

# Land use has global and systematic effects on local zoonotic host communities

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## Main text

Changes to biodiversity under human land use (e.g. agriculture, urbanisation) are increasingly recognised to influence zoonotic disease risk and emergence in humans<sup>1,2</sup>, but the underlying ecological processes that interact to drive infection risk are not well understood. In particular, it has been widely hypothesised that systematic differences in species' resilience to anthropogenic impacts, linked to traits, life-histories and phylogeny, might result in habitat disturbance causing predictable changes in the diversity and species composition of potential zoonotic hosts<sup>3,4</sup>. Here, we analyse 6801 ecological assemblages and 376 host species worldwide to show that land use has global and systematic effects on local zoonotic host communities. Controlling for research effort and survey methods, known wildlife hosts of human-shared pathogens and parasites comprise a significantly greater proportion of site-level species richness (18%–72% increase) and total abundance (21%–144% increase) in intensively-used secondary, human-managed and urban ecosystems than in nearby undisturbed habitats. This effect varies in magnitude among mammalian and avian taxonomic orders. In particular, rodent, bat and passerine bird zoonotic host species increasingly dominate disturbed assemblages, which may be one factor underpinning the global importance of these taxa as zoonotic disease reservoirs. Crucially, we further show that mammal species that harbour more pathogens overall (either human-shared or non-human shared) are more likely to occur in managed ecosystems, suggesting that our overall results may be mediated by ecological or life-history traits that influence both host status and human-tolerance<sup>5,6</sup>. Our results suggest that global changes in mode and intensity of land use are creating growing interfaces between people, livestock and wildlife reservoirs of zoonotic disease, and highlight the need to prioritise wildlife and human disease surveillance in agriculturally intensifying and urbanising habitats.

Anthropogenic environmental change impacts many dimensions of human health and wellbeing, including the incidence and emergence of zoonotic and vector-borne diseases<sup>1</sup>. Although large-scale research into environmental drivers of disease has mostly focused on climate, there is growing consensus that land use change (conversion of natural habitats to agricultural, urban or otherwise anthropogenic ecosystems) is a globally-significant mediator of human infection risk and disease emergence<sup>2,3</sup>. Land use change directly and indirectly

drives biodiversity loss, turnover and homogenisation (including through biological invasions and rare species losses)<sup>7,8</sup>, modifies landscape structure in ways that modulate epidemiological processes (e.g. fragmentation<sup>9</sup>, resource provisioning<sup>10</sup>) and can increase human-wildlife contact (e.g. via agricultural practices or hunting)<sup>1</sup>. These processes interact to influence transmission dynamics in reservoir and vector communities and ultimately spillover risk to humans<sup>11,12</sup>, with land use change implicated in driving both endemic (e.g. trypanosomiasis<sup>13</sup>, malaria<sup>14</sup>) and epidemic (e.g. Nipah<sup>15</sup>, West Nile<sup>16</sup>) zoonoses. However, the complexity of these systems (Extended Data 1) has made it difficult to identify whether land use has consistent effects on the ecological factors underpinning zoonotic disease risk<sup>2</sup>, a critical knowledge gap given the extensive global land change anticipated this century<sup>17</sup>.

Although there is broad evidence for regulatory effects of local species diversity on pathogen transmission<sup>18</sup>, such effects are not universal: higher disease risk in depauperate assemblages has been observed for some disease systems (e.g. *Borrelia*<sup>19</sup>, West Nile<sup>16</sup>, *Ribeiroia*<sup>6</sup>) but not others. One ecological factor underlying these inconsistencies may be differences in host species sensitivity to human pressures<sup>4</sup>. It is often proposed that more effective zoonotic host species might be generally more likely to persist in disturbed ecosystems, since certain trait profiles (e.g. ‘fast’ life-histories, higher population densities) correlate to both reservoir status and reduced extirpation risk in several vertebrate taxa<sup>20,21</sup>. Alternatively, any such tendencies might be taxonomically or geographically idiosyncratic: for example, mammals that are more closely phylogenetically-related to humans are more likely to be zoonotic reservoirs<sup>22</sup>, but may also be highly variable in their sensitivity to human impacts<sup>20</sup>. Reservoir host responses to disturbance have been investigated in certain taxa (e.g. primates<sup>23</sup>) and disease systems<sup>13,19</sup>, but to date there has been no comprehensive analysis of the effects of land use on zoonotic host diversity and species composition.

Here, we use a global dataset of 6801 ecological assemblages derived from the Projecting Responses of Ecological Diversity in Changing Terrestrial Systems (PREDICTS) biodiversity database<sup>24</sup>, to test whether land use has systematic effects on the zoonotic potential of local wildlife communities. We identified records of wildlife hosts of known human pathogens and endoparasites (henceforth ‘*pathogens*’) within PREDICTS using a comprehensive host-pathogen associations database, and classified species as zoonotic hosts (henceforth ‘*hosts*’) based on evidence of association with at least one human-shared pathogen (Methods). PREDICTS compiles >3.2 million species records from 666 studies that sampled biodiversity across land use gradients, enabling global comparisons of local assemblages in primary vegetation (minimally-disturbed baseline) to nearby secondary

(recovering from past disturbance), managed (cropland, pasture, plantation) and urban sites, of varying use intensities (here, minimal or substantial-use), sampled using the same protocols<sup>24</sup>. We identified records of 376 host species in a dataset of 6801 survey sites from 184 studies across 6 continents, with a taxonomic distribution broadly representative of known zoonotic host diversity (Figure 1, Supp. Tables 1-2; Methods). We compared host responses to land use to those of all other species at the same locations (‘non-hosts’, approximating the response of background biodiversity; n=6512 species), using Bayesian mixed-effects models to control for study methods and sampling design (Methods). Pathogen detection is sensitive to research effort, such that some poorly studied species might be misclassified as non-hosts. We account for this uncertainty in our models using a bootstrap approach, with each iteration transitioning a proportion of non-host species to host status, with species-level transition rates determined by both publication effort and taxonomic order (Supp. Text 1, Extended Data 2). All parameter estimates are obtained across each full bootstrap ensemble (Methods).

We first estimated the effects of land use type and intensity on two community metrics: site-level host species richness (number of host species; related to potential pathogen richness) and host total abundance (total number of host individuals; a more epidemiologically-relevant metric related to opportunities for transmission)<sup>25</sup>. Both host richness and total abundance either persist or increase in response to land use, against a background of consistent declines in all other (non-host) species in human-dominated habitats (Figure 2a-b). Together these changes lead to hosts comprising an increasing proportion of ecological assemblages in secondary, managed and urban land (Figure 2c-d, Supp. Tables 3-5). Notably, land use intensity has clear positive effects on community zoonotic potential both within and between land use types, with largest increases in substantial-use secondary and managed (posterior median: +18-21% host proportion richness, +21-26% proportion abundance) and urban sites (+62-72% proportion richness, +136-144% proportion abundance; but with higher uncertainty due to sparser sampling). These results are robust to testing for sensitivity to random study-level variability (Extended Data 3a), geographical biases in data coverage<sup>24</sup> (Extended Data 3b), and strictness of host status definition (Extended Data 4). The last of these is crucial to understanding disease risk, since species capable of being infected by a given pathogen may not contribute substantially to transmission dynamics or zoonotic spillover risk. We therefore repeated analyses for mammals only (the major reservoirs of zoonoses globally) with reservoir status strictly-defined as an association with at least one zoonotic agent (aetiologic agent of a specific

human disease with a known animal reservoir), based on pathogen detection, isolation or confirmed reservoir status (143 host species, 2026 sites, 63 studies). Overall trends remain consistent, although with notably stronger effects on host proportion of total abundance (+42-52% in secondary and managed land), and weaker effects on host richness that may reflect underlying variability in responses between mammal taxa (Extended Data 4).

To examine the possibility of such taxonomic variability in host responses to land use, we separately analysed mean land use effects on species-level occurrence and abundance of zoonotic host (strictly-defined) and non-host species, for several mammalian (Carnivora, Cetartiodactyla, Chiroptera, Primates, Rodentia) and avian orders (Passeriformes, Psittaciformes) that are well-sampled in PREDICTS and harbour the majority of known zoonoses (Methods). We again used bootstrapping to account for host status uncertainty, and predicted abundance change using a hurdle model-based approach to account for zero-inflation (combining separately-fitted occurrence and zero-truncated abundance models; Extended Data 5). Within most orders, non-host species tend to decline more strongly in response to land disturbance than host species, but with substantial between-order variation in the direction and clarity of effects (Figure 3, Supp. Table 6). Notably, within passerine birds, bats and rodents, hosts and non-hosts show clear divergent responses to land use, with host species abundances on average increasing (+14-96% Passeriformes, +45% Chiroptera, +52% Rodentia) while non-host abundances decline (-28-43% Passeriformes, -13% Chiroptera, -53% Rodentia) in human-dominated relative to primary sites (Figure 3). Although such a tendency has been observed in some disease systems, our results suggest this is a more general phenomenon in these taxa, which may contribute to numerous documented links between anthropogenic ecosystems and bat-, rodent- and bird-borne emerging infections (e.g. corona-, henipa-, arena- and flaviviruses, *Borrelia* spp.)<sup>15,16,19</sup>. In contrast, primate and carnivore host responses are not clearly distinguishable from overall species declines in these orders, consistent with past studies that found no consistent links between land disturbance and disease in primates<sup>23</sup> and highlighting the importance of ecotonal or edge habitats as human-primate epidemiological interfaces<sup>14</sup> (although sparser urban sampling means that the urban adaptations of certain primates, such as macaques, are likely underrepresented).

The differing responses of host and non-host species may be mediated by covariance between traits influencing both host status and human-tolerance<sup>26</sup>, but could also reflect histories of human-wildlife contact and coevolution of shared pathogens<sup>11</sup>. If the former is the case we hypothesise that harbouring a higher number of pathogens overall (richness of either zoonotic or non-zoonotic pathogens; a metric often correlated to species traits<sup>27</sup>), would be

associated with more positive species responses to land use. We tested this across all mammals in our dataset (due to more complete pathogen data availability than for other taxa; 546 species, 1950 sites), here controlling for species-level differences in research effort by analysing residual pathogen richness not explained by publication effort (Methods, Extended Data 6). We find that increasing pathogen richness positively predicts species' probability of occurrence in managed but not primary sites, a result that is consistent for either human-shared or non-human-shared pathogens (no documented infection of either people or domestic animals; Extended Data 7, [Supp. Table 7](#)). This suggests that the net increase in zoonotic host diversity in disturbed sites is at least partly trait-mediated: in particular, species traits associated with a faster pace-of-life are often correlated both with reservoir status and infection outcomes<sup>5,26</sup> (potentially owing to life-history trade-offs between reproductive rate and immune investment<sup>28</sup>) and with population resilience to anthropogenic pressures<sup>20</sup>. [A trait-mediated explanation is also supported by our finding that differential host and non-host species responses to land use are most clearly detected when comparing across large clades with a wide diversity of life-histories, such as rodents, passerines and, notably, mammals overall \(Extended Data 5\). In contrast, generally longer-lived, large-bodied clades \(e.g. primates, carnivores\) show more idiosyncratic or negative responses to landscape disturbance \(Figure 3\).](#)

Overall, our results indicate that the homogenising impacts of land use on biodiversity globally<sup>8</sup> have produced systematic changes to local zoonotic host communities, which may be one factor underpinning links between human-disturbed ecosystems and disease emergence. By leveraging site-level survey data, our analyses reflect community changes at the epidemiologically-relevant local landscape scale<sup>21</sup>, negating the need to ignore community interactions or generalise ecological processes to coarser spatial scales (a typical limitation of global studies [that can confound or mask biodiversity-disease relationships](#)<sup>29</sup>). Our results reflect zoonotic potential, since proximity to reservoir hosts is not necessarily sufficient for zoonotic spillover<sup>30</sup>, and emergent disease risk will depend on contextual factors (e.g. pathogen prevalence, intermediate host/vector populations, landscape structure, human socioeconomics) that may synergistically or antagonistically affect contact and transmission dynamics<sup>11</sup>. Nonetheless, land use also predictably impacts other factors that can amplify within- and cross-species transmission<sup>31</sup> (e.g. resource provisioning<sup>10</sup>, vector diversity<sup>32</sup>), and increases potential for human-wildlife contact<sup>12</sup>: for example, human populations are consistently higher at disturbed sites in our dataset (Extended Data 8). Global expansions of agricultural and urban land forecast for the coming decades, predominantly in

low- and middle-income countries<sup>17</sup>, thus have the potential to create growing risk interfaces for zoonotic pathogen exposure. In particular, the large effect sizes but sparser data availability for urban ecosystems (especially for mammals; Extended Data 4) highlight a key knowledge gap for anticipating urbanisation effects on public health and biodiversity. Our findings therefore strongly support calls to enhance proactive human and animal surveillance within agricultural, pastoral and urbanising ecosystems<sup>33,34</sup>, and highlight the need to incorporate disease-related costs into future land use and conservation planning.

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## End notes

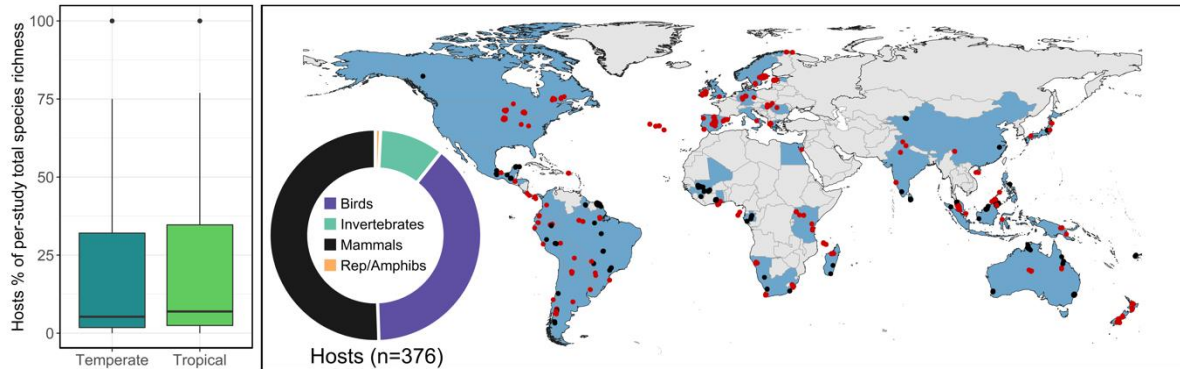
**Supplementary Information** is available in the accompanying document.

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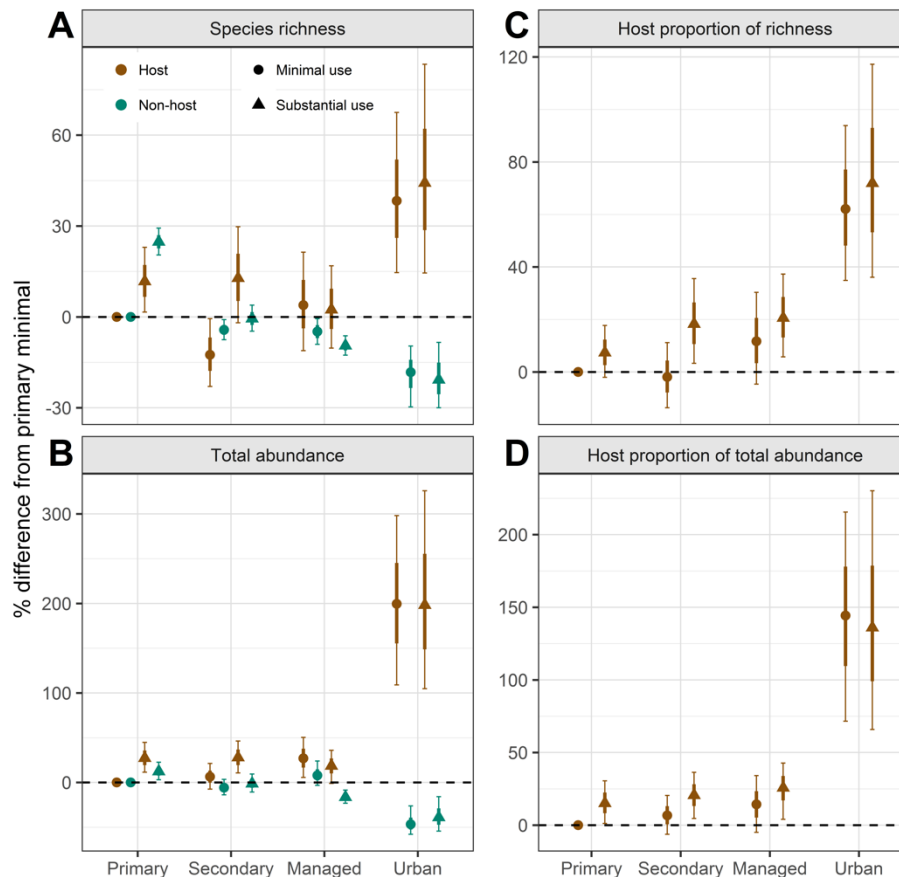
**Author contributions:** RG, DWR, KEJ and TN conceived and designed the study, [CAD contributed to the design of statistical analyses](#), RG collated and processed the data and led and conducted the analyses with DWR, KQC and TN, and all authors contributed to writing the manuscript.

**Author information:** The authors declare no competing interests.

## Figures

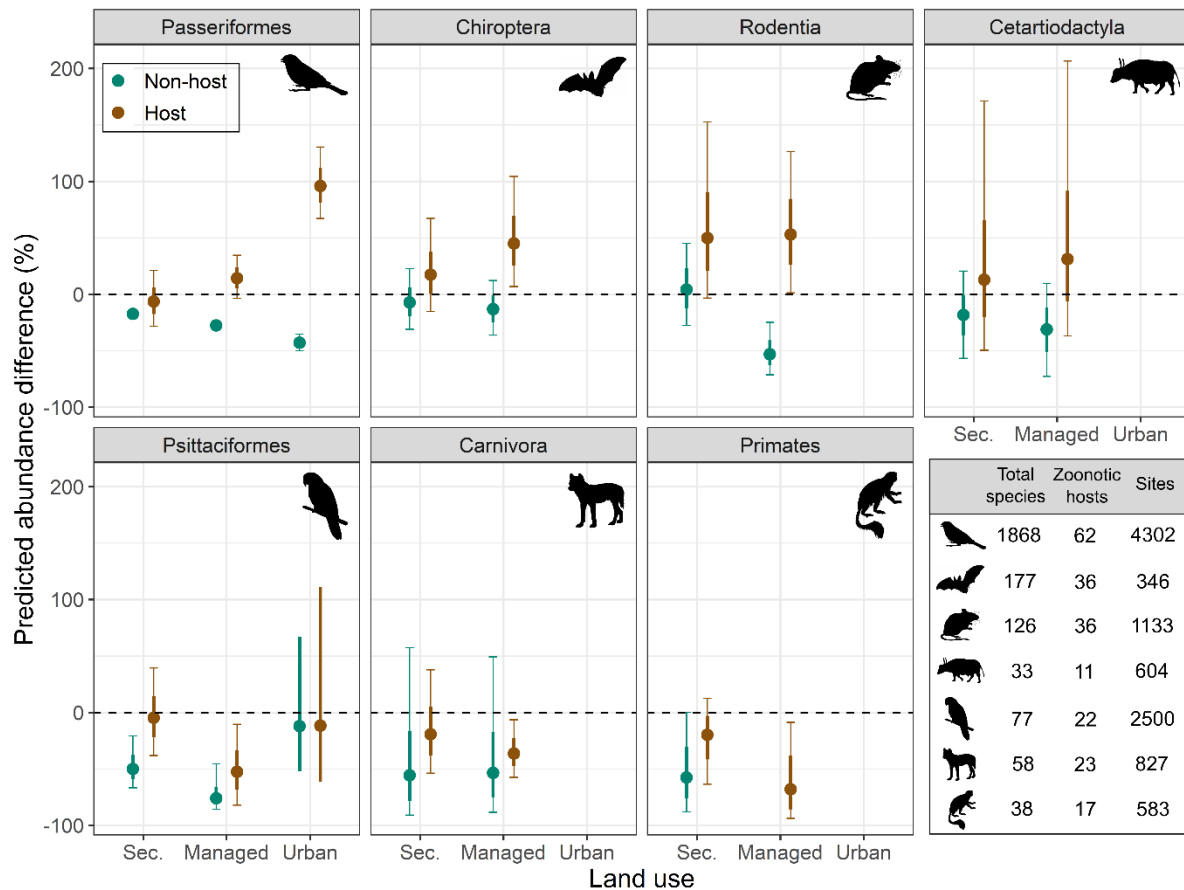


**Figure 1: Combined ecological communities and zoonotic host species dataset.** Points show the geographical locations of surveyed assemblages (n=6801 sites), with mammal survey locations in black and all other sites in red, and countries containing sites shaded in blue. Inset chart shows the taxonomic distribution of hosts of human-shared pathogens (birds, invertebrates, mammals, reptiles and amphibians; see Methods). Boxplot summarises, across all studies (n=184), host species richness as a percentage of the total per-study sampled richness, split across temperate and tropical biomes.



**Figure 2: Effects of land use on site-level host species richness and total abundance.**

Points, wide and narrow error bars show modelled percentage difference in diversity metrics (posterior marginal median, 67% and 95% quantile ranges respectively, across 1000 bootstrap models) relative to a baseline of primary land under minimal use (dashed line) (n=6801 sites: primary (1423 and 1457 for minimal and substantial use, respectively), secondary (1044, 629), managed (565, 1314), urban (136, 233)). Models are of species richness (A) and total abundance (B) of host species and of all other (non-host) species, and of hosts as a proportion of total site-level richness and abundance (C-D). Point shape denotes land use intensity (minimal or substantial) and colour denotes host (brown) or non-host (green). All posterior estimates were calculated across an ensemble of 1000 bootstrapped models, each with a proportion of non-hosts probabilistically transitioned to host status (median 121, range 90–150; Extended Data 2) to account for variability in species-level research effort (Methods, Supp. Text 1). Models also included fixed effects for human population density and random effects for study methods and biome (Methods). Parameter estimates represent averaged effect sizes across multiple studies with differing survey methods and taxonomic focus, so do not have an absolute numerical interpretation.



**Figure 3: Effects of land use on species abundance of mammalian and avian zoonotic hosts and non-hosts.** Points, wide and narrow error bars show average difference in species abundance (posterior median, 67% and 95% quantile ranges respectively, across 500 bootstrap models) in secondary (Sec.), managed and urban sites relative to a primary land baseline (dashed line), across all host (brown) and non-host (green) species in each mammalian or avian order. For mammals, zoonotic host status was defined strictly (direct pathogen detection, isolation or confirmed reservoir status), and urban sites were excluded owing to sparse urban sampling (only 2 studies; additionally, no non-host primates were recorded in managed land, and urban 95% quantile range for Psittaciformes is not shown due to high uncertainty). Abundance differences were predicted using a hurdle model approach (by combining estimates from separately-fitted occurrence and zero-truncated abundance models; see Extended Data 5, Methods). The inset table show per-order numbers of species in the dataset (between 8% and 35% of total described species in each order), known zoonotic hosts (prior to bootstrap), and sampled sites.

## Methods

We combined a global database of ecological assemblages (Projecting Responses of Ecological Diversity In Changing Terrestrial Systems, PREDICTS)<sup>24</sup> with data on host-pathogen and host-parasite associations, to create a global, spatially-explicit dataset of local zoonotic host diversity. We define pathogens and parasites (henceforth '*pathogens*') as including bacteria, viruses, protozoa, helminths and fungi (excluding ectoparasites). PREDICTS contains species records compiled from 666 published studies that sampled local biodiversity across land use type and intensity gradients, allowing global space-for-time analysis of land use effects on local species assemblages (i.e. comparison between sites with natural vegetation considered to be a baseline). We analysed relative differences in wildlife host community metrics (zoonotic host species richness and abundance) between undisturbed (primary) land and nearby sites under varying degrees of anthropogenic disturbance. We subsequently conducted further analyses to examine how host species responses to land use vary across different mammalian and avian orders, and to test whether mammal pathogen richness (including both human and non-human pathogens) covaries with tolerance to land use.

## Datasets

*Ecological community and land use data.* Each of the >3.2 million records in PREDICTS is a per-species, per-site measure of either occurrence (including absences) or abundance, alongside metadata on site location, land use type and use intensity. The database provides as representative a sample as possible of local biodiversity responses to human pressure, containing 47,000 species in a taxonomic distribution broadly proportional to the numbers of described species in major terrestrial taxonomic groups<sup>24</sup>. We first pre-processed PREDICTS following previous studies<sup>7</sup>: records collected during multiple sampling events at one survey site (e.g. multiple transects) were combined into a single site record, and for studies whose methods were sensitive to sampling effort (e.g. area sampled), species abundances were adjusted to standardise sampling effort across all sites within each study, by assuming a linear relationship between sampling effort and recorded abundance measures (both following ref.<sup>7</sup>). Our analyses of species occurrence and richness are therefore based on discrete count data, whereas abundances are pseudo-continuous (counts adjusted for survey effort). Due to the multi-source structure of PREDICTS (multiple studies with differing methods and scope),



the absolute species richness and abundance measures are non-comparable between studies<sup>24</sup>, so our analyses necessarily measure relative differences across land use classes.

*Host-pathogen association data.* We compiled animal host-pathogen associations from several source databases, to provide as comprehensive a dataset as possible of zoonotic host species and their pathogens: the Enhanced Infectious Diseases (EID2) database<sup>35</sup>; the Global Mammal Parasite Database v2.0 (GMPD2) which collates records of parasites of cetartiodactyls, carnivores and primates<sup>36</sup>; Plourde *et al.*'s reservoir hosts database<sup>37</sup>; Olival *et al.*'s mammal-virus associations database<sup>22</sup>; and Han *et al.*'s rodent zoonotic reservoirs database<sup>38</sup> augmented with pathogen data from the Global Infectious Disease and Epidemiology Network (GIDEON) (Supp. Table 8). We harmonised species names across all databases, excluding instances where either hosts or pathogens could not be classified to species level. To prevent erroneous matches due to misspelling or taxonomic revision, all host species synonyms were accessed from Catalogue Of Life using 'taxize' v.0.8.9<sup>39</sup>. Combined, the dataset contained 20,382 associations between 3883 animal host species and 5694 pathogen species.

Each source database applies different methods and taxonomic scope. EID2 defines associations broadly, based on evidence of a cargo species being found in association with a carrier (host) species, rather than strict evidence of a pathogenic relationship or reservoir status<sup>35</sup>. The other 4 databases were developed using targeted searches of literature and/or surveillance reports, focus mainly on mammals, and provide more specific information on strength of evidence for host status (either serology, pathogen detection/isolation, and/or evidence of acting as reservoir for cross-species transmission). We therefore harmonised definitions of host-pathogen associations across the full combined database. Across all animal taxa we broadly defined associations based on any documented evidence (cargo-carrier or stronger, i.e. including all datasets). Additionally, for mammals only (due to more comprehensive pathogen data availability), we were able to define two further tiers based on progressively stronger evidence: firstly, serological or stronger evidence of infection, and secondly, either direct pathogen detection, isolation or reservoir status. Across all pathogens, we also harmonised definitions of zoonotic status. Each pathogen was classified as *human-shared* if recorded as infecting humans within either one of the source host-pathogen databases or an external human pathogens list collated from multiple sources (Supp. Table 8). Because the source datasets contain some organisms that infect humans and animals rarely or opportunistically, or that may not strictly be zoonotic (e.g. pathogens with an environmental

or anthroponotic reservoir), pathogens were also more specifically defined as *zoonotic agents* (aetiologic agent of a specific human disease with a known animal reservoir) if classed as such in GIDEON, Wertheim *et al.*'s Atlas of Human Infectious Diseases<sup>40</sup> or Taylor *et al.*'s human pathogens database<sup>41</sup>.

*Combined datasets of hosts and land use.* We combined PREDICTS with the compiled host-pathogen database by matching records by species binomial, and each species record was given a binary classification of 'host' or 'non-host' of human-shared pathogens. We adopted a two-tiered definition of host status, to examine the impact of making more or less conservative assumptions about the likelihood of a species contributing to pathogen transmission dynamics and spillover to humans. Firstly, we defined host status broadly: as any species with an association with at least one human-shared pathogen (as defined above), which for mammals must be based on serological or stronger evidence of infection (henceforth referred to as the '*full dataset*'). 177 studies in PREDICTS contained host species matches (190 mammals, 146 birds, 1 reptile, 2 amphibians, 37 invertebrates, listed in Supp. Table 1). Secondly, since mammals are the predominant reservoirs of both endemic and emerging zoonotic infections due to their phylogenetic proximity to humans<sup>42,43</sup>, we also defined mammal species as zoonotic reservoir hosts based on stricter criteria: an association with at least one zoonotic agent (as defined above) which must be based on direct pathogen detection, isolation or confirmed reservoir status (henceforth referred to as '*mammal reservoirs subset*'). Within PREDICTS, 63 studies contained host matches based on this narrower definition (143 mammal reservoir hosts; Extended Data 4, Supp. Table 1).

Prior to analysis, we filtered PREDICTS to include only studies that sampled taxa relevant to zoonotic transmission, since the full database includes many studies with a different taxonomic scope (e.g. plants or non-vector invertebrates)<sup>24</sup>. We retained all studies that sampled any mammal or bird species, as these groups are the main reservoir hosts of zoonoses. For all other taxa, given that zoonoses and their hosts occur globally, we made the more conservative assumption that studies with no sampled hosts represent false absences (i.e. resulting from study aims and methodology) rather than true absences (i.e. no hosts are present), and only included studies with at least one host match in one sampled site in community models. This resulted in a final dataset of 530,161 records from 6801 sites in 184 studies (full dataset) and 51,801 records from 2066 sites within 66 studies (mammal reservoirs dataset; including mammal studies only) (Figure 1). Some host records were of arthropod vectors, but as these are a small proportion of records (around 2%; Supp. Table 1)

we generically refer to all matched species as '*hosts*'. By matching on species binomial we assume that pathogens are equally likely to occur anywhere within their hosts' geographical range; evidence from terrestrial mammal orders suggests that this assumption is reasonable globally<sup>44,45</sup>. Although overlooking geographical variation in pathogen occurrence, pathogen geographical distributions are poorly understood and subject to change, making it difficult to define geographical constraints on host status.

We aggregated land use classes in PREDICTS to ensure a more even distribution of sampled sites. We assigned each survey site's land use type to one of four categories: primary vegetation, secondary vegetation, managed ecosystems (plantation forest, pasture and cropland) and urban. Land use intensity was assigned to either minimal, substantial (combining light and intense use), or cannot decide (the latter were excluded from models). Original use intensity definitions<sup>7</sup> reflect gradation of potential human impacts within land use types; for example urban sites range from minimal (villages, large managed green spaces) to high intensity (impervious with few green areas). Land use categories simplify complex landscape processes, so our aggregation might mask subtle differences in disturbance mode and intensity. However, although some local studies have found differences in zoonotic host abundance and pathogen prevalence between different management regimes<sup>46</sup>, we had no *a priori* reason to hypothesise differences between managed ecosystem types globally. Study regions were categorised as temperate or tropical, following ref.<sup>47</sup>.

## Statistical analysis

Accounting for species-level differences in pathogen discovery effort. The probability of identifying zoonotic pathogens within a species is strongly influenced by effort, meaning that poorly-studied species in our data could be falsely classified as non-hosts. Since research effort might also positively correlate with species' abundance in anthropogenic landscapes, accounting for this uncertainty is crucial. In statistical models we therefore consider host status (and derived metrics such as host richness) to be an uncertain variable, by assuming that all known hosts in our dataset are true hosts (true positives), and that non-hosts comprise a mixture of true non-hosts and an unknown number of misclassified species. We propagate this uncertainty into all model estimates using a bootstrapping approach, in which each iteration transitions a proportion of non-host species to host status with a probability influenced by research effort and taxonomic group (with poorly-researched species in taxonomic orders known to host more zoonoses having the highest transition rates; Extended Data 2, Supp. Text 1).

We estimate disease-related research effort using species publication counts extracted from the PubMed biomedical database (1950–2018) for every species within our dataset ( $n=7285$ ; Extended Data 2c), following other studies in disease macroecology in which publication effort often explains much of the variation in response variables<sup>22,48</sup>. Across 100 randomly-sampled mammal species from PREDICTS, PubMed publication counts were highly correlated to those from Web of Science and Google Scholar (both Pearson  $r = 0.93$ ), indicating robustness to choice of publications database. Using publication counts directly to index species misclassification probability is problematic, since the relationship between publication effort and host status is both nonlinear (e.g. due to positive feedback, where pathogen detection drives increasing research towards a species or taxon) and taxon-specific (e.g. because some taxa are more intensely targeted for surveillance). We therefore calculate a trait-free approximation of false classification probability for non-host species (detailed in Supp. Text 1) by assuming, first, that a species' relative likelihood of being a zoonotic host is proportional to the number of known hosts in the same taxonomic order (i.e. a poorly-studied primate is more likely to be a zoonotic host than a poorly-studied moth), and second, that confidence in non-host status accrues and saturates with increasing publication effort (following the cumulative curve of publication effort for known hosts within the same order; Extended Data 2a-b). Therefore, under-researched mammals, followed by birds, have the highest estimated false classification probabilities, but with substantial variation among mammalian and avian orders (Extended Data 2d-e).

Since data constraints prevent direct observation of how host detections accrue with discovery effort, our trait-free approximation leverages current knowledge of the distribution of zoonotic hosts and publication effort across broad taxonomic groups, and thus might over- or underestimate absolute host potential in any particular species. For example, because species traits and research effort are autocorrelated, our assumption that all non-host species per taxonomic group are equally likely to host zoonoses may conservatively overestimate host potential in less-researched species: many ecological traits that make species more likely to be poorly-studied (e.g. lower population density, smaller range size<sup>49,50</sup>) would often be expected to reduce their relative importance in multi-host pathogen systems<sup>51</sup>. Nonetheless, our approach is sufficient to address our study's main confounding factor, i.e. the potential for biased distribution of research across land use types and biomes globally.

*Community models of host species richness and total abundance.* All modelling was conducted using mixed-effects regression in a Bayesian inference framework (Integrated

Nested Laplace Approximation (INLA)<sup>52</sup>. We aggregated ecological communities data to site-level by calculating the per-site species richness (number of species) and total abundance (total number of sampled individuals, adjusted for survey effort) of host and non-host species. Land use type and intensity were combined into a categorical variable with 8 factor levels (type+intensity, for 4 types and 2 intensity levels). During model selection we considered fixed effects for land use and log-transformed 2015 human population density extracted from CIESIN (because synanthropic species diversity might respond to changes in human population density independently of land use; Extended Data 8). All models included random intercept for study to account for between-study variation, and we additionally considered random intercepts for spatial block within study (to account for the local spatial arrangement of sites), site ID (to account for overdispersion caused by site-level differences)<sup>7</sup> and biome (as defined in PREDICTS).

We modelled the effects of land use on the richness and total abundance of host and non-host species separately, using a Poisson likelihood (log-link) to model species richness (discrete counts). Since abundance data were continuous following adjustment for survey effort, we followed other PREDICTS studies<sup>7</sup> and modelled log-transformed abundance with a Gaussian likelihood; log-transformation both reduces overdispersion and harmonises interpretation of the fixed effects with the species richness models (i.e. both measure relative changes in geometric mean diversity from primary land under minimal use). We also modelled the effects of land use on host richness and abundance as a proportion of overall site-level sampled species richness or abundance, by including log total species richness as an offset in Poisson models, and log total abundance as a continuous fixed effect (effectively an offset) in abundance models.

For each response variable we first selected among candidate model structures, comparing all combinations of random effects with all fixed effects included, and subsequently comparing all possible fixed effects combinations using the best-fitting random effects structure. In all cases we selected among models using the Bayesian pointwise diagnostic metric Watanabe-Akaike Information Criterion (WAIC)<sup>53</sup> (Supp. Table 3-4). The final models were subsequently checked for fit and adherence to model assumptions, including testing for spatial autocorrelation in residuals (Extended Data 9). We then bootstrapped each final model for 1000 iterations to incorporate research effort. For each iteration, each non-host species was randomly transitioned to host status as a Bernoulli trial with success probability  $p$  equal to estimated false negative probability (as described above; Supp. Text 1, Extended Data 2), all community response variables were recalculated, the

model was fitted and 2500 samples were drawn from the approximated joint posterior distribution. We then calculated posterior marginal parameter estimates (median and quantile ranges) across all samples from the bootstrap ensemble (Figure 2, Supp. Table 5). Between 90 and 150 non-host species (median 121) were selected to transition per iteration, increasing the total number of hosts by 24–40% (median 32%; Extended Data 2e). Because study coverage is heterogeneous globally, we subjected the full model ensembles to random and geographical cross-validation (Extended Data 3). We also conducted the same modelling procedure using only the strictly-defined mammal reservoirs subset (Extended Data 4).

*Species-level estimates of land use effects on mammalian and avian zoonotic hosts.* Because aggregate community diversity metrics may mask important variation between taxonomic groups, we separately modelled the average effects of land use type on the occupancy and abundance of all hosts and non-hosts of zoonotic agents within five mammalian (Carnivora, Cetartiodactyla, Chiroptera, Primates, Rodentia) and two avian orders (Passeriformes, Psittaciformes). For mammals we defined zoonotic host status strictly (pathogen detection, isolation or confirmed reservoir status, as described above) and excluded urban sites due to sparse urban sampling for mammals in PREDICTS (only 2 studies). All models included an interaction term between land use type and zoonotic host status (host or non-host) and random intercepts for each species-study combination and for taxonomic family (to account for gross phylogenetic differences). We again accounted for variable research effort per species as described above, fitting 500 models per order, and calculating posterior marginal estimates across samples drawn from the whole ensemble (Supp. Table 6).

Abundance data were overdispersed and zero-inflated due to the high proportion of absence records (i.e. sites where species were not found despite being sampled for). We therefore used a hurdle model-based approach<sup>54</sup> to estimate effects of land use on abundance, by separately fitting occurrence models (presence-absence; binomial likelihood, logit-link) to the complete dataset for each mammalian order, and zero-truncated abundance models (ZTA, log-abundance with Gaussian likelihood) to the dataset with absences removed (Extended Data 5). Mean differences in abundance across land uses are then calculated as the product of the proportional differences in predicted occurrence probability and ZTA relative to primary land<sup>54</sup>. We used posterior samples from paired occurrence (transformed to probability scale) and ZTA models (transformed to linear scale) to calculate a distribution of hurdle predictions separately for each bootstrap iteration (i.e. with the same non-hosts reclassified). We then summarised predicted changes per land use type across samples from the entire bootstrap

ensemble (median and quantile ranges; Figure 3). Due to the complex nested structure of PREDICTS, our hurdle predictions assume independence between occurrence and ZTA processes<sup>54</sup>, so do not formally account for the possibility of covariance at random effects (species or family) level. For clarity, we therefore show the contributions of each separate model for each order (Extended Data 5, Supp. Table 6). In most orders, and when fitting models across all mammal species, land use often appears to act most consistently on species occurrence, with more variable effects on ZTA, suggesting that the independence assumption may be broadly reasonable at this global and cross-taxa scale.

#### *Relationship between pathogen richness and responses to land use across mammal species.*

Pathogen richness (the number of pathogens hosted by a species) is a widely-analysed trait in disease macroecology, with both overall pathogen richness, shared pathogen richness (i.e. number of pathogens shared between focal species) and zoonotic pathogen richness often correlated to species traits such as intrinsic population density, life history strategy and geographic range size<sup>5,22,27,55</sup>. If human-disturbed landscapes systematically select for species trait profiles that facilitate host status, we might expect to observe positive responses to land use in species with higher richness of either human-shared or non human-shared pathogens<sup>23</sup>. We tested this hypothesis for mammals, due to availability of much more comprehensive pathogen data than for other taxa, by analysing the relationship between species pathogen richness and probability of occurrence across three land use types (primary, secondary and managed; urban sites excluded due to limited sampling).

Within the subset of PREDICTS studies that sampled for mammals, containing 26,569 records of 546 mammal species (1950 sites, 66 studies), we used the host-pathogen association dataset to calculate, firstly, each mammal species' richness of human-shared pathogens, and secondly its richness of pathogens with no evidence of infecting either humans or domestic animals (‘non human-shared’), defining associations based on serological evidence or stronger. Of the 546 mammals, 190 species had at least one known human-shared pathogen (human-shared pathogen richness mean 1.92, sd 6.07) and 96 species had at least one non human-shared pathogen (non human-shared pathogen richness mean 0.81, sd 4.16). We account for research effort differently than in the binary host status models above, since pathogen richness is a continuous variable that is influenced by magnitude of effort (i.e. more effort would be expected to increase the number of detected pathogens; Extended Data 6b-c). Therefore, we account for effort by estimating per-species residual pathogen richness not explained by publication effort (i.e. the difference between observed



pathogen richness and expected pathogen richness given publication effort and taxonomic group). To do this, we modelled the effect of publication effort on pathogen richness (discrete counts) separately for human-shared and non human-shared pathogens, using a Poisson likelihood with a continuous fixed effect of log-publications and random intercepts and slopes for each mammalian Order and Family (to account for broad taxonomic differences in host-pathogen ecology between orders<sup>22</sup>). We fitted the model to data from all mammal species in our host-pathogen database (n=780) and predicted expected mean pathogen richness for all mammals in PREDICTS. We calculated residuals from observed values for these species (Extended Data 6), which we expect represent trait-mediated variation, given the evidence that mammal pathogen richness covaries with species traits after accounting for phylogeny and research effort<sup>22</sup>.

We then modelled the relationship between residual pathogen richness (scaled to mean 0, sd 1) and species probability of occurrence across land use types, separately for human-shared and non-human-shared pathogens (Extended Data 7). Species occurrence was modelled using a binomial (logit-link) likelihood, with fixed effects for the interaction between residual pathogen richness and land use type, and random intercepts for species, order, study and spatial block within study. As with prior analyses, models were checked for fit and adherence to assumptions. Pathogen surveillance in animals is often focused on species of zoonotic concern, meaning that pathogen inventories (especially of non-human-shared pathogens) may be more complete for some taxonomic groups than others. We therefore tested model sensitivity to separately fitting models containing, firstly, only species from the four most comprehensively-sampled mammalian orders for parasites and pathogens (Primates, Cetartiodactyla, Perissodactyla and Carnivora; the focal taxa of the Global Mammal Parasite Database<sup>36</sup>), and secondly, species from all other mammal orders. We also tested for sensitivity to uncertainty in the publications-pathogen richness relationship, by separately fitting the land use model to 400 sets of residuals derived using posterior samples from the fitted publication effort model (Extended Data 6g-h), and summarising parameters across the full ensemble. Fixed effects directions and strength of evidence were consistent across all models (Supp. Table 7). Data processing and analyses were conducted in R v. 3.4.1<sup>56</sup>, with model inference conducted in R-INLA<sup>52</sup>.

**Data availability:** All code and data used in this analysis (where not freely available online) are archived at Figshare (<https://figshare.com/s/cede860bb9c77abf23ff>; doi: 10.6084/m9.figshare.7624289), and data sources are listed in Supp. Table 8.



## Extended Data

### List of extended data items

Extended Data 1: Conceptual framework for the effects of land use change on zoonotic disease transmission.

Extended Data 2: Approximating research effort bias for non-host species within the PREDICTS dataset.

Extended Data 3: Random (study-level) and geographical cross-validation of community models (full dataset).

Extended Data 4: Effects of land use on site-level mammalian reservoir host species richness and total abundance.

Extended Data 5: Effects of land use on occurrence and zero-truncated abundance (abundance given presence) of mammalian and avian hosts and non-hosts of zoonotic agents.

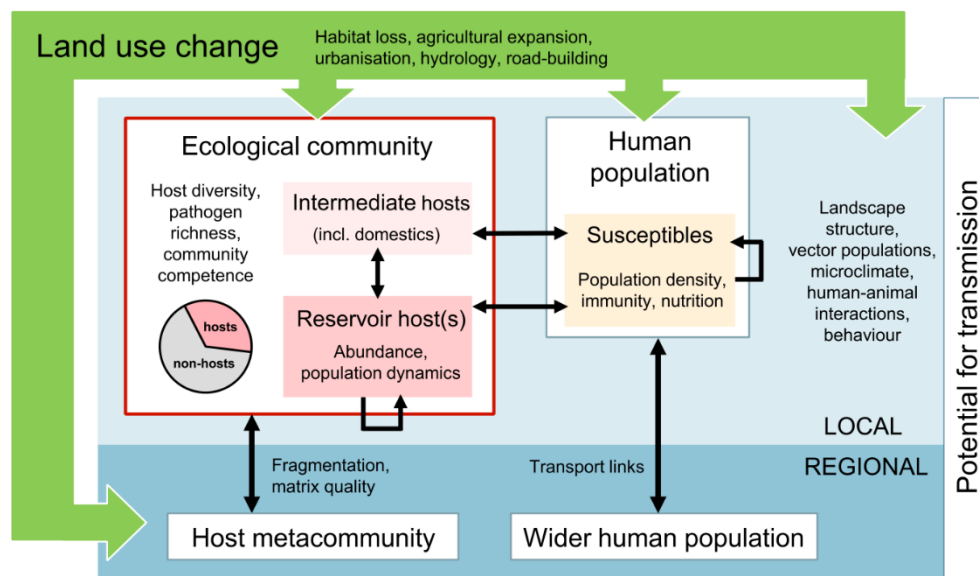
Extended Data 6: Residual human-shared and non-human-shared pathogen richness across mammals.

Extended Data 7: Effects of land use on the relationship between mammal pathogen richness and species occurrence probability.

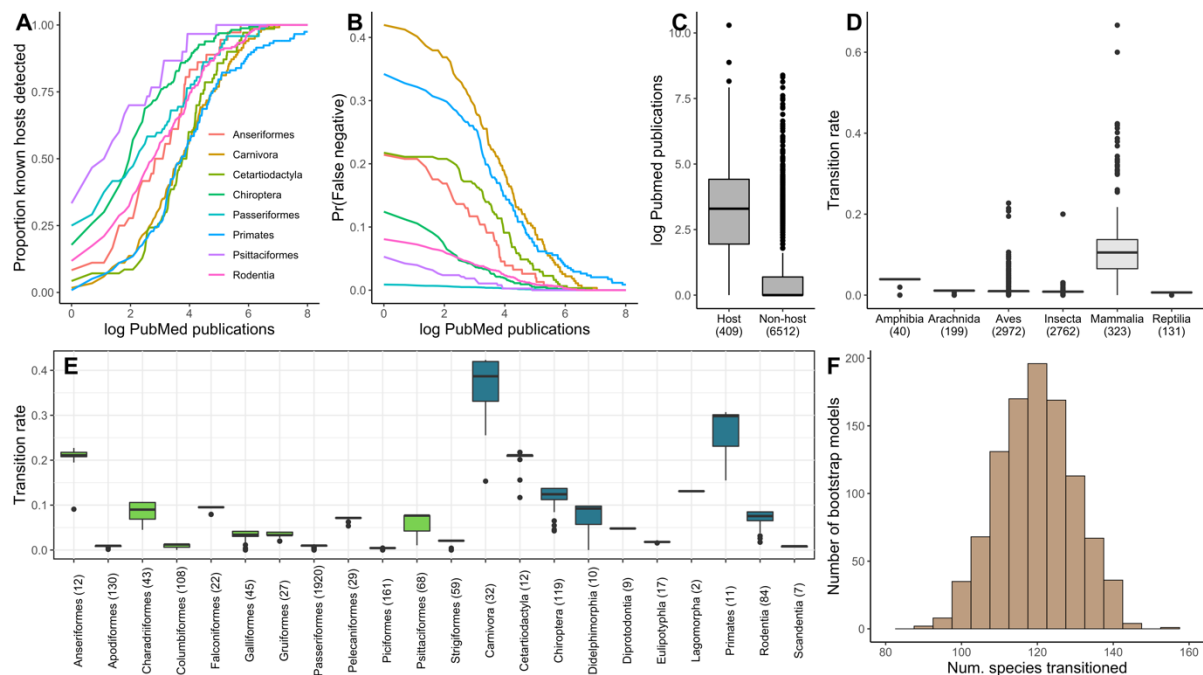
Extended Data 8: Differences in human population density between land use types, for all sites within the full dataset.

Extended Data 9: Diagnostic plots for all community models (full dataset and mammal reservoirs subset).

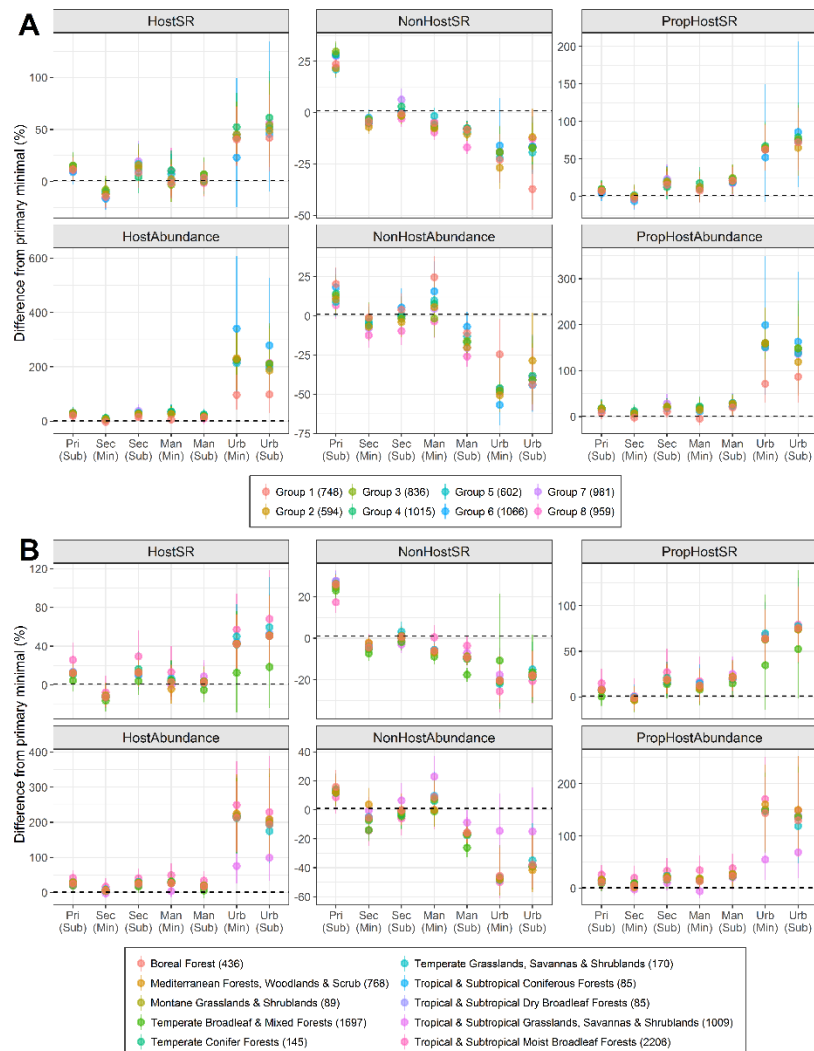
**Extended Data 1: Conceptual framework for the effects of land use change on zoonotic disease transmission.** Pathogen transmission between potential hosts is shown as black arrows. Land use change (green driver) acts on both ecological community composition, including humans (white boxes) and on environmental features that influence contact and transmission both locally (light blue box) and at broader geographical scales (dark blue box). These processes occur within a broader socio-ecological system context also influenced by additional environmental (e.g. climatic), socioeconomic and demographic factors. Unpicking the relative influence of these different processes on disease outcomes is challenging in local disease system studies, where multiple processes may be acting on pathogen prevalence and transmission intensity. The aim of this analysis was therefore to specifically examine, at a global scale, the effects of land use change on the composition of the potential host community (excluding domestic species), denoted below by the red box.



**Extended Data 2: Approximating research effort bias for non-host species within the PREDICTS dataset.** For all non-host species, we approximated the likelihood of false classification given research effort (i.e. probability of being a host, but not detected), based on the distribution of publication effort across known zoonotic hosts within the same taxonomic order (Supp. Text 1). Line graphs show, for several orders, the cumulative curve of publication counts for known zoonotic hosts (A; shown on log-scale), and approximated false classification probability, which declines and asymptotes with increasing levels of research effort (B) (line colours denote taxonomic order). Boxplots show the distribution of PubMed publications for all host and non-host species in PREDICTS (C), and false classification probabilities (used as bootstrap transition rates) for all non-hosts per taxonomic class in PREDICTS (D), and per key mammalian and avian order (E) (bracketed numbers denote number of non-host species per-group; boxes show median and interquartile range, whiskers show values within 1.5\*IQR from quartile). Histogram shows the number of non-host species transitioned to host status for each of 1000 bootstrapped models of the full dataset (F; median 121, 95% quantile range 102–142).

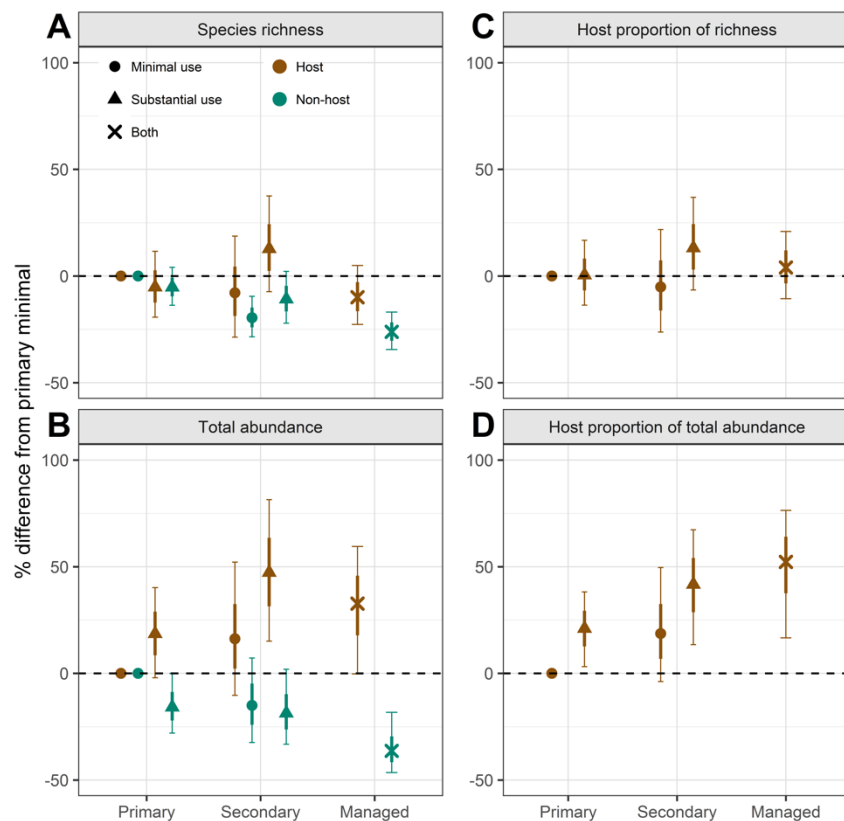


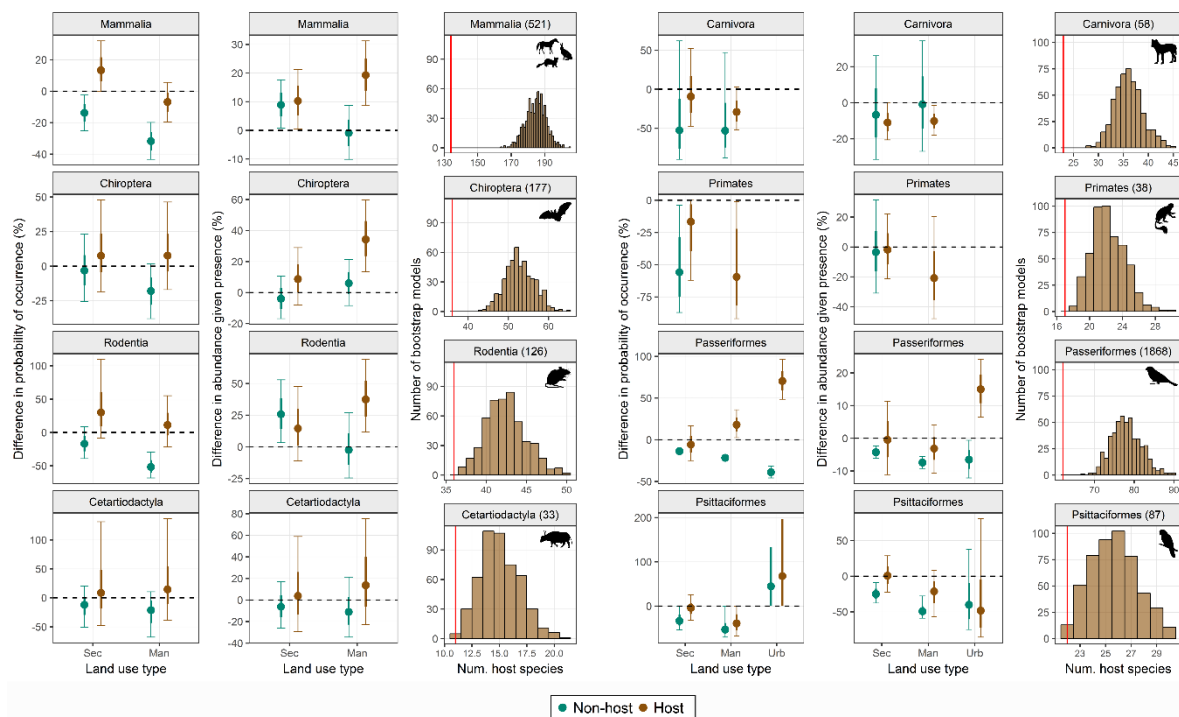
**Extended Data 3: Random (study-level) and geographical cross-validation of community models (full dataset).** We tested the sensitivity of fixed effects estimates to both random and geographically-structured (biome-level) subsampling. For random tests we fitted 8 hold-out models, excluding all sites from 12.5% of studies at a time (mean 12.5% of total sites excluded per model, range 4%-19%; results in A). For geographical tests we fitted 14 hold-out models, with each excluding all sites from one biome (mean 7% of sites excluded per model, range 0.07%-32%; results in B). Points and error bars show posterior marginal parameter distributions for each hold-out model (median and 95% quantile range, with colour denoting hold-out group or biome), calculated across samples from 500 bootstrap iterations per-model to account for variable research effort across species. Directionality and evidence for fixed-effects estimates are robust to both tests, suggesting that our results are not driven by data from any particular subset of studies or regions. Urban parameters are however the most sensitive to exclusion of data, likely due to the relatively sparse representation of urban vertebrate diversity in the PREDICTS database (17 studies in our full dataset).



**Extended Data 4: Effects of land use on site-level mammalian reservoir host species**

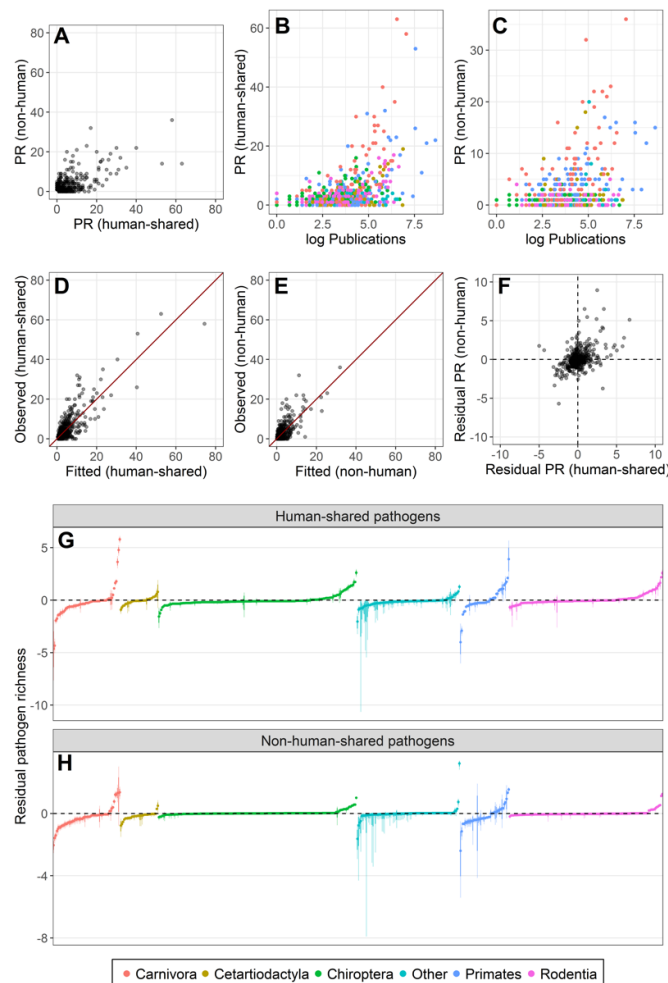
**richness and total abundance.** Points, wide and narrow error bars show differences in diversity metrics from primary minimal use baseline (posterior marginal median, 67% and 95% quantile ranges respectively, across 1000 bootstrap models). Models are of species richness (A) and total abundance (B) of reservoir host and all other (non-host) species, and of hosts as a proportion of site-level richness (C) and total abundance (D). For managed and urban sites, use intensities were combined to improve evenness of sampling (n=2026 sites from 63 studies: primary (589 and 572 for minimal and substantial use respectively), secondary (144, 257), managed (348) and urban (116)). Posterior estimates were calculated across an ensemble of 1000 bootstrapped models (median 51, range 38–62 non-hosts transitioned to host status, i.e. increasing host number by 28–46%) (Methods). Urban sites results show the same trend as the full dataset (Figure 2), but are not visualised due to wide uncertainty: 88.7% (-2.1, 252.3) proportion richness, 307% (78.8, 500.7) proportion abundance (posterior median and 95% quantile range; see Supp. Table 4). Point shape indicates use intensity (minimal, substantial or both combined) and colour indicates host (brown) or non-host (green). Reservoir species are listed in Supp. Table 1 (mammal species listed as ‘Detection/reservoir’ in the ‘Evidence of host status’ column).



**Extended Data 5: Effects of land use on occurrence and zero-truncated abundance****(abundance given presence) of mammalian and avian hosts and non-hosts of zoonotic****agents.** Each row of three plots shows the results of species-level modelling for each of 5mammalian and 2 avian orders, and for mammals overall. Points, wide and narrow error barsshow average difference in species occurrence probability (left column) and zero-truncatedabundance (ZTA; middle column) (posterior median, 67% and 95% quantile ranges across500 and 750 bootstrap iterations, for each order and all mammals respectively). Differencesare shown in secondary (Sec), managed and urban sites relative to a primary land baseline(dashed line), across all host (brown) and non-host (green) species. Histograms show, foreach taxonomic group, the distribution of host species counts across all bootstrap models (i.e.after reclassifying non-hosts) compared to current number of known hosts (red vertical line),and the total number of species included in models (brackets in plot title). Estimates fromoccupancy and ZTA models (Supp. Table 6) were combined, assuming independence ofprocesses, to give the hurdle predictions in Figure 3. Mammal reservoir status was definedbased on strict criteria (pathogen detection or isolation), and the full list of host speciesincluded in these estimates is provided in Supp. Table 1 (scored ‘1’ in the ‘zoonotic agenthost’ column).

**Extended Data 6: Residual human-shared and non human-shared pathogen richness**

**across mammals.** Distribution of human and non human-shared pathogen richness (A) and relationship to publication counts (B-C) are shown for mammals in our host-pathogen association dataset (n=780 species; points represent species shaded by Order, associations defined on serological or stronger evidence). Observed versus fitted plots (D-E) show where observed deviates from expected pathogen richness given log-publications and taxonomic group (Poisson likelihood with random intercepts and slopes for Order and Family; slope estimates for log-publications are similar for both human and non human-shared pathogens,  $\beta$  of 0.298 and 0.248 respectively). We used the fitted models to predict expected pathogen richness for mammals in PREDICTS (n=546) and derived residuals from observed values (shown in F), which were used in land use models (Extended Data 7). Calculating per-species residual quantile ranges across 2500 posterior parameter samples shows that within-species residual variance is generally small relative to residual size (G-H, points and error-bars show posterior median, 67% and 95% intervals, scaled to unit variance), and land use model results are robust to including this uncertainty (Methods, Supp. Table 7).



**Extended Data 7: Effects of land use on the relationship between mammal species**

**pathogen richness and occurrence probability.** Points and error bars show intercept (A-B)

and slope parameters (C-D) of the relationship between residual pathogen richness (scaled to

mean 0 and unit variance) and mammal species occurrence probability (on the log odds scale;

median and 95% credible interval). Intercept parameters represent the average occurrence

probability of a species with residual pathogen richness of 0 (i.e. with average pathogen

richness given research effort and taxonomy), and slope parameters represent the change in

occurrence probability for one scaled unit (standard deviation) increase in residual pathogen

richness ([Extended Data 6g-h](#)). Intercept and slope parameters for primary and secondary

land measure the differences relative to managed land (i.e. delta-intercept or delta-slope; B,

D). Plotted lines show these relationships on the probability scale (E-F), showing the median

(black line), 67% (dark shading) and 95% (light shading) quantile range, based on 3000

samples from the joint posterior distribution. For both human-shared and non human-shared

pathogens, there is a positive relationship between a species' residual pathogen richness and

its probability of occurrence in human-managed land. For human-shared pathogens, the

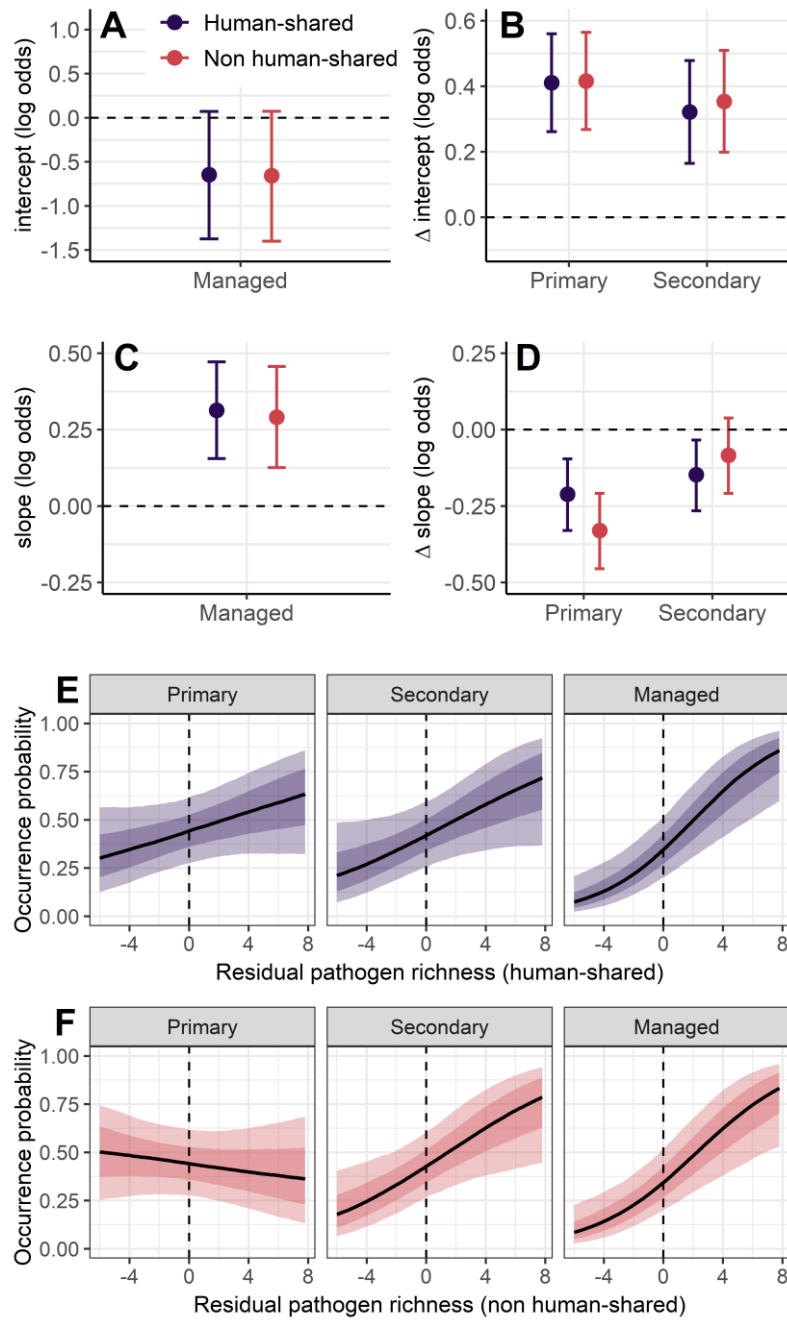
strength of this relationship (slope parameter) is significantly larger in managed sites than in

both primary and secondary land, and for non human-shared pathogens significantly larger in

managed than in primary land (D; slopes for primary land not significantly different from 0).

Full model summaries [and results of sensitivity analyses](#) are in Supp. Table 7.





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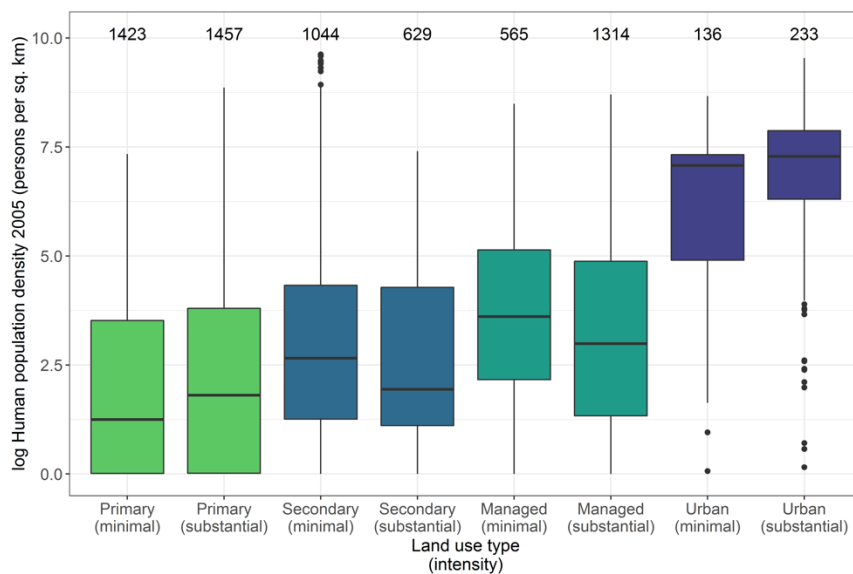
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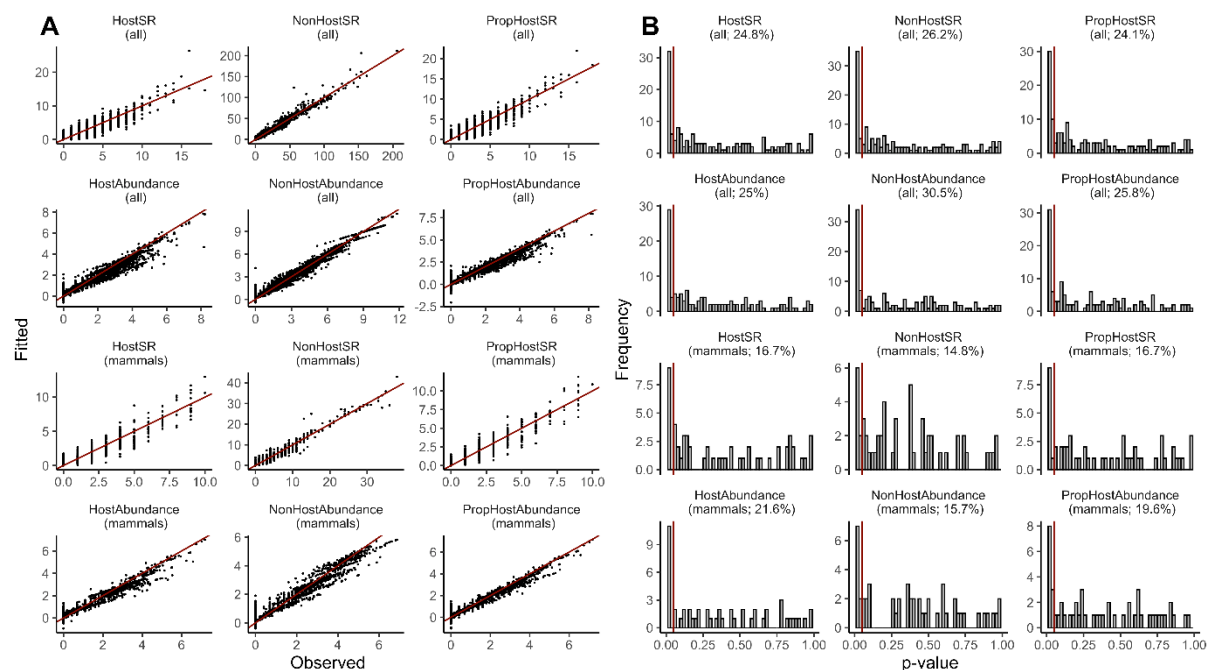
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**Extended Data 8: Differences in human population density between land use types, for all sites within the full dataset.** Boxplots show the distribution of log-transformed human population density, separated by land use type and intensity, across all sites included in community models (n=6801; boxes coloured by land use type, numbers denote the number of sites in each category). Human population density estimates were extracted from CIESIN Gridded Population of the World 4, for 2005, the median year of studies included in the dataset. Per-site log human density estimates were considered as fixed effects in community models of host diversity, since human-tolerant or synanthropic species might respond to human population change independently of land use (Methods).



**Extended Data 9: Diagnostic plots for all community models (full dataset and mammal reservoirs subset).** Species richness counts were modelled with a Poisson likelihood, and abundance (adjusted counts) were log-transformed and modelled with a Gaussian likelihood (see Methods). Plot titles refer to model response variables: species richness (SR), total abundance (Abundance), for hosts, non-hosts, and for hosts as a proportion of the community (Prop). Points in (A) show observed data against model-fitted values, and the red line shows the expectation if observed equals fitted. We also tested for spatial autocorrelation of residuals across all sites within each study, with histograms (B) showing the distribution of per-study Moran's  $I$   $p$ -values (indicating significance of spatial autocorrelation among sites within that study) for each model. Numbers in brackets are the percentage of studies that contained significant spatial autocorrelation ( $p < 0.05$ , shown as a red line). Overall, spatial autocorrelation was fairly low across the dataset (statistically significant in 14%-30% of studies across the different models, with maximum 26% for models with host metrics as response variables). Residuals and statistics were derived from a single fitted model including community mean false classification probability as a linear covariate to account for research effort (with known hosts given a false classification probability of 0), rather than the full bootstrap ensemble.



**ENDS**