

The coevolutionary mosaic of bat betacoronavirus emergence risk

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Driven by the need to understand the ecological factors involved in the emergence of *Betacoronavirus* (the genus causing SARS, MERS, and COVID-19 in human) through bat hosts, we develop an approach to the assessment of spillover risk based on the Geographic Mosaic Theory of Coevolution. In doing so, we provide a global mapping of the spillover risk posed by betacoronaviruses, reflecting the fact that this risk is best understood through the multi-faceted prism of ecological and evolutionary mechanisms. Our framework reveals that host richness alone, although a component of viral hazard, is not a sufficiently integrative predictor of risk. We offer alternative insights based on viral sharing, host compositional uniqueness, and host phylogenetic diversity and phylogeographic regions. By comparing our aggregated measure of risk to a proxy for human density, namely the proportion of each pixel that is covered by urban or built land, we provide a synthetic risk map, allowing the identification of hotspots where the bat-betacoronavirus interaction network may facilitate the emergence of novel viruses and their spillover into human populations.

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Disease emergence is complex, and is driven not only by animal-human contact, but also by the underlying evolutionary dynamics in viral reservoirs.¹ Although host richness is often used as a superficial proxy for spillover risk,^{2,3} these approaches oversimplify the relevant interspecific heterogeneity in immunology, behavior, and other traits, and therefore overlook unique host pools that allow for the rapid evolution of highly divergent viruses.⁴ In the case of generalist pathogens like betacoronaviruses, there is conceptual and empirical support to the idea that these community-level mechanisms are even more important,⁵ particularly given that cross-species transmission may, as a rule, structure viral evolution more than co-divergence with hosts.⁶ This creates a disconnect between coevolutionary theory and most existing ecological frameworks for mapping spillover risk.

The geographic mosaic theory of coevolution (GMTC) attempts to explicitly connect microevolutionary dynamics to the macroecology and biogeography of symbiotic interactions.⁷ The GMTC posits that co-evolutionary processes among pairs⁸ or complexes⁹ of species are structured in space by the rippling effects of abiotic conditions onto evolutionary mechanisms, giving rise to fragmented systems with different ecologies over large spatial extents.¹⁰ The GMTC predicts a spatial fragmentation of coevolutionary dynamics under the joint action of three processes:¹¹ coevolutionary hot- and coldspots, which appear when the intensity of *interaction* (in terms of reciprocal fitness consequences) varies spatially; selection mosaics, wherein the intensity of *selection* varies across space, driven by both the biotic complexity of

the community (locally diverse hosts and viruses are more biotically complex) and the local favorability of the environment;¹² and trait remixing, which occurs when coevolutionary dynamics change when community-level *functional traits* change through meta-community dynamics.

Here, we apply the GMTC to explore and explain the global biogeography of betacoronaviruses, the group that includes SARS-CoV, MERS-CoV, and SARS-CoV-2. In their bat reservoirs, coronaviruses evolve through a mix of host jumps, recombination among disparate lineages, and, to a lesser degree, co-divergence with their hosts—²a mix of mechanisms that creates a complex and nonlinear relationship between host diversity and viral emergence. Working from a recently published database of bat hosts of betacoronaviruses, we test whether spatial structure in bat-betacoronavirus coevolution is identifiable at a global scale. Aiming to explain these patterns, we develop a generalized framework for applying the GMTC to host-virus interactions, with a specific emphasis on the potential to create independent coevolutionary dynamics (and therefore spatial fragmentation in risk) through heterogeneity. We develop a trivariate risk assessment system that connects each GMTC mechanism to a quantifiable aspect of host-virus interactions: (i) viral sharing rates in host communities, representing the strength of potential interaction between viruses and any one host (i.e., places where viruses undergo constant host switching may be coevolutionary coldspots); (ii) the phylogenetic diversity of hosts, as a proxy for variation in the immunological mechanisms that antagonize viruses (i.e., the selection mosaic); and (iii) the local uniqueness of the bat community, representing the potential for viruses to be exposed to novel host traits (e.g., variation in receptor sequences). Together, we argue that these can be used to identify and map the evolutionary drivers that—in conjunction with transmission processes (e.g., viral prevalence in reservoirs and animal-human contact rates)—determine disease emergence risk.

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Results and Discussion

1.1. Bat and betacoronavirus biogeography are broadly consistent Most previous work has assumed that the presence or richness of key groups of bat hosts are predictive of coronavirus diversity.^{2,3} Projecting bat and betacoronavirus phylogeny over space (fig. 1), we find support for the idea that bat community assembly is directly responsible for a global mosaic of viral evolution. The distinct groupings (represented by different colors, symbolizing positions in a subspace formed by the first two phylogenetic principal components) are essentially equivalent between the two groups, and can be coarsely delineated as (1) south and southeast Asia; (2) east Asia (including northern China), west Asia, and the Mediterranean coast; (3) Eurasia above a northing of 40; and (4) Africa and Latin America. In some cases, this diverges from expectations about coronavirus biogeography: for example, previous work has rarely flagged India as a region of interest, but for both bats and betacoronaviruses, the subcontinent falls into the same regions as the southeast Asian peninsula (and indeed, the region is home to known bat hosts of multiple betacoronavirus subgenera, including nobecoviruses, sarbecoviruses, and merbecoviruses).³

Overall, these results suggest that the boundaries of bat and betacoronavirus biogeographic regions are largely consistent. This may be surprising, given that cross-species transmission may play a stronger role in coronavirus diversification than cospeciation—²a property that would theoretically allow for substantial broad divergence in their biogeography. However, host jumps at the family level or higher are relatively rare and significant events in coronavirus evolutionary history;^{2,13} as a result, the mosaic of betacoronavirus phylogeography is assembled from a set of overlapping smaller coevolutionary systems, superimposed in space and filtered by the importance of different subgroups in local host communities. For example, the most speciose and cosmopolitan family of bats, the vesper bats (*Vespertilionidae*), are considered the primary hosts of the subgenus *Merbécovirus* (MERS-like viruses);^{3,13} but in the Americas, where merbecoviruses are the only lineage present, they have only been found in other bat taxa (e.g., *Molossidae*, *Phyllostomidae*).^{14–17} At the coarsest scale, these heterogeneities are lost, and betacoronavirus biogeography tracks the deep rifts in bat evolutionary history—but within broad regions, the component coevolutionary systems may have very different dynamics.

1.2. Hotspots of bat and betacoronavirus biodiversity are distinct Bats, the second most diverse groups of mammals, are found worldwide; gradients in their species richness generally track broader patterns of mammal diversity, with a striking Neotropical hotspot (especially in the Amazon basin) and a secondary hotspot centered in the southeast Asian peninsula. These hotspots of bat diversity are



Figure 1 Phylogeographic regions of bats (top) and viruses (bottom) based on the joint analysis of their occurrence and phylogenetic relatedness. The different colors show tendencies to separate alongside the first two components of a PCoA. Note that the PCoA for the bats and viruses are independent, and so cannot be compared directly – that being said, the regions can be compared across maps.

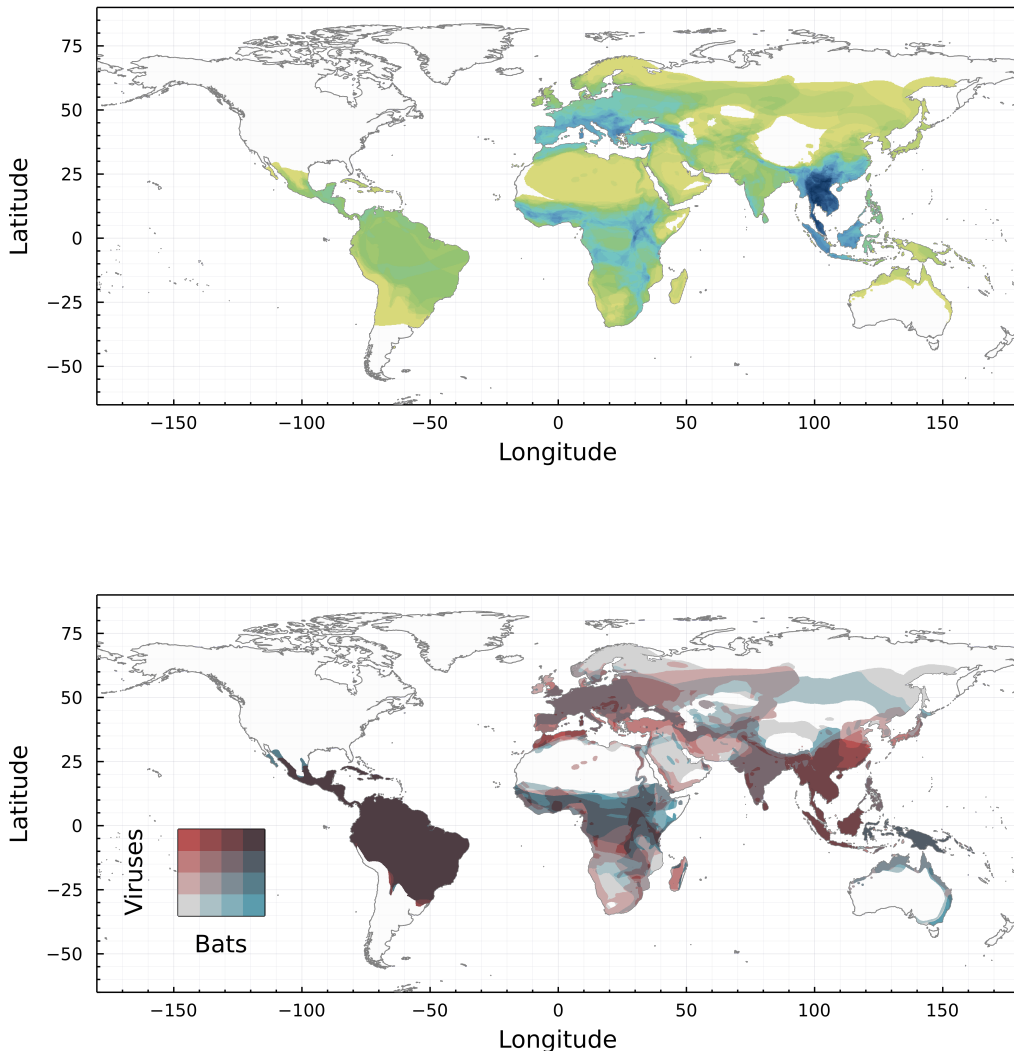


Figure 2 Top panel: relative diversity of known bat hosts of betacoronaviruses. This map shows that the region with the largest number of possible hosts is South-Eastern Asia. Bottom panel: congruence between the evolutionary distinctiveness of the hosts (grey to blue) and the viruses (grey to red).

generally presumed to be hotspots of viral adaptive radiation, and therefore areas of concern for human health.^{2,18} However, the hotspots of known bat betacoronavirus hosts show a distinct pattern, with primary hotspots (both in terms of size and higher values) of host richness situated in southeast Asia, parts of southern Europe, and to a lesser extent parts of Africa in the -25-0 range of latitudes (fig. 2; top). Although hundreds of species likely host undiscovered betacoronaviruses, machine learning predictions have suggested that these undiscovered reservoirs should follow the same diversity gradient.¹⁹ In principle, these hotspots of locally-diverse, virus-rich bat communities should drive more adaptive diversification in their viruses.

However, we find that the global pattern of betacoronavirus phylogenetic distinctiveness is quite distinct from both bat host richness and phylogenetic distinctiveness (fig. 2; bottom). In contrast to the sparsity of Neotropical betacoronavirus hosts, South and Central America have the most evolutionary distinct hosts *and* viruses, followed by secondary hotspots in southeast Asia and the Rift Valley region have mostly distinct viruses. Some degree of sampling bias may contribute to these patterns: for example, the Neotropics are one of the places where the fewest bat betacoronavirus sequences have been generated (cite2), resulting in a sparser phylogenetic tree, and artificially inflating distinctiveness; conversely, disproportionate research effort in eastern China²⁰ may have led to a more complete inventory of the local diversity of coronaviruses, again inflating these metrics relative to underlying patterns. Even

accounting for these potential biases, though, there is obvious heterogeneity in betacoronavirus evolutionary distinctiveness that is distinct from overall bat diversity.

Overall, these patterns recapitulate the evolutionary history of both the order Chiroptera and the genus *Betacoronavirus*. Horseshoe bats (Rhinolophidae) include the reservoirs of the SARS-like viruses (sub-genus *Sarbecovirus*), the group of pandemic threats that have been of the greatest interest to researchers¹³ (and so have been sampled most intensively).²⁰ The hotspots of host richness and viral diversity in southeast Asia—both of which are disproportionately high, considering the global landscape of bat species richness—are almost entirely driven by viral adaptive radiation through host switching within this clade^{3,19}. In contrast, the Neotropical hotspot of viral distinctiveness is driven by isolation by host vicariance. Out of the four main groups of betacoronaviruses, only merbecoviruses have been found in animals in the Americas—an introduction that is generally presumed to be ancient.^{3,21} While comparatively understudied, New World merbecoviruses have been found in the ghost-faced bats (Mormoopidae), Neotropical leaf-nosed bats (Phyllostomidae), and free-tailed bats (Molossidae).^{14–17} The former two groups and a clade of the latter are endemic to the Neotropics, while the explosive adaptive radiations of the phyllostomids are responsible for the hotspot of bat diversity in the Amazon.²² Together, these clades of New World bats play host to a distinct regime of betacoronavirus coevolution.

1.3. Coevolutionary regimes structure evolutionary risk of zoonotic emergence The existence of well-defined cophylogenetic regions suggests that the bat-betacoronavirus system is spatially fragmented enough to create differential coevolutionary processes; in turn, the coevolutionary mosaic contributes to heterogeneity in emergence risk. These ideas are, respectively, supported by the existence of hotspots of viral uniqueness and the diverse origins of human betacoronaviruses. Together, these ideas point to a predictable relationship between host community structure and coevolutionary pressure: phylogeographic structure in bat hosts—and their diverse immune strategies—²³ creates a landscape of selective pressure; the trajectory of viruses' coevolutionary response is, in turn, constrained by the opportunities they have for either coevolutionary specialization or diversification through host jumps and recombination.

Based on the geographic mosaic theory of coevolution, we developed a trivariate map of three facets of coevolutionary pressure (see Methods): (1) *host phylogenetic diversity*: a high diversity of evolutionary histories should expose viruses to more variation in host immune traits; (2) *host community uniqueness*: exposure to greater host trait heterogeneity can drive viral diversification, and coevolving with more unique host communities should create more unique branches of viral evolution; and (3) propensity for *viral sharing*: frequent cross-species transmission may act as a buffer on selective pressure, while lower rates of exchange may enable more simultaneous trajectories of viral specialization to coexist within a given community. We combine global maps of all three to generate a map of coevolutionary regimes, where close colors represent similar risks, and paler pixels represent overall higher risk (fig. 3). We find that these regions do not neatly overlap with those defined in fig. 1 or fig. 2, reinforcing the notion that local-scale coevolutionary mosaics can form within cophylogenetic regions.

The greatest evolutionary potential for zoonotic emergence exists where pathogen pools have a high genetic diversity and high propensity for cross-species transmission. In our framework, emergence risk is therefore maximized under higher phylogenetic diversity (viruses are exposed to different host clades), higher host uniqueness (viruses are experiencing novel, heterogeneous host traits combinations), and low to medium viral sharing (host-virus pairs can coevolve independently, but divergent viruses may still have opportunities for recombination). In fig. 3, this corresponds to yellow areas (dynamics dominated by low viral sharing, with equal contributions of selection mosaics and trait remixing; southeast Asia, and the Indian sub-continent), green-yellow areas (dynamics with low viral sharing but dominated by the selection mosaic effect of host diversity; sub-Saharan Africa), and red-yellow areas (dynamics with low viral sharing but dominated by trait remixing in host communities; the Middle East). Translating this axis of variation back into a univariate risk map (fig. 4) highlights that this evolutionary landscape has a striking correspondence to regions where zoonotic betacoronaviruses have previously emerged.

Compared to approaches that map emergence risk based only on the number of known bat hosts of betacoronaviruses, our framework suggests regions where high viral sharing dominates coevolutionary dynamics—such as Latin America, or Eurasia above a northing of 30—would pose less of a relative risk of zoonotic emergence. Nevertheless, areas of high host uniqueness coupled with high viral sharing (red-to-pink in fig. 3) could create hotspots facilitated by viral codivergence. Our framework identifies Madagascar, where most bat species are endemic following evolutionary divergence from sister species in both African and Asian continents,²⁴ as one such hotspot; interestingly, a recent study²⁵ reported a novel and highly divergent lineage of nobecoviruses from Madagascar-endemic pteropid bat species

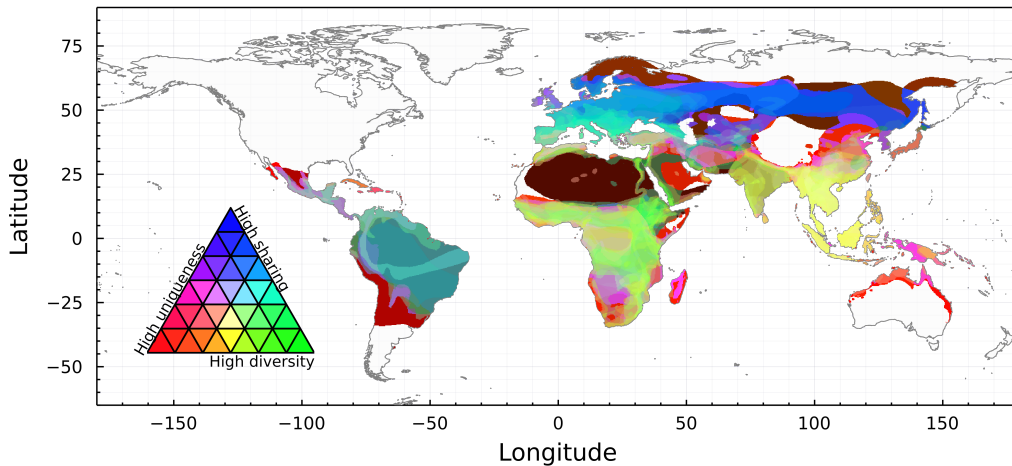


Figure 3 Trivariate additive mapping of the components of risk in the red/green/blue, where high virus sharing is encoded in the blue channel, host phylogenetic diversity in the green channel, and compositional uniqueness in the red channel. The main driver of emergence risk (possibility of spatially overlapped but coevolutionarily independent host-viral dynamics) corresponds to low viral sharing, i.e. pixels around yellow.

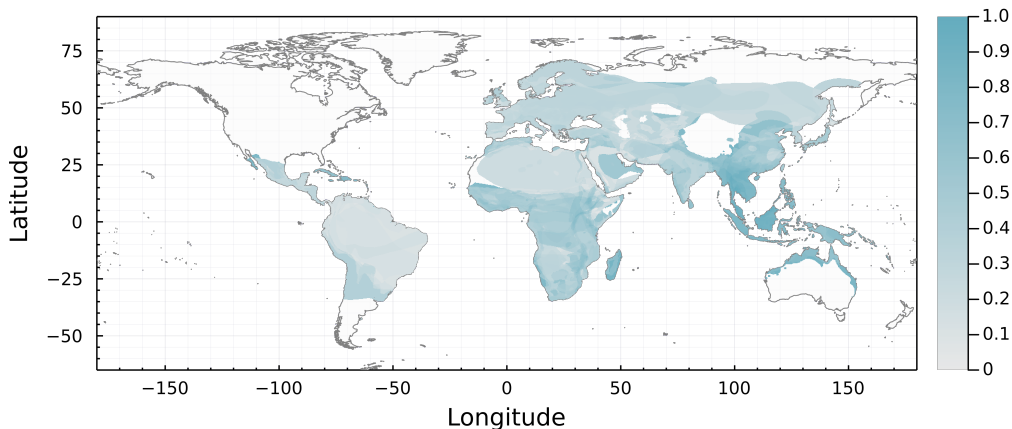


Figure 4 Extraction of a measure of *Betacoronavirus* emergence risk from bat hosts based on the colorimetric space from fig. 3. The risk is a composite measure of the color value and angular distance to the yellow hue, as defined in the methods, ranged in the unit space. Based on these analyses, South-Eastern Asia, Madagascar, the Middle-East, and Africa below the Sahara desert have the highest relative risk of emergence.

(*Pteropus rufus* and *Rousettus madagascariensis*), again supporting the predictive power of the coevolutionary framework.

1.4. Human landscapes filter the geography of emergence risk The relationship between the underlying pathogen pool and emergence risk is mediated by both human-wildlife interfaces (the probability of spillover) and opportunities for onward transmission (the probability that spillovers become epidemics)¹. As a proxy for both, we finally overlaid the risk component from the composite map (see above) with the proportion of built land, as a proxy for a mix of habitat disturbance, potential for bat synanthropy or contact with bridge hosts like livestock,^{26,27} and human population density and connectivity^{1,28,29} (fig. 5). Accounting for these factors, most of South America and Europe are at comparatively lower risk, as—although densely populated—settlements tend to be in areas with lower potential risk. Conversely, regions like Malaysia and the northern coast of Australia have a high evolutionary risk component, but should represent a relatively lower effective risk due to low human density. However, southeast Asia, the Indian subcontinent, and scattered hotspots in sub-Saharan Africa are at high risk due to the overlap between human populations and natural opportunities for cross-species transmission of betacoronaviruses.

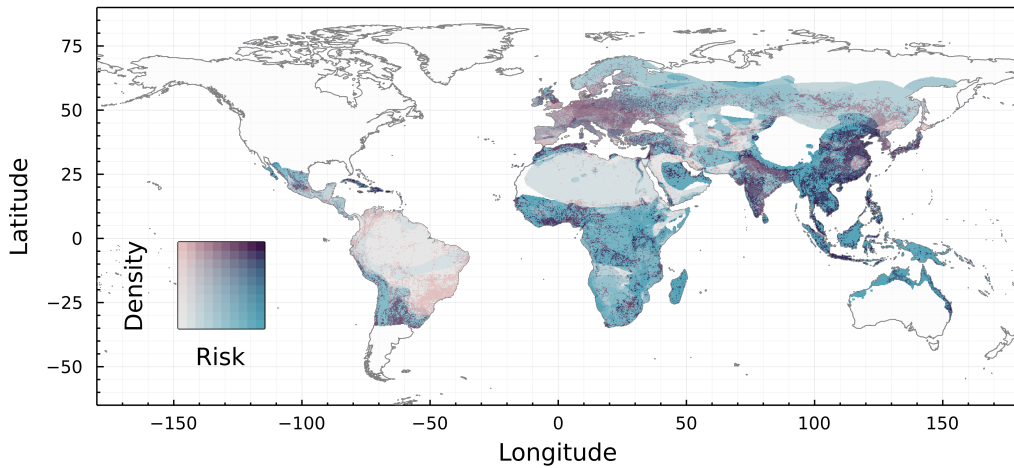


Figure 5 Overlap of the percent of each pixel occupied by urbanized structures, representing the degree of settlement, on the spillover risk map (where the risk comes only from wildlife, and ignores multi-hosts chains of transmissions including non-bats hosts). Darker pixels correspond to more risk, in that the GMTC-derived risk of fig. 4 is high *and* the pixel is densely occupied by human populations. This approach increases the relative risk of several regions in Africa, and highlights the risk in India, southeast China, and the Arabian peninsula where areas of high to moderate risk overlap with areas of denser population.

Reassuringly, these predictions correspond to the geographic origins of the three bat-origin coronaviruses that have recently emerged in human populations. While available information puts the spillover of SARS-CoV-2 in a live animal market in Wuhan, China, the ultimate origin of the virus is almost certainly in a divergent lineage of sarbecoviruses from the Indochinese peninsula that was poorly characterized prior to the pandemic.^{30–32} Similarly, the SARS-CoV outbreak began in Guangdong province in 2002, reaching humans through small carnivore bridge hosts, but was eventually traced back to a set of likely progenitor viruses found in cave-dwelling horseshoe bats in Yunnan province;³³ nearby, antibody evidence has indicated human exposure to SARS-like viruses.³⁴ MERS-CoV was originally detected in Saudi Arabia, accompanied by a nearly identical virus sequenced from an Egyptian tomb bat (*Taphozous perforatus*),³⁵ but is widespread in camels in East Africa and the Middle East, and may have reached its bridge host decades earlier than originally supposed;³⁶ as a result, the geography of the original bat-to-camel transmission is still widely regarded as uncertain. All of these are broadly consistent with the risk factors we identify. Notably, India and west Africa are additional hotspots that have yet to experience the emergence of a bat coronavirus into human populations, but may still be at risk—particularly given known gaps in bat surveillance,²⁰ and a dense population in both regions with global connectivity. In any of these regions, surveillance on viral reservoirs can be paired with targeted monitoring of high-risk human populations (i.e., those with regular wildlife contact)³⁷ for maximum impact.

2

Conclusion

Bats are important reservoir hosts for different classes of microorganisms, many of which a threat to human health.^{38,39} Chiropterans emerged around 64 million years ago and are one of the most diverse mammalian orders, with an estimated richness of more than 1400 species.^{40,41} They exhibit a broad variety of habitat use, behaviour, and feeding strategies, putting them at key positions in the delivery and provisioning of several ecosystem services, tied to important ecosystem-derived benefits to human.⁴² For example, bats are an essential component of many seed-dispersal networks.⁴³ Over two-thirds of bats are known to be either obligate or facultative insectivores, therefore actively contributing for agricultural pest control,^{44,45} and vectors of pathogens that put a risk on human health.^{46,47} Because bats are globally distributed and have a long evolutionary history, phylogeographic and biogeographic approaches are required to shed light on the contemporary distribution of coevolutionary processes between bats and the pathogens they host. Not all areas in which bats, viruses, and human are co-occurring are facing a risk of spillover towards human populations, and the areas in which this risk exist may not be facing risks of the same nature and magnitude.

Here, we propose a simple framework with broad explanatory power that helps contextualize discoveries like highly divergent nobecoviruses in Madagascar and the previously-neglected adaptive radiation

of sarbecoviruses outside of southern China and throughout southeast Asia. In doing so, it advances ecological theory beyond the current state of the art for global maps of emergence risk. For example, previous studies that have used host richness as proxy have predicted a high diversity of unsampled bat viruses,¹⁸ bat coronaviruses,² and even specifically betacoronaviruses¹⁹ in both the Amazon and southeast Asia. While we find that both regions are characterized by highly divergent host and viral communities, our framework identifies key differences between the regions. We find that Latin America is a hotspot of both host and viral distinctiveness, suggesting that this branch of the bat-betacoronavirus complex may be undergoing independent evolutionary dynamics from the rest of the global pool, but with limited potential for viral diversification—a finding that is supported by previous work indicating a higher rate of codivergence in Latin America.² In contrast, in southeast Asia, host richness and viral distinctiveness are high but sharing is low; this suggests a different type of evolutionary dynamics that could generate high local diversity of viruses through host switching and viral recombination (see e.g.,¹³ as well as the discovery of recombinant viruses that share genetic material from both the SARS-CoV and SARS-CoV-2 branches of the Sarbecovirus lineage).⁴⁸ Both of these regions are priority areas for sampling, especially given predictions that they contain many bat hosts of undiscovered betacoronaviruses.^{19,20} However, both the evolutionary and ecological aspects of emergence risk are likely higher in southeast Asia—a fact that will only become more relevant, as bats track shifting climates and exchange viruses with other species, creating a hotspot of cross-species transmission unique to the region.⁴⁹

The diversity and diversification potential of bats responds to anthropogenic factors others than shifting climates.⁵⁰ Land use changes could significantly decrease bat suitability, notably through effects on diet and availability of habitats.⁵¹ As our results establish that the diversification of bats betacoronaviruses happens on top of processes affecting hosts, biogeographic variation in human population density and anthropogenic disturbances may feed into co-evolutionary dynamics. Increase in humans-hosts contacts also increase the risk of emergence of novel diseases,⁵² so does the changes in landscape connectivity at local/regional scales.⁵³ This represents a challenge for both conservation strategies and disease ecology: some areas can a high emergence risk and more potential for the acquisition of zoonotic viruses through bat-human encounters.⁵⁴ In particular, the challenge ahead lies in the need to quantify actual exposure (and risk) accounting for several transmission scenarios, including both direct and indirect bat - human interactions, and feeding back into the provision of ecosystem services by bats.

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Methods

3.1. Known *Betacoronavirus* hosts We downloaded the data on bats hosts of *Betacoronavirus* from <https://www.viralemergence.org/betacov> on Apr. 2022,¹⁹ and filtered it to “known” hosts (established before the emergence of SARS-CoV-2) and “novel” hosts (confirmed through sampling and competence assays since the initial data collection). The original database was assembled by a combination of data mining and literature surveys, including automated alerts on the “bats” and “coronavirus” keywords to identify novel empirical evidence of bats-betacoronaviruses associations; this yielded a total of 126 known hosts, 47 of which were novel hosts.

3.2. Bat occurrences We downloaded the rangemap of every current bat species that was classified as an empirically documented host of *Betacoronavirus* from the previous step, according to recent IUCN data.⁵⁵ The range maps were subsequently rasterized using the `rasterize` function from GDAL⁵⁶ at a resolution of approximately 100kmx100km. For every pixel in the resulting raster where at least one bat host of *Betacoronavirus* was present, we extract the species pool (list of all known bat hosts), which was used to calculate the following risk assessment components: bat phylogenetic diversity, bat compositional uniqueness, and predicted viral sharing risk.

3.3. Bat phylogenetic diversity For every pixel, we measured Faith’s Phylogenetic Diversity⁵⁷ based on a recent synthetic tree with robust time calibration, covering about 6000 mammalian species.⁵⁸ Faith’s PD measures the sum of unique branches from an arbitrary root to a set of tips, and comparatively larger values indicate a more phylogenetic diverse species pool. We measured phylogenetic diversity starting from the root of the entire tree (and not from Chiroptera); this bears no consequences on the resulting values, since all branches leading up to Chiroptera are only counted one per species pool, and (as we explain when describing the assembly of the composite risk map), all individual risk components are ranged in [0,1]. This measure incorporates a richness component, which we chose not to correct for; the interpretation of the phylogenetic diversity is therefore a weighted species richness, that accounts for phylogenetic over/under-dispersal in some places.

3.4. Bat compositional uniqueness For every species pool, we measured its Local Contribution to Beta-Diversity;⁵⁹ LCBD works from a species-data matrix (traditionally noted as \mathbf{Y}), where species are rows and sites are columns, and a value of 1 indicates occurrence. We extracted the \mathbf{Y} matrix assuming that every pixel represents a unique location, and following best practices⁶⁰ transformed it using Hellinger’s distance to account for unequal bat richness at different pixels. The correction of raw community data is particularly important for two reasons: first, it prevents the artifact of richer sites having higher importance; second, it removes the effect of overall species richness, which is already incorporated in the phylogenetic diversity component. High values of LCBD indicate that the pixel has a community that is on average more dissimilar in species composition than what is expected knowing the entire matrix, i.e. a more unique community. Recent results by⁶¹ shows that LCBD measures are robust with regards to spatial scale, and are therefore applicable at the global scale.

3.5. Viral sharing between hosts For all bat hosts of *Betacoronavirus*, we extracted their predicted viral sharing network, generated from a previously published generalized additive mixed model of virus sharing by a tensor function of phylogenetic distance and geographic range overlap across mammals.⁶² This network stores pairwise values of viral community similarity. To project viral sharing values into a single value for every pixel, we averaged the pairwise scores. High values of the average sharing propensity means that this specific extant bat assemblage is likely to be proficient at exchanging viruses.

3.6. Composite risk map To visualize the aggregated risk at the global scale, we combine the three individual risk components (phylogenetic diversity, compositional uniqueness, and viral sharing) using an additive color model.⁶³ In this approach, every risk component gets assigned a component in the RGB color model (phylogenetic diversity is green, compositional uniqueness is red, and viral sharing is blue). In order to achieve a valid RGB measure, all components are re-scaled to the [0,1] interval, so that a pixel with no sharing, no phylogenetic diversity, and no compositional uniqueness is black, and a pixel with maximal values for each is white. This additive model conveys both the intensity of the

overall risk, but also the nature of the risk as colors diverge towards combinations of values for three risk components. Out of the possible combinations, the most risky in terms of rapid diversification and spillover potential is high phylogenetic diversity and low viral sharing,⁶⁴ in that this allows multiple independent host-virus coevolutionary dynamics to take place in the same location. In the colorimetric space, this correspond to yellow – because the HSV space is more amenable to calculations for feature extraction,⁶⁵ we measured the risk level by calculating the angular distance of the hue of each pixel to a reference value of 60 (yellow), and weighted this risk level by the value component. Specifically, given a pixel with colorimetric coordinates (h, s, v) , its ranged weighted risk value is

$$v \times \left[1 - \frac{|\text{atan}(\cos(\text{rad}(h)), \sin(\text{rad}(h))) - X|}{2\pi} \right],$$

where X is $\text{atan}(\cos(\text{rad}(60)), \sin(\text{rad}(60)))$, a constant approximately equal to 0.5235.

3.7. Viral phyogeography and evolutionary diversification To next represent phyogeography of betacoronaviruses in bats, we aggregated and analyzed betacoronavirus sequence data. We used the following query to pull all *Betacoronavirus* sequence data from the GenBank Nucleotide database except SARS-CoV-2; (“Betacoronavirus”[Organism] OR betacoronavirus[All Fields]) NOT (“Severe acute respiratory syndrome coronavirus 2”[Organism] OR sars-cov-2[All Fields]). We added a single representative sequence for SARS-CoV-2 and manually curated to remove sequences without the RNA-dependent RNA polymerase (RdRp) sequence or that contained words indicating recombinant or laboratory strains including “patent”, “mutant”, “GFP”, and “recombinant”. We filtered over-represented taxa including betacoronavirus 1, hCoV-OC43, Middle East respiratory syndrome coronavirus, Murine hepatitis virus, and hCoV-HKU1. Curated betacoronavirus RdRp sequences were then aligned using MAFFT⁶⁶ v1.4.0 (Algorithm FFT-NS-2, Scoring matrix 200PAM / k=2, gap open penalty 1.53m offset value 0.123) and a maximum likelihood tree reconstructed in IQ-TREE⁶⁷ v1.6.12 with ModelFinder⁶⁸ ultrafast bootstrap approximation⁶⁹ with a general time reversible model with empirical base frequencies and the 5-discrete-rate-category FreeRaye model of nucleotide substitution (GTR+F+R5).

We first tested the hypothesis that hotspots of viral diversification would track hotspots of bat diversification. To do so, we plotted the number of known bat hosts (specifically only those included in the phylogeny, so there was a 1:1 correspondence between data sources) against the “mean evolutionary distinctiveness” of the associated viruses. To calculate this, we derived the fair proportions evolutionary distinctiveness⁷⁰ for each of the viruses in the tree, then averaged these at the bat species level, projected these values onto their geographic distributions, and averaged across every bat found in a given pixel. As such, this can be thought of as a map of the mean evolutionary distinctiveness of the known viral community believed to be associated with a particular subset of bats present.

3.8. Co-distribution of hosts and viral hotspots Subsequently, we tested the hypothesis that the biogeography of bat betacoronaviruses should track the biogeography of their hosts. To test this idea, we loosely adapted a method from,^{71,72} who proposed a phylogenetic method for the delineation of animal biogeographic regions. In their original method, a distance matrix - where each row or column represents a geographic raster’s grid cell, and the dissimilarity values are the “beta diversity similarity” of their community assemble - undergoes non-metric multidimensional scaling (NMDS); the first two axes of the NMDS are projected geographically using a four-color bivariate map. Here, we build on this idea with an entirely novel methodology. First, we measure the phylogenetic distance between the different viruses in the betacoronaviruses tree by using the cophenetic function in ape;⁷³ subsequently, we take a principal components analysis of that distance matrix (readily interchangeable for NMDS in this case) to project the viral tree into an n-dimensional space. We then take the first two principal components and, as with the evolutionary distinctiveness analysis, aggregated these to a mean host value and projected them using a four-color bivariate map.

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