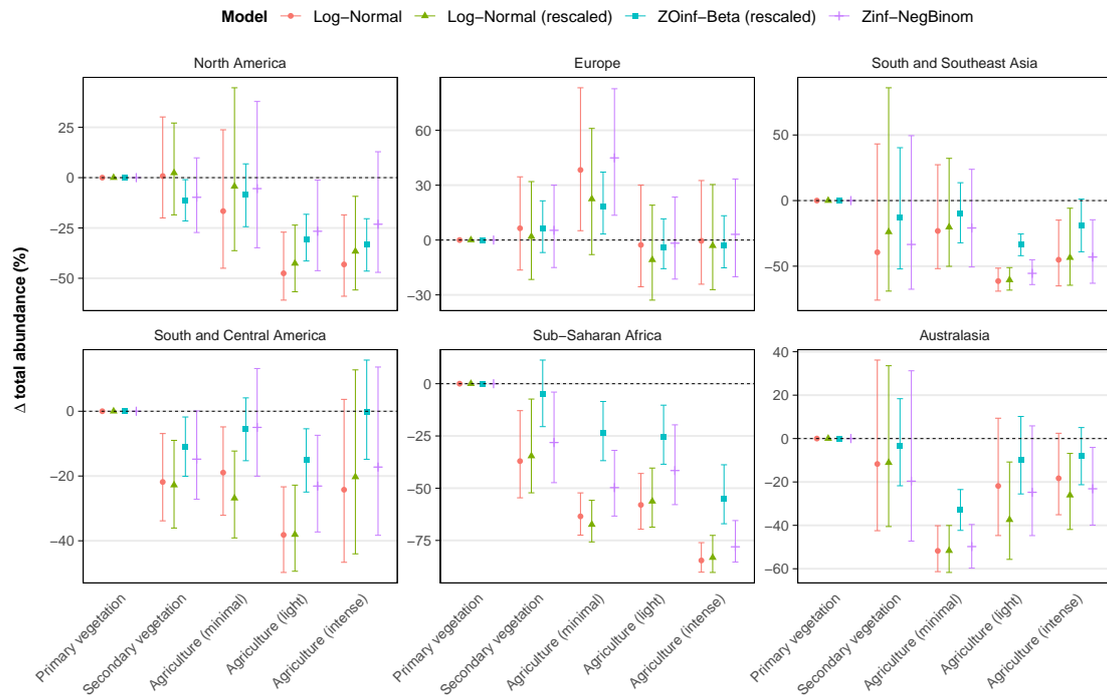


Figure S20: Comparison of model responses of abundance per land use and region as percentage change from primary vegetation. Points indicate median modelled responses and error bars represent 95% credible intervals. Shifted from left-to-right: Rescaled abundance modelled with a zero-and-one inflated beta distribution (red), log-transformed rescaled abundance modelled with a Gaussian distribution (green), and log-transformed total abundance (unscaled) modelled with a Gaussian distribution (blue). Although there is approximate agreement between the models, responses from the beta model tend to be more conservative.

B.7 Sensitivity analysis

To test the robustness of the regional responses to agricultural land use shown in Figure 1, we conducted a sensitivity analysis on the model coefficients by removing individual studies from the model data and refitting the models of species richness and total abundance (Fig. S21 and Fig. S22). Due to the high computational cost of refitting Bayesian `brms` models, we used the R package `glmmTMB` (Brooks et al., 2017) to fit these models. Major changes in coefficients due to the removal of a study were further investigated but disregarded if the changes were not consistent across land uses and/or if the study in question corresponded to the largest study in that region (in which case large changes in coefficients are to be expected). For example, whilst the Australasian study `CC1_2007_Ewers 1` (New Zealand) results to substantially more negative diversity change across minimal-, light- and intense-use agriculture, this study is the largest PREDICTS study in Australasia and was therefore not removed from the analysis. This does suggest, however, that diversity declines in agricultural landscapes in Australasia may be less pronounced in Australia than in New Zealand.

For Europe, a consistent change in coefficients of species richness and total abundance was observed for the study `SC1.2011_Meijer 1`, which corresponds to one of two arthropod studies in the PREDICTS database that stem from the Azores islands. Due to the disproportionate impact of the Azores on European biodiversity responses, their geographical isolation as a small island group in the Atlantic ocean, and their substantial distance from the European mainland, we decided to remove the Azores records from the analysis. To be consistent, we removed studies from islands of similar size from other regions, which resulted in the removal of studies from Puerto Rico (South and Central America) and Comoros (Subsaharan Africa).

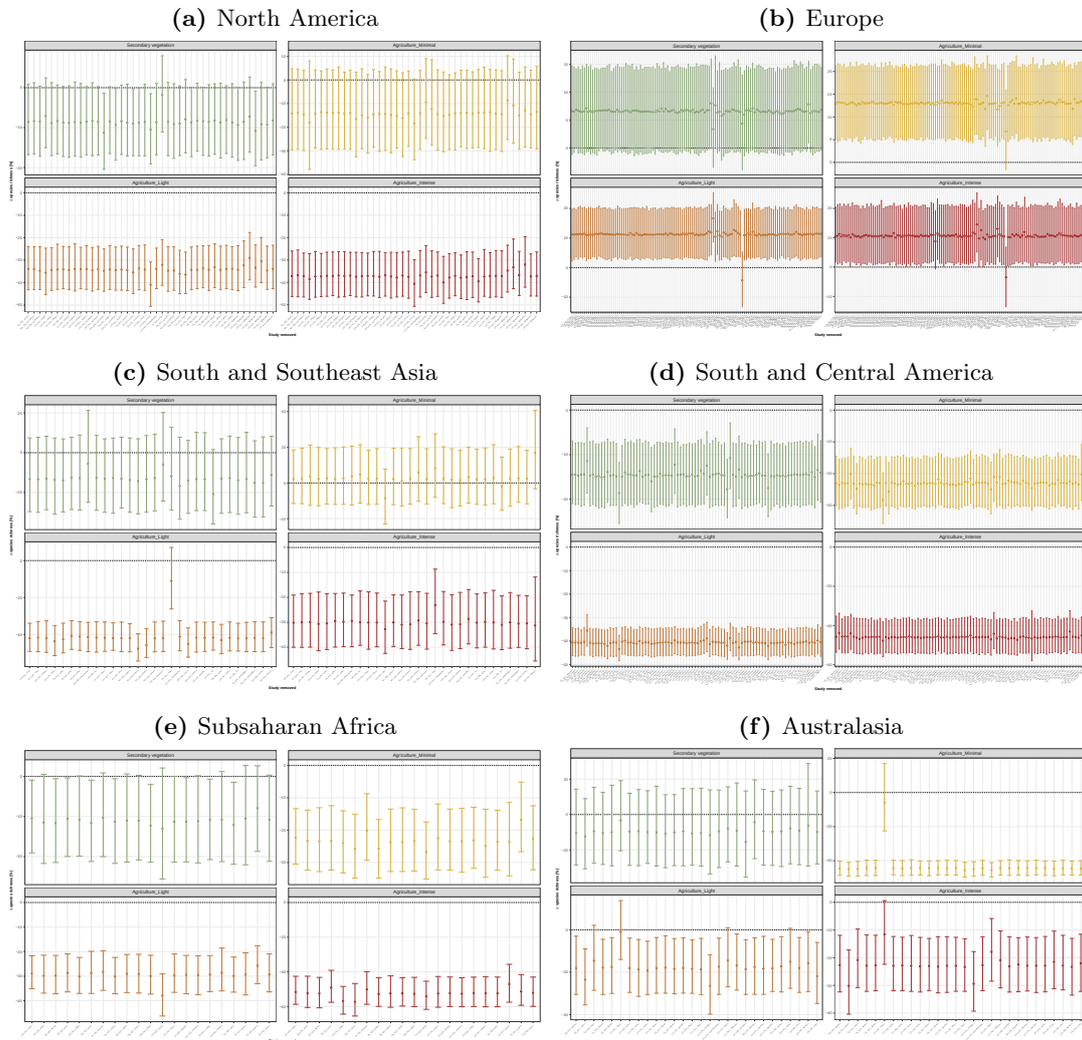


Figure S21: Sensitivity test of individual studies (through their individual removal from the dataset) on the regional model coefficients of species richness in secondary vegetation and minimal-, light- and intense-use agriculture, relative to primary vegetation. Points indicate median modelled responses and error bars represent 95% confidence intervals. In Europe, we observe a consistent bias across land use coefficients for the study `SC1_2011_Meijer 1` (located in the Azores island group), which results in substantially more positive coefficients of diversity change if the study is included.

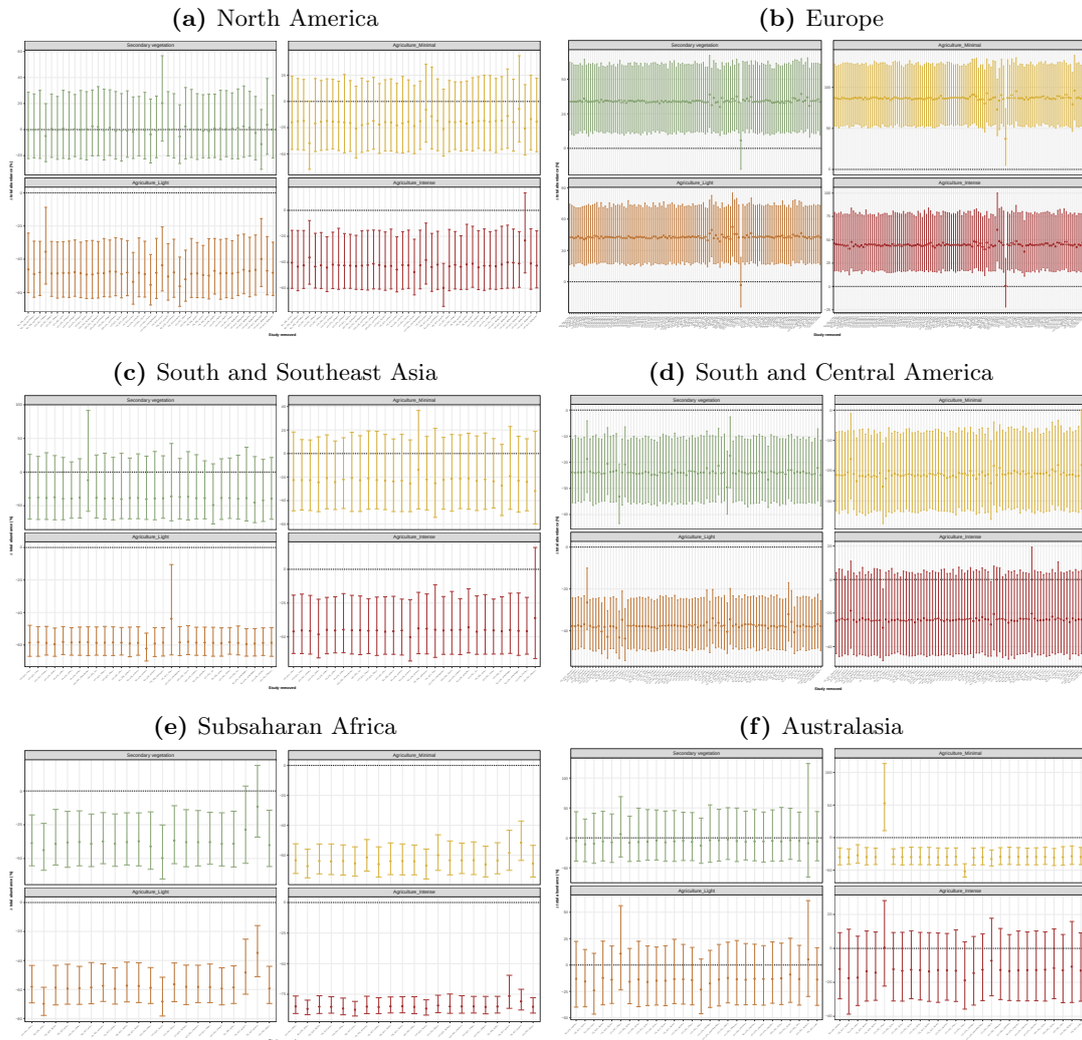


Figure S22: Sensitivity test of individual studies (through their individual removal from the dataset) on the regional model coefficients of total abundance in secondary vegetation and minimal-, light- and intense-use agriculture, relative to primary vegetation. Points indicate median modelled responses and error bars represent 95% confidence intervals. In Europe, we observe a consistent bias across land use coefficients for the study *SC1_2011_Meijer_1* (located in the Azores island group), which results in substantially more positive coefficients of diversity change if the study is included.

B.8 Influence of soil-core samples

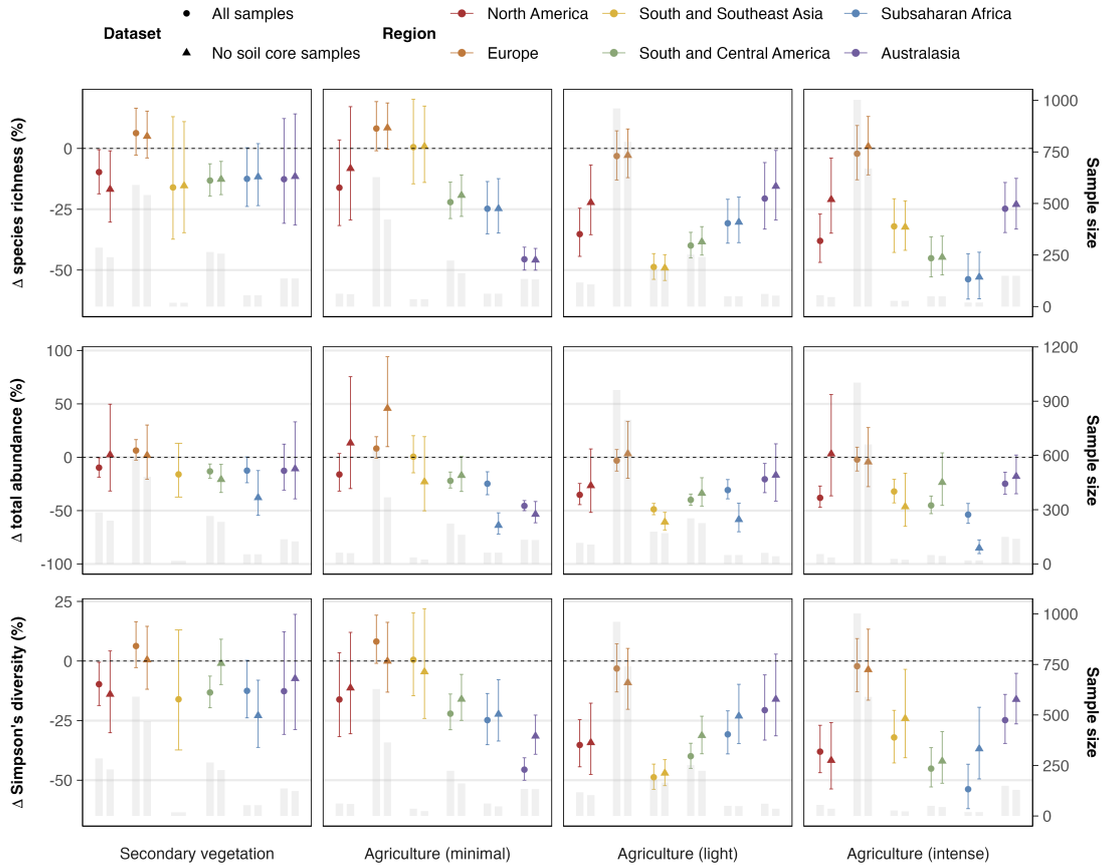


Figure S23: Comparison of the PREDICTS dataset used in this study with and without soil-core samples, comparing regional responses of species richness, total abundance and Simpson's diversity index to agricultural land use, relative to primary vegetation. Points indicate median modelled responses and error bars represent 95% confidence intervals. The figure shows that the removal of soil core samples does not substantially change diversity responses, with the exception that the response of total abundance in sub-Saharan Africa is more negative without soil core samples.

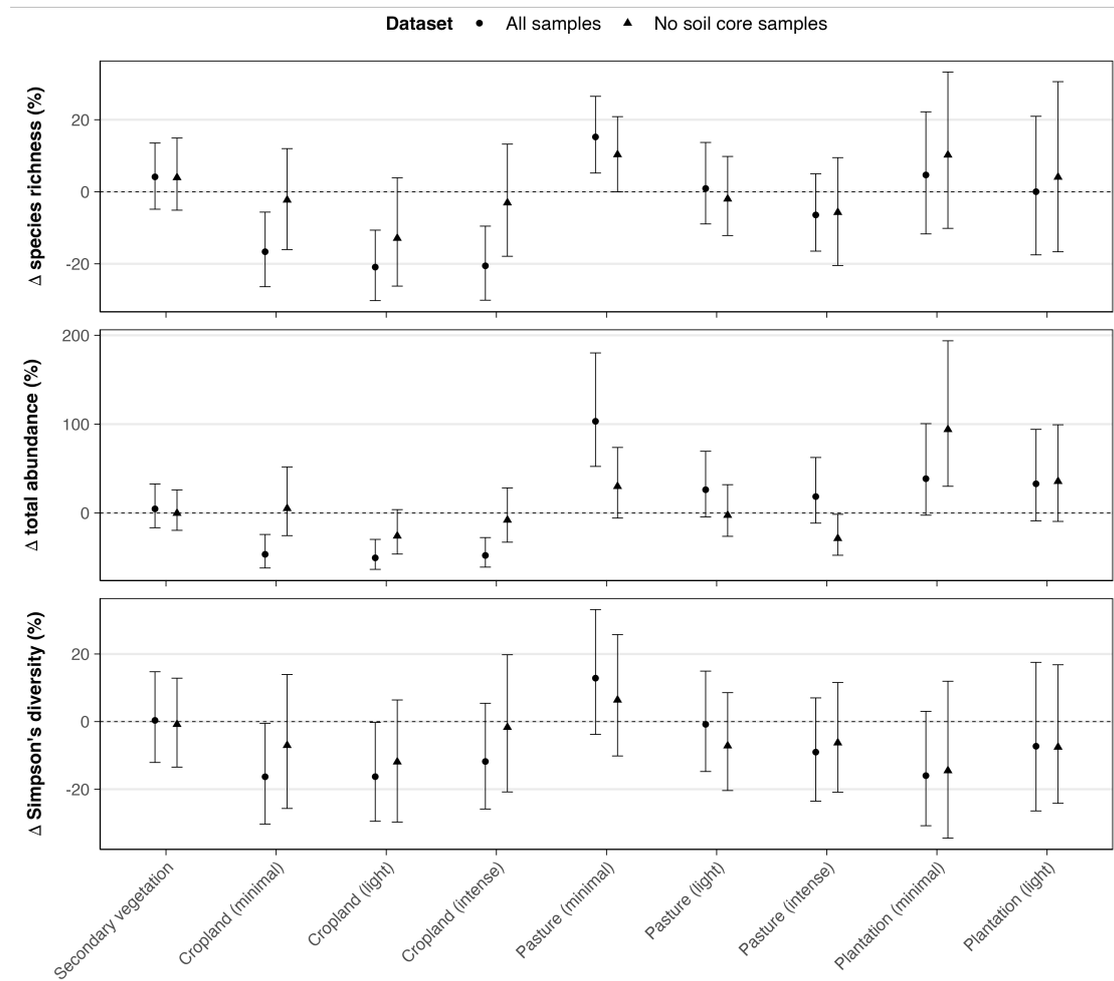


Figure S24: Comparison of the PREDICTS dataset used in this study with and without soil-core samples, comparing European responses of species richness, total abundance and Simpson's diversity index relative to primary vegetation, per individual land-use type and intensity. Points indicate median modelled responses for Europe and error bars represent 95% confidence intervals. The figure shows that the removal of soil core samples does not substantially change diversity responses, although responses in cropland are slightly less negative without core samples.

References

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- He, H. S., DeZonia, B. E., and Mladenoff, D. J. (Oct. 2000). “An aggregation index (AI) to quantify spatial patterns of landscapes”. In: *Landscape Ecology* 15.7, pp. 591–601. ISSN: 1572-9761. DOI: 10.1023/A:1008102521322.
- Hudson, L. N. et al. (2017). “The database of the PREDICTS (Projecting Responses of Ecological Diversity In Changing Terrestrial Systems) project”. In: *Ecology and Evolution* 7.1, pp. 145–188. DOI: <https://doi.org/10.1002/ece3.2579>.

Supplementary Information B

Model Fits and Diagnostics Plots

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A Regional responses

The following model diagnostic plots refer to the comparison of regional responses of biodiversity in primary vegetation, secondary vegetation and agricultural land use (Fig. 1).

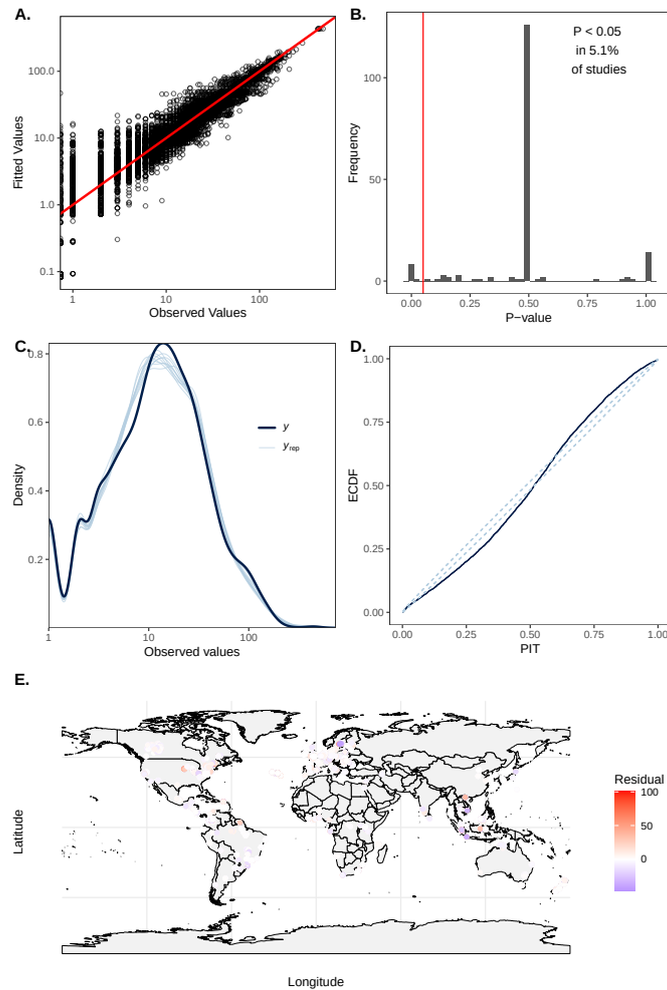


Figure M1: Model diagnostic plots for the zero-inflated negative binomial model of global species richness responses to land use: $\text{Richness} \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

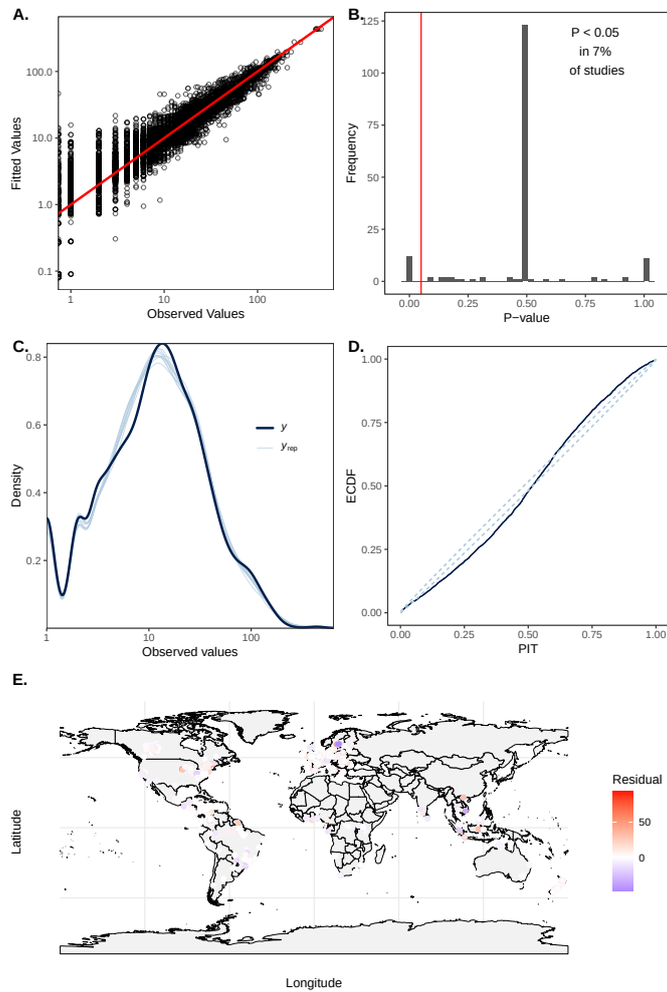


Figure M2: Model diagnostic plots for the zero-inflated negative binomial model of regional species richness responses to land use: $\text{Richness} \sim \text{LandUse} \times \text{Region} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M1: Posterior summary of the global model of species richness (Fig. M1).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ UI2 + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 7604) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 341) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.24 | 0.05 | 1.15 | 1.35 | 1.00 | 1224 | 2943 |
| ~SS:SSB (Number of levels: 1696) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.21 | 0.01 | 0.19 | 0.24 | 1.00 | 5120 | 9570 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.84 | 0.07 | 2.70 | 2.97 | 1.01 | 632 | 1234 |
| zi_Intercept | -5.37 | 0.22 | -5.84 | -4.97 | 1.00 | 24669 | 13954 |
| UI2Secondaryvegetation | -0.13 | 0.02 | -0.17 | -0.09 | 1.00 | 13895 | 15118 |
| UI2Agriculture_Minimal | -0.22 | 0.02 | -0.27 | -0.18 | 1.00 | 15282 | 14596 |
| UI2Agriculture_Light | -0.33 | 0.02 | -0.37 | -0.28 | 1.00 | 11134 | 13619 |
| UI2Agriculture_Intense | -0.36 | 0.03 | -0.41 | -0.31 | 1.00 | 12246 | 14491 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| shape | 14.80 | 0.55 | 13.74 | 15.92 | 1.00 | 14225 | 14438 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.905 (95% CI [0.894, 0.912]) | | | | | | | |
| Marginal R2: 0.005 (95% CI [0.003, 0.007]) | | | | | | | |

Table M2: Posterior summary of the regional model of species richness (Fig. M2).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ UI2 * Region + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 7188) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 323) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.27 | 0.05 | 1.17 | 1.38 | 1.00 | 3432 | 6393 |
| ~SS:SSB (Number of levels: 1632) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.21 | 0.01 | 0.19 | 0.23 | 1.00 | 5617 | 10817 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.49 | 0.12 | 2.25 | 2.73 | 1.00 | 1216 | 2423 |
| zi_Intercept | -5.32 | 0.22 | -5.79 | -4.92 | 1.00 | 40384 | 13614 |
| UI2Secondaryvegetation | 0.06 | 0.05 | -0.03 | 0.15 | 1.00 | 5995 | 11716 |
| UI2Agriculture_Minimal | 0.08 | 0.05 | -0.01 | 0.16 | 1.00 | 5938 | 10724 |
| UI2Agriculture_Light | -0.04 | 0.05 | -0.14 | 0.07 | 1.00 | 4923 | 9310 |
| UI2Agriculture_Intense | -0.03 | 0.06 | -0.14 | 0.08 | 1.00 | 5053 | 9350 |
| RegionNorthAmerica | 0.37 | 0.24 | -0.08 | 0.84 | 1.00 | 1701 | 3614 |
| RegionSouthandSoutheastAsia | 0.90 | 0.27 | 0.37 | 1.43 | 1.00 | 2123 | 3712 |
| RegionSouthandCentralAmerica | 0.39 | 0.19 | 0.02 | 0.75 | 1.00 | 1388 | 3233 |
| RegionSubsaharanAfrica | 0.21 | 0.31 | -0.40 | 0.82 | 1.00 | 2536 | 4568 |
| RegionAustralasia | 0.36 | 0.28 | -0.18 | 0.90 | 1.00 | 2166 | 3794 |
| UI2Secondaryvegetation:RegionNorthAmerica | -0.16 | 0.07 | -0.31 | -0.02 | 1.00 | 9631 | 13407 |
| UI2Agriculture_Minimal:RegionNorthAmerica | -0.25 | 0.12 | -0.48 | -0.02 | 1.00 | 16968 | 16148 |

```

UI2Agriculture_Light:RegionNorthAmerica      -0.40    0.09   -0.58   -0.22  1.00   10111  14428
UI2Agriculture_Intense:RegionNorthAmerica    -0.45    0.10   -0.65   -0.25  1.00   11462  14597
UI2Secondaryvegetation:RegionSouthandSoutheastAsia -0.23    0.15   -0.52    0.06  1.00   22859  15124
UI2Agriculture_Minimal:RegionSouthandSoutheastAsia -0.07    0.09   -0.25    0.12  1.00   15094  15125
UI2Agriculture_Light:RegionSouthandSoutheastAsia -0.63    0.08   -0.78   -0.48  1.00    7978  13286
UI2Agriculture_Intense:RegionSouthandSoutheastAsia -0.36    0.10   -0.55   -0.17  1.00   11207  13845
UI2Secondaryvegetation:RegionSouthandCentralAmerica -0.20    0.06   -0.32   -0.08  1.00    7910  13236
UI2Agriculture_Minimal:RegionSouthandCentralAmerica -0.32    0.07   -0.45   -0.19  1.00    9353  14920
UI2Agriculture_Light:RegionSouthandCentralAmerica -0.47    0.07   -0.61   -0.34  1.00    7024  12012
UI2Agriculture_Intense:RegionSouthandCentralAmerica -0.58    0.09   -0.76   -0.39  1.00    9220  13961
UI2Secondaryvegetation:RegionSubsaharanAfrica -0.18    0.09   -0.35   -0.01  1.00   12434  15682
UI2Agriculture_Minimal:RegionSubsaharanAfrica -0.36    0.09   -0.53   -0.19  1.00   12645  15148
UI2Agriculture_Light:RegionSubsaharanAfrica -0.33    0.09   -0.49   -0.16  1.00    9166  14359
UI2Agriculture_Intense:RegionSubsaharanAfrica -0.74    0.12   -0.97   -0.50  1.00   13992  14771
UI2Secondaryvegetation:RegionAustralasia -0.19    0.14   -0.47    0.07  1.00   17174  15374
UI2Agriculture_Minimal:RegionAustralasia -0.68    0.06   -0.80   -0.56  1.00    9210  13255
UI2Agriculture_Light:RegionAustralasia -0.19    0.10   -0.39    0.01  1.00   10897  14221
UI2Agriculture_Intense:RegionAustralasia -0.26    0.09   -0.43   -0.09  1.00    8904  13779

Further Distributional Parameters:
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
shape  17.37    0.74   15.96   18.86 1.00   16775   15686

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
and Tail_ESS are effective sample size measures, and Rhat is the potential
scale reduction factor on split chains (at convergence, Rhat = 1).

# Bayesian R2 with Compatibility Interval

Conditional R2: 0.918 (95% CI [0.908, 0.925])
Marginal R2: 0.025 (95% CI [0.006, 0.058])

```

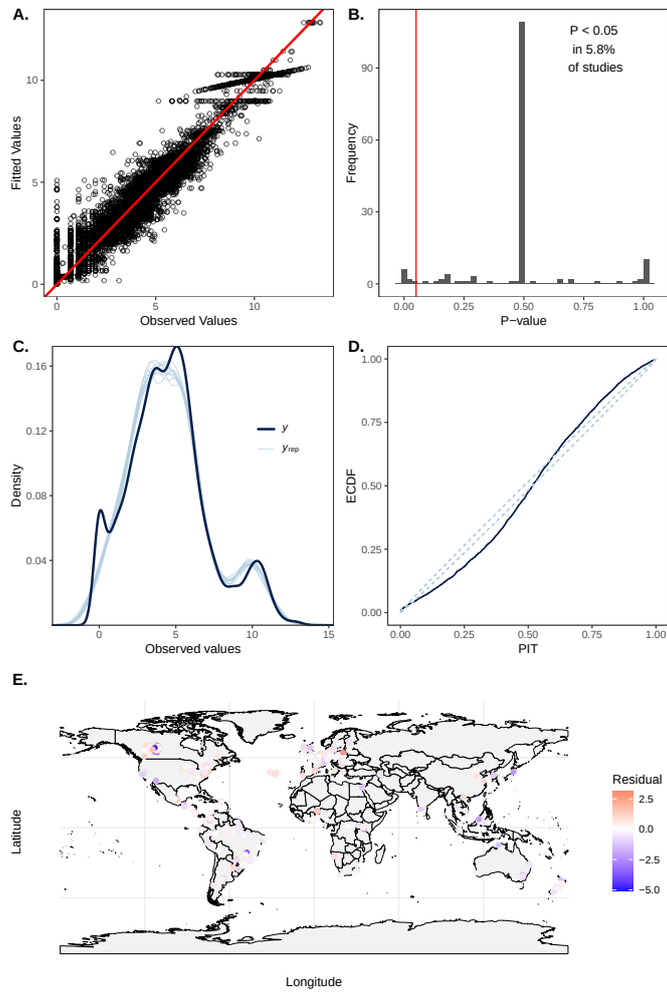


Figure M3: Model diagnostic plots for the Gaussian model of global total abundance responses to land use: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

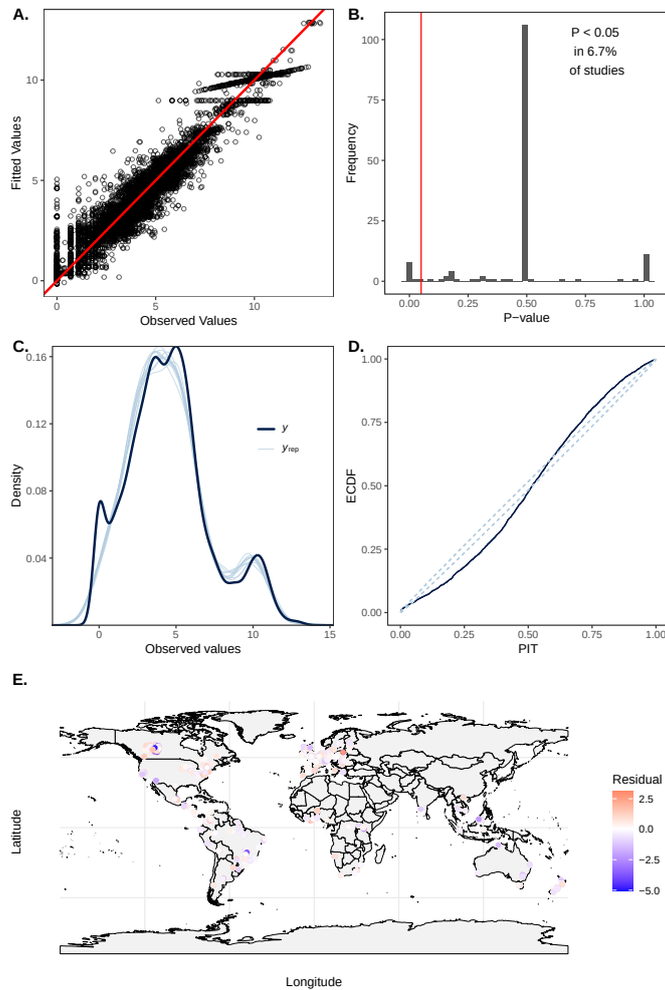


Figure M4: Model diagnostic plots for the Gaussian model of regional total abundance responses to land use: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} \times \text{Region} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M3: Posterior summary of the global model of total abundance (Fig. M3).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ UI2 + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 7122) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 314) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 2.11 | 0.09 | 1.94 | 2.29 | 1.00 | 2258 | 4282 |
| ~SS:SSB (Number of levels: 1624) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.42 | 0.02 | 0.37 | 0.47 | 1.00 | 5368 | 10618 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 5.26 | 0.12 | 5.02 | 5.51 | 1.01 | 1174 | 2780 |
| UI2Secondaryvegetation | -0.24 | 0.05 | -0.33 | -0.15 | 1.00 | 23249 | 17202 |
| UI2Agriculture_Minimal | -0.27 | 0.05 | -0.36 | -0.18 | 1.00 | 23482 | 17854 |
| UI2Agriculture_Light | -0.46 | 0.05 | -0.55 | -0.37 | 1.00 | 18742 | 17183 |
| UI2Agriculture_Intense | -0.45 | 0.05 | -0.55 | -0.34 | 1.00 | 20184 | 15549 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.82 | 0.01 | 0.81 | 0.84 | 1.00 | 18357 | 14190 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.905 (95% CI [0.903, 0.907]) | | | | | | | |
| Marginal R2: 0.005 (95% CI [0.003, 0.006]) | | | | | | | |

Table M4: Posterior summary of the regional model of total abundance (Fig. M4).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ UI2 * Region + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 6751) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 297) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 2.11 | 0.09 | 1.93 | 2.29 | 1.00 | 2086 | 4042 |
| ~SS:SSB (Number of levels: 1561) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.43 | 0.02 | 0.38 | 0.47 | 1.00 | 5203 | 9227 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 4.51 | 0.23 | 4.04 | 4.96 | 1.00 | 964 | 1878 |
| UI2Secondaryvegetation | 0.07 | 0.13 | -0.18 | 0.31 | 1.00 | 2325 | 4719 |
| UI2Agriculture_Minimal | 0.32 | 0.14 | 0.05 | 0.59 | 1.00 | 2335 | 5283 |
| UI2Agriculture_Light | -0.02 | 0.13 | -0.29 | 0.24 | 1.00 | 2058 | 4385 |
| UI2Agriculture_Intense | 0.00 | 0.14 | -0.28 | 0.27 | 1.00 | 2092 | 4468 |
| RegionNorthAmerica | 0.45 | 0.41 | -0.35 | 1.27 | 1.00 | 1158 | 2590 |
| RegionSouthandSoutheastAsia | 1.76 | 0.52 | 0.74 | 2.76 | 1.01 | 1341 | 3162 |
| RegionSouthandCentralAmerica | 0.97 | 0.34 | 0.32 | 1.65 | 1.00 | 1169 | 2742 |
| RegionSubsaharanAfrica | 0.66 | 0.54 | -0.41 | 1.70 | 1.00 | 1630 | 3831 |
| RegionAustralasia | 2.03 | 0.51 | 1.05 | 3.05 | 1.00 | 1467 | 3164 |
| UI2Secondaryvegetation:RegionNorthAmerica | -0.06 | 0.17 | -0.39 | 0.28 | 1.00 | 3383 | 7314 |
| UI2Agriculture_Minimal:RegionNorthAmerica | -0.51 | 0.24 | -0.97 | -0.05 | 1.00 | 4427 | 9051 |
| UI2Agriculture_Light:RegionNorthAmerica | -0.62 | 0.21 | -1.03 | -0.22 | 1.00 | 3433 | 7454 |
| UI2Agriculture_Intense:RegionNorthAmerica | -0.55 | 0.23 | -0.99 | -0.11 | 1.00 | 3902 | 8626 |
| UI2Secondaryvegetation:RegionSouthandSoutheastAsia | -0.57 | 0.45 | -1.47 | 0.31 | 1.00 | 13338 | 14827 |
| UI2Agriculture_Minimal:RegionSouthandSoutheastAsia | -0.58 | 0.28 | -1.14 | -0.02 | 1.00 | 6903 | 11526 |

| | | | | | | | |
|---|-----------------|-----------|----------|----------|---------|----------|----------|
| UI2Agriculture_Light:RegionSouthandSoutheastAsia | -0.92 | 0.17 | -1.26 | -0.58 | 1.00 | 3069 | 6629 |
| UI2Agriculture_Intense:RegionSouthandSoutheastAsia | -0.61 | 0.28 | -1.16 | -0.07 | 1.00 | 5940 | 10905 |
| UI2Secondaryvegetation:RegionSouthandCentralAmerica | -0.31 | 0.15 | -0.61 | -0.02 | 1.00 | 2922 | 6142 |
| UI2Agriculture_Minimal:RegionSouthandCentralAmerica | -0.54 | 0.16 | -0.86 | -0.22 | 1.00 | 2980 | 6595 |
| UI2Agriculture_Light:RegionSouthandCentralAmerica | -0.46 | 0.17 | -0.79 | -0.13 | 1.00 | 2805 | 6150 |
| UI2Agriculture_Intense:RegionSouthandCentralAmerica | -0.28 | 0.22 | -0.72 | 0.15 | 1.00 | 4031 | 8593 |
| UI2Secondaryvegetation:RegionSubsaharanAfrica | -0.53 | 0.21 | -0.94 | -0.11 | 1.00 | 4588 | 9345 |
| UI2Agriculture_Minimal:RegionSubsaharanAfrica | -1.32 | 0.20 | -1.71 | -0.93 | 1.00 | 3860 | 8551 |
| UI2Agriculture_Light:RegionSubsaharanAfrica | -0.84 | 0.21 | -1.25 | -0.43 | 1.00 | 3742 | 8848 |
| UI2Agriculture_Intense:RegionSubsaharanAfrica | -1.85 | 0.26 | -2.35 | -1.33 | 1.00 | 5248 | 10283 |
| UI2Secondaryvegetation:RegionAustralasia | -0.19 | 0.26 | -0.69 | 0.31 | 1.00 | 5753 | 9603 |
| UI2Agriculture_Minimal:RegionAustralasia | -1.05 | 0.18 | -1.40 | -0.71 | 1.00 | 3413 | 7710 |
| UI2Agriculture_Light:RegionAustralasia | -0.24 | 0.22 | -0.66 | 0.19 | 1.00 | 4013 | 8438 |
| UI2Agriculture_Intense:RegionAustralasia | -0.21 | 0.18 | -0.56 | 0.16 | 1.00 | 2959 | 6459 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.81 | 0.01 | 0.80 | 0.83 | 1.00 | 13326 | 12249 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| | Conditional R2: | 0.910 | (95% CI | [0.908, | 0.912]) | | |
| | Marginal R2: | 0.053 | (95% CI | [0.023, | 0.086]) | | |

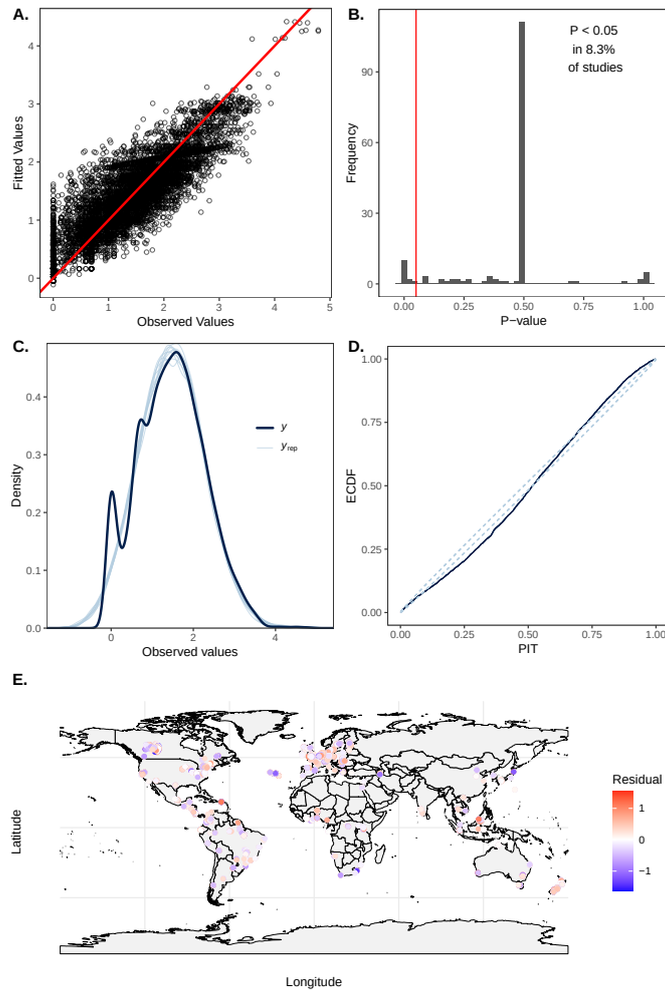


Figure M5: Model diagnostic plots for the Gaussian model of global Simpson's diversity responses to land use: $\text{Log}(\text{Simpson}) \sim \text{LandUse} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

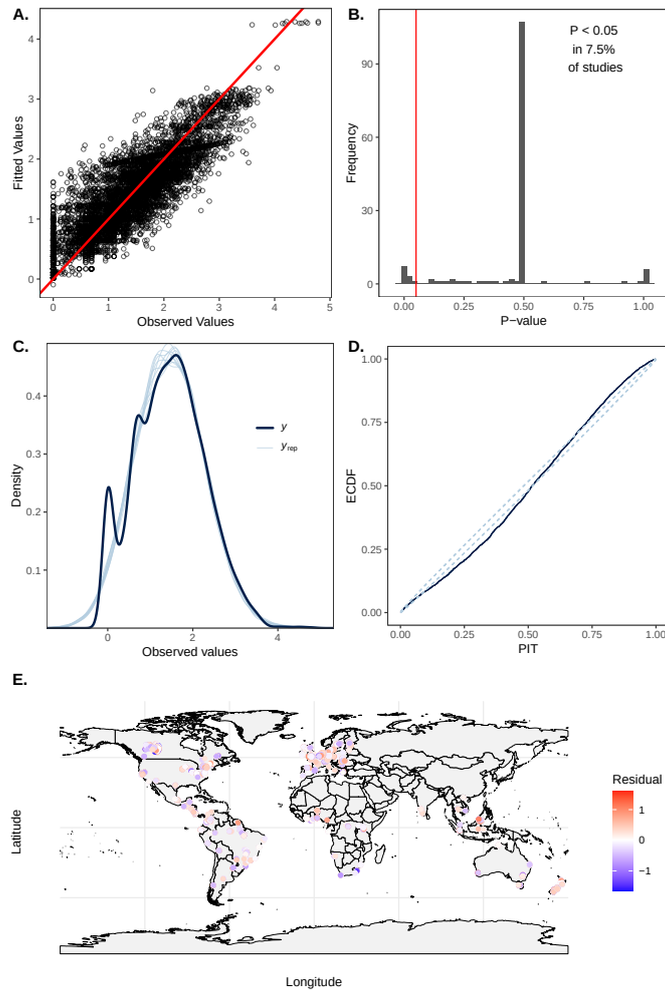


Figure M6: Model diagnostic plots for the Gaussian model of regional Simpson's diversity responses to land use: $\text{Log}(\text{Simpson}) \sim \text{LandUse} \times \text{Region} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M5: Posterior summary of the global model of Simpson's diversity index (Fig. M5).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: $\text{LogSimp} \sim \text{UI2} + (1 \mid \text{SS}/\text{SSB})$ | | | | | | | |
| Data: model_data (Number of observations: 6726) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 314) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.72 | 0.03 | 0.66 | 0.78 | 1.00 | 4243 | 7967 |
| ~SS:SSB (Number of levels: 1589) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.21 | 0.01 | 0.18 | 0.23 | 1.00 | 5581 | 10057 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.70 | 0.04 | 1.61 | 1.79 | 1.00 | 2570 | 5044 |
| UI2Secondaryvegetation | -0.14 | 0.03 | -0.19 | -0.09 | 1.00 | 19685 | 15914 |
| UI2Agriculture_Minimal | -0.28 | 0.03 | -0.33 | -0.23 | 1.00 | 19862 | 16476 |
| UI2Agriculture_Light | -0.37 | 0.03 | -0.42 | -0.32 | 1.00 | 16391 | 14549 |
| UI2Agriculture_Intense | -0.38 | 0.03 | -0.44 | -0.32 | 1.00 | 19441 | 16033 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.45 | 0.00 | 0.44 | 0.46 | 1.00 | 17170 | 14736 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.694 (95% CI [0.685, 0.702]) | | | | | | | |
| Marginal R2: 0.035 (95% CI [0.027, 0.043]) | | | | | | | |

Table M6: Posterior summary of the regional model of Simpson's diversity index (Fig. M6).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: $\text{LogSimp} \sim \text{UI2} * \text{Region} + (1 \mid \text{SS}/\text{SSB})$ | | | | | | | |
| Data: model_data (Number of observations: 6355) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 297) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.73 | 0.03 | 0.67 | 0.80 | 1.00 | 4843 | 8369 |
| ~SS:SSB (Number of levels: 1526) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.20 | 0.01 | 0.18 | 0.23 | 1.00 | 6174 | 11261 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.54 | 0.09 | 1.36 | 1.73 | 1.00 | 2414 | 4768 |
| UI2Secondaryvegetation | 0.00 | 0.07 | -0.13 | 0.13 | 1.00 | 3621 | 6816 |
| UI2Agriculture_Minimal | -0.03 | 0.07 | -0.17 | 0.11 | 1.00 | 3522 | 7354 |
| UI2Agriculture_Light | -0.09 | 0.07 | -0.23 | 0.04 | 1.00 | 3137 | 6415 |
| UI2Agriculture_Intense | -0.10 | 0.07 | -0.25 | 0.04 | 1.00 | 3273 | 6664 |
| RegionNorthAmerica | 0.20 | 0.16 | -0.11 | 0.50 | 1.00 | 2848 | 5792 |
| RegionSouthandSoutheastAsia | 0.44 | 0.19 | 0.07 | 0.81 | 1.00 | 3897 | 7016 |
| RegionSouthandCentralAmerica | 0.01 | 0.13 | -0.24 | 0.26 | 1.00 | 2785 | 5903 |
| RegionSubsaharanAfrica | 0.08 | 0.20 | -0.31 | 0.47 | 1.00 | 3546 | 7044 |
| RegionAustralasia | -0.05 | 0.19 | -0.43 | 0.33 | 1.00 | 3409 | 6850 |
| UI2Secondaryvegetation:RegionNorthAmerica | -0.12 | 0.09 | -0.30 | 0.06 | 1.00 | 5193 | 9339 |
| UI2Agriculture_Minimal:RegionNorthAmerica | -0.16 | 0.13 | -0.41 | 0.09 | 1.00 | 7145 | 12140 |
| UI2Agriculture_Light:RegionNorthAmerica | -0.49 | 0.11 | -0.71 | -0.28 | 1.00 | 5699 | 10021 |
| UI2Agriculture_Intense:RegionNorthAmerica | -0.63 | 0.12 | -0.87 | -0.39 | 1.00 | 6236 | 11806 |
| UI2Secondaryvegetation:RegionSouthandSoutheastAsia | -0.15 | 0.25 | -0.64 | 0.33 | 1.00 | 21751 | 15055 |
| UI2Agriculture_Minimal:RegionSouthandSoutheastAsia | -0.01 | 0.15 | -0.31 | 0.29 | 1.00 | 10156 | 13999 |

| | | | | | | | |
|---|---|-----------|----------|----------|------|----------|----------|
| UI2Agriculture_Light:RegionSouthandSoutheastAsia | -0.54 | 0.09 | -0.72 | -0.36 | 1.00 | 4695 | 8612 |
| UI2Agriculture_Intense:RegionSouthandSoutheastAsia | -0.18 | 0.15 | -0.47 | 0.11 | 1.00 | 9234 | 11576 |
| UI2Secondaryvegetation:RegionSouthandCentralAmerica | -0.01 | 0.08 | -0.17 | 0.15 | 1.00 | 4591 | 8688 |
| UI2Agriculture_Minimal:RegionSouthandCentralAmerica | -0.17 | 0.09 | -0.34 | 0.00 | 1.00 | 4591 | 9232 |
| UI2Agriculture_Light:RegionSouthandCentralAmerica | -0.25 | 0.09 | -0.42 | -0.07 | 1.00 | 4334 | 9006 |
| UI2Agriculture_Intense:RegionSouthandCentralAmerica | -0.45 | 0.12 | -0.68 | -0.22 | 1.00 | 6293 | 11740 |
| UI2Secondaryvegetation:RegionSubsaharanAfrica | -0.26 | 0.11 | -0.48 | -0.03 | 1.00 | 7884 | 12780 |
| UI2Agriculture_Minimal:RegionSubsaharanAfrica | -0.21 | 0.11 | -0.43 | -0.00 | 1.00 | 6023 | 11434 |
| UI2Agriculture_Light:RegionSubsaharanAfrica | -0.17 | 0.11 | -0.39 | 0.05 | 1.00 | 5808 | 11481 |
| UI2Agriculture_Intense:RegionSubsaharanAfrica | -0.36 | 0.14 | -0.64 | -0.09 | 1.00 | 8289 | 12953 |
| UI2Secondaryvegetation:RegionAustralasia | -0.07 | 0.14 | -0.34 | 0.20 | 1.00 | 9137 | 12772 |
| UI2Agriculture_Minimal:RegionAustralasia | -0.35 | 0.09 | -0.53 | -0.16 | 1.00 | 4956 | 10133 |
| UI2Agriculture_Light:RegionAustralasia | -0.09 | 0.12 | -0.33 | 0.15 | 1.00 | 6697 | 11970 |
| UI2Agriculture_Intense:RegionAustralasia | -0.07 | 0.10 | -0.27 | 0.12 | 1.00 | 4895 | 10045 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.44 | 0.00 | 0.43 | 0.45 | 1.00 | 19210 | 14800 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| | Conditional R2: 0.707 (95% CI [0.699, 0.716]) | | | | | | |
| | Marginal R2: 0.056 (95% CI [0.036, 0.084]) | | | | | | |

B Regional responses per land use type and intensity

The following model diagnostic plots refer to the comparison of global and Europe-specific responses to individual land use types and intensities, for the three taxonomic groups Arthropoda, Chordata and Plantae (Fig. 2).

B.1 Arthropoda

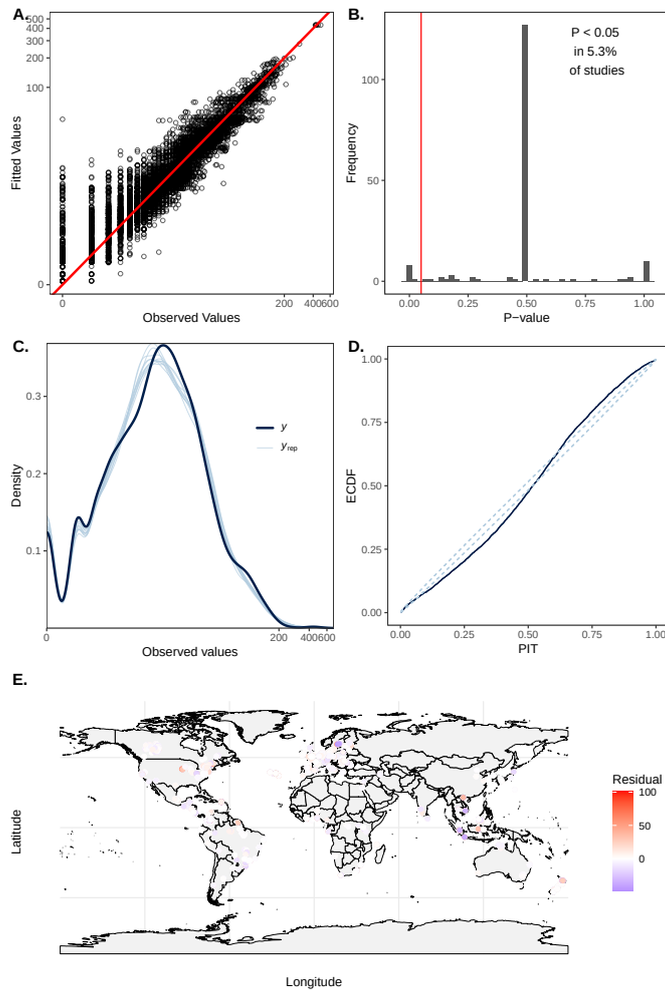


Figure M7: Model diagnostic plots for the zero-inflated negative binomial model of global species richness responses to land use: $\text{Richness} \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

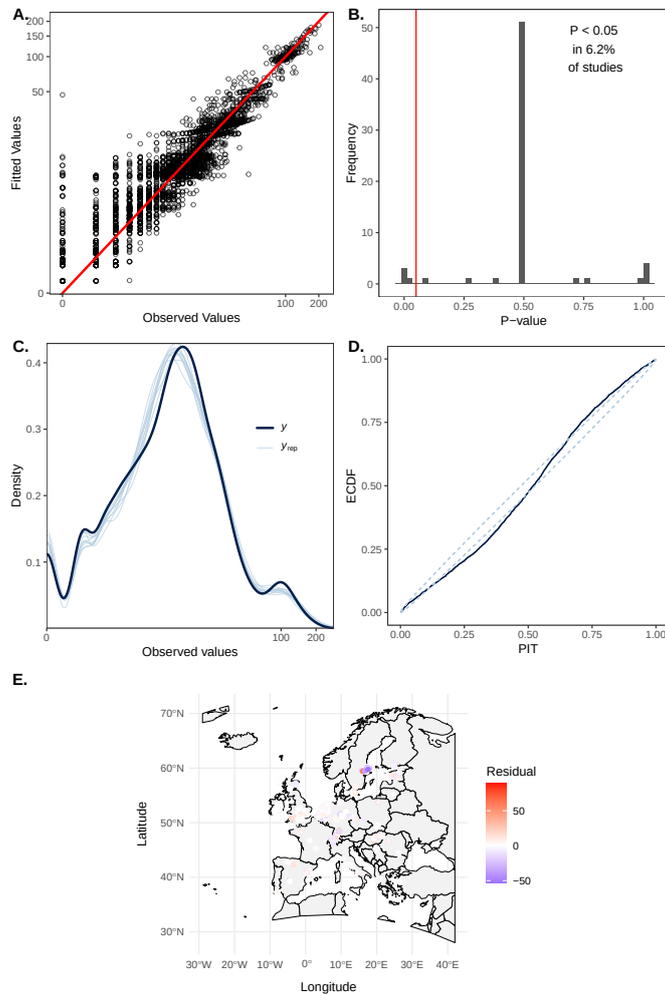


Figure M8: Model diagnostic plots for the zero-inflated negative binomial model of species richness responses to land use in Europe: $\text{Richness} \sim \text{LandUse} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M7: Posterior summary of the global model of species richness (Fig. M7).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ LU + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 7604) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 341) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.25 | 0.05 | 1.16 | 1.35 | 1.00 | 1678 | 3445 |
| ~SS:SSB (Number of levels: 1696) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.19 | 0.01 | 0.17 | 0.22 | 1.00 | 5479 | 10314 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.85 | 0.07 | 2.71 | 2.99 | 1.01 | 565 | 1502 |
| zi_Intercept | -5.34 | 0.22 | -5.79 | -4.94 | 1.00 | 32271 | 14439 |
| LUSecondaryvegetation | -0.13 | 0.02 | -0.17 | -0.08 | 1.00 | 16543 | 15550 |
| LUCropland_Minimal | -0.32 | 0.04 | -0.40 | -0.24 | 1.00 | 18845 | 16656 |
| LUCropland_Light | -0.50 | 0.04 | -0.57 | -0.43 | 1.00 | 12102 | 14779 |
| LUCropland_Intense | -0.54 | 0.04 | -0.61 | -0.47 | 1.00 | 12150 | 14509 |
| LUPasture_Minimal | -0.22 | 0.03 | -0.27 | -0.17 | 1.00 | 19322 | 16125 |
| LUPasture_Light | -0.25 | 0.03 | -0.30 | -0.19 | 1.00 | 13961 | 14473 |
| LUPasture_Intense | -0.39 | 0.03 | -0.46 | -0.32 | 1.00 | 13757 | 15129 |
| LUPlantationforest_Minimal | -0.10 | 0.04 | -0.18 | -0.02 | 1.00 | 24793 | 16319 |
| LUPlantationforest_Light | -0.37 | 0.03 | -0.44 | -0.30 | 1.00 | 21402 | 15936 |
| LUPlantationforest_Intense | -0.10 | 0.04 | -0.18 | -0.01 | 1.00 | 18243 | 16009 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| shape | 15.01 | 0.56 | 13.94 | 16.15 | 1.00 | 17382 | 13094 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.903 (95% CI [0.892, 0.911]) | | | | | | | |
| Marginal R2: 0.008 (95% CI [0.005, 0.011]) | | | | | | | |

Table M8: Posterior summary of the model of species richness for Europe (Fig. M8).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ LU + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 3412) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 118) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.37 | 0.09 | 1.20 | 1.57 | 1.00 | 3463 | 6871 |
| ~SS:SSB (Number of levels: 897) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.15 | 0.01 | 0.12 | 0.18 | 1.00 | 6293 | 11021 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.56 | 0.13 | 2.30 | 2.83 | 1.00 | 1405 | 2502 |
| zi_Intercept | -5.07 | 0.27 | -5.66 | -4.59 | 1.00 | 38999 | 13164 |
| LUSecondaryvegetation | 0.04 | 0.05 | -0.05 | 0.13 | 1.00 | 14463 | 16164 |
| LUCropland_Minimal | -0.18 | 0.06 | -0.30 | -0.06 | 1.00 | 16937 | 16715 |
| LUCropland_Light | -0.23 | 0.06 | -0.36 | -0.10 | 1.00 | 11119 | 14853 |
| LUCropland_Intense | -0.23 | 0.06 | -0.35 | -0.11 | 1.00 | 11474 | 14306 |
| LUPasture_Minimal | 0.14 | 0.05 | 0.05 | 0.24 | 1.00 | 14487 | 16566 |

| | | | | | | | |
|----------------------------|-------|------|-------|------|------|-------|-------|
| LUPasture_Light | 0.02 | 0.05 | -0.09 | 0.12 | 1.00 | 11248 | 14082 |
| LUPasture_Intense | -0.06 | 0.06 | -0.18 | 0.05 | 1.00 | 10769 | 13479 |
| LUPlantationforest_Minimal | 0.04 | 0.09 | -0.13 | 0.21 | 1.00 | 23539 | 16593 |
| LUPlantationforest_Light | 0.01 | 0.10 | -0.19 | 0.20 | 1.00 | 31103 | 16369 |
| LUPlantationforest_Intense | 0.72 | 0.08 | 0.57 | 0.88 | 1.00 | 13519 | 15080 |

Further Distributional Parameters:

| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
|-------|----------|-----------|----------|----------|------|----------|----------|
| shape | 19.38 | 1.39 | 16.83 | 22.30 | 1.00 | 14984 | 14998 |

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1).

Bayesian R2 with Compatibility Interval

Conditional R2: 0.917 (95% CI [0.908, 0.924])

Marginal R2: 0.007 (95% CI [0.003, 0.012])

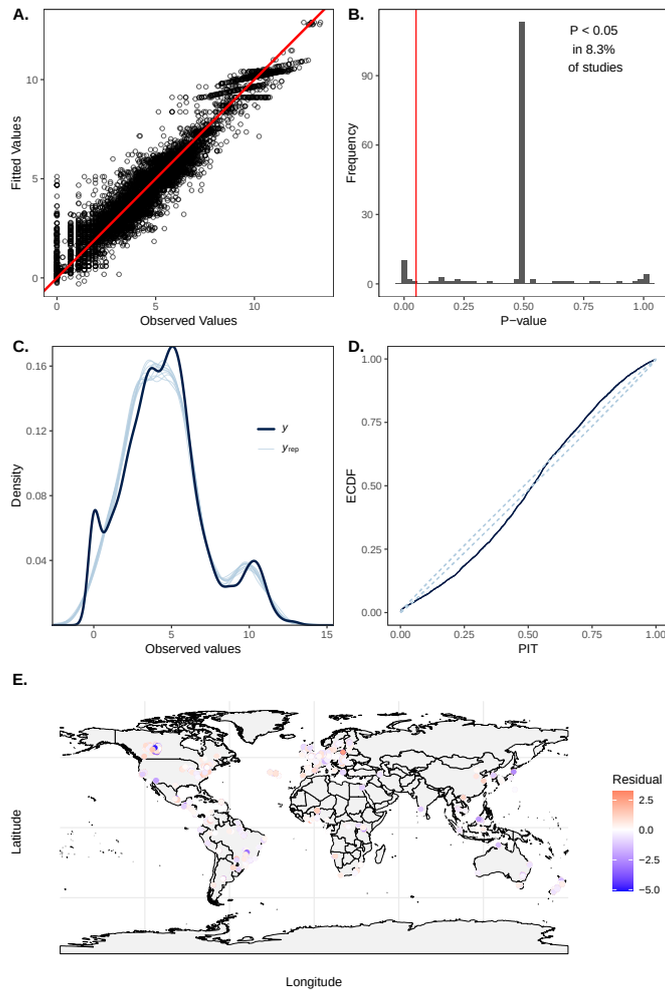


Figure M9: Model diagnostic plots for the Gaussian model of global total abundance responses to land use: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

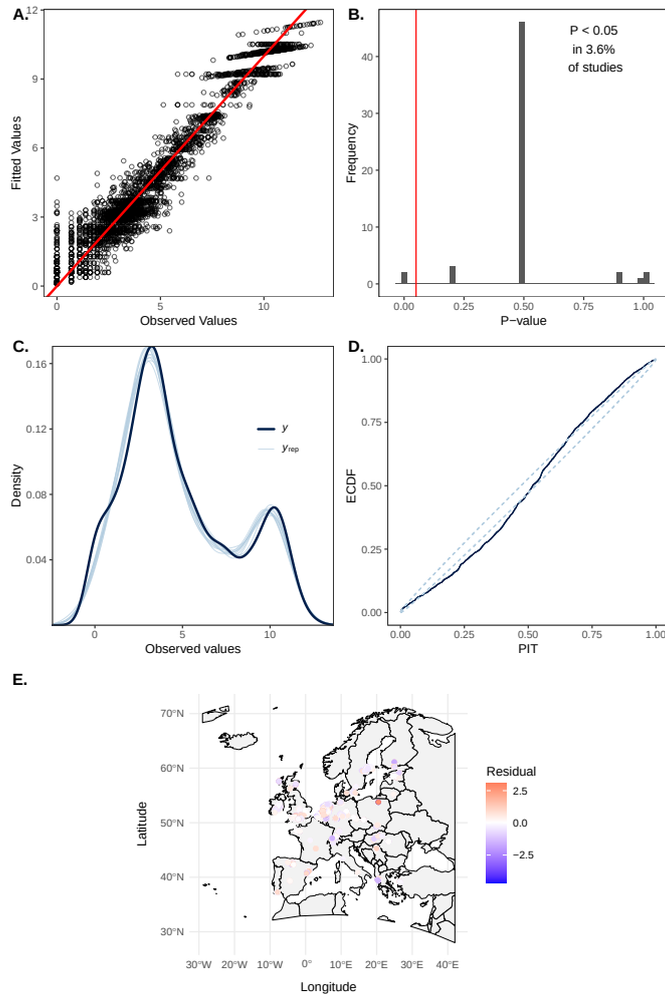


Figure M10: Model diagnostic plots for the Gaussian model of total abundance responses to land use in Europe: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M9: Posterior summary of the global model of total abundance (Fig. M9).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ LU + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 7122) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 314) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 2.09 | 0.09 | 1.92 | 2.26 | 1.00 | 1809 | 4331 |
| ~SS:SSB (Number of levels: 1624) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.41 | 0.02 | 0.37 | 0.45 | 1.00 | 5925 | 11912 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 5.30 | 0.12 | 5.06 | 5.54 | 1.01 | 897 | 2327 |
| LUSecondaryvegetation | -0.22 | 0.05 | -0.31 | -0.13 | 1.00 | 13832 | 14589 |
| LUCropland_Minimal | -0.80 | 0.07 | -0.95 | -0.66 | 1.00 | 17671 | 16103 |
| LUCropland_Light | -0.91 | 0.07 | -1.05 | -0.76 | 1.00 | 11117 | 14650 |
| LUCropland_Intense | -0.93 | 0.07 | -1.08 | -0.78 | 1.00 | 12122 | 13991 |
| LUPasture_Minimal | -0.07 | 0.06 | -0.18 | 0.05 | 1.00 | 17254 | 15105 |
| LUPasture_Light | -0.20 | 0.06 | -0.32 | -0.08 | 1.00 | 13161 | 14900 |
| LUPasture_Intense | -0.30 | 0.07 | -0.44 | -0.16 | 1.00 | 13665 | 16240 |
| LUPlantationforest_Minimal | -0.00 | 0.09 | -0.18 | 0.17 | 1.00 | 22838 | 15678 |
| LUPlantationforest_Light | -0.55 | 0.07 | -0.68 | -0.41 | 1.00 | 21637 | 16108 |
| LUPlantationforest_Intense | 0.06 | 0.10 | -0.13 | 0.25 | 1.00 | 23689 | 15508 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.81 | 0.01 | 0.80 | 0.83 | 1.00 | 19874 | 14735 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.907 (95% CI [0.905, 0.909]) | | | | | | | |
| Marginal R2: 0.018 (95% CI [0.014, 0.022]) | | | | | | | |

Table M10: Posterior summary of the model of total abundance for Europe (Fig. M10).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ LU + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 3211) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 111) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 2.13 | 0.15 | 1.85 | 2.46 | 1.00 | 1759 | 3174 |
| ~SS:SSB (Number of levels: 876) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.31 | 0.03 | 0.25 | 0.38 | 1.00 | 4148 | 8416 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 4.67 | 0.24 | 4.20 | 5.14 | 1.00 | 1088 | 2121 |
| LUSecondaryvegetation | 0.06 | 0.13 | -0.18 | 0.30 | 1.00 | 3822 | 8141 |
| LUCropland_Minimal | -0.61 | 0.16 | -0.93 | -0.29 | 1.00 | 4391 | 9601 |
| LUCropland_Light | -0.68 | 0.16 | -0.99 | -0.35 | 1.00 | 3066 | 6797 |
| LUCropland_Intense | -0.62 | 0.15 | -0.92 | -0.32 | 1.00 | 3286 | 7139 |
| LUPasture_Minimal | 0.72 | 0.15 | 0.42 | 1.02 | 1.00 | 3597 | 8666 |
| LUPasture_Light | 0.24 | 0.14 | -0.04 | 0.52 | 1.00 | 3130 | 7309 |
| LUPasture_Intense | 0.18 | 0.15 | -0.11 | 0.48 | 1.00 | 3091 | 6657 |
| LUPlantationforest_Minimal | 0.34 | 0.19 | -0.03 | 0.70 | 1.00 | 6741 | 11355 |
| LUPlantationforest_Light | 0.29 | 0.20 | -0.09 | 0.68 | 1.00 | 11575 | 15096 |

```
LUPlantationforest_Intense 1.26 0.22 0.83 1.69 1.00 5209 10657
```

```
Further Distributional Parameters:
```

```
Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS  
sigma 0.79 0.01 0.77 0.82 1.00 12548 13361
```

```
Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS  
and Tail_ESS are effective sample size measures, and Rhat is the potential  
scale reduction factor on split chains (at convergence, Rhat = 1).
```

```
# Bayesian R2 with Compatibility Interval
```

```
Conditional R2: 0.938 (95% CI [0.937, 0.940])  
Marginal R2: 0.025 (95% CI [0.019, 0.032])
```

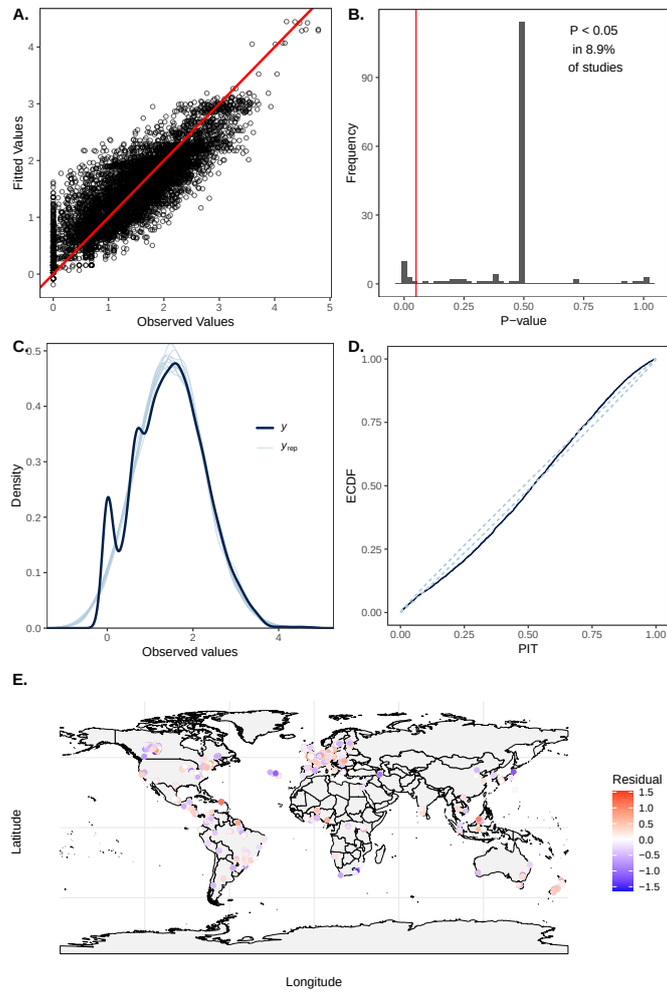


Figure M11: Model diagnostic plots for the Gaussian model of global Simpson's diversity responses to land use: $\text{Log}(\text{Simpson}) \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

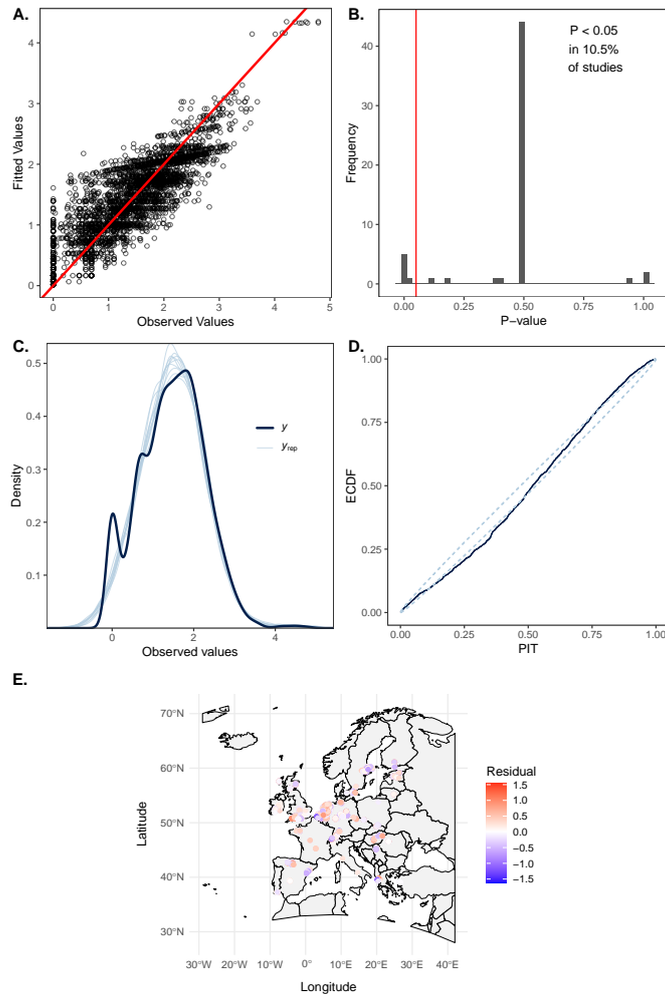


Figure M12: Model diagnostic plots for the Gaussian model of Simpson's diversity responses to land use in Europe: $\text{Log}(\text{Simpson}) \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M11: Posterior summary of the global model of Simpson's diversity index (Fig. M11).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogSimp ~ LU + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 6726) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 314) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.72 | 0.03 | 0.66 | 0.79 | 1.00 | 2370 | 4781 |
| ~SS:SSB (Number of levels: 1589) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.20 | 0.01 | 0.18 | 0.22 | 1.00 | 4678 | 9207 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.70 | 0.05 | 1.61 | 1.79 | 1.01 | 1301 | 2503 |
| LUSecondaryvegetation | -0.14 | 0.03 | -0.19 | -0.09 | 1.00 | 11008 | 13901 |
| LUCropland_Minimal | -0.37 | 0.04 | -0.46 | -0.29 | 1.00 | 11418 | 14441 |
| LUCropland_Light | -0.45 | 0.04 | -0.53 | -0.37 | 1.00 | 8191 | 11489 |
| LUCropland_Intense | -0.45 | 0.04 | -0.54 | -0.37 | 1.00 | 8241 | 12190 |
| LUPasture_Minimal | -0.23 | 0.03 | -0.30 | -0.17 | 1.00 | 12631 | 14704 |
| LUPasture_Light | -0.30 | 0.03 | -0.37 | -0.24 | 1.00 | 9823 | 13644 |
| LUPasture_Intense | -0.40 | 0.04 | -0.48 | -0.32 | 1.00 | 10096 | 14376 |
| LUPlantationforest_Minimal | -0.24 | 0.05 | -0.34 | -0.14 | 1.00 | 16751 | 14853 |
| LUPlantationforest_Light | -0.43 | 0.04 | -0.50 | -0.35 | 1.00 | 14275 | 14196 |
| LUPlantationforest_Intense | -0.20 | 0.05 | -0.30 | -0.09 | 1.00 | 15825 | 16041 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.45 | 0.00 | 0.44 | 0.46 | 1.00 | 14353 | 14046 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.694 (95% CI [0.686, 0.702]) | | | | | | | |
| Marginal R2: 0.044 (95% CI [0.034, 0.055]) | | | | | | | |

Table M12: Posterior summary of the model of Simpson's diversity index for Europe (Fig. M12).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogSimp ~ LU + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 3032) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 111) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.79 | 0.06 | 0.68 | 0.91 | 1.00 | 6265 | 9955 |
| ~SS:SSB (Number of levels: 853) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.19 | 0.02 | 0.15 | 0.23 | 1.00 | 5619 | 10698 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.55 | 0.10 | 1.35 | 1.75 | 1.00 | 4361 | 8155 |
| LUSecondaryvegetation | 0.00 | 0.07 | -0.13 | 0.14 | 1.00 | 11446 | 15568 |
| LUCropland_Minimal | -0.18 | 0.09 | -0.36 | -0.01 | 1.00 | 12156 | 15524 |
| LUCropland_Light | -0.18 | 0.09 | -0.36 | -0.01 | 1.00 | 9570 | 13592 |
| LUCropland_Intense | -0.13 | 0.09 | -0.30 | 0.04 | 1.00 | 9566 | 12947 |
| LUPasture_Minimal | 0.12 | 0.08 | -0.05 | 0.28 | 1.00 | 10640 | 14536 |
| LUPasture_Light | -0.01 | 0.08 | -0.17 | 0.14 | 1.00 | 9529 | 13244 |
| LUPasture_Intense | -0.10 | 0.08 | -0.26 | 0.06 | 1.00 | 9469 | 13533 |
| LUPlantationforest_Minimal | -0.17 | 0.11 | -0.39 | 0.04 | 1.00 | 18237 | 16782 |

| | | | | | | | |
|----------------------------|-------|------|-------|------|------|-------|-------|
| LUPlantationforest_Light | -0.08 | 0.11 | -0.29 | 0.15 | 1.00 | 27318 | 16927 |
| LUPlantationforest_Intense | 0.40 | 0.12 | 0.16 | 0.64 | 1.00 | 13332 | 15760 |

Further Distributional Parameters:

| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
|-------|----------|-----------|----------|----------|------|----------|----------|
| sigma | 0.44 | 0.01 | 0.43 | 0.45 | 1.00 | 18178 | 14228 |

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1).

Bayesian R2 with Compatibility Interval

Conditional R2: 0.690 (95% CI [0.677, 0.703])
 Marginal R2: 0.020 (95% CI [0.010, 0.032])

B.2 Chordata

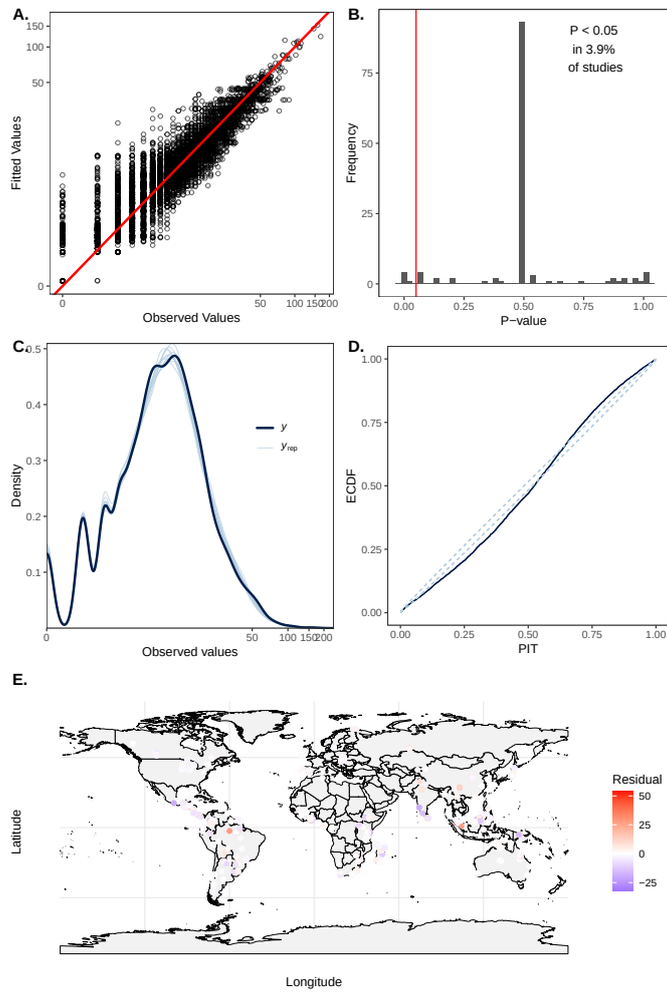


Figure M13: Model diagnostic plots for the zero-inflated negative binomial model of global species richness responses to land use: $\text{Richness} \sim \text{LandUse} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

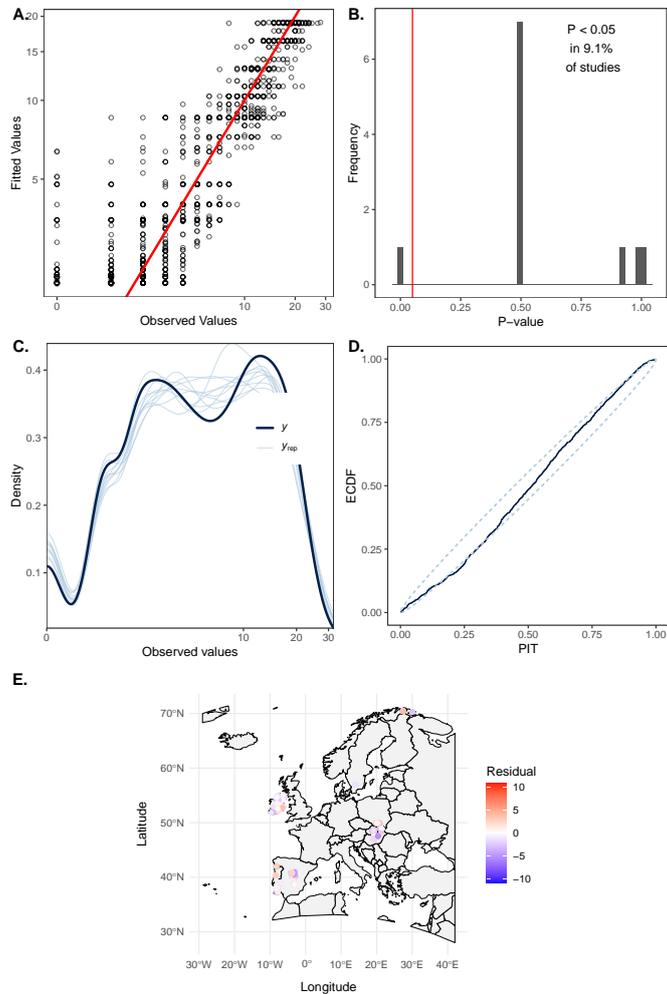


Figure M14: Model diagnostic plots for the zero-inflated negative binomial model of species richness responses to land use in Europe: $\text{Richness} \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M13: Posterior summary of the global model of species richness (Fig. M13).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ LU + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 7842) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 224) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.07 | 0.06 | 0.97 | 1.19 | 1.00 | 2378 | 5275 |
| ~SS:SSB (Number of levels: 855) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.22 | 0.01 | 0.19 | 0.24 | 1.00 | 6180 | 11195 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.44 | 0.07 | 2.30 | 2.59 | 1.01 | 1006 | 2371 |
| zi_Intercept | -8.88 | 1.25 | -11.88 | -7.05 | 1.00 | 23800 | 11167 |
| LUSecondaryvegetation | -0.09 | 0.02 | -0.12 | -0.05 | 1.00 | 24133 | 16956 |
| LUCropland_Minimal | -0.17 | 0.03 | -0.23 | -0.11 | 1.00 | 33215 | 16169 |
| LUCropland_Light | -0.20 | 0.03 | -0.27 | -0.13 | 1.00 | 36759 | 15186 |
| LUCropland_Intense | -0.23 | 0.10 | -0.42 | -0.04 | 1.00 | 37110 | 15235 |
| LUPasture_Minimal | -0.24 | 0.04 | -0.32 | -0.17 | 1.00 | 30795 | 16130 |
| LUPasture_Light | -0.31 | 0.03 | -0.37 | -0.26 | 1.00 | 30864 | 16740 |
| LUPasture_Intense | -0.19 | 0.05 | -0.29 | -0.09 | 1.00 | 22248 | 16565 |
| LUPlantationforest_Minimal | -0.07 | 0.03 | -0.12 | -0.02 | 1.00 | 23920 | 16106 |
| LUPlantationforest_Light | -0.22 | 0.02 | -0.26 | -0.18 | 1.00 | 24204 | 16937 |
| LUPlantationforest_Intense | -0.46 | 0.04 | -0.53 | -0.39 | 1.00 | 30199 | 15965 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| shape | 31.42 | 1.87 | 27.94 | 35.32 | 1.00 | 31326 | 14006 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.855 (95% CI [0.848, 0.862]) | | | | | | | |
| Marginal R2: 0.011 (95% CI [0.007, 0.015]) | | | | | | | |

Table M14: Posterior summary of the model of species richness for Europe (Fig. M14).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ LU + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 1157) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 15) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.69 | 0.17 | 0.43 | 1.08 | 1.00 | 8141 | 11199 |
| ~SS:SSB (Number of levels: 113) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.29 | 0.06 | 0.18 | 0.42 | 1.00 | 5098 | 10136 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.33 | 0.20 | 1.94 | 2.72 | 1.00 | 6423 | 8930 |
| zi_Intercept | -5.26 | 0.97 | -7.81 | -4.05 | 1.00 | 9163 | 7633 |
| LUSecondaryvegetation | -0.15 | 0.04 | -0.23 | -0.07 | 1.00 | 18047 | 15462 |
| LUCropland_Minimal | 0.19 | 0.09 | 0.01 | 0.36 | 1.00 | 23136 | 15474 |
| LUCropland_Intense | -0.22 | 0.12 | -0.46 | 0.02 | 1.00 | 28209 | 15648 |
| LUPasture_Minimal | -0.55 | 0.13 | -0.80 | -0.29 | 1.00 | 21141 | 15364 |
| LUPasture_Light | -0.32 | 0.08 | -0.47 | -0.17 | 1.00 | 19260 | 16543 |

| | | | | | | | |
|----------------------------|-------|------|-------|-------|------|-------|-------|
| LUPasture_Intense | -0.01 | 0.31 | -0.63 | 0.56 | 1.00 | 32256 | 13428 |
| LUPlantationforest_Minimal | 0.00 | 0.06 | -0.11 | 0.11 | 1.00 | 19519 | 16181 |
| LUPlantationforest_Light | -0.23 | 0.05 | -0.33 | -0.13 | 1.00 | 22194 | 15704 |
| LUPlantationforest_Intense | -0.61 | 0.07 | -0.74 | -0.48 | 1.00 | 24867 | 16728 |

Further Distributional Parameters:

| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
|-------|---------------|-----------------|----------|-------------|------|----------|----------|
| shape | 6322954526.55 | 290719243411.97 | 487.10 | 13141967.76 | 1.00 | 14777 | 8381 |

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1).

Bayesian R2 with Compatibility Interval

Conditional R2: 0.850 (95% CI [0.837, 0.861])
 Marginal R2: 0.083 (95% CI [0.029, 0.158])

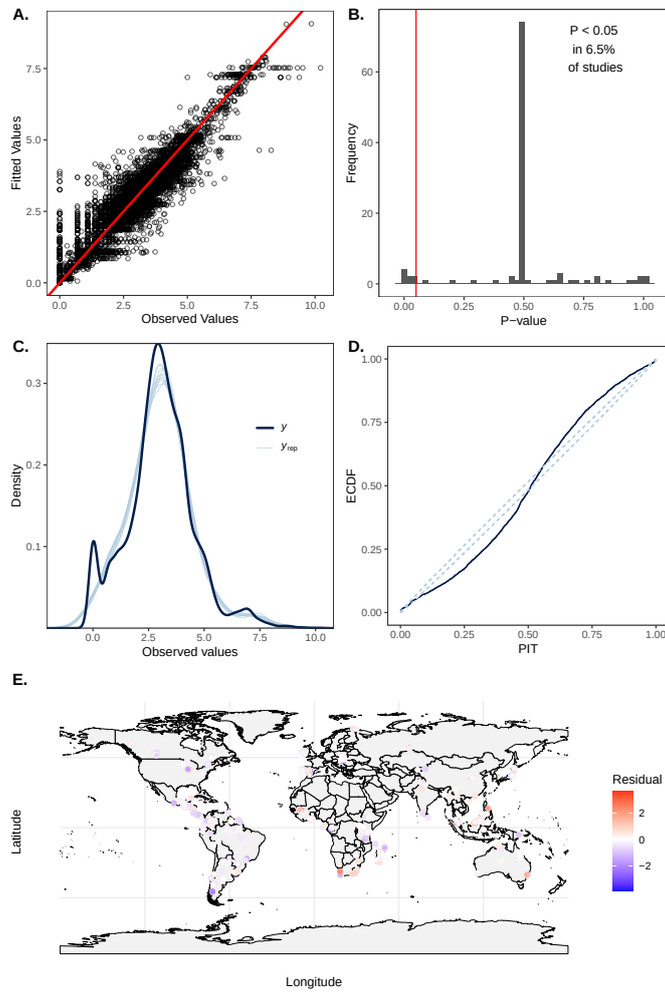


Figure M15: Model diagnostic plots for the Gaussian model of global total abundance responses to land use: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

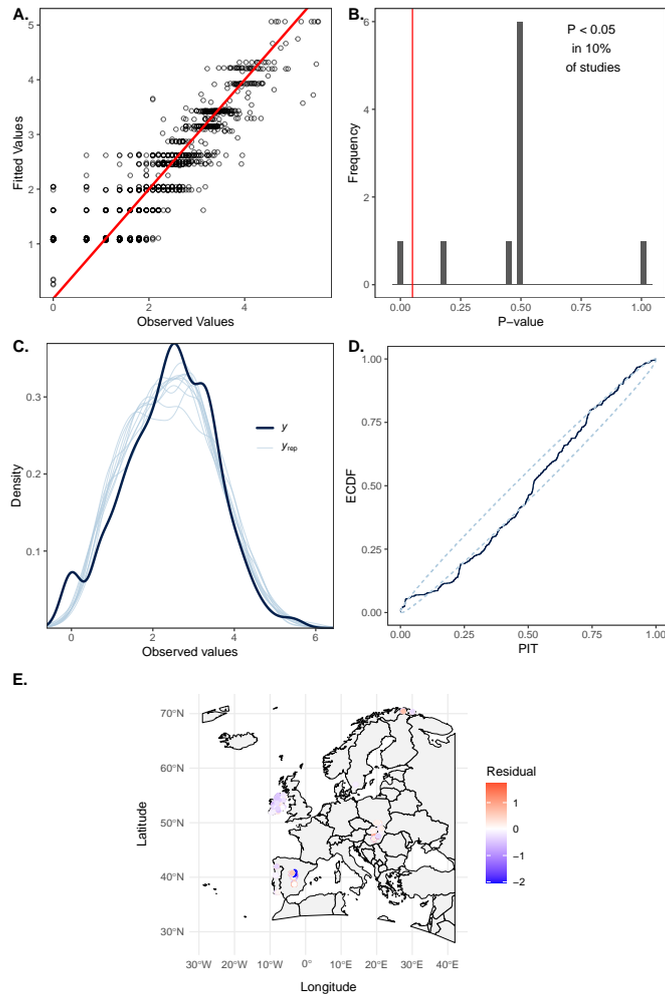


Figure M16: Model diagnostic plots for the Gaussian model of total abundance responses to land use in Europe: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M15: Posterior summary of the global model of total abundance (Fig. M15).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ LU + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 7007) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 199) | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.73 | 0.09 | 1.56 | 1.92 | 1.00 | 2471 | 5066 |
| ~SS:SSB (Number of levels: 751) | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.33 | 0.02 | 0.29 | 0.37 | 1.00 | 6843 | 12431 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 3.83 | 0.12 | 3.59 | 4.07 | 1.01 | 983 | 2029 |
| LUSsecondaryvegetation | -0.04 | 0.03 | -0.10 | 0.02 | 1.00 | 20326 | 16038 |
| LUCropland_Minimal | 0.01 | 0.05 | -0.08 | 0.11 | 1.00 | 32767 | 15044 |
| LUCropland_Light | -0.20 | 0.06 | -0.32 | -0.07 | 1.00 | 36154 | 15222 |
| LUCropland_Intense | -0.04 | 0.09 | -0.23 | 0.15 | 1.00 | 31540 | 15829 |
| LUPasture_Minimal | -0.12 | 0.06 | -0.25 | 0.00 | 1.00 | 30492 | 16358 |
| LUPasture_Light | -0.52 | 0.05 | -0.61 | -0.43 | 1.00 | 29056 | 15774 |
| LUPasture_Intense | -0.13 | 0.10 | -0.32 | 0.06 | 1.00 | 25215 | 15414 |
| LUPlantationforest_Minimal | -0.02 | 0.04 | -0.10 | 0.07 | 1.00 | 23613 | 17202 |
| LUPlantationforest_Light | -0.24 | 0.04 | -0.31 | -0.17 | 1.00 | 22643 | 16276 |
| LUPlantationforest_Intense | -0.26 | 0.05 | -0.36 | -0.15 | 1.00 | 28902 | 15481 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.60 | 0.01 | 0.59 | 0.61 | 1.00 | 30488 | 14143 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.838 (95% CI [0.834, 0.841]) | | | | | | | |
| Marginal R2: 0.008 (95% CI [0.006, 0.010]) | | | | | | | |

Figure M17: Posterior summary of the model of total abundance for Europe (Fig. M16).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ LU + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 966) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 14) | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.41 | 0.31 | 0.02 | 1.14 | 1.03 | 160 | 71 |
| ~SS:SSB (Number of levels: 35) | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.13 | 0.16 | 0.86 | 1.47 | 1.01 | 2123 | 7896 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 3.36 | 0.28 | 2.80 | 3.90 | 1.01 | 3048 | 5019 |
| LUSsecondaryvegetation | -0.27 | 0.08 | -0.43 | -0.11 | 1.02 | 255 | 758 |
| LUCropland_Minimal | 0.16 | 0.11 | -0.06 | 0.37 | 1.01 | 1120 | 13231 |
| LUCropland_Intense | -0.30 | 0.12 | -0.53 | -0.08 | 1.01 | 386 | 7667 |
| LUPasture_Minimal | -0.34 | 0.18 | -0.71 | 0.01 | 1.01 | 1155 | 7831 |
| LUPasture_Light | -0.43 | 0.11 | -0.65 | -0.20 | 1.00 | 5675 | 8496 |
| LUPasture_Intense | 0.04 | 0.55 | -1.03 | 1.13 | 1.00 | 16908 | 13144 |
| LUPlantationforest_Minimal | 0.10 | 0.10 | -0.09 | 0.29 | 1.02 | 219 | 179 |
| LUPlantationforest_Light | -0.28 | 0.11 | -0.48 | -0.03 | 1.03 | 125 | 25 |
| LUPlantationforest_Intense | -0.92 | 0.13 | -1.16 | -0.68 | 1.03 | 151 | 74 |

```
Further Distributional Parameters:
      Estimate Est.Error l-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sigma    0.51      0.01    0.49    0.54 1.02     267    185

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
and Tail_ESS are effective sample size measures, and Rhat is the potential
scale reduction factor on split chains (at convergence, Rhat = 1).

# Bayesian R2 with Compatibility Interval

Conditional R2: 0.790 (95% CI [0.777, 0.800])
Marginal R2: 0.051 (95% CI [0.036, 0.070])
```

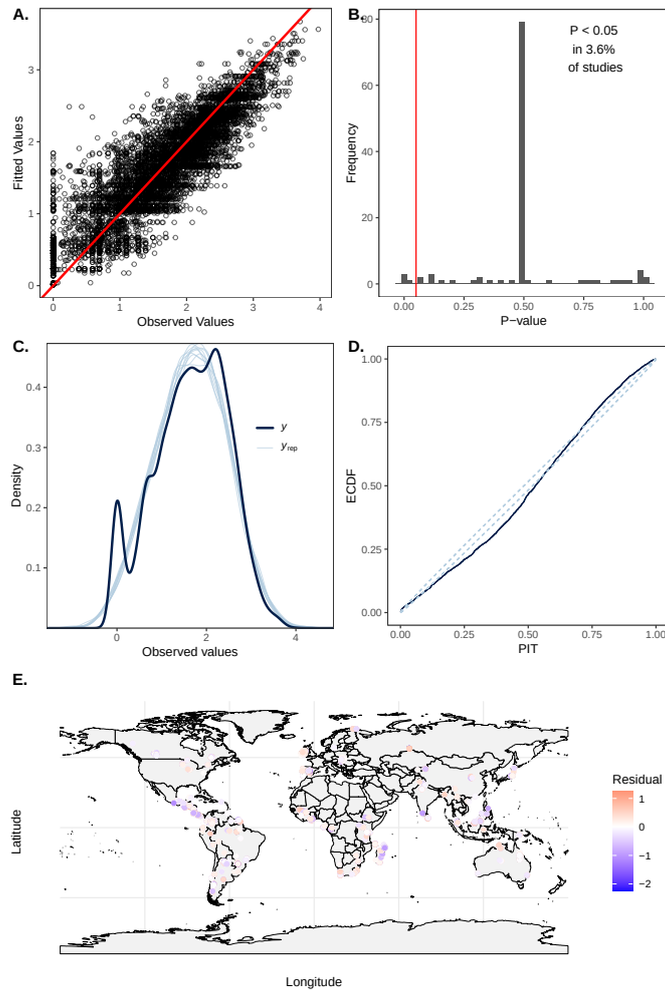


Figure M18: Model diagnostic plots for the Gaussian model of global Simpson's diversity responses to land use: $\text{Log}(\text{Simpson}) \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

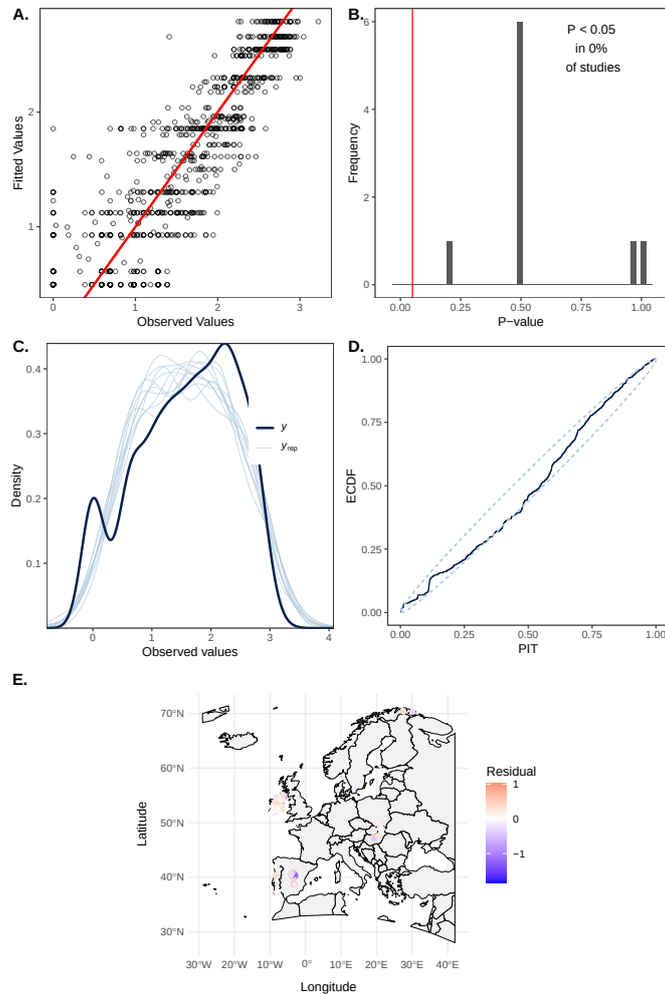


Figure M19: Model diagnostic plots for the Gaussian model of Simpson's diversity responses to land use in Europe: $\text{Log}(\text{Simpson}) \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Figure M20: Posterior summary of the global model of Simpson's diversity index (Fig. M18).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogSimp ~ LU + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 6708) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 199) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.82 | 0.05 | 0.73 | 0.91 | 1.00 | 5216 | 9227 |
| ~SS:SSB (Number of levels: 742) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.22 | 0.01 | 0.20 | 0.25 | 1.00 | 7211 | 11805 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.80 | 0.06 | 1.68 | 1.92 | 1.00 | 2555 | 5477 |
| LUSecondaryvegetation | -0.11 | 0.02 | -0.15 | -0.07 | 1.00 | 28142 | 18557 |
| LUCropland_Minimal | -0.18 | 0.03 | -0.24 | -0.11 | 1.00 | 39875 | 17040 |
| LUCropland_Light | -0.30 | 0.04 | -0.39 | -0.22 | 1.00 | 45355 | 15905 |
| LUCropland_Intense | -0.25 | 0.07 | -0.38 | -0.12 | 1.00 | 41140 | 16051 |
| LUPasture_Minimal | -0.31 | 0.05 | -0.40 | -0.23 | 1.00 | 38275 | 16438 |
| LUPasture_Light | -0.27 | 0.03 | -0.34 | -0.21 | 1.00 | 35881 | 15628 |
| LUPasture_Intense | -0.33 | 0.06 | -0.46 | -0.20 | 1.00 | 33476 | 16888 |
| LUPlantationforest_Minimal | -0.11 | 0.03 | -0.17 | -0.05 | 1.00 | 31383 | 16441 |
| LUPlantationforest_Light | -0.18 | 0.03 | -0.23 | -0.13 | 1.00 | 30372 | 17262 |
| LUPlantationforest_Intense | -0.36 | 0.04 | -0.43 | -0.28 | 1.00 | 36510 | 16790 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.41 | 0.00 | 0.40 | 0.42 | 1.00 | 30952 | 14920 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.749 (95% CI [0.743, 0.755]) | | | | | | | |
| Marginal R2: 0.019 (95% CI [0.014, 0.024]) | | | | | | | |

Figure M21: Posterior summary of the model of Simpson's diversity index for Europe (Fig. M19).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogSimp ~ LU + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 923) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 14) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.51 | 0.19 | 0.12 | 0.90 | 1.00 | 5069 | 3798 |
| ~SS:SSB (Number of levels: 33) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.30 | 0.11 | 0.08 | 0.53 | 1.00 | 3666 | 5649 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.04 | 0.17 | 1.70 | 2.38 | 1.00 | 11948 | 13019 |
| LUSecondaryvegetation | -0.24 | 0.06 | -0.37 | -0.12 | 1.00 | 13029 | 14983 |
| LUCropland_Minimal | 0.13 | 0.09 | -0.05 | 0.31 | 1.00 | 16062 | 16549 |
| LUCropland_Intense | -0.36 | 0.09 | -0.54 | -0.17 | 1.00 | 17482 | 15810 |
| LUPasture_Minimal | -0.80 | 0.15 | -1.09 | -0.50 | 1.00 | 20455 | 15872 |
| LUPasture_Light | -0.49 | 0.09 | -0.67 | -0.30 | 1.00 | 17064 | 16462 |
| LUPasture_Intense | -0.37 | 0.45 | -1.26 | 0.51 | 1.00 | 31683 | 14779 |
| LUPlantationforest_Minimal | -0.05 | 0.08 | -0.20 | 0.10 | 1.00 | 14322 | 16492 |
| LUPlantationforest_Light | -0.32 | 0.07 | -0.47 | -0.18 | 1.00 | 20009 | 16533 |

```

LUPlantationforest_Intense  -0.46    0.09   -0.64   -0.28  1.00   25689   14591

Further Distributional Parameters:
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sigma    0.40    0.01    0.39    0.42  1.00   29767   14504

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
and Tail_ESS are effective sample size measures, and Rhat is the potential
scale reduction factor on split chains (at convergence, Rhat = 1).

# Bayesian R2 with Compatibility Interval

Conditional R2: 0.769 (95% CI [0.756, 0.782])
Marginal R2: 0.057 (95% CI [0.036, 0.082])

```

C Compositional dissimilarity

Here we provide an example model diagnostics plot from one of the 100 models fitted on different permutations of compositional dissimilarity between natural vegetation and other land uses (Fig. 4). These diagnostics are from the model of total dissimilarity, but diagnostics from the turnover and nestedness components are similar.

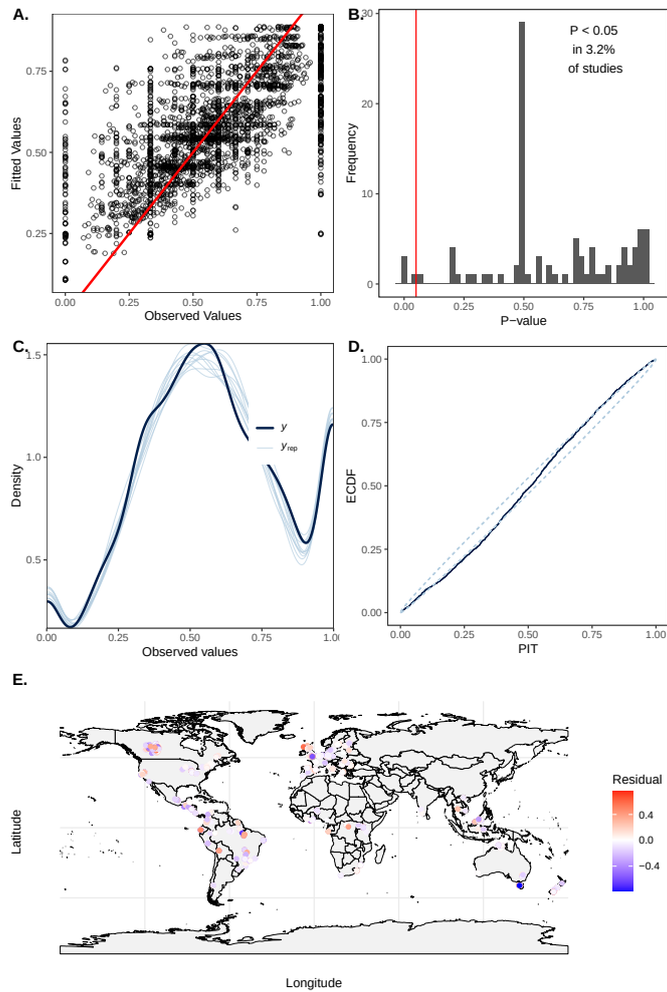


Figure M22: Model diagnostic plots for the zero-and-one inflated beta model of total compositional dissimilarity (Sørensen's Index): $\text{Dissimilarity} \sim \text{LandUse} * \text{Region} + (1|SS)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

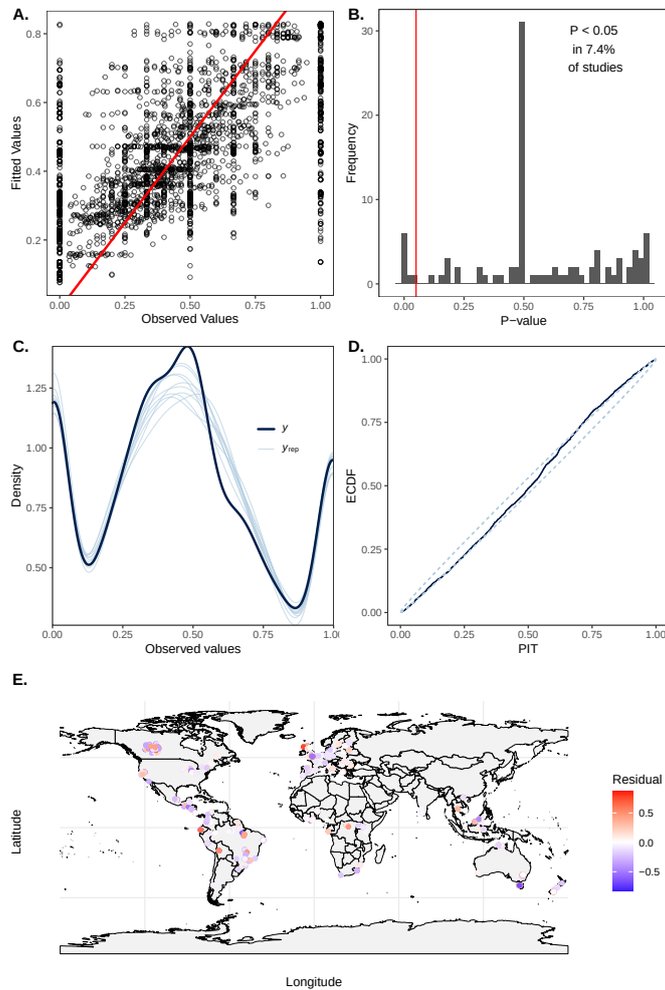


Figure M23: Model diagnostic plots for the zero-and-one inflated beta model of species turnover (Sørensen's Index): $\text{Turnover} \sim \text{LandUse} * \text{Region} + (1|SS)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

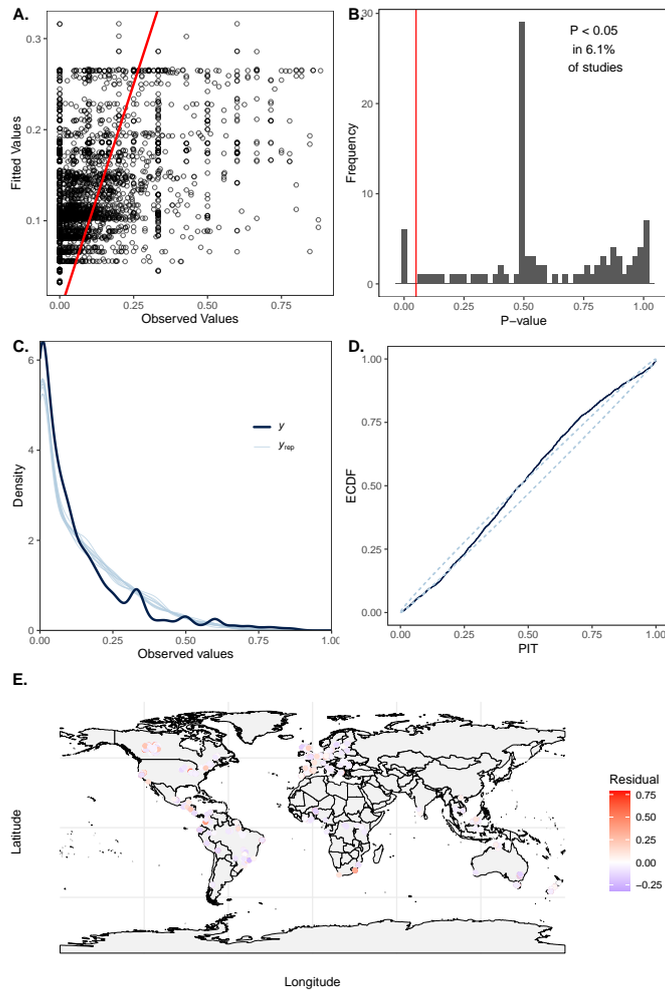


Figure M24: Model diagnostic plots for the zero-and-one inflated beta model of species nestedness (Sørensen's Index): $Nestedness \sim LandUse * Region + (1|SS)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M16: Posterior summary of the model of total compositional dissimilarity (Fig. M22).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_one_inflated_beta | | | | | | | |
| Links: mu = logit; phi = log; zoi = logit; coi = logit | | | | | | | |
| Formula: Dissimilarity ~ LU * Region + Dist_geo + (1 SS) | | | | | | | |
| phi ~ 1 | | | | | | | |
| zoi ~ (1 SS) | | | | | | | |
| coi ~ (1 SS) | | | | | | | |
| Data: model_data (Number of observations: 2858) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 187) | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.55 | 0.04 | 0.48 | 0.63 | 1.00 | 4093 | 8313 |
| sd(zoi_Intercept) | 2.80 | 0.34 | 2.22 | 3.54 | 1.00 | 2997 | 5966 |
| sd(coi_Intercept) | 3.55 | 0.78 | 2.31 | 5.37 | 1.00 | 5005 | 8751 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | -0.27 | 0.09 | -0.44 | -0.10 | 1.00 | 1995 | 4476 |
| phi_Intercept | 2.43 | 0.03 | 2.38 | 2.49 | 1.00 | 21620 | 13743 |
| zoi_Intercept | -3.53 | 0.36 | -4.30 | -2.88 | 1.00 | 2651 | 5135 |
| coi_Intercept | 1.92 | 0.58 | 0.86 | 3.14 | 1.00 | 3270 | 6305 |
| LUagriculture_Minimal | 0.75 | 0.09 | 0.58 | 0.93 | 1.00 | 5485 | 9451 |
| LUagriculture_Light | 0.28 | 0.14 | 0.01 | 0.55 | 1.00 | 3876 | 7036 |
| LUagriculture_Intense | 0.55 | 0.19 | 0.18 | 0.92 | 1.00 | 4280 | 9516 |
| RegionNorthAmerica | 0.40 | 0.14 | 0.11 | 0.68 | 1.00 | 2085 | 4501 |
| RegionSouthandSoutheastAsia | -0.29 | 0.19 | -0.67 | 0.09 | 1.00 | 3473 | 6496 |
| RegionSouthandCentralAmerica | -0.17 | 0.13 | -0.42 | 0.09 | 1.00 | 2518 | 4834 |
| RegionSubsaharanAfrica | -0.07 | 0.19 | -0.44 | 0.30 | 1.00 | 3234 | 5406 |
| RegionAustralasia | 0.27 | 0.19 | -0.10 | 0.64 | 1.00 | 2043 | 4714 |
| Dist_geo | 0.12 | 0.02 | 0.08 | 0.16 | 1.00 | 12869 | 15537 |
| LUagriculture_Minimal:RegionNorthAmerica | -0.73 | 0.15 | -1.03 | -0.43 | 1.00 | 9854 | 13640 |
| LUagriculture_Light:RegionNorthAmerica | -0.15 | 0.20 | -0.54 | 0.24 | 1.00 | 5883 | 10883 |
| LUagriculture_Intense:RegionNorthAmerica | -0.19 | 0.29 | -0.75 | 0.38 | 1.00 | 6873 | 12517 |
| LUagriculture_Minimal:RegionSouthandSoutheastAsia | -0.40 | 0.32 | -1.04 | 0.23 | 1.00 | 9962 | 13721 |
| LUagriculture_Light:RegionSouthandSoutheastAsia | 0.40 | 0.16 | 0.08 | 0.72 | 1.00 | 4684 | 8863 |
| LUagriculture_Intense:RegionSouthandSoutheastAsia | 0.70 | 0.36 | -0.01 | 1.42 | 1.00 | 9138 | 12652 |
| LUagriculture_Minimal:RegionSouthandCentralAmerica | -0.53 | 0.12 | -0.78 | -0.29 | 1.00 | 7520 | 11689 |
| LUagriculture_Light:RegionSouthandCentralAmerica | 0.49 | 0.18 | 0.14 | 0.84 | 1.00 | 4797 | 9274 |
| LUagriculture_Intense:RegionSouthandCentralAmerica | 0.13 | 0.25 | -0.35 | 0.63 | 1.00 | 5399 | 11412 |
| LUagriculture_Minimal:RegionSubsaharanAfrica | -0.64 | 0.16 | -0.94 | -0.32 | 1.00 | 8290 | 12643 |
| LUagriculture_Light:RegionSubsaharanAfrica | -0.44 | 0.22 | -0.86 | -0.02 | 1.00 | 5837 | 11520 |
| LUagriculture_Intense:RegionSubsaharanAfrica | -0.30 | 0.27 | -0.83 | 0.22 | 1.00 | 6129 | 11011 |
| LUagriculture_Minimal:RegionAustralasia | -0.45 | 0.16 | -0.75 | -0.13 | 1.00 | 10221 | 13590 |
| LUagriculture_Light:RegionAustralasia | 0.14 | 0.29 | -0.43 | 0.70 | 1.00 | 7867 | 12357 |
| LUagriculture_Intense:RegionAustralasia | -0.41 | 0.24 | -0.88 | 0.06 | 1.00 | 5722 | 10523 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.434 (95% CI [0.412, 0.457]) | | | | | | | |
| Marginal R2: 0.102 (95% CI [0.071, 0.136]) | | | | | | | |

Table M17: Posterior summary of the model of species turnover (Fig. M23).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_one_inflated_beta | | | | | | | |
| Links: mu = logit; phi = log; zoi = logit; coi = logit | | | | | | | |
| Formula: Dissimilarity ~ LU * Region + Dist_geo + (1 SS) | | | | | | | |
| phi ~ 1 | | | | | | | |
| zoi ~ (1 SS) | | | | | | | |
| coi ~ (1 SS) | | | | | | | |
| Data: model_data (Number of observations: 2858) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 187) | | | | | | | |
| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.54 | 0.04 | 0.46 | 0.62 | 1.00 | 5435 | 9801 |

| | | | | | | | |
|-------------------|------|------|------|------|------|------|------|
| sd(zoi_Intercept) | 3.20 | 0.33 | 2.61 | 3.91 | 1.00 | 4454 | 8184 |
| sd(coi_Intercept) | 2.51 | 0.40 | 1.85 | 3.38 | 1.00 | 4760 | 8903 |

Regression Coefficients:

| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
|--|----------|-----------|----------|----------|------|----------|----------|
| Intercept | -0.62 | 0.09 | -0.80 | -0.44 | 1.00 | 3074 | 5555 |
| phi_Intercept | 2.37 | 0.03 | 2.30 | 2.43 | 1.00 | 24875 | 14479 |
| zoi_Intercept | -2.38 | 0.32 | -3.07 | -1.80 | 1.00 | 2460 | 5087 |
| coi_Intercept | -0.92 | 0.34 | -1.63 | -0.29 | 1.00 | 3072 | 6288 |
| LUAgriculture_Minimal | 0.93 | 0.09 | 0.75 | 1.12 | 1.00 | 8512 | 12063 |
| LUAgriculture_Light | 0.42 | 0.16 | 0.11 | 0.73 | 1.00 | 5473 | 9255 |
| LUAgriculture_Intense | 0.57 | 0.25 | 0.08 | 1.06 | 1.00 | 5535 | 8982 |
| RegionNorthAmerica | 0.61 | 0.15 | 0.31 | 0.90 | 1.00 | 3038 | 5513 |
| RegionSouthandSoutheastAsia | -0.27 | 0.20 | -0.67 | 0.11 | 1.00 | 5332 | 8952 |
| RegionSouthandCentralAmerica | -0.16 | 0.13 | -0.42 | 0.11 | 1.00 | 4132 | 8688 |
| RegionSubsaharanAfrica | 0.01 | 0.20 | -0.37 | 0.39 | 1.00 | 4863 | 8355 |
| RegionAustralasia | 0.33 | 0.19 | -0.04 | 0.70 | 1.00 | 3903 | 6910 |
| Dist_geo | 0.10 | 0.02 | 0.06 | 0.14 | 1.00 | 17213 | 16189 |
| LUAgriculture_Minimal:RegionNorthAmerica | -0.76 | 0.19 | -1.12 | -0.39 | 1.00 | 12820 | 14461 |
| LUAgriculture_Light:RegionNorthAmerica | -0.38 | 0.23 | -0.83 | 0.07 | 1.00 | 7942 | 12577 |
| LUAgriculture_Intense:RegionNorthAmerica | -0.53 | 0.35 | -1.21 | 0.16 | 1.00 | 7998 | 11420 |
| LUAgriculture_Minimal:RegionSouthandSoutheastAsia | -0.75 | 0.34 | -1.42 | -0.09 | 1.00 | 13998 | 14496 |
| LUAgriculture_Light:RegionSouthandSoutheastAsia | -0.04 | 0.18 | -0.40 | 0.31 | 1.00 | 6188 | 11190 |
| LUAgriculture_Intense:RegionSouthandSoutheastAsia | 0.66 | 0.40 | -0.12 | 1.46 | 1.00 | 8792 | 13027 |
| LUAgriculture_Minimal:RegionSouthandCentralAmerica | -0.82 | 0.14 | -1.10 | -0.54 | 1.00 | 11913 | 14634 |
| LUAgriculture_Light:RegionSouthandCentralAmerica | 0.29 | 0.20 | -0.10 | 0.68 | 1.00 | 6654 | 11079 |
| LUAgriculture_Intense:RegionSouthandCentralAmerica | 0.18 | 0.30 | -0.41 | 0.76 | 1.00 | 6632 | 11044 |
| LUAgriculture_Minimal:RegionSubsaharanAfrica | -0.98 | 0.19 | -1.36 | -0.61 | 1.00 | 13361 | 15589 |
| LUAgriculture_Light:RegionSubsaharanAfrica | -0.58 | 0.28 | -1.13 | -0.05 | 1.00 | 9439 | 13802 |
| LUAgriculture_Intense:RegionSubsaharanAfrica | -0.57 | 0.33 | -1.23 | 0.08 | 1.00 | 7501 | 11545 |
| LUAgriculture_Minimal:RegionAustralasia | -0.74 | 0.16 | -1.05 | -0.42 | 1.00 | 14440 | 14477 |
| LUAgriculture_Light:RegionAustralasia | -0.03 | 0.31 | -0.64 | 0.59 | 1.00 | 11471 | 13452 |
| LUAgriculture_Intense:RegionAustralasia | -0.61 | 0.32 | -1.23 | 0.01 | 1.00 | 6772 | 10432 |

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1).

Bayesian R2 with Compatibility Interval

Conditional R2: 0.368 (95% CI [0.344, 0.391])
Marginal R2: 0.074 (95% CI [0.049, 0.104])

Table M18: Posterior summary of the model of species nestedness (Fig. M24).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_one_inflated_beta | | | | | | | |
| Links: mu = logit; phi = log; zoi = logit; coi = logit | | | | | | | |
| Formula: Dissimilarity ~ LU * Region + Dist_geo + (1 SS) | | | | | | | |
| phi ~ 1 | | | | | | | |
| zoi ~ (1 SS) | | | | | | | |
| coi ~ (1 SS) | | | | | | | |
| Data: model_data (Number of observations: 2858) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 187) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.50 | 0.04 | 0.42 | 0.58 | 1.00 | 6790 | 10891 |
| sd(zoi_Intercept) | 1.75 | 0.18 | 1.43 | 2.12 | 1.00 | 5031 | 9780 |
| sd(coi_Intercept) | 0.82 | 0.65 | 0.03 | 2.43 | 1.00 | 12742 | 10078 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | -1.68 | 0.09 | -1.86 | -1.49 | 1.00 | 4749 | 8736 |
| phi_Intercept | 2.01 | 0.03 | 1.95 | 2.08 | 1.00 | 27789 | 13724 |
| zoi_Intercept | -1.96 | 0.19 | -2.34 | -1.61 | 1.00 | 4504 | 8444 |
| coi_Intercept | -7.52 | 1.46 | -11.03 | -5.37 | 1.00 | 25369 | 10802 |
| LUAgriculture_Minimal | -0.09 | 0.12 | -0.33 | 0.16 | 1.00 | 8612 | 12507 |
| LUAgriculture_Light | -0.11 | 0.18 | -0.48 | 0.25 | 1.00 | 6803 | 11065 |
| LUAgriculture_Intense | 0.07 | 0.25 | -0.43 | 0.55 | 1.00 | 7242 | 11493 |
| RegionNorthAmerica | -0.09 | 0.15 | -0.39 | 0.20 | 1.00 | 4888 | 9063 |
| RegionSouthandSoutheastAsia | -0.23 | 0.21 | -0.65 | 0.18 | 1.00 | 8372 | 11862 |
| RegionSouthandCentralAmerica | 0.01 | 0.14 | -0.27 | 0.28 | 1.00 | 6240 | 10677 |
| RegionSubsaharanAfrica | 0.01 | 0.19 | -0.38 | 0.38 | 1.00 | 7448 | 12146 |
| RegionAustralasia | -0.08 | 0.19 | -0.45 | 0.30 | 1.00 | 5723 | 10142 |
| Dist_geo | -0.01 | 0.03 | -0.07 | 0.04 | 1.00 | 18472 | 16433 |
| LUAgriculture_Minimal:RegionNorthAmerica | -0.14 | 0.22 | -0.58 | 0.29 | 1.00 | 14692 | 14910 |
| LUAgriculture_Light:RegionNorthAmerica | 0.18 | 0.28 | -0.37 | 0.71 | 1.00 | 9861 | 14863 |
| LUAgriculture_Intense:RegionNorthAmerica | 0.87 | 0.39 | 0.09 | 1.63 | 1.00 | 10321 | 15146 |

```

LUAgriculture_Minimal:RegionSouthandSoutheastAsia -0.06 0.46 -0.97 0.80 1.00 17060 15998
LUAgriculture_Light:RegionSouthandSoutheastAsia 0.87 0.22 0.44 1.30 1.00 8162 13206
LUAgriculture_Intense:RegionSouthandSoutheastAsia -0.32 0.59 -1.57 0.76 1.00 16648 14728
LUAgriculture_Minimal:RegionSouthandCentralAmerica 0.00 0.18 -0.34 0.35 1.00 12097 15015
LUAgriculture_Light:RegionSouthandCentralAmerica 0.41 0.23 -0.05 0.87 1.00 8131 11671
LUAgriculture_Intense:RegionSouthandCentralAmerica -0.25 0.33 -0.90 0.41 1.00 9525 14377
LUAgriculture_Minimal:RegionSubsaharanAfrica 0.50 0.22 0.07 0.93 1.00 12767 15703
LUAgriculture_Light:RegionSubsaharanAfrica 0.46 0.30 -0.12 1.05 1.00 10156 13478
LUAgriculture_Intense:RegionSubsaharanAfrica 0.16 0.37 -0.58 0.89 1.00 10221 14098
LUAgriculture_Minimal:RegionAustralasia 0.29 0.22 -0.14 0.71 1.00 15343 14689
LUAgriculture_Light:RegionAustralasia -0.46 0.42 -1.32 0.35 1.00 15268 14034
LUAgriculture_Intense:RegionAustralasia 0.13 0.32 -0.49 0.78 1.00 9619 13227

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
and Tail_ESS are effective sample size measures, and Rhat is the potential
scale reduction factor on split chains (at convergence, Rhat = 1).

# Bayesian R2 with Compatibility Interval

Conditional R2: 0.155 (95% CI [0.134, 0.177])
Marginal R2: 0.036 (95% CI [0.019, 0.061])

```

D Effects of range extent, habitat specificity and non-native species

D.1 Range extent

The following model diagnostic plots refer to the comparison of biodiversity responses in primary vegetation, secondary vegetation and agricultural land use between species with different range extents, comparing Europe and North America (Fig. 5).

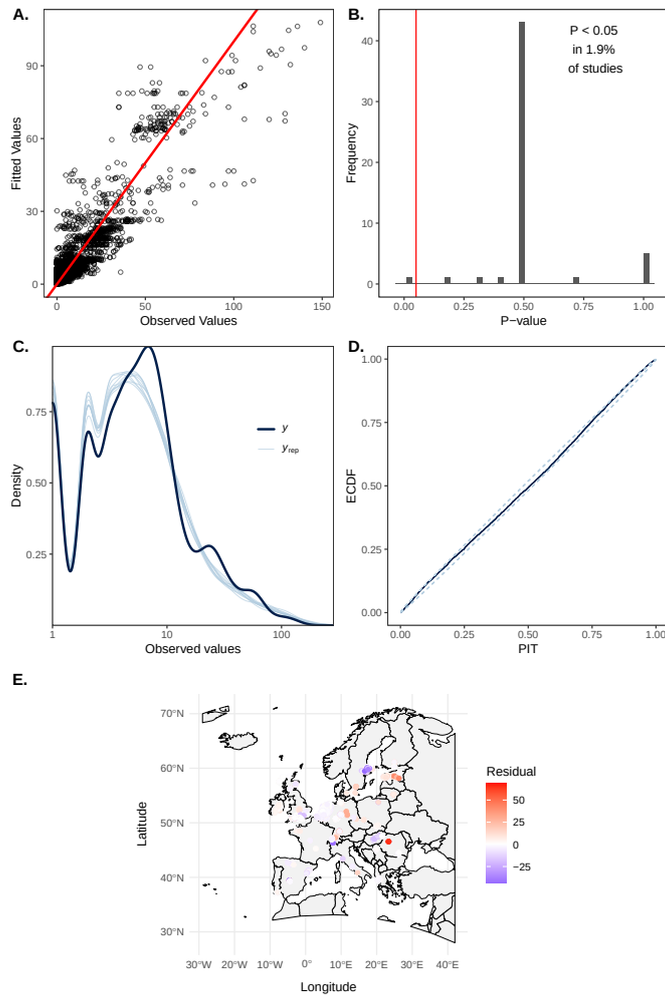


Figure M25: Model diagnostic plots for the zero-inflated negative binomial model of species richness responses to land use and range extent in Europe: $\text{Richness} \sim \text{LandUse} \times \text{RangeExtent} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

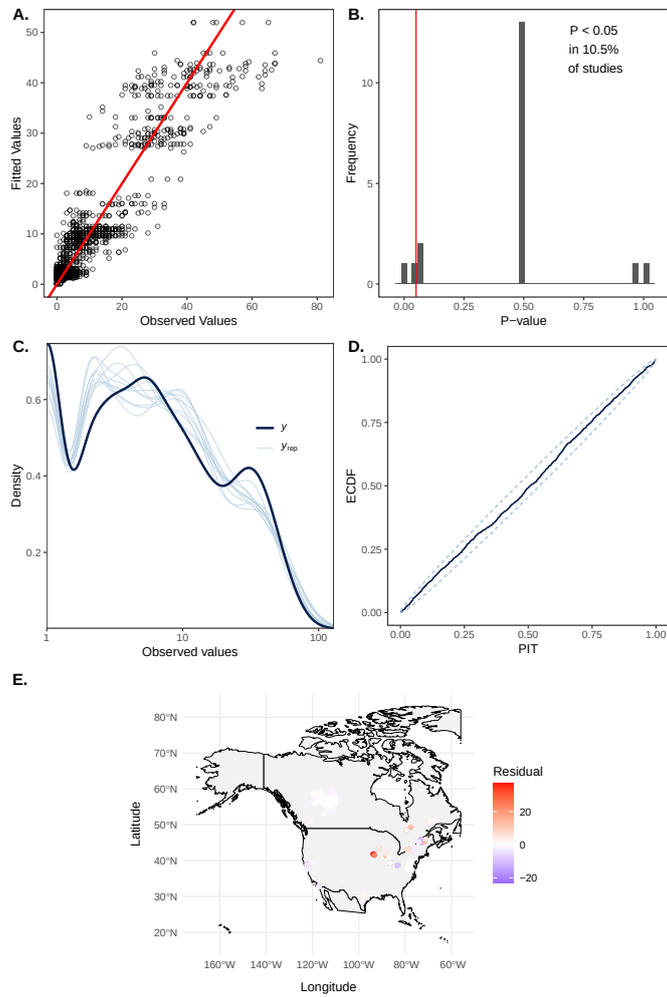


Figure M26: Model diagnostic plots for the zero-inflated negative binomial model of species richness responses to land use and range extent in North America: $\text{Richness} \sim \text{LandUse} \times \text{RangeExtent} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M19: Posterior summary of the model of species richness for Europe (Fig. M25).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ UI2 * Quants + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 6114) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 105) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.26 | 0.09 | 1.10 | 1.45 | 1.00 | 1177 | 2323 |
| ~SS:SSB (Number of levels: 861) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.16 | 0.02 | 0.12 | 0.20 | 1.00 | 3016 | 5291 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.89 | 0.14 | 1.61 | 2.17 | 1.01 | 661 | 1581 |
| zi_Intercept | -3.28 | 0.14 | -3.59 | -3.02 | 1.00 | 12695 | 6953 |
| UI2Secondaryvegetation | -0.69 | 0.09 | -0.87 | -0.51 | 1.00 | 3734 | 6049 |
| UI2Agriculture_Minimal | -0.45 | 0.08 | -0.60 | -0.30 | 1.00 | 3485 | 5708 |
| UI2Agriculture_Light | -0.70 | 0.09 | -0.88 | -0.53 | 1.00 | 3136 | 5116 |
| UI2Agriculture_Intense | -0.57 | 0.09 | -0.75 | -0.39 | 1.00 | 3378 | 5739 |
| Quants2 | -0.09 | 0.07 | -0.23 | 0.05 | 1.00 | 2739 | 5151 |
| UI2Secondaryvegetation:Quants2 | 1.19 | 0.08 | 1.02 | 1.36 | 1.00 | 3108 | 5742 |
| UI2Agriculture_Minimal:Quants2 | 0.87 | 0.08 | 0.71 | 1.04 | 1.00 | 3263 | 5836 |
| UI2Agriculture_Light:Quants2 | 1.08 | 0.08 | 0.92 | 1.24 | 1.00 | 3002 | 5300 |
| UI2Agriculture_Intense:Quants2 | 0.88 | 0.08 | 0.73 | 1.04 | 1.00 | 3031 | 5730 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| shape | 4.57 | 0.21 | 4.16 | 5.00 | 1.00 | 9437 | 7300 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.800 (95% CI [0.780, 0.820]) | | | | | | | |
| Marginal R2: 0.037 (95% CI [0.021, 0.056]) | | | | | | | |

Table M20: Posterior summary of the model of species richness for North America (Fig. ??).

| BRMS Summary | | | | | | | |
|--|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ UI2 * Quants + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 1697) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 36) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.51 | 0.19 | 1.19 | 1.95 | 1.00 | 1859 | 3630 |
| ~SS:SSB (Number of levels: 120) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.11 | 0.04 | 0.04 | 0.18 | 1.00 | 2555 | 3301 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.50 | 0.26 | 1.00 | 2.02 | 1.00 | 819 | 1833 |
| zi_Intercept | -6.00 | 1.13 | -8.81 | -4.46 | 1.00 | 7993 | 4232 |
| UI2Secondaryvegetation | -0.21 | 0.07 | -0.35 | -0.07 | 1.00 | 9649 | 7390 |
| UI2Agriculture_Minimal | -0.70 | 0.15 | -0.99 | -0.41 | 1.00 | 8618 | 7219 |
| UI2Agriculture_Light | -0.45 | 0.12 | -0.69 | -0.22 | 1.00 | 9699 | 7985 |
| UI2Agriculture_Intense | -0.54 | 0.15 | -0.84 | -0.26 | 1.00 | 9776 | 8143 |
| Quants2 | 0.12 | 0.04 | 0.04 | 0.20 | 1.00 | 8959 | 7692 |
| UI2Secondaryvegetation:Quants2 | 0.23 | 0.06 | 0.10 | 0.35 | 1.00 | 10205 | 7734 |

| | | | | | | | |
|--------------------------------|------|------|-------|------|------|-------|------|
| UI2Agriculture_Minimal:Quants2 | 0.92 | 0.15 | 0.62 | 1.20 | 1.00 | 11101 | 7846 |
| UI2Agriculture_Light:Quants2 | 0.17 | 0.13 | -0.09 | 0.43 | 1.00 | 11960 | 8060 |
| UI2Agriculture_Intense:Quants2 | 0.15 | 0.15 | -0.15 | 0.45 | 1.00 | 12104 | 7908 |

Further Distributional Parameters:

| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
|-------|----------|-----------|----------|----------|------|----------|----------|
| shape | 8.13 | 0.78 | 6.73 | 9.81 | 1.00 | 12088 | 7071 |

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1).

Bayesian R2 with Compatibility Interval

Conditional R2: 0.848 (95% CI [0.831, 0.861])
 Marginal R2: 0.005 (95% CI [0.001, 0.012])

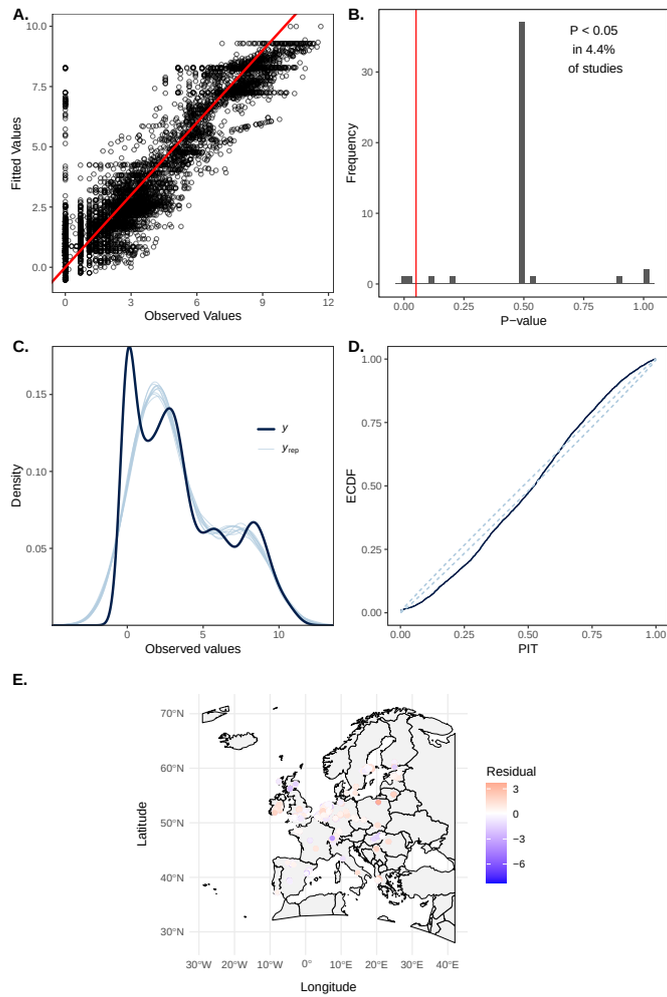


Figure M27: Model diagnostic plots for the Gaussian model of total abundance responses to land use and range extent in Europe: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} \times \text{RangeExtent} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

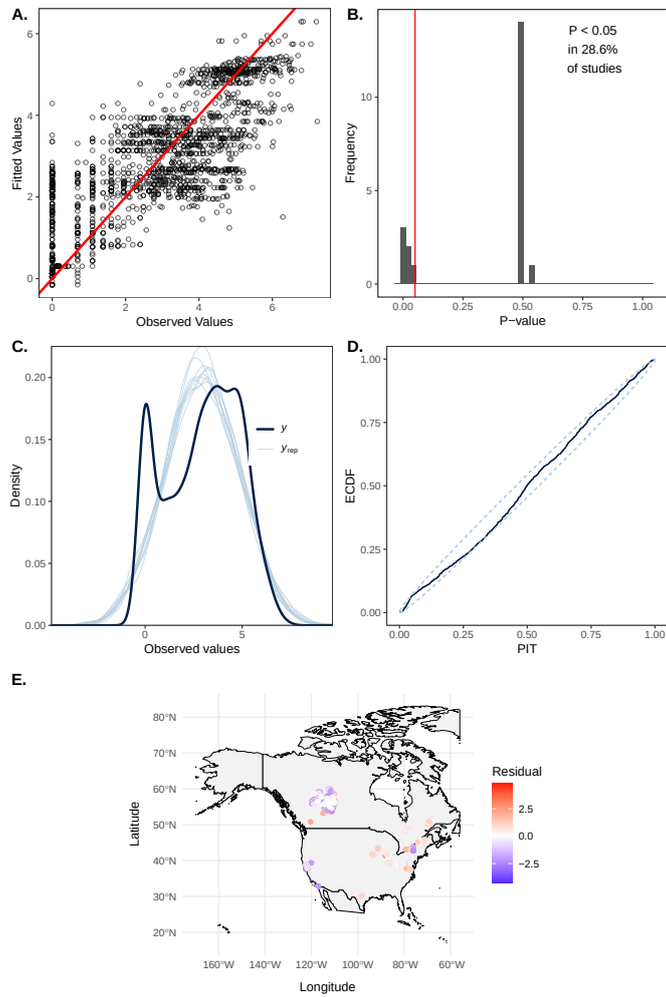


Figure M28: Model diagnostic plots for the Gaussian model of total abundance responses to land use and range extent in North America: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} \times \text{RangeExtent} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M21: Posterior summary of the model of total abundance for Europe (Fig. M27).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ UI2 * Quants + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 5741) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 99) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.91 | 0.16 | 1.63 | 2.24 | 1.00 | 1443 | 3048 |
| ~SS:SSB (Number of levels: 847) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.73 | 0.05 | 0.63 | 0.83 | 1.00 | 2434 | 4230 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.83 | 0.29 | 2.25 | 3.41 | 1.00 | 1576 | 3417 |
| UI2Secondaryvegetation | 0.00 | 0.23 | -0.45 | 0.45 | 1.00 | 2925 | 5271 |
| UI2Agriculture_Minimal | 0.08 | 0.23 | -0.38 | 0.52 | 1.00 | 2880 | 5322 |
| UI2Agriculture_Light | -0.25 | 0.24 | -0.71 | 0.22 | 1.00 | 2818 | 4950 |
| UI2Agriculture_Intense | -0.17 | 0.24 | -0.65 | 0.30 | 1.00 | 2845 | 4790 |
| Quants2 | 0.62 | 0.17 | 0.27 | 0.95 | 1.00 | 2344 | 4293 |
| UI2Secondaryvegetation:Quants2 | 0.34 | 0.20 | -0.04 | 0.73 | 1.00 | 2802 | 4873 |
| UI2Agriculture_Minimal:Quants2 | 0.42 | 0.19 | 0.05 | 0.80 | 1.00 | 2596 | 4812 |
| UI2Agriculture_Light:Quants2 | 0.58 | 0.18 | 0.22 | 0.94 | 1.00 | 2619 | 4778 |
| UI2Agriculture_Intense:Quants2 | 0.33 | 0.18 | -0.03 | 0.69 | 1.00 | 2496 | 5031 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 1.27 | 0.01 | 1.25 | 1.30 | 1.00 | 6912 | 7135 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.825 (95% CI [0.820, 0.830]) | | | | | | | |
| Marginal R2: 0.030 (95% CI [0.026, 0.034]) | | | | | | | |

Table M22: Posterior summary of the model of total abundance for North America (Fig. M28).

| BRMS Summary | | | | | | | |
|--|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ UI2 * Quants + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 1677) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 35) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.66 | 0.22 | 1.29 | 2.15 | 1.00 | 3236 | 5349 |
| ~SS:SSB (Number of levels: 119) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.38 | 0.07 | 0.25 | 0.52 | 1.00 | 3299 | 6328 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 3.39 | 0.30 | 2.81 | 3.97 | 1.00 | 2321 | 4106 |
| UI2Secondaryvegetation | -0.28 | 0.14 | -0.55 | -0.01 | 1.00 | 13088 | 7844 |
| UI2Agriculture_Minimal | -0.62 | 0.23 | -1.08 | -0.16 | 1.00 | 12505 | 7993 |
| UI2Agriculture_Light | -0.77 | 0.22 | -1.20 | -0.35 | 1.00 | 13961 | 8184 |
| UI2Agriculture_Intense | -1.04 | 0.30 | -1.63 | -0.45 | 1.00 | 15351 | 8048 |
| Quants2 | -0.08 | 0.09 | -0.24 | 0.09 | 1.00 | 12837 | 8193 |
| UI2Secondaryvegetation:Quants2 | 0.42 | 0.13 | 0.15 | 0.67 | 1.00 | 13569 | 8593 |
| UI2Agriculture_Minimal:Quants2 | 0.92 | 0.24 | 0.46 | 1.38 | 1.00 | 17378 | 7338 |
| UI2Agriculture_Light:Quants2 | 0.18 | 0.25 | -0.30 | 0.66 | 1.00 | 15600 | 8403 |
| UI2Agriculture_Intense:Quants2 | 0.58 | 0.37 | -0.14 | 1.29 | 1.00 | 15842 | 7552 |

```

Further Distributional Parameters:
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sigma    1.23      0.02    1.18    1.27 1.00   16728    7006

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
and Tail_ESS are effective sample size measures, and Rhat is the potential
scale reduction factor on split chains (at convergence, Rhat = 1).

# Bayesian R2 with Compatibility Interval

Conditional R2: 0.581 (95% CI [0.559, 0.600])
Marginal R2: 0.023 (95% CI [0.011, 0.037])

```

D.2 Habitat specificity

The following model diagnostic plots refer to the comparison of biodiversity responses in primary vegetation, secondary vegetation and agricultural land use between species with different levels of habitat specificity, comparing Europe and North America (Fig. 5).

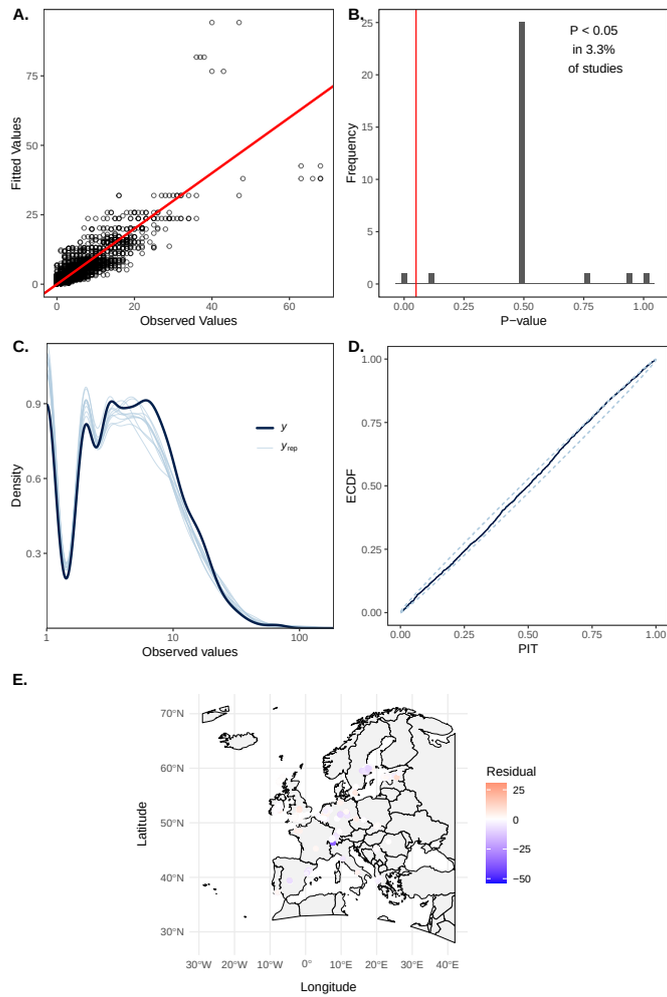


Figure M29: Model diagnostic plots for the zero-inflated negative binomial model of species richness responses to land use and habitat specificity in Europe: $\text{Richness} \sim \text{LandUse} \times \text{Habitat-Specificity} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

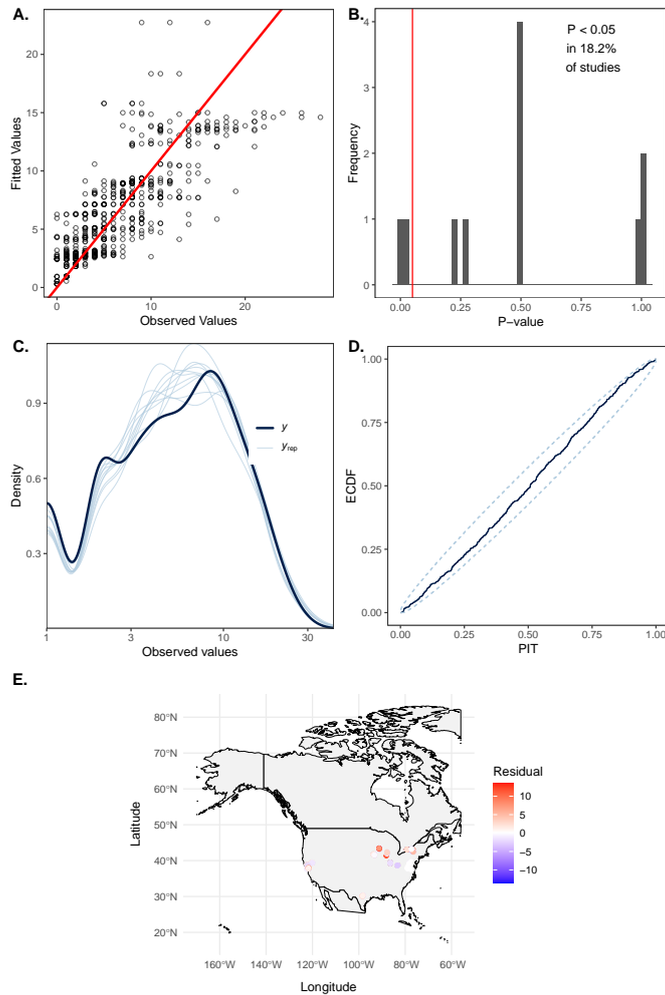


Figure M30: Model diagnostic plots for the zero-inflated negative binomial model of species richness responses to land use and habitat specificity in North America: $Richness \sim LandUse \times HabitatSpecificity + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M23: Posterior summary of the model of species richness for Europe (Fig. M29).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ UI2 * Quants + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 3389) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 59) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.11 | 0.11 | 0.91 | 1.34 | 1.00 | 1537 | 3371 |
| ~SS:SSB (Number of levels: 280) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.23 | 0.03 | 0.17 | 0.29 | 1.00 | 2857 | 5086 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.28 | 0.19 | 0.91 | 1.64 | 1.00 | 1047 | 2295 |
| zi_Intercept | -6.30 | 1.14 | -9.18 | -4.75 | 1.00 | 10361 | 5057 |
| UI2Secondaryvegetation | -0.37 | 0.13 | -0.62 | -0.12 | 1.00 | 4233 | 6135 |
| UI2Agriculture_Minimal | -0.49 | 0.12 | -0.74 | -0.24 | 1.00 | 3928 | 6321 |
| UI2Agriculture_Light | -0.49 | 0.14 | -0.76 | -0.21 | 1.00 | 3252 | 5350 |
| UI2Agriculture_Intense | -0.54 | 0.14 | -0.82 | -0.26 | 1.00 | 3262 | 5604 |
| Quants2 | -0.35 | 0.09 | -0.53 | -0.17 | 1.00 | 2811 | 4701 |
| UI2Secondaryvegetation:Quants2 | 0.93 | 0.10 | 0.73 | 1.13 | 1.00 | 3133 | 5345 |
| UI2Agriculture_Minimal:Quants2 | 1.25 | 0.11 | 1.05 | 1.46 | 1.00 | 3205 | 5745 |
| UI2Agriculture_Light:Quants2 | 1.11 | 0.10 | 0.92 | 1.31 | 1.00 | 3028 | 4783 |
| UI2Agriculture_Intense:Quants2 | 1.19 | 0.10 | 1.00 | 1.39 | 1.00 | 3015 | 4851 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| shape | 11.23 | 0.99 | 9.46 | 13.32 | 1.00 | 13751 | 7153 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.757 (95% CI [0.747, 0.767]) | | | | | | | |
| Marginal R2: 0.044 (95% CI [0.023, 0.073]) | | | | | | | |

Table M24: Posterior summary of the model of species richness for North America (Fig. ??).

| BRMS Summary | | | | | | | |
|--|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Species_richness ~ UI2 * Quants + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 605) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 19) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.08 | 0.22 | 0.74 | 1.62 | 1.00 | 2260 | 4434 |
| ~SS:SSB (Number of levels: 76) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.11 | 0.06 | 0.01 | 0.22 | 1.00 | 1986 | 2904 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 1.74 | 0.26 | 1.22 | 2.27 | 1.00 | 1189 | 2643 |
| zi_Intercept | -5.68 | 1.15 | -8.65 | -4.11 | 1.00 | 8700 | 4197 |
| UI2Secondaryvegetation | 0.05 | 0.10 | -0.15 | 0.26 | 1.00 | 8320 | 7457 |
| UI2Agriculture_Minimal | -0.48 | 0.22 | -0.92 | -0.05 | 1.00 | 7286 | 6680 |
| UI2Agriculture_Light | -0.60 | 0.16 | -0.92 | -0.29 | 1.00 | 6719 | 7571 |
| UI2Agriculture_Intense | -0.85 | 0.19 | -1.23 | -0.47 | 1.00 | 7873 | 7677 |
| Quants2 | -0.08 | 0.09 | -0.25 | 0.09 | 1.00 | 5066 | 6650 |
| UI2Secondaryvegetation:Quants2 | -0.34 | 0.10 | -0.54 | -0.14 | 1.00 | 5425 | 6772 |

| | | | | | | | |
|--------------------------------|------|------|-------|------|------|------|------|
| UI2Agriculture_Minimal:Quants2 | 0.33 | 0.21 | -0.08 | 0.75 | 1.00 | 6831 | 6774 |
| UI2Agriculture_Light:Quants2 | 0.32 | 0.18 | -0.04 | 0.68 | 1.00 | 6379 | 7347 |
| UI2Agriculture_Intense:Quants2 | 0.57 | 0.20 | 0.17 | 0.97 | 1.00 | 7772 | 7352 |

Further Distributional Parameters:

| | Estimate | Est.Error | l-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
|-------|----------|-----------|----------|----------|------|----------|----------|
| shape | 17.33 | 3.92 | 11.49 | 26.63 | 1.00 | 10655 | 6845 |

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1).

Bayesian R2 with Compatibility Interval

Conditional R2: 0.646 (95% CI [0.610, 0.677])
 Marginal R2: 0.040 (95% CI [0.007, 0.102])

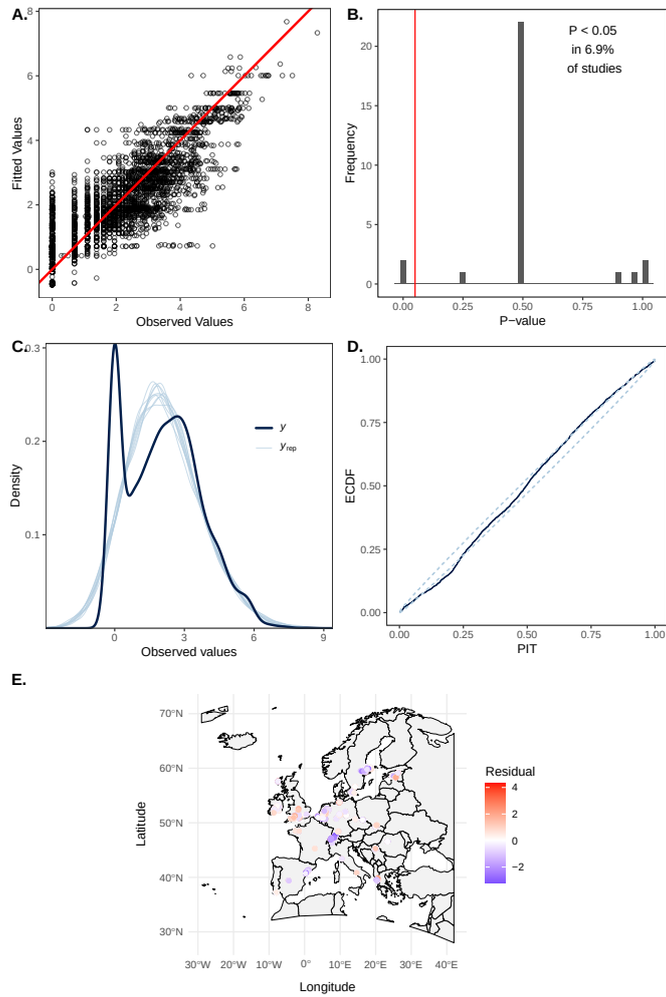


Figure M31: Model diagnostic plots for the Gaussian model of total abundance responses to land use and habitat specificity in Europe: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} \times \text{HabitatSpecificity} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

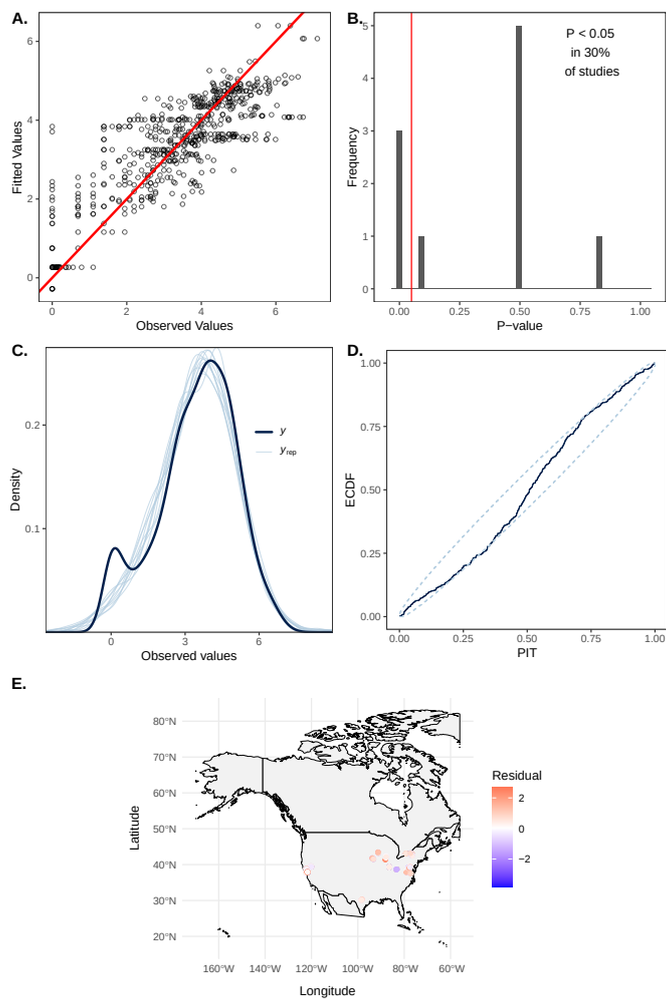


Figure M32: Model diagnostic plots for the Gaussian model of total abundance responses to land use and habitat specificity in North America: $\text{Log}(\text{Abundance}+1) \sim \text{LandUse} \times \text{Habitat-Specificity} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Figure M33: Posterior summary of the model of total abundance for Europe (Fig. M31).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ UI2 * Quants + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 3233) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 55) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.52 | 0.16 | 1.25 | 1.87 | 1.00 | 1681 | 3195 |
| ~SS:SSB (Number of levels: 273) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.29 | 0.04 | 0.21 | 0.37 | 1.00 | 3096 | 5738 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 2.22 | 0.29 | 1.63 | 2.80 | 1.00 | 1315 | 2060 |
| UI2Secondaryvegetation | -0.52 | 0.22 | -0.96 | -0.10 | 1.00 | 4192 | 5436 |
| UI2Agriculture_Minimal | -0.28 | 0.21 | -0.70 | 0.12 | 1.00 | 3990 | 6087 |
| UI2Agriculture_Light | -0.17 | 0.23 | -0.62 | 0.28 | 1.00 | 3442 | 5745 |
| UI2Agriculture_Intense | -0.21 | 0.23 | -0.66 | 0.23 | 1.00 | 3310 | 5244 |
| Quants2 | -0.53 | 0.15 | -0.83 | -0.23 | 1.00 | 3410 | 5373 |
| UI2Secondaryvegetation:Quants2 | 1.75 | 0.17 | 1.41 | 2.09 | 1.00 | 3749 | 6309 |
| UI2Agriculture_Minimal:Quants2 | 1.95 | 0.17 | 1.61 | 2.30 | 1.00 | 3700 | 6197 |
| UI2Agriculture_Light:Quants2 | 1.50 | 0.16 | 1.18 | 1.82 | 1.00 | 3498 | 5653 |
| UI2Agriculture_Intense:Quants2 | 1.66 | 0.16 | 1.34 | 1.98 | 1.00 | 3529 | 5571 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.88 | 0.01 | 0.86 | 0.91 | 1.00 | 14456 | 7432 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.693 (95% CI [0.683, 0.704]) | | | | | | | |
| Marginal R2: 0.126 (95% CI [0.114, 0.138]) | | | | | | | |

Figure M34: Posterior summary of the model of total abundance for North America (Fig. M32).

| BRMS Summary | | | | | | | |
|--|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: LogAbund ~ UI2 * Quants + (1 SS/SSB) | | | | | | | |
| Data: model_data (Number of observations: 585) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 18) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.48 | 0.32 | 0.96 | 2.22 | 1.00 | 4449 | 5656 |
| ~SS:SSB (Number of levels: 75) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.57 | 0.10 | 0.39 | 0.78 | 1.00 | 3372 | 5937 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 3.63 | 0.41 | 2.82 | 4.45 | 1.00 | 4221 | 5356 |
| UI2Secondaryvegetation | 0.13 | 0.24 | -0.33 | 0.60 | 1.00 | 10628 | 7513 |
| UI2Agriculture_Minimal | 0.32 | 0.48 | -0.63 | 1.26 | 1.00 | 12634 | 7358 |
| UI2Agriculture_Light | -1.03 | 0.32 | -1.66 | -0.41 | 1.00 | 11549 | 7636 |
| UI2Agriculture_Intense | -1.68 | 0.43 | -2.52 | -0.86 | 1.00 | 12092 | 8025 |
| Quants2 | 0.37 | 0.19 | 0.01 | 0.73 | 1.00 | 8829 | 7665 |
| UI2Secondaryvegetation:Quants2 | -0.70 | 0.23 | -1.14 | -0.25 | 1.00 | 9704 | 8146 |
| UI2Agriculture_Minimal:Quants2 | -0.65 | 0.47 | -1.58 | 0.27 | 1.00 | 11400 | 7260 |
| UI2Agriculture_Light:Quants2 | 0.22 | 0.36 | -0.48 | 0.92 | 1.00 | 11246 | 8224 |
| UI2Agriculture_Intense:Quants2 | 1.64 | 0.51 | 0.66 | 2.65 | 1.00 | 12925 | 8343 |

```
Further Distributional Parameters:
      Estimate Est.Error l-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sigma    0.98    0.03    0.92    1.04 1.00    12818    7423

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
and Tail_ESS are effective sample size measures, and Rhat is the potential
scale reduction factor on split chains (at convergence, Rhat = 1).

# Bayesian R2 with Compatibility Interval

Conditional R2: 0.630 (95% CI [0.595, 0.663])
Marginal R2: 0.061 (95% CI [0.032, 0.101])
```

D.3 Non-native species proportion

The following model diagnostic plots refer to the comparison of the proportions of non-native species in primary vegetation, secondary vegetation and agricultural land use in Europe and North America (Fig. 5).

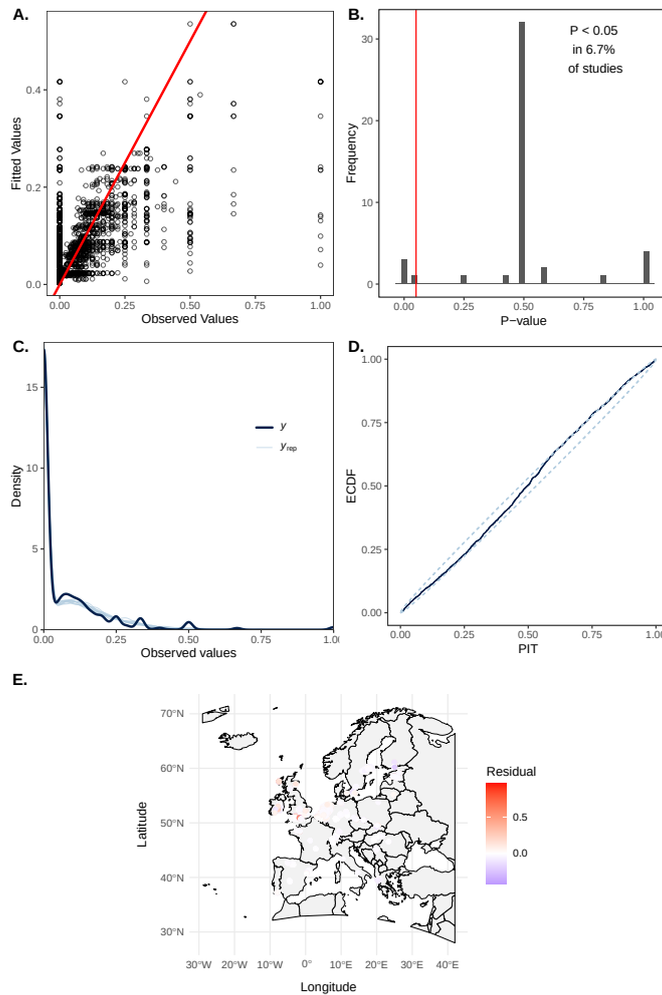


Figure M35: Model diagnostic plots for the zero-and-one inflated beta model of non-native species proportions of richness P_{SR} in Europe: $P_{SR} \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Due to the zero-and-one inflated structure of the data, deviations of PIT values from uniformity are to be expected and do not necessarily indicate poor model fit. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

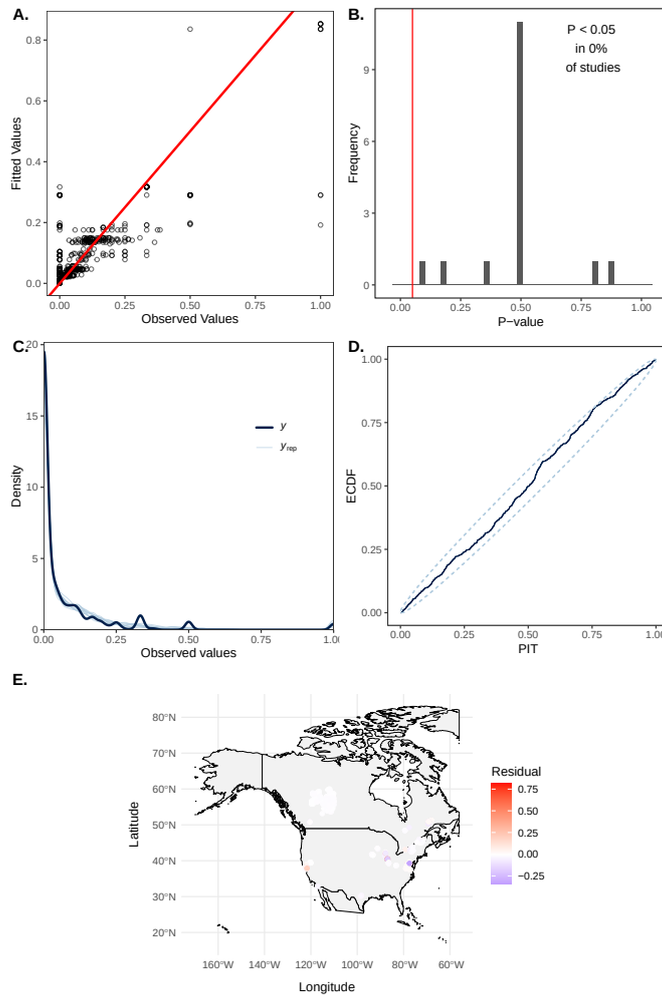


Figure M36: Model diagnostic plots for the zero-and-one-inflated beta model of non-native proportions of species richness P_{SR} in North America: $P_{SR} \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Due to the zero-and-one inflated structure of the data, deviations of PIT values from uniformity are to be expected and do not necessarily indicate poor model fit. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M25: Posterior summary of the model of species richness for Europe (Fig. M35).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_one_inflated_beta | | | | | | | |
| Links: mu = logit; phi = log; zoi = logit; coi = logit | | | | | | | |
| Formula: SiteSpeciesProps ~ UI2 + (1 SS/SSB) | | | | | | | |
| phi ~ 1 | | | | | | | |
| zoi ~ 1 + (1 SS) | | | | | | | |
| coi ~ 1 + (1 SS) | | | | | | | |
| Data: model_data (Number of observations: 2837) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 99) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.86 | 0.08 | 0.73 | 1.02 | 1.00 | 3693 | 7103 |
| sd(zoi_Intercept) | 3.66 | 0.47 | 2.86 | 4.70 | 1.00 | 5324 | 9853 |
| sd(coi_Intercept) | 2.37 | 0.59 | 1.43 | 3.75 | 1.00 | 10014 | 12712 |
| ~SS:SSB (Number of levels: 823) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.08 | 0.04 | 0.01 | 0.14 | 1.00 | 3649 | 4537 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | -1.82 | 0.16 | -2.14 | -1.52 | 1.00 | 3927 | 7542 |
| phi_Intercept | 3.44 | 0.05 | 3.35 | 3.53 | 1.00 | 20882 | 15297 |
| zoi_Intercept | -0.61 | 0.40 | -1.42 | 0.18 | 1.00 | 1819 | 3867 |
| coi_Intercept | -6.22 | 0.80 | -8.01 | -4.90 | 1.00 | 14550 | 13338 |
| UI2Secondaryvegetation | -0.05 | 0.13 | -0.31 | 0.22 | 1.00 | 9089 | 12602 |
| UI2Agriculture_Minimal | -0.00 | 0.14 | -0.27 | 0.28 | 1.00 | 8772 | 12633 |
| UI2Agriculture_Light | 0.17 | 0.14 | -0.11 | 0.45 | 1.00 | 8197 | 12246 |
| UI2Agriculture_Intense | 0.11 | 0.15 | -0.17 | 0.40 | 1.00 | 8079 | 11756 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.353 (95% CI [0.309, 0.402]) | | | | | | | |
| Marginal R2: 0.003 (95% CI [2.961e-04, 0.008]) | | | | | | | |

Table M26: Posterior summary of the model of species richness for North America (Fig. ??).

| BRMS Summary | | | | | | | |
|--|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_one_inflated_beta | | | | | | | |
| Links: mu = logit; phi = log; zoi = logit; coi = logit | | | | | | | |
| Formula: SiteSpeciesProps ~ UI2 + (1 SS/SSB) | | | | | | | |
| phi ~ 1 | | | | | | | |
| zoi ~ 1 + (1 SS) | | | | | | | |
| coi ~ 1 + (1 SS) | | | | | | | |
| Data: model_data (Number of observations: 832) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 35) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.06 | 0.19 | 0.77 | 1.51 | 1.00 | 2071 | 3682 |
| sd(zoi_Intercept) | 5.19 | 1.14 | 3.42 | 7.83 | 1.00 | 2794 | 4846 |
| sd(coi_Intercept) | 3.81 | 1.26 | 2.05 | 6.81 | 1.00 | 5529 | 6003 |
| ~SS:SSB (Number of levels: 119) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.08 | 0.05 | 0.00 | 0.19 | 1.00 | 2601 | 2917 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | -2.31 | 0.25 | -2.82 | -1.82 | 1.00 | 996 | 2041 |
| phi_Intercept | 4.00 | 0.08 | 3.85 | 4.16 | 1.00 | 10248 | 7498 |
| zoi_Intercept | 1.01 | 0.85 | -0.61 | 2.80 | 1.00 | 1244 | 2462 |
| coi_Intercept | -6.23 | 1.36 | -9.37 | -3.99 | 1.00 | 6056 | 5299 |
| UI2Secondaryvegetation | 0.30 | 0.12 | 0.07 | 0.54 | 1.00 | 6543 | 7357 |
| UI2Agriculture_Minimal | 0.48 | 0.20 | 0.08 | 0.86 | 1.00 | 4748 | 7035 |
| UI2Agriculture_Light | 0.53 | 0.22 | 0.10 | 0.95 | 1.00 | 3773 | 5904 |

```
UI2Agriculture_Intense 0.53 0.27 -0.00 1.04 1.00 4984 6886
```

```
Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS  
and Tail_ESS are effective sample size measures, and Rhat is the potential  
scale reduction factor on split chains (at convergence, Rhat = 1).
```

```
# Bayesian R2 with Compatibility Interval
```

```
Conditional R2: 0.658 (95% CI [0.552, 0.715])
```

```
Marginal R2: 0.002 (95% CI [7.614e-08, 0.012])
```

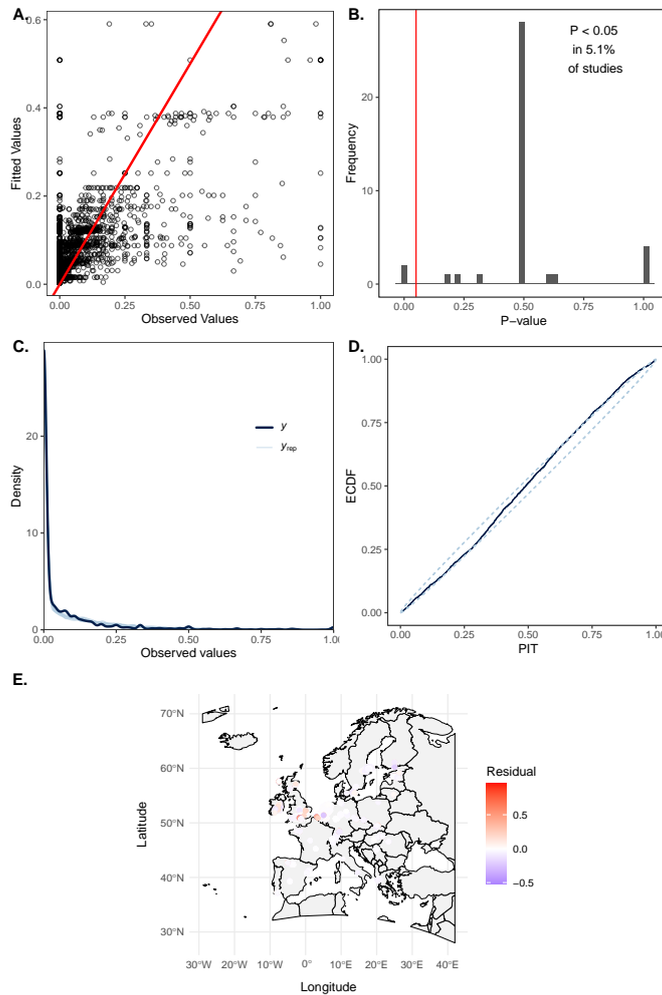


Figure M37: Model diagnostic plots for the zero-and-one-inflated beta model of non-native proportions of total abundance P_{TA} in Europe: $P_{TA} \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Due to the zero-and-one inflated structure of the data, deviations of PIT values from uniformity are to be expected and do not necessarily indicate poor model fit. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

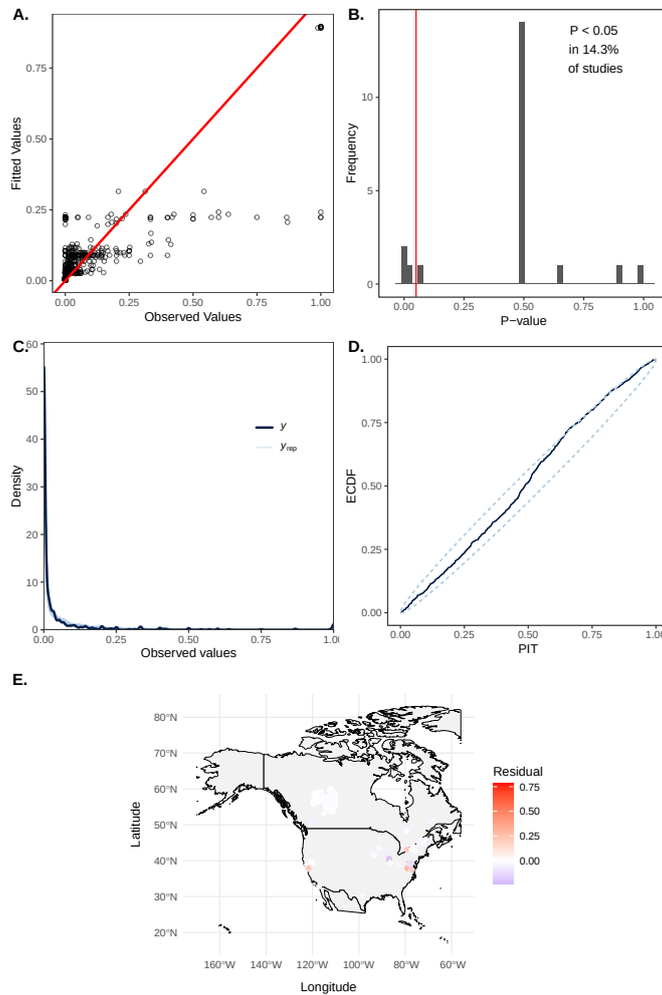


Figure M38: Model diagnostic plots for the zero-and-one-inflated beta model of non-native proportions of total abundance P_{TA} in North America: $P_{TA} \sim \text{LandUse} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Due to the zero-and-one inflated structure of the data, deviations of PIT values from uniformity are to be expected and do not necessarily indicate poor model fit. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M27: Posterior summary of the model of total abundance for Europe (Fig. M37).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_one_inflated_beta | | | | | | | |
| Links: mu = logit; phi = log; zoi = logit; coi = logit | | | | | | | |
| Formula: SiteTotAbunProps ~ UI2 + (1 SS/SSB) | | | | | | | |
| phi ~ 1 | | | | | | | |
| zoi ~ (1 SS) | | | | | | | |
| coi ~ (1 SS) | | | | | | | |
| Data: model_data (Number of observations: 2837) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 99) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.02 | 0.10 | 0.85 | 1.23 | 1.00 | 6129 | 11007 |
| sd(zoi_Intercept) | 3.66 | 0.47 | 2.86 | 4.71 | 1.00 | 5835 | 10593 |
| sd(coi_Intercept) | 2.36 | 0.59 | 1.42 | 3.72 | 1.00 | 10034 | 13639 |
| ~SS:SSB (Number of levels: 823) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.18 | 0.06 | 0.05 | 0.29 | 1.00 | 3358 | 4339 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | -1.88 | 0.23 | -2.33 | -1.43 | 1.00 | 6490 | 10653 |
| phi_Intercept | 2.35 | 0.05 | 2.26 | 2.45 | 1.00 | 13682 | 14449 |
| zoi_Intercept | -0.62 | 0.41 | -1.45 | 0.15 | 1.00 | 2006 | 4348 |
| coi_Intercept | -6.22 | 0.80 | -8.05 | -4.91 | 1.00 | 15486 | 13607 |
| UI2Secondaryvegetation | -0.09 | 0.21 | -0.51 | 0.33 | 1.00 | 9832 | 13033 |
| UI2Agriculture_Minimal | -0.14 | 0.22 | -0.57 | 0.30 | 1.00 | 9582 | 12465 |
| UI2Agriculture_Light | 0.12 | 0.23 | -0.32 | 0.56 | 1.00 | 8993 | 11493 |
| UI2Agriculture_Intense | 0.07 | 0.23 | -0.37 | 0.53 | 1.00 | 8918 | 12233 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.364 (95% CI [0.313, 0.412]) | | | | | | | |
| Marginal R2: 0.004 (95% CI [1.334e-04, 0.010]) | | | | | | | |

Table M28: Posterior summary of the model of total abundance for North America (Fig. M38).

| BRMS Summary | | | | | | | |
|--|----------|-----------|----------|----------|------|----------|----------|
| Family: zero_one_inflated_beta | | | | | | | |
| Links: mu = logit; phi = log; zoi = logit; coi = logit | | | | | | | |
| Formula: SiteTotAbunProps ~ UI2 + (1 SS/SSB) | | | | | | | |
| phi ~ 1 | | | | | | | |
| zoi ~ 1 + (1 SS) | | | | | | | |
| coi ~ 1 + (1 SS) | | | | | | | |
| Data: model_data (Number of observations: 832) | | | | | | | |
| Draws: 4 chains, each with iter = 5000; warmup = 2500; thin = 1; | | | | | | | |
| total post-warmup draws = 10000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 35) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 1.50 | 0.29 | 1.04 | 2.17 | 1.00 | 2935 | 5413 |
| sd(zoi_Intercept) | 5.14 | 1.13 | 3.40 | 7.78 | 1.00 | 2627 | 5391 |
| sd(coi_Intercept) | 3.75 | 1.18 | 2.01 | 6.55 | 1.00 | 6608 | 6724 |
| ~SS:SSB (Number of levels: 119) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.25 | 0.16 | 0.01 | 0.56 | 1.00 | 1010 | 3282 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | -2.29 | 0.39 | -3.06 | -1.53 | 1.00 | 1712 | 3326 |
| phi_Intercept | 2.61 | 0.10 | 2.41 | 2.82 | 1.00 | 2468 | 4435 |
| zoi_Intercept | 0.98 | 0.85 | -0.66 | 2.72 | 1.00 | 1435 | 2792 |
| coi_Intercept | -6.22 | 1.34 | -9.24 | -4.01 | 1.00 | 7007 | 7286 |
| UI2Secondaryvegetation | 0.10 | 0.23 | -0.36 | 0.57 | 1.00 | 7274 | 6367 |
| UI2Agriculture_Minimal | -0.15 | 0.40 | -0.94 | 0.63 | 1.00 | 4981 | 7290 |
| UI2Agriculture_Light | 0.68 | 0.43 | -0.14 | 1.55 | 1.00 | 3742 | 5370 |

```
UI2Agriculture_Intense 0.37 0.58 -0.80 1.49 1.00 4899 6503
```

```
Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS  
and Tail_ESS are effective sample size measures, and Rhat is the potential  
scale reduction factor on split chains (at convergence, Rhat = 1).
```

```
# Bayesian R2 with Compatibility Interval
```

```
Conditional R2: 0.576 (95% CI [0.440, 0.648])  
Marginal R2: 0.003 (95% CI [2.082e-06, 0.027])
```

E Abundance model comparison

The following model diagnostic plots refer to the comparison of four possible models of total abundance: The Gaussian model of $\log(\text{total abundance}+1)$, the Gaussian model of $\log(\text{rescaled abundance}+0.01)$, the zero-and-one-inflated beta model of rescaled abundance, and the zero-inflated negative binomial model of (rounded) total abundance (Fig. S20).

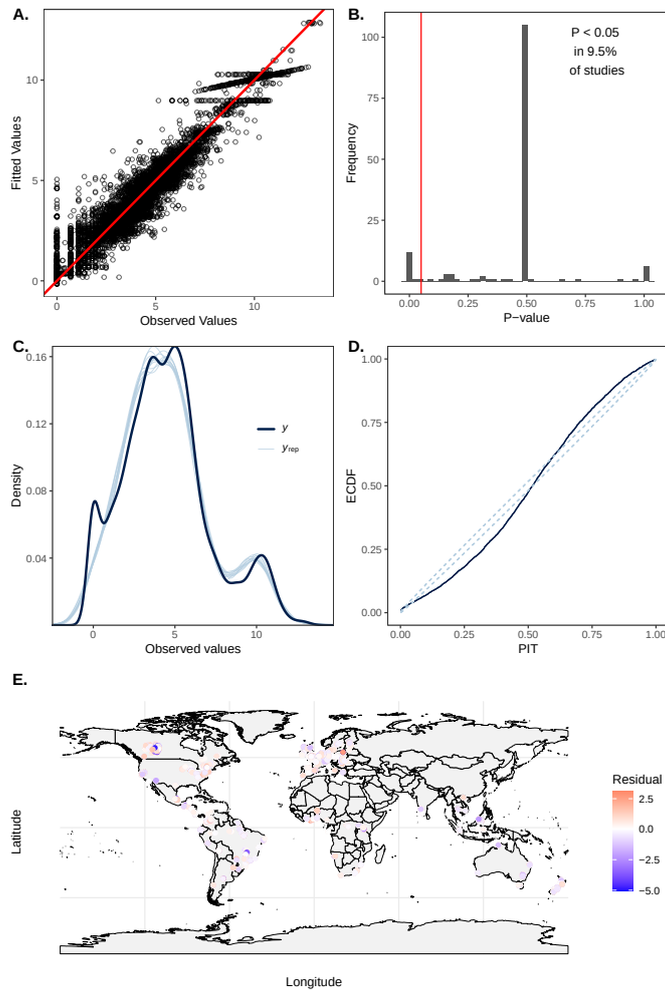


Figure M39: Model diagnostic plots for the Gaussian model of $\log(\text{total abundance}+1) \sim \text{LandUse} \times \text{Region} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

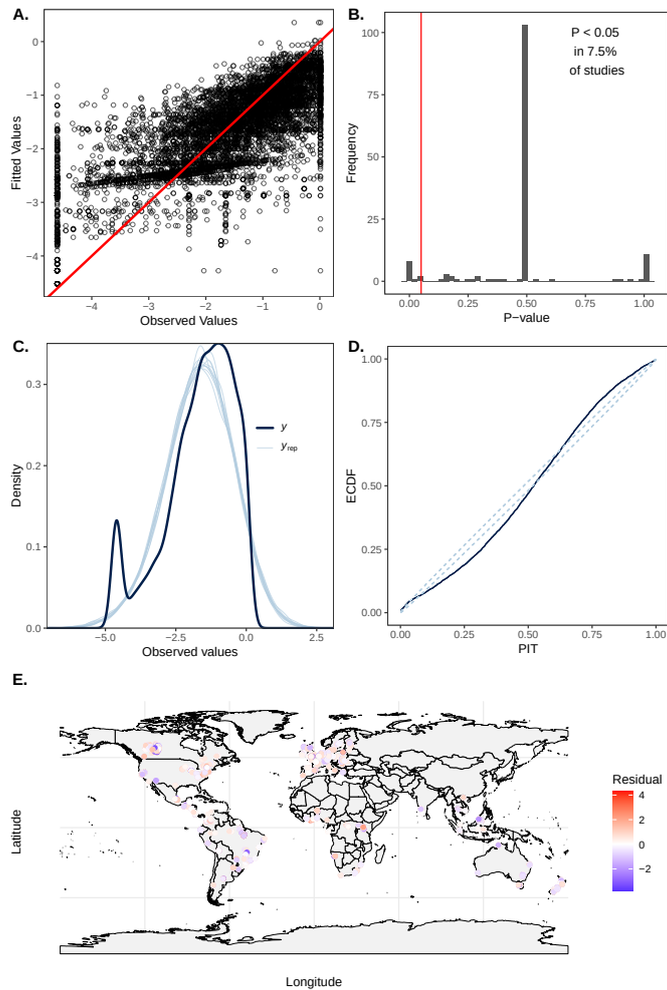


Figure M40: Model diagnostic plots for the Gaussian model of $\log(\text{rescaled total abundance} + 0.01) \sim \text{LandUse} \times \text{Region} + (1|\text{SS}/\text{SSB})$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

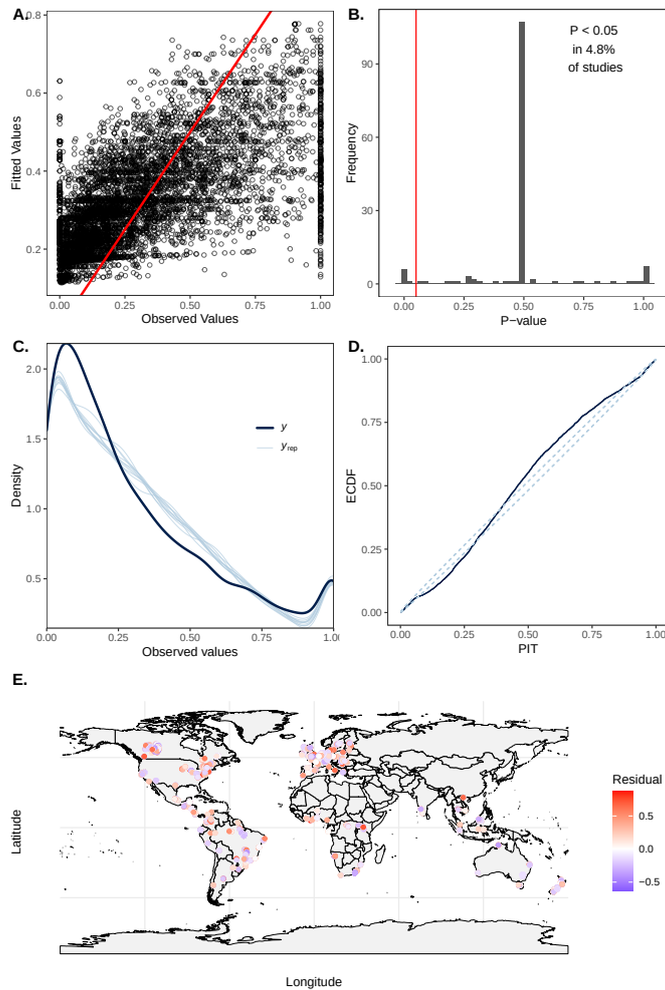


Figure M41: Model diagnostic plots for the beta distributed model of rescaled total abundance $\sim \text{LandUse} \times \text{Region} + (1|SS/SSB)$. Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

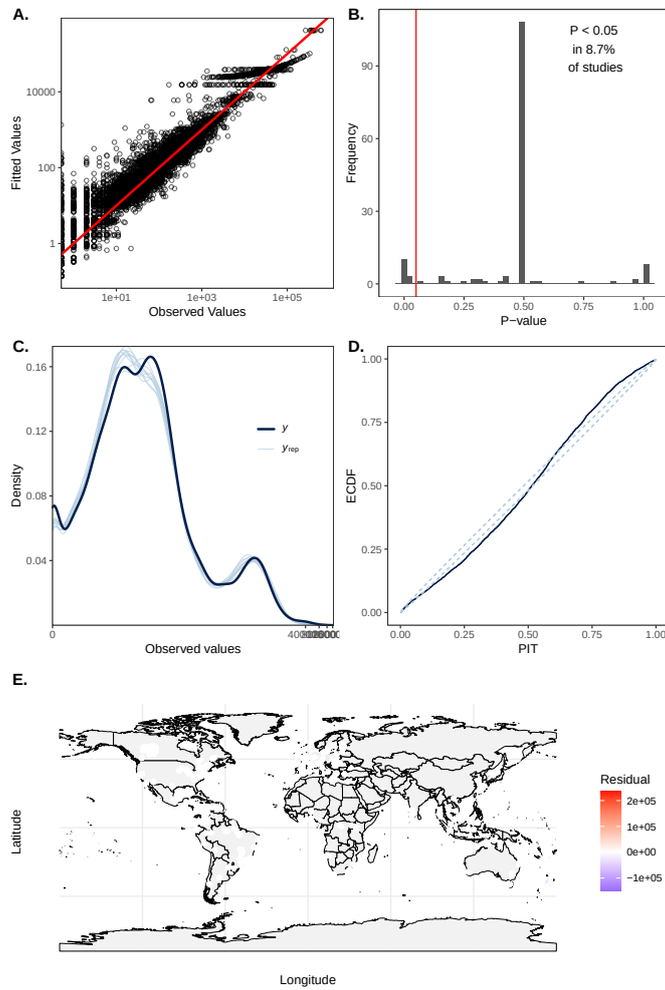


Figure M42: Model diagnostic plots for the zero-inflated negative binomial model of total abundance \sim LandUse \times Region + (1|SS/SSB). Model diagnostics show (a) fitted versus observed values, (b) spatial autocorrelation test results, (c) density plot of predictions and observed values, (d) ECDF of PIT values, and (e) spatial plot of model residuals. Map lines delineate study areas and do not necessarily depict accepted national boundaries.

Table M29: Posterior summary of the Gaussian model of $\log(\text{total abundance}+1)$ (Fig. M39).

| BRMS Summary | | | | | | | |
|---|----------|-----------|----------|----------|------|----------|----------|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: $\text{LogAbund} \sim \text{UI2} * \text{Region} + (1 \text{SS}/\text{SSB})$ | | | | | | | |
| Data: model_data (Number of observations: 6751) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 297) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 2.11 | 0.09 | 1.93 | 2.29 | 1.00 | 2233 | 4433 |
| ~SS:SSB (Number of levels: 1561) | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sd(Intercept) | 0.42 | 0.02 | 0.38 | 0.47 | 1.00 | 5133 | 9603 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 4.52 | 0.24 | 4.06 | 4.99 | 1.01 | 966 | 2347 |
| UI2Secondaryvegetation | 0.06 | 0.12 | -0.19 | 0.30 | 1.00 | 2496 | 5894 |
| UI2Agriculture_Minimal | 0.32 | 0.14 | 0.05 | 0.58 | 1.00 | 2570 | 5728 |
| UI2Agriculture_Light | -0.03 | 0.13 | -0.29 | 0.23 | 1.00 | 2325 | 4971 |
| UI2Agriculture_Intense | -0.01 | 0.14 | -0.28 | 0.26 | 1.00 | 2398 | 4976 |
| RegionNorthAmerica | 0.44 | 0.41 | -0.36 | 1.24 | 1.00 | 1343 | 3448 |
| RegionSouthandSoutheastAsia | 1.77 | 0.52 | 0.76 | 2.79 | 1.00 | 1781 | 4232 |
| RegionSouthandCentralAmerica | 0.97 | 0.34 | 0.32 | 1.64 | 1.00 | 1185 | 2812 |
| RegionSubsaharanAfrica | 0.65 | 0.54 | -0.43 | 1.72 | 1.00 | 1690 | 3616 |
| RegionAustralasia | 2.02 | 0.51 | 1.03 | 3.01 | 1.00 | 1869 | 3936 |
| UI2Secondaryvegetation:RegionNorthAmerica | -0.05 | 0.17 | -0.38 | 0.29 | 1.00 | 3651 | 7859 |
| UI2Agriculture_Minimal:RegionNorthAmerica | -0.50 | 0.23 | -0.96 | -0.04 | 1.00 | 4489 | 10349 |
| UI2Agriculture_Light:RegionNorthAmerica | -0.61 | 0.20 | -1.00 | -0.21 | 1.00 | 3960 | 8045 |
| UI2Agriculture_Intense:RegionNorthAmerica | -0.54 | 0.23 | -0.99 | -0.09 | 1.00 | 4577 | 9581 |
| UI2Secondaryvegetation:RegionSouthandSoutheastAsia | -0.56 | 0.46 | -1.45 | 0.34 | 1.00 | 14235 | 14128 |
| UI2Agriculture_Minimal:RegionSouthandSoutheastAsia | -0.57 | 0.28 | -1.13 | -0.01 | 1.00 | 7906 | 11350 |
| UI2Agriculture_Light:RegionSouthandSoutheastAsia | -0.91 | 0.17 | -1.24 | -0.57 | 1.00 | 3462 | 7930 |
| UI2Agriculture_Intense:RegionSouthandSoutheastAsia | -0.60 | 0.27 | -1.13 | -0.07 | 1.00 | 7160 | 12236 |
| UI2Secondaryvegetation:RegionSouthandCentralAmerica | -0.30 | 0.15 | -0.60 | -0.01 | 1.00 | 3134 | 7552 |
| UI2Agriculture_Minimal:RegionSouthandCentralAmerica | -0.53 | 0.16 | -0.84 | -0.21 | 1.00 | 3191 | 6550 |
| UI2Agriculture_Light:RegionSouthandCentralAmerica | -0.45 | 0.17 | -0.78 | -0.11 | 1.00 | 3014 | 6697 |
| UI2Agriculture_Intense:RegionSouthandCentralAmerica | -0.27 | 0.22 | -0.70 | 0.16 | 1.00 | 4503 | 9359 |
| UI2Secondaryvegetation:RegionSubsaharanAfrica | -0.52 | 0.21 | -0.92 | -0.11 | 1.00 | 5173 | 10073 |
| UI2Agriculture_Minimal:RegionSubsaharanAfrica | -1.31 | 0.20 | -1.70 | -0.93 | 1.00 | 4194 | 8757 |
| UI2Agriculture_Light:RegionSubsaharanAfrica | -0.83 | 0.21 | -1.24 | -0.43 | 1.00 | 4461 | 9178 |
| UI2Agriculture_Intense:RegionSubsaharanAfrica | -1.84 | 0.26 | -2.35 | -1.32 | 1.00 | 6165 | 10745 |
| UI2Secondaryvegetation:RegionAustralasia | -0.18 | 0.25 | -0.67 | 0.31 | 1.00 | 6298 | 11106 |
| UI2Agriculture_Minimal:RegionAustralasia | -1.04 | 0.17 | -1.38 | -0.70 | 1.00 | 3667 | 7811 |
| UI2Agriculture_Light:RegionAustralasia | -0.23 | 0.21 | -0.64 | 0.20 | 1.00 | 4588 | 9173 |
| UI2Agriculture_Intense:RegionAustralasia | -0.20 | 0.18 | -0.55 | 0.16 | 1.00 | 3443 | 7870 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| sigma | 0.81 | 0.01 | 0.80 | 0.83 | 1.00 | 14469 | 15031 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.910 (95% CI [0.908, 0.912]) | | | | | | | |
| Marginal R2: 0.053 (95% CI [0.023, 0.086]) | | | | | | | |

Table M30: Posterior summary of the Gaussian model of $\log(\text{rescaled total abundance}+0.01)$ (Fig. M40).

| BRMS Summary | | | | | | | |
|--|--|--|--|--|--|--|--|
| Family: gaussian | | | | | | | |
| Links: mu = identity | | | | | | | |
| Formula: $\text{LogAbund}_{\text{RS}} \sim \text{UI2} * \text{Region} + (1 \text{SS}/\text{SSB})$ | | | | | | | |
| Data: model_data (Number of observations: 6751) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |

```

~SS (Number of levels: 297)
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sd(Intercept)    0.68    0.04    0.61    0.76 1.00    5181    9106

~SS:SSB (Number of levels: 1561)
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sd(Intercept)    0.35    0.02    0.31    0.40 1.00    5112    9455

Regression Coefficients:
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
Intercept                -1.18    0.13    -1.45    -0.92 1.00    2258    3108
UI2Secondaryvegetation    0.02    0.12    -0.22    0.27 1.00    2563    5095
UI2Agriculture_Minimal    0.19    0.13    -0.07    0.46 1.00    2434    4905
UI2Agriculture_Light    -0.11    0.13    -0.36    0.14 1.00    2262    4034
UI2Agriculture_Intense   -0.03    0.13    -0.29    0.23 1.00    2309    4754
RegionNorthAmerica        0.24    0.19    -0.13    0.62 1.00    3095    5839
RegionSouthandSoutheastAsia 0.50    0.23    0.05    0.95 1.00    3685    6731
RegionSouthandCentralAmerica 0.48    0.17    0.14    0.82 1.00    2797    4730
RegionSubsaharanAfrica    0.24    0.23    -0.21    0.69 1.00    3778    6672
RegionAustralasia         0.27    0.23    -0.20    0.72 1.00    3278    6084
UI2Secondaryvegetation:RegionNorthAmerica -0.00    0.17    -0.33    0.33 1.00    3634    7799
UI2Agriculture_Minimal:RegionNorthAmerica -0.23    0.23    -0.68    0.23 1.00    4786    9142
UI2Agriculture_Light:RegionNorthAmerica  -0.42    0.20    -0.81    -0.03 1.00    3715    8045
UI2Agriculture_Intense:RegionNorthAmerica -0.41    0.22    -0.85    0.03 1.00    4336    8467
UI2Secondaryvegetation:RegionSouthandSoutheastAsia -0.28    0.46    -1.18    0.61 1.00    13505   14160
UI2Agriculture_Minimal:RegionSouthandSoutheastAsia -0.40    0.29    -0.97    0.17 1.00    6874    10764
UI2Agriculture_Light:RegionSouthandSoutheastAsia -0.79    0.17    -1.11    -0.46 1.00    3361    7216
UI2Agriculture_Intense:RegionSouthandSoutheastAsia -0.53    0.28    -1.08    0.02 1.00    6186    11951
UI2Secondaryvegetation:RegionSouthandCentralAmerica -0.28    0.15    -0.58    0.02 1.00    3229    6762
UI2Agriculture_Minimal:RegionSouthandCentralAmerica -0.50    0.16    -0.81    -0.18 1.00    3013    6357
UI2Agriculture_Light:RegionSouthandCentralAmerica -0.35    0.16    -0.68    -0.03 1.00    3048    5583
UI2Agriculture_Intense:RegionSouthandCentralAmerica -0.19    0.22    -0.62    0.24 1.00    4697    10107
UI2Secondaryvegetation:RegionSubsaharanAfrica -0.43    0.21    -0.85    -0.02 1.00    5188    9537
UI2Agriculture_Minimal:RegionSubsaharanAfrica -1.26    0.20    -1.65    -0.87 1.00    4095    9015
UI2Agriculture_Light:RegionSubsaharanAfrica  -0.69    0.21    -1.09    -0.28 1.00    4415    8407
UI2Agriculture_Intense:RegionSubsaharanAfrica -1.62    0.27    -2.14    -1.10 1.00    6054    10261
UI2Secondaryvegetation:RegionAustralasia -0.13    0.24    -0.60    0.33 1.00    5677    10170
UI2Agriculture_Minimal:RegionAustralasia -0.89    0.18    -1.24    -0.55 1.00    3557    7447
UI2Agriculture_Light:RegionAustralasia -0.34    0.22    -0.77    0.08 1.00    4575    7755
UI2Agriculture_Intense:RegionAustralasia -0.26    0.18    -0.61    0.09 1.00    3559    8249

Further Distributional Parameters:
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sigma    0.86    0.01    0.85    0.88 1.00    14528   15033

Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
and Tail_ESS are effective sample size measures, and Rhat is the potential
scale reduction factor on split chains (at convergence, Rhat = 1).

# Bayesian R2 with Compatibility Interval

Conditional R2: 0.501 (95% CI [0.487, 0.515])
Marginal R2: 0.049 (95% CI [0.035, 0.066])

```

Table M31: Posterior summary of the zero-and-one-inflated beta model of rescaled total abundance (Fig. M41).

```

----- BRMS Summary -----
Family: zero_one_inflated_beta
Links: mu = logit
Formula: Total_abundance_RS ~ UI2 * Region + (1 | SS/SSB)
Data: model_data (Number of observations: 6751)
Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1;
      total post-warmup draws = 20000

Multilevel Hyperparameters:
~SS (Number of levels: 297)
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sd(Intercept)    0.75    0.04    0.67    0.84 1.00    4549    9393

~SS:SSB (Number of levels: 1561)
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
sd(Intercept)    0.27    0.03    0.22    0.32 1.00    3996    7372

Regression Coefficients:
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
Intercept                -0.47    0.13    -0.73    -0.21 1.00    1688    3468
UI2Secondaryvegetation    0.11    0.12    -0.13    0.35 1.00    2066    5330
UI2Agriculture_Minimal    0.32    0.13    0.06    0.59 1.00    1922    4727
UI2Agriculture_Light    -0.07    0.13    -0.32    0.18 1.00    1710    4437

```

| | | | | | | | |
|---|-----------|----------|----------|-------|----------|----------|-------|
| UI2Agriculture_Intense | -0.05 | 0.13 | -0.31 | 0.22 | 1.00 | 1897 | 4840 |
| RegionNorthAmerica | 0.37 | 0.20 | -0.02 | 0.75 | 1.00 | 2349 | 4799 |
| RegionSouthandSoutheastAsia | 0.48 | 0.24 | 0.02 | 0.95 | 1.00 | 3261 | 7079 |
| RegionSouthandCentralAmerica | 0.37 | 0.18 | 0.02 | 0.72 | 1.00 | 2206 | 4742 |
| RegionSubsaharanAfrica | 0.32 | 0.24 | -0.15 | 0.81 | 1.00 | 3285 | 6285 |
| RegionAustralasia | 0.11 | 0.24 | -0.37 | 0.59 | 1.00 | 2518 | 5593 |
| UI2Secondaryvegetation:RegionNorthAmerica | -0.35 | 0.17 | -0.68 | -0.02 | 1.00 | 2903 | 6589 |
| UI2Agriculture_Minimal:RegionNorthAmerica | -0.50 | 0.23 | -0.94 | -0.05 | 1.00 | 3311 | 7741 |
| UI2Agriculture_Light:RegionNorthAmerica | -0.61 | 0.20 | -1.00 | -0.22 | 1.00 | 3093 | 6468 |
| UI2Agriculture_Intense:RegionNorthAmerica | -0.70 | 0.22 | -1.15 | -0.27 | 1.00 | 3553 | 8131 |
| UI2Secondaryvegetation:RegionSouthandSoutheastAsia | -0.36 | 0.61 | -1.54 | 0.84 | 1.00 | 12235 | 13832 |
| UI2Agriculture_Minimal:RegionSouthandSoutheastAsia | -0.55 | 0.29 | -1.12 | 0.01 | 1.00 | 5523 | 12328 |
| UI2Agriculture_Light:RegionSouthandSoutheastAsia | -0.70 | 0.17 | -1.02 | -0.38 | 1.00 | 2749 | 7334 |
| UI2Agriculture_Intense:RegionSouthandSoutheastAsia | -0.39 | 0.28 | -0.95 | 0.15 | 1.00 | 5791 | 11955 |
| UI2Secondaryvegetation:RegionSouthandCentralAmerica | -0.34 | 0.16 | -0.65 | -0.04 | 1.00 | 2633 | 6429 |
| UI2Agriculture_Minimal:RegionSouthandCentralAmerica | -0.44 | 0.17 | -0.77 | -0.11 | 1.00 | 2558 | 6418 |
| UI2Agriculture_Light:RegionSouthandCentralAmerica | -0.25 | 0.17 | -0.58 | 0.08 | 1.00 | 2382 | 6235 |
| UI2Agriculture_Intense:RegionSouthandCentralAmerica | 0.04 | 0.22 | -0.39 | 0.46 | 1.00 | 3855 | 8212 |
| UI2Secondaryvegetation:RegionSubsaharanAfrica | -0.21 | 0.21 | -0.62 | 0.19 | 1.00 | 4585 | 11349 |
| UI2Agriculture_Minimal:RegionSubsaharanAfrica | -0.83 | 0.22 | -1.26 | -0.41 | 1.00 | 3989 | 8197 |
| UI2Agriculture_Light:RegionSubsaharanAfrica | -0.48 | 0.22 | -0.92 | -0.06 | 1.00 | 3917 | 8852 |
| UI2Agriculture_Intense:RegionSubsaharanAfrica | -1.33 | 0.28 | -1.89 | -0.80 | 1.00 | 5693 | 11814 |
| UI2Secondaryvegetation:RegionAustralasia | -0.17 | 0.24 | -0.66 | 0.31 | 1.00 | 4408 | 9969 |
| UI2Agriculture_Minimal:RegionAustralasia | -1.01 | 0.18 | -1.36 | -0.67 | 1.00 | 3076 | 7756 |
| UI2Agriculture_Light:RegionAustralasia | -0.11 | 0.23 | -0.55 | 0.34 | 1.00 | 3946 | 10268 |
| UI2Agriculture_Intense:RegionAustralasia | -0.10 | 0.19 | -0.47 | 0.27 | 1.00 | 2968 | 8007 |
| Further Distributional Parameters: | | | | | | | |
| Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS | |
| phi | 5.30 | 0.10 | 5.10 | 5.51 | 1.00 | 12121 | 13779 |
| zoi | 0.10 | 0.00 | 0.10 | 0.11 | 1.00 | 28769 | 13844 |
| coi | 0.44 | 0.02 | 0.40 | 0.47 | 1.00 | 29953 | 14009 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| Conditional R2: 0.276 (95% CI [0.261, 0.291]) | | | | | | | |
| Marginal R2: 0.037 (95% CI [0.024, 0.053]) | | | | | | | |

Table M32: Posterior summary of the zero-inflated negative binomial model of total abundance (Fig. M42).

| BRMS Summary | | | | | | | |
|---|-----------|-----------|----------|----------|----------|----------|----------|
| Family: zero_inflated_negbinomial | | | | | | | |
| Links: mu = log; zi = logit | | | | | | | |
| Formula: Total_abundance ~ UI2 * Region + (1 SS/SSB) | | | | | | | |
| zi ~ 1 | | | | | | | |
| Data: model_data (Number of observations: 6751) | | | | | | | |
| Draws: 4 chains, each with iter = 10000; warmup = 5000; thin = 1; | | | | | | | |
| total post-warmup draws = 20000 | | | | | | | |
| Multilevel Hyperparameters: | | | | | | | |
| ~SS (Number of levels: 297) | | | | | | | |
| Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS | |
| sd(Intercept) | 2.22 | 0.10 | 2.03 | 2.42 | 1.00 | 4298 | 8353 |
| ~SS:SSB (Number of levels: 1561) | | | | | | | |
| Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS | |
| sd(Intercept) | 0.46 | 0.02 | 0.41 | 0.50 | 1.00 | 5834 | 11529 |
| Regression Coefficients: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| Intercept | 4.69 | 0.24 | 4.22 | 5.16 | 1.00 | 2112 | 4289 |
| zi_Intercept | -5.01 | 0.20 | -5.44 | -4.65 | 1.00 | 42844 | 14598 |
| UI2Secondaryvegetation | 0.05 | 0.11 | -0.17 | 0.26 | 1.00 | 5945 | 10031 |
| UI2Agriculture_Minimal | 0.36 | 0.12 | 0.12 | 0.61 | 1.00 | 5853 | 10513 |
| UI2Agriculture_Light | -0.02 | 0.12 | -0.26 | 0.21 | 1.00 | 5263 | 9737 |
| UI2Agriculture_Intense | 0.03 | 0.13 | -0.23 | 0.28 | 1.00 | 5501 | 9538 |
| RegionNorthAmerica | 0.43 | 0.43 | -0.42 | 1.28 | 1.00 | 2397 | 4680 |
| RegionSouthandSoutheastAsia | 1.81 | 0.54 | 0.73 | 2.86 | 1.00 | 3775 | 7503 |
| RegionSouthandCentralAmerica | 0.92 | 0.35 | 0.23 | 1.63 | 1.00 | 2409 | 4666 |
| RegionSubsaharanAfrica | 0.39 | 0.57 | -0.70 | 1.51 | 1.00 | 3323 | 6706 |
| RegionAustralasia | 2.18 | 0.53 | 1.13 | 3.23 | 1.00 | 3323 | 6724 |
| UI2Secondaryvegetation:RegionNorthAmerica | -0.15 | 0.15 | -0.44 | 0.15 | 1.00 | 8749 | 12652 |
| UI2Agriculture_Minimal:RegionNorthAmerica | -0.42 | 0.23 | -0.87 | 0.03 | 1.00 | 10964 | 14597 |
| UI2Agriculture_Light:RegionNorthAmerica | -0.29 | 0.20 | -0.69 | 0.11 | 1.00 | 9436 | 14590 |
| UI2Agriculture_Intense:RegionNorthAmerica | -0.29 | 0.23 | -0.75 | 0.16 | 1.00 | 10962 | 14894 |

| | | | | | | | |
|---|--|-----------|----------|----------|------|----------|----------|
| UI2Secondaryvegetation:RegionSouthandSoutheastAsia | -0.44 | 0.43 | -1.26 | 0.42 | 1.00 | 30970 | 14724 |
| UI2Agriculture_Minimal:RegionSouthandSoutheastAsia | -0.60 | 0.25 | -1.10 | -0.10 | 1.00 | 14664 | 14294 |
| UI2Agriculture_Light:RegionSouthandSoutheastAsia | -0.79 | 0.16 | -1.10 | -0.48 | 1.00 | 7728 | 13338 |
| UI2Agriculture_Intense:RegionSouthandSoutheastAsia | -0.60 | 0.25 | -1.08 | -0.11 | 1.00 | 12850 | 14578 |
| UI2Secondaryvegetation:RegionSouthandCentralAmerica | -0.21 | 0.14 | -0.47 | 0.06 | 1.00 | 7210 | 12627 |
| UI2Agriculture_Minimal:RegionSouthandCentralAmerica | -0.42 | 0.15 | -0.72 | -0.12 | 1.00 | 6972 | 11225 |
| UI2Agriculture_Light:RegionSouthandCentralAmerica | -0.25 | 0.15 | -0.55 | 0.06 | 1.00 | 6940 | 12150 |
| UI2Agriculture_Intense:RegionSouthandCentralAmerica | -0.21 | 0.20 | -0.61 | 0.18 | 1.00 | 9453 | 13869 |
| UI2Secondaryvegetation:RegionSubsaharanAfrica | -0.39 | 0.20 | -0.77 | -0.00 | 1.00 | 11222 | 14507 |
| UI2Agriculture_Minimal:RegionSubsaharanAfrica | -1.06 | 0.20 | -1.44 | -0.67 | 1.00 | 9881 | 14039 |
| UI2Agriculture_Light:RegionSubsaharanAfrica | -0.51 | 0.20 | -0.91 | -0.11 | 1.00 | 9719 | 14239 |
| UI2Agriculture_Intense:RegionSubsaharanAfrica | -1.52 | 0.25 | -2.00 | -1.03 | 1.00 | 11939 | 14456 |
| UI2Secondaryvegetation:RegionAustralasia | -0.25 | 0.26 | -0.76 | 0.28 | 1.00 | 17905 | 14256 |
| UI2Agriculture_Minimal:RegionAustralasia | -1.06 | 0.16 | -1.37 | -0.74 | 1.00 | 8043 | 12915 |
| UI2Agriculture_Light:RegionAustralasia | -0.25 | 0.21 | -0.65 | 0.15 | 1.00 | 10260 | 14579 |
| UI2Agriculture_Intense:RegionAustralasia | -0.29 | 0.18 | -0.64 | 0.05 | 1.00 | 7916 | 12310 |
| Further Distributional Parameters: | | | | | | | |
| | Estimate | Est.Error | 1-95% CI | u-95% CI | Rhat | Bulk_ESS | Tail_ESS |
| shape | 2.03 | 0.04 | 1.94 | 2.11 | 1.00 | 19526 | 15571 |
| Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS and Tail_ESS are effective sample size measures, and Rhat is the potential scale reduction factor on split chains (at convergence, Rhat = 1). | | | | | | | |
| # Bayesian R2 with Compatibility Interval | | | | | | | |
| | Conditional R2: 0.790 (95% CI [0.673, 0.839]) | | | | | | |
| | Marginal R2: 0.000 (95% CI [7.755e-06, 4.241e-04]) | | | | | | |