1 Origin and cross-species transmission of bat coronaviruses in China

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18 Abstract

- 19 Bats are presumed reservoirs of diverse coronaviruses (CoVs) including progenitors of Severe Acute
- 20 Respiratory Syndrome (SARS)-CoV and SARS-CoV-2, the causative agent of COVID-19. However, the
- 21 evolution and diversification of these coronaviruses remains poorly understood. We used a Bayesian
- 22 statistical framework and sequence data from all known bat-CoVs (including 630 novel CoV sequences)

to study their macroevolution, cross-species transmission, and dispersal in China. We find that host-

- switching was more frequent and across more distantly related host taxa in alpha- than beta-CoVs, and
- 25 more highly constrained by phylogenetic distance for beta-CoVs. We show that inter-family and -genus
- switching is most common in Rhinolophidae and the genus *Rhinolophus*. Our analyses identify the host
- 27 taxa and geographic regions that define hotspots of CoV evolutionary diversity in China that could help
- 28 target bat-CoV discovery for proactive zoonotic disease surveillance. Finally, we present a phylogenetic
- analysis suggesting a likely origin for SARS-CoV-2 in *Rhinolophus* spp. bats.

31 Introduction

32 Coronaviruses (CoVs) are RNA viruses causing respiratory and enteric diseases with varying

33 pathogenicity in humans and animals. All CoVs known to infect humans are zoonotic, or of animal origin, with many thought to originate in bat hosts^{1,2}. Due to their large genome size (the largest non-34 35 segmented RNA viral genome), frequent recombination and high genomic plasticity, CoVs are prone to cross-species transmission and are able to rapidly adapt to new hosts^{1,3}. This phenomenon is thought to 36 have led to the emergence of a number of CoVs affecting livestock and human health⁴⁻⁹. Three of these 37 38 causing significant outbreaks originated in China during the last two decades. Severe Acute Respiratory 39 Syndrome (SARS)-CoV emerged first in humans in Guangdong province, southern China, in 2002 and 40 spread globally, causing fatal respiratory infections in close to 800 people¹⁰⁻¹². Subsequent investigations identified horseshoe bats (genus Rhinolophus) as the natural reservoirs of SARS-related CoVs and the 41 likely origin of SARS-CoV¹³⁻¹⁶. In 2016, Swine Acute Diarrhea Syndrome (SADS)-CoV caused the death of 42 over 25,000 pigs in farms within Guangdong province¹⁷. This virus appears to have originated within 43 *Rhinolophus* spp. bats, and belongs to the HKU2-CoV clade previously detected in bats in the region¹⁷⁻¹⁹. 44 45 In 2019, a novel coronavirus (SARS-CoV-2) caused an outbreak of respiratory illness (COVID-19) first 46 detected in Wuhan, Hubei province, China, which has since become a pandemic. This emerging human virus is closely related to SARS-CoV, and also appears to have originated in horseshoe bats^{20,21} - with its 47 full genome 96% similar to a viral sequence reported from *Rhinolophus affinis*²⁰. Closely related 48 sequences were also identified in Malayan pangolins^{22,23}. 49

A growing body of research has identified bats as the evolutionary sources of SARS- and Middle East Respiratory Syndrome (MERS)-CoVs ^{13,14,24-26}, and as the source of progenitors for the human CoVs, NL63 and 229E^{27,28}. The emergence of SARS-CoV-2 further underscores the importance of bat-origin CoVs to global health, and understanding their origin and cross-species transmission is a high priority for

54	pandemic preparedness ^{20,29} . Bats harbor the largest diversity of CoVs among mammals and two CoV						
55	genera, alpha- and beta-CoVs ($lpha$ - and eta -CoVs), have been widely detected in bats from most regions of						
56	the world ^{30,31} . Bat-CoV diversity seems to be correlated with host taxonomic diversity globally, the						
57	highest CoV diversity being found in areas with the highest bat species richness ³² . Host switching of						
58	viruses over evolutionary time is an important mechanism driving the evolution of bat coronaviruses in						
59	nature and appears to vary geographically ^{32,33} . However, detailed analyses of host-switching have been						
60	hampered by incomplete or opportunistic sampling, typically with relatively low numbers of viral						
61	sequences from any given region ³⁴ .						
62	China has a rich bat fauna, with more than 100 described bat species and several endemic species						
63	representing both the Palearctic and Indo-Malay regions ³⁵ . Its situation at the crossroads of two						
64	zoogeographic regions heightens China's potential to harbor a unique and distinctive CoV diversity.						
65	Since the emergence of SARS-CoV in 2002, China has been the focus of an intense viral surveillance and						
66	a large number of diverse bat-CoVs has been discovered in the region ³⁶⁻⁴⁴ . However, the macroevolution						
67	of CoVs in their bat hosts in China and their cross-species transmission dynamics remain poorly						
68	understood.						
69	In this study, we analyze an extensive field-collected dataset of bat-CoV sequences from across China.						
70	We use a phylogeographic Bayesian statistical framework to reconstruct virus transmission history						
71	between different bat host species and virus spatial spread over evolutionary time. Our objectives were						
72	to compare the macroevolutionary patterns of $lpha-$ and $eta-$ CoVs and identify the hosts and geographical						
73	regions that act as centers of evolutionary diversification for bat-CoVs in China. These analyses aim to						
74	improve our understanding of how CoVs evolve, diversify, circulate among, and transmit between bat						
75	families and genera to help identify bat hosts and regions where the risk of CoV spillover is the highest.						

76 Results

77 Taxonomic and geographic sampling

78	We generated 630 partial sequences (440 nt) of the RNA-dependent RNA polymerase (RdRp) gene from
79	bat rectal swabs collected in China and added 608 bat-CoV and eight pangolin CoV sequences from
80	China available in GenBank or GISAID to our datasets (list of GenBank and GISAID accession numbers
81	available in Supplementary Note 1). For each CoV genus, two datasets were created: one including all
82	bat-CoV sequences with known host (host dataset) and one including all bat-CoV sequences with known
83	sampling location at the province level (geographic dataset). To create a geographically discrete
84	partitioning scheme that was more ecologically relevant than administrative borders for our
85	phylogeographic reconstructions, we defined six zoogeographic regions within China by clustering
86	provinces with similar mammalian diversity using hierarchical clustering ⁴⁵ (see Methods): South western
87	region (SW), Northern region (NO), Central northern region (CN), Central region (CE), Southern region
88	(SO) and Hainan island (HI) (Fig. 1 and Supplementary Fig. 1).
89	Our host datasets included 701 $lpha$ -CoV sequences (353 new sequences, including 102 new SADSr-CoV
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97 Ancestral hosts and cross-species transmission

98 We used a Bayesian discrete phylogeographic approach implemented in BEAST⁴⁶ to reconstruct the

ancestral host of each node in the phylogenetic tree using bat host family as a discrete character state.

100	The phylogenetic reconstructions for α -CoVs in China suggest an evolutionary origin within rhinolophid
101	and vespertilionid bats (Fig. 2A). The first α -CoV lineage to diverge historically corresponds to the
102	subgenus Rhinacovirus (L1), originating within rhinolophid bats, and includes sequences related to
103	HKU2-CoV and SADS-CoV (Supplementary Fig. 2). Then several lineages, labelled L2 to L7, emerged from
104	vespertilionid bats (Fig. 2A). The subgenus Decacovirus (L2) includes sequences mostly associated with
105	the Rhinolophidae and Hipposideridae and related to HKU10-CoV (Supplementary Fig. 3), while the
106	subgenera Myotacovirus (L3) and Pedacovirus (L5) as well as an unidentified lineage (L4) include CoVs
107	mainly from vespertilionid bats and related to HKU6-, HKU10-, and 512-CoVs (Supplementary Fig. 4-5).
108	Finally, a well-supported node comprises the subgenera Nyctacovirus (L6) from vespertilionid bats and
109	Minunacovirus (L7) from miniopterid bats, and includes HKU7-, HKU8-, 1A-, and 1B-CoVs
110	(Supplementary Fig. 6). These seven $lpha$ -CoV lineages are mostly associated with a single host family but
111	each also included several sequences identified from other bat families (Fig. 2A, Supplementary Fig. 2-6
112	and Supplementary Table 1), suggesting frequent cross-species transmission events have occurred
113	among bats. Ancestral host reconstructions based on the random data subset, to normalize sampling
114	effort, gave very similar results with rhinolophids and vespertilionids being the most likely ancestral
115	hosts of most α -CoV lineages too (Supplementary Fig. 7A). However, the topology of the tree based on
116	the random subset was slightly different as the lineage L5 was paraphyletic.
117	Chinese β -CoVs likely originated from vespertilionid and rhinolophid bats (Fig. 2B). The MCC tree was
118	clearly structured into four main lineages: Merbecovirus (Lineage C), including MERS-related (MERSr-)
119	CoVs, HKU4- and HKU5-CoVs and strictly restricted to vespertilionid bats (Supplementary Fig. 8);
120	Nobecovirus (lineage D), originating from pteropodid bats and corresponding to HKU9-CoV
121	(Supplementary Fig. 9); Hibecovirus (lineage E) comprising sequences isolated in hipposiderid bats
122	(Supplementary Fig. 10) and Sarbecovirus (Lineage B) including sequences related to HKU3- and SARS-
123	related (SARSr-) CoVs originating in rhinolophid bats (Supplementary Fig. 11). We show that SARS-CoV-2

124 forms a divergent clade within Sarbecovirus and is most closely related to viruses sampled from 125 Rhinolophus malayanus and R. affinis and from Malayan pangolins (Manis javanica) (Fig. 3). Similar tree 126 topology and ancestral host inference were obtained with the random subset (Supplementary Fig. 7B). 127 We used a Bayesian Stochastic Search Variable Selection (BSSVS) procedure⁴⁷ to identify viral host 128 switches (transmission over evolutionary time) between bat families and genera that occurred along the 129 branches of the MCC annotated tree and calculated Bayesian Factor (BF) to estimate the significance of 130 these switches (Fig. 4). We identified nine highly supported (BF > 10) inter-family host switches for α -131 CoVs and three for β -CoVs (Fig. 4A and 4B). These results are robust over a range of sample sizes, with 132 seven of these nine switches for α -CoVs and the exact same three host switches for β -CoVs having 133 strong BF support (BF > 10) when analyzing our random subset (Supplementary Tables 2 and 3). To 134 quantify the magnitude of these host switches, we estimated the number of host switching events (Markov jumps)^{48,49} along the significant inter-family switches (Fig. 4C and 4D) and estimated the rate of 135 136 inter-family host switching events per unit of time for each CoV genus. The rate of inter-family host 137 switching events was five times higher in the evolutionary history of α - (0.010 host switches/unit time) 138 than β -CoVs (0.002 host switches/unit time) in China. For α -CoVs, host switching events from the 139 Rhinolophidae and the Miniopteridae were greater than from other bat families while rhinolophids were 140 the highest donor family for β -CoVs. The Rhinolophidae and the Vespertilionidae for α -CoVs and the 141 Hipposideridae for β -CoVs received the highest numbers of switching events (Fig. 4C and 4D). When 142 using the random dataset, similar results were obtained for β -CoVs while rhinolophids were the highest 143 donor family for α -CoVs (Supplementary Tables 4 and 5).

At the genus level, we identified 20 highly supported inter-genus host switches for α-CoVs, 17 of them
were also highly significant using the random subset (Fig. 5A and Supplementary Table 6). Sixteen highly
supported inter-genus switches were identified for β-CoVs (Fig. 5B). Similar results were obtained for
the random β-CoV subset (Supplementary Table 7). Most of the significant cross-genus CoV switches for

148 α -CoVs, 15 of 20 (75%), were between genera in different bat families, while this proportion was only 6 149 of 16 (37.5%) for β -CoVs. The estimated rate of inter-genus host switching events (Markov jumps) was 150 similar for α - (0.014 host switches/unit time) and β -CoVs (0.014 host switches/unit time). For α -CoVs, 151 *Rhinolophus* and *Miniopterus* were the greatest donor genera and *Rhinolophus* was the greatest receiver 152 (Supplementary Table 8). For β -CoVs, *Rousettus* was the greatest donor and *Eonycteris* the greatest 153 receiver genus (Supplementary Table 9).

154 CoV spatiotemporal dispersal in China

155 We used our Bayesian discrete phylogeographic model with zoogeographic regions as character states 156 to reconstruct the spatiotemporal dynamics of CoV dispersal in China. Eleven and seven highly 157 significant (BF > 10) dispersal routes within China were identified for α - and β -CoVs, respectively (Fig. 6). 158 Seven and five of these dispersal routes, respectively, remained significant when using our random 159 subsets (Supplementary Tables 10 and 11). The Rhinacovirus lineage (L1) that includes HKU2- and SADS-160 CoV likely originated in the SO region while all other α -CoV lineages historically arose in SW China and 161 spread to other regions before several dispersal events from SO and NO in all directions (Fig. 6A and 162 Supplementary Fig. 12). A roughly similar pattern of α -CoV dispersal was obtained using the random 163 subset (Supplementary Tables 10 and 12).

The oldest inferred dispersal movements for β-CoVs occurred among the SO and SW regions (Fig. 6B).
The SO region was the likely origin of *Merbecovirus* (Lineage C, including HKU4- and HKU5-CoV) and *Sarbecovirus* subgenera (Lineage B, including HKU3- and SARSr-CoVs) while the *Nobecovirus* (lineage D,
including HKU9-CoV) and *Hibecovirus* (lineage E) subgenera originated in SW China (Supplementary Fig.
12). Then several dispersal movements likely originated from SO and CE (Fig. 6B). More recent
southward dispersal from NO was observed. Similar spatiotemporal dispersal patterns were observed
using the random subset of β-CoVs (Supplementary Tables 11 and 13).

171	The estimated rate of migration events per unit of time along these significant dispersal routes was
172	more than two times higher for α - (0.026 host switches/unit time) than β -CoVs (0.011 host switches/unit
173	time) and SO was the region involved in the greatest total number of migration events for both α - and β -
174	CoVs. SO had the highest number of outbound and inbound migration events for α -CoVs (Fig. 6C and
175	Supplementary Table 12). For β -CoVs, the highest number of outbound migration events was estimated
176	to be from NO and SO while SO and SW had the highest numbers of inbound migration events (Fig. 6D
177	and Supplementary Table 13).
178	Phylogenetic diversity
179	In order to identify the hotspots of CoV phylogenetic diversity in China and evaluate phylogenetic
180	clustering of CoVs, we calculated the Mean Phylogenetic Distance (MPD) and the Mean Nearest Taxon
181	Distance (MNTD) statistics ⁵⁰ and their standardized effect size (SES).

182 We found significant and negative SES MPD values, indicating significant phylogenetic clustering, within 183 all bat families and genera for both α - and β -CoVs, except within the Aselliscus and Tylonycteris for α -184 CoVs (Fig. 7A and 7B). Negative and mostly significant SES MNTD values, reflecting phylogenetic 185 structure closer to the tips, were also observed within most bat families and genera for α - and β -CoVs 186 but we found non-significant positive SES MNTD value for vespertilionid bats, and particularly for those 187 in the *Pipistrellus* genus, for β -CoVs (Fig. 7A and 7B). In general, we observed lower phylogenetic 188 diversity for β - than α -CoVs within all bat families and most genera when looking at SES MPD, but the 189 difference in the level of diversity between α - and β -CoVs is less important when looking at SES MNTD 190 (Fig. 7). These results suggest stronger basal clustering (reflected by larger SES MPD values) for β -CoVs 191 than α -CoVs, indicating stronger host structuring effect and phylogenetic conservatism for β -CoVs. Very 192 similar results were obtained with the random subsets for both α - and β -CoVs (Supplementary Tables 193 14-21).

194 We found negative and mostly significant values of MPD and MNTD (Fig. 7C and Supplementary Tables 195 22-25) indicating significant phylogenetic clustering of CoV lineages in bat communities within the same 196 zoogeographic region. However, SES MPD values for α -CoVs in SW were positive (significant for the 197 random subset) indicating a greater evolutionary diversity of CoVs in that region than others (Fig. 7 and 198 Supplementary Tables 22-25). We used a linear regression analysis to assess the relationship between 199 CoV phylogenetic diversity and bat species richness in China and determine if bat richness is a significant 200 predictor of bat-CoV diversity and evolution. α -CoV phylogenetic diversity (MPD) was not significantly 201 correlated to total bat species richness or sampled bat species richness in zoogeographic regions or 202 provinces (Supplementary Table 26). Non-significant correlations between bat species richness and β -203 CoV phylogenetic diversity were also observed at the zoogeographic region level (Supplementary Table 204 27). However, a significant correlation was observed between sampled bat species richness and β -CoV 205 phylogenetic diversity at the province level (Supplementary Table 27). Similar results were obtained 206 when using the random subsets (Supplementary Tables 26 and 27). These findings suggest that bat host 207 diversity is not the main driver of CoV diversity in China and that other ecological or biogeographic 208 factors may influence this diversity. We observed higher CoV diversity than expected in several southern 209 or central provinces (Hainan, Guangxi, Hunan) given their underlying total or sampled bat diversity 210 (Supplementary Fig. 13 and 14).

We also assessed patterns of CoV phylogenetic turnover/differentiation among Chinese zoogeographic
regions and bat host families by measuring the inter-region and inter-host values of MPD (equivalent to
a measure of phylogenetic β-diversity) and their SES. We found positive inter-family SES MPD values,
except between Pteropodidae and Hipposideridae for α-CoVs and between Rhinolophidae and
Hipposideridae for β-CoVs (Fig. 8A and 8B and Supplementary Tables 28 and 29), suggesting higher
phylogenetic differentiation of CoVs among most bat families than among random communities. Our
phylo-ordination based on inter-family MPD values indicated that α-CoVs from vespertilionids and

218	miniopterids, and from hipposiderids and pteropodids; as well as eta -CoVs from rhinolophids and
219	hipposiderids are phylogenetically closely related (Fig. 8A and 8B). We also observed strong
220	phylogenetic turnover between α -CoV strains from rhinolophids and from miniopterids and all other bat
221	families, and between β -CoV strains from vespertilionids and all other bat families (Supplementary
222	Tables 28 and 29). Phylo-ordination among bat genera based on inter-genus MPD confirmed these
223	results and indicated that CoV strains from genera belonging to the same bat family were mostly more
224	closely related to each other than to genera from other families (Fig. 8C and 8D and Supplementary
225	Tables 30 and 31).
226	We observed high and positive inter-region SES MPD values between SW/HI and all other regions,
227	suggesting that these two regions host higher endemic diversity (Fig. 9 and Supplementary Tables 32
228	and 31). Negative inter-region SES MPD values suggested that the phylogenetic turnover among other
229	regions was less important than expected among random communities. Our phylo-ordination among
230	zoogeographic regions also reflected the high phylogenetic turnover and deep evolutionary

distinctiveness of both α - and β -CoVs from SW and HI regions (Fig. 9 and Supplementary Tables 32 and

232 33). Similar results were obtained using the random subset (Supplementary Tables 32 and 33).

233 Mantel tests

234 Mantel tests revealed a positive and significant correlation between CoV genetic differentiation (F_{ST}) and

235 geographic distance matrices, both with and without provinces including fewer than four viral

236 sequences, for α- (r = 0.25, p = 0.0097; r = 0.32, p = 0.0196; respectively) and β-CoVs (r = 0.22, p =

237 0.0095; r = 0.23, p = 0.0336; respectively). We also detected a positive and highly significant correlation

- between CoV genetic differentiation (F_{ST}) and their host phylogenetic distance matrices, both with and
- without genera including fewer than four viral sequences, for β -CoVs (r = 0.41, p = 0; r = 0.39, p = 0
- 240 0.0012; respectively) but not for α -CoVs (r = -0.13, p = 0.8413; r = 0.02, p = 0.5019; respectively).

241 Discussion

242 Our phylogenetic analysis shows a high diversity of CoVs from bats sampled in China, with most bat 243 genera included in this study (10/16) infected by both α - and β -CoVs. In our phylogenetic analysis that 244 includes all known bat-CoVs from China, we find that SARS-CoV-2 is likely derived from a clade of viruses 245 originating in horseshoe bats (Rhinolophus spp.). The geographic location of this origin appears to be 246 Yunnan province. However, it is important to note that: 1) our study collected and analyzed samples 247 solely from China; 2) many sampling sites were close to the borders of Myanmar and Lao PDR; and 3) 248 most of the bats sampled in Yunnan also occur in these countries, including R. affinis and R. malayanus, the species harboring the CoVs with highest RdRp sequence identity to SARS-CoV-2^{20,21}. For these 249 250 reasons, we cannot rule out an origin for the clade of viruses that are progenitors of SARS-CoV-2 that is 251 outside China, and within Myanmar, Lao PDR, Vietnam or another Southeast Asian country. Additionally, 252 our analysis shows that the virus RmYN02 from *R. malayanus*, which is characterized by the insertion of 253 multiple amino acids at the junction site of the S1 and S2 subunits of the Spike (S) protein, belongs to 254 the same clade as both RaTG13 and SARS-CoV-2, providing further support for the natural origin of SARS-CoV-2 in *Rhinolophus* spp. bats in the region^{20,21}. Finally, while our analysis shows that the RdRp 255 256 sequences of coronaviruses from the Malayan pangolin are closely related to SARS-CoV-2 RdRp, analysis 257 of full genomes of these viruses suggest that these terrestrial mammals are less likely to be the origin of SARS-CoV-2 than *Rhinolophus* spp. bats^{22,23}. 258

This analysis also demonstrates that a significant amount of cross-species transmission has occurred among bat hosts over evolutionary time. Our Bayesian phylogeographic inference and analysis of host switching showed varying levels of viral connectivity among bat hosts and allowed us to identify significant host transitions that appear to have occurred during bat-CoV evolution in China.

263 We found that bats in the family Rhinolophidae (horseshoe bats) played a key role in the evolution and 264 cross-species transmission history of α -CoVs. The family Rhinolophidae and the genus Rhinolophus were 265 involved in more inter-family and inter-genus highly significant host switching of α -CoVs than any other 266 family or genus. They were the greatest receivers of α -CoV host switching events and second greatest 267 donors after Miniopteridae/Miniopterus. The Rhinolophidae, together with the Hipposideridae, also 268 played an important role in the evolution of β -CoVs, being at the origin of most inter-family host 269 switching events. Chinese horseshoe bats are characterized by a distinct and evolutionary divergent α -270 CoV diversity, while their β -CoV diversity is similar to that found in the Hipposideridae. The 271 Rhinolophidae comprises a single genus, Rhinolophus, and is the most speciose bat family after the Vespertilionidae in China⁵¹, with 20 known species, just under a third of global *Rhinolophus* diversity, 272 mostly in Southern China³⁵. This family likely originated in Asia^{52,53}, but some studies suggest an African 273 origin^{54,55}. Rhinolophid fossils from the middle Eocene (38 - 47.8 Mya) have been found in China, 274 suggesting a westward dispersal of the group from eastern Asia to Europe⁵⁶. The ancient likely origin of 275 276 the Rhinolophidae in Asia and China in particular may explain the central role they played in the 277 evolution and diversification of bat-CoVs in this region, including SARSr-CoVs, MERS-cluster CoVs, and 278 SADSr-CoVs, which contain important human and livestock pathogens. Horseshoe bats are known to share roosts with genera from all other bat families in this study⁵⁷, which may also favor CoV cross-279 species transmission from and to rhinolophids³⁴. A global meta-analysis showing higher rates of viral 280 sharing among co-roosting cave bats supports this finding⁵⁸. 281

Vespertilionid and miniopterid bats (largely within the *Myotis* and *Miniopterus* genera) also appear to
 have been involved in several significant host switches during α-CoV evolution. However, no significant
 transition from vespertilionid bats was identified for β-CoVs and these bats exhibit a divergent β-CoV
 diversity compared to other bat families. Vespertilionid and miniopterid bats are characterized by strong
 basal phylogenetic clustering but high recent CoV diversification rates, indicating a more rapid

evolutionary radiation of CoVs in these bat hosts. At the genus level, similar findings were observed for
the genera *Myotis*, *Pipistrellus* and *Miniopterus*.

289 A significant correlation between geographic distance and genetic differentiation of both α - and β -CoVs 290 has been detected, even if only a relatively small proportion of the variance is explained by geographic 291 distance. We also revealed a significant effect of host phylogeny on β -CoV evolution while it had a 292 minimal effect on α -CoV diversity. Contrary to the α -CoV phylogeny, the basal phylogenetic structure of 293 β -CoVs mirrored the phylogeny of their bat hosts, with a clear distinction between the Yangochiroptera, 294 encompassing the Vespertilionidae and Miniopteridae, and the Yinpterochiroptera, which includes the 295 megabat family Pteropodidae and the microbat families Rhinolophidae and Hipposideridae, as evidenced in recent bat phylogenies^{52,59}. These findings suggest a profound co-macroevolutionary 296 297 process between β -CoVs and their bat hosts, even if host switches also occurred throughout their 298 evolution as our study showed. The phylogenetic structure of α -CoVs, with numerous and closely related 299 lineages identified in the Vespertilionidae and Miniopteridae, contrasts with the β -CoV 300 macroevolutionary pattern and suggests α -CoVs have undergone an adaptive radiation in these two 301 Yangochiroptera families. Our BSSVS procedure and Markov jump estimates revealed higher 302 connectivity, both qualitatively and quantitatively, among bat families and genera in the α -CoV cross-303 species transmission history. Larger numbers of highly significant host transitions and higher rates of 304 switching events along these pathways were inferred for α - than β -CoVs, especially at the host family 305 level. These findings suggest that α -CoVs are able to switch hosts more frequently and between more 306 distantly related taxa, and that phylogenetic distance among hosts represents a higher constraint on 307 host switches for β - than α -CoVs. This is supported by more frequent dispersal events in the evolution of 308 α - than β -CoVs in China.

Variation in the extent of host jumps between α and β-CoVs within the same hosts in the same
 environment may be due to virus-specific factors such as differences in receptor usage between α- and

311	β -CoVs ⁶⁰⁻⁶² . Coronaviruses use a large diversity of receptors, and their entry into host cells is mediated
312	by the spike protein with an ectodomain consisting of a receptor-binding subunit S1 and a membrane-
313	fusion subunit S2 ⁶³ . However, despite differences in the core structure of their S1 receptor binding
314	domains (RBD), several $lpha$ - and eta -CoV species are able to recognize and bind to the same host
315	receptors ⁶⁴ . Other factors such as mutation rate, recombination potential, or replication rate might also
316	be involved in differences in host switching potential between α - and β -CoVs. A better understanding of
317	receptor usage and other biological characteristics of these bat-CoVs may help predict their cross-
318	species transmission and zoonotic potential.
319	We also found that some bat genera were infected by a single CoV genus: <i>Miniopterus</i> (Miniopteridae)
320	and <i>Murina</i> (Vespertilionidae) carried only α -CoVs, while <i>Cynopterus</i> , <i>Eonycteris</i> , <i>Megaerops</i>
321	(Pteropodidae) and <i>Pipistrellus</i> (Vespertilionidae) hosted only β -CoVs. This was found despite using the
322	same conserved pan-CoV PCR assays for all specimens screened and it can't be explained by differences
323	in sampling effort for these genera (Supplementary Table 1): for example, >250 α -CoV sequences but no
324	β -CoV were discovered in <i>Miniopterus</i> bats in China during our recent fieldwork. These migratory bats,
325	which seem to have played a key role in the evolution of α -CoVs, share roosts with several other bat
326	genera hosting β -CoVs in China ⁵⁷ , suggesting high likelihood of being exposed to β -CoVs. Biological or
327	ecological properties of miniopterid bats may explain this observation and clearly warrant further
328	investigation.
329	Our Bayesian ancestral reconstructions revealed the importance of South western and Southern China
330	as centers of diversification for both α - and β -CoVs. These two regions are hotspots of CoV phylogenetic
331	diversity, harboring evolutionarily old and phylogenetically diverse lineages of $lpha-$ and $eta-$ CoVs. South

332 western China acted as a refugium during Quaternary glaciation for numerous plant and animal species

- including several bat species, such as *Rhinolophus affinis*⁶⁵, *Rhinolophus sinicus*⁶⁶, *Myotis davidii*⁶⁷, and
- 334 *Cynopterus sphinx*⁶⁸. The stable and long-term persistence of bats and other mammals throughout the

Quaternary may explain the deep macroevolutionary diversity of bat-CoVs in these regions⁶⁹. Several highly significant and ancient CoV dispersal routes from these two regions have been identified in this study. Other viruses, such as the Avian Influenza A viruses H5N6, H7N9 and H5N1, also likely originated in South western and Southern Chinese regions^{70,71}.

339 Our findings suggest that bat host diversity is not the main driver of CoV diversity in China and that 340 other ecological or biogeographic factors may influence this diversity. Overall, there were no significant 341 correlations between CoV phylogenetic diversity and bat species diversity (total or sampled) for each 342 province or biogeographic region, apart from a weak correlation between β -CoV phylogenetic diversity 343 and the number of bat species sampled at the province level. Yet, we observed higher than expected 344 phylogenetic diversity in several southern provinces (Hainan, Guangxi, Hunan). These results and main 345 conclusions are consistent and robust even when we account for geographic biases in sampling effort by 346 analyzing random subsets of the data.

347 Despite being the most exhaustive study of bat-CoVs in China, this study had several limitations that 348 must be taken into consideration when interpreting our results. First, only partial RdRp sequences were 349 generated in this study and used in our phylogenetic analysis as the non-invasive samples (rectal 350 swabs/feces) collected in this study prevented us from generating longer sequences in many cases. The 351 *RdRp* gene is a suitable marker for this kind of study as it reflects vertical ancestry and is less prone to recombination than other regions of the CoV genome such as the spike protein gene^{16,72}. While using 352 353 long sequences is always preferable, our phylogenetic trees are well supported and their topology consistent with trees obtained using longer sequences or whole genomes^{30,73}. Second, most sequences 354 355 in this study were obtained by consensus PCR using primers targeting highly conserved regions. Even if 356 this broadly reactive PCR assay designed to detect widely variant CoVs has proven its ability to detect a large diversity of CoVs in a wide diversity of bats and mammals^{30,74-77}, we may not rule out that some 357

bat-CoV variants remained undetected. Using deep sequencing techniques would allow to detect this
unknown and highly divergent diversity.

360 In this study, we identified the host taxa and geographic regions that together define hotspots of CoV 361 phylogenetic diversity and centers of diversification in China. These findings may provide a strategy for 362 targeted discovery of bat-borne CoVs of zoonotic or livestock infection potential, and for early detection of bat-CoV outbreaks in livestock and people, as proposed elsewhere⁷⁸. Our results suggest that future 363 364 sampling and viral discovery should target two hotspots of CoV diversification in Southern and South 365 western China in particular, as well as neighboring countries where similar bat species live. These 366 regions are characterized by a subtropical to tropical climate; dense, growing and rapidly urbanizing populations of people; a high degree of poultry and livestock production; and other factors which may 367 promote cross-species transmission and disease emergence⁷⁸⁻⁸⁰. Additionally, faster rates of evolution in 368 369 the tropics have been described for other RNA viruses which could favor cross-species transmission of RNA viruses in these regions⁸¹. Both SARS-CoV and SADS-CoV emerged in this region, and several bat 370 371 SARSr-CoVs with high zoonotic potential have recently been reported from there, although the dynamics of their circulation in wild bat populations remain poorly understood^{16,61}. Importantly, the closest known 372 relative of SARS-CoV-2, a SARS-related virus, was found in a *Rhinolophus* sp. bat in this region²⁰, 373 374 although it is important to note that our survey was limited to China, and that the bat hosts of this virus 375 also occur in nearby Myanmar and Lao PDR. The significant public health and food security implications 376 of these outbreaks reinforces the need for enhanced, targeted sampling and discovery of novel CoVs. 377 Because intensive sampling has not, to our knowledge, been undertaken in countries bordering 378 southern China, these surveys should be extended to include Myanmar, Lao PDR, and Vietnam, and 379 perhaps across southeast Asia. Our finding that Rhinolophus spp. are most likely to be involved in host-380 switching events makes them a key target for future longitudinal surveillance programs, but surveillance

targeted the genera *Hipposideros* and *Aselliscus* may also be fruitful as they share numerous β-CoVs
 with *Rhinolophus* bats.

383 In the aftermath of the SARS-CoV and MERS-CoV outbreaks, β -CoVs have been the main focus of bat-CoV studies in China, Africa, and Europe^{17,32,36,61,82}. However, we have shown that α -CoVs have a higher 384 propensity to switch host within their natural bat reservoirs, and therefore also have a high cross-385 386 species transmission potential and risk of spillover. This is exemplified by the recent emergence of SADS-387 CoV in pigs in Guangdong province¹⁷. Two human α -CoVs, NL63 and 229E, also likely originated in bats^{27,28}, reminding us that past spillover events from bat species can readily be established in the 388 389 human population. Future work discovering and characterizing the biological properties of bat α -CoVs 390 may therefore be of potential value for public and livestock health. Our study, and recent analysis of viral discovery rates⁸³, suggest that a substantially wider sampling and discovery net will be required to 391 392 capture the complete diversity of coronaviruses in their natural hosts and assess their potential for 393 cross-species transmission. The bat genera Rhinolophus, Hipposideros, Myotis and Miniopterus, all 394 involved in numerous naturally-occurring host switches throughout α -CoV evolution, should be a 395 particular target for α -CoV discovery in China and across southeast Asia, with *in vitro* and experimental 396 characterization to better understand their potential to infect people or livestock and cause disease.

397 Methods

398 Bat sampling

Bat oral and rectal swabs and fecal pellets were collected from 2010 to 2015 in numerous Chinese
provinces (Anhui, Beijing, Guangdong, Guangxi, Guizhou, Hainan, Henan, Hubei, Hunan, Jiangxi, Macau,
Shanxi, Sichuan, Yunnan, and Zhejiang). Fecal pellets were collected from tarps placed below bat
colonies. Bats were captured using mist nets at their roost site or feeding areas. Each captured bat was
stored into a cotton bag, all sampling was non-lethal and bats were released at the site of capture

404 immediately after sample collection. A wing punch was also collected for barcoding purport	se. Bat-
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405 handling methods were approved by Tufts University IACUC committee (proposal #G2017-32) and

406 Wuhan Institute of Virology Chinese Academy of Sciences IACUC committee (proposal WIVA05201705).

407 Samples were stored in viral transport medium at -80°C directly after collection.

408 RNA extraction and PCR screening

- 409 RNA was extracted from 200 μl swab rectal samples or fecal pellets with the High Pure Viral RNA Kit
- 410 (Roche) following the manufacturer's instructions. RNA was eluted in 50 μl elution buffer and stored at -
- 411 80°C. A one-step hemi-nested RT-PCR (Invitrogen) was used to detect coronavirus RNA using a set of
- 412 primers targeting a 440-nt fragment of the *RdRp* gene and optimized for bat-CoV detection (CoV-FWD3:

413 GGTTGGGAYTAYCCHAARTGTGA; CoV-RVS3: CCATCATCASWYRAATCATCATA; CoV-FWD4/Bat:

414 GAYTAYCCHAARTGTGAYAGAGC)⁸⁴. For the first round PCR, the amplification was performed as follows:

415 50°C for 30 min, 94°C for 2 min, followed by 40 cycles consisting of 94°C for 20 sec, 50°C for 30 sec, 68°C

416 for 30 sec, and a final extension step at 68°C for 5 min. For the second round PCR, the amplification was

417 performed as follows: 94°C for 2 min followed by 40 cycles consisting of 94°C for 20 sec, 59°C for 30 sec,

418 72°C for 30 sec, and a final extension step at 72°C for 7 min. PCR products were gel purified and

- 419 sequenced with an ABI Prism 3730 DNA analyzer (Applied Biosystems, USA). PCR products with low
- 420 concentration or bad sequencing quality were cloned into pGEM-T Easy Vector (Promega) for
- 421 sequencing. Positive results detected in bat genera that were not known to harbor a specific CoV lineage

422 previously were repeated a second time (PCR + sequencing) as a confirmation. Species identifications

423 from the field were also confirmed and re-confirmed by cytochrome (cytb) DNA barcoding using DNA

424 extracted from the feces or swabs⁸⁵. Only viral detection and barcoding results confirmed at least twice

425 were included in this study.

426 Sequence data

427 We also added bat-CoV RdRp sequences from China available in GenBank to our dataset. All sequences 428 for which sampling year and host or sampling location information was available either in GenBank 429 metadata or in the original publication were included (as of March 15, 2018). Our final datasets include 430 630 sequences generated for this study and 616 sequences from GenBank or GISAID (list of GenBank 431 and GISAID accession numbers available in Supplementary Note 1, and Supplementary Tables 34 and 432 35). Nucleotide sequences were aligned using MUSCLE and trimmed to 360 base pair length to reduce 433 the proportion of missing data in the alignments. All phylogenetic analyses were performed on both the 434 complete data and random subset, and for α - and β -CoVs separately.

435 Defining zoogeographic regions in China

436 Hierachical clustering was used to define zoogeographic regions within China by clustering provinces with similar mammalian diversity⁴⁵. Hierarchical cluster analysis classifies several objects into small 437 438 groups based on similarities between them. To do this, we created a presence/absence matrix of all extant terrestrial mammals present in China using data from the IUCN spatial database⁸⁶ and generated 439 440 a cluster dendrogram using the function *hclust* with average method of the R package stats. Hong Kong 441 and Macau were included within the neighboring Guangdong province. We then visually identified 442 geographically contiguous clusters of provinces for which CoV sequences are available (Fig. 1 and 443 Supplementary Fig. 1).

We identified six zoogeographic regions within China based on the similarity of the mammal community in these provinces: South western region (SW; Yunnan province), Northern region (NO; Xizang, Gansu, Jilin, Anhui, Henan, Shandong, Shaanxi, Hebei and Shanxi provinces and Beijing municipality), Central northern region (CN; Sichuan and Hubei provinces), Central region (CE; Guangxi, Guizhou, Hunan, Jiangxi and Zhejiang provinces), Southern region (SO; Guangdong and Fujian provinces, Hong Kong, Macau and Taiwan), and Hainan island (HI). Hunan and Jiangxi, clustering with the SO provinces in our dendrogram,

450 were included within the central region to create a geographically contiguous Central cluster

451 (Supplementary Fig. 1). These six zoogeographic regions are very similar to the biogeographic regions

452 traditionally recognized in China⁸⁷. The three β -CoV sequences from HI were included in the SO region to

453 avoid creating a cluster with a very small number of sequences.

454 Model selection and phylogenetic analysis

455 Bayesian phylogenetic analysis were performed in BEAST 1.8.4⁴⁶. Sampling years were used as tip dates.

456 Preliminary analysis were run to select the best fitting combination of substitution models (HKY/GTR),

457 codon partition scheme, molecular clock (strict/lognormal uncorrelated relaxed clock) and coalescent

458 models (constant population size/exponential growth/GMRF Bayesian Skyride). Model combinations

459 were compared and the best fitting model was selected using a modified Akaike information criterion

460 (AICM) implemented in Tracer 1.6^{88} . We also used TEMPEST⁸⁹ to assess the temporal structure within

461 our α - and β -CoV datasets. TEMPEST showed that both datasets did not contain sufficient temporal

462 information to accurately estimate substitution rates or time to the most recent common ancestor

463 (TMRCA). Therefore we used a fixed substitution rate of 1.0 for all our BEAST analysis.

All subsequent BEAST analysis were performed under the best fitting model including a HKY substitution model with two codons partitions ((1+2), 3), a strict molecular clock and a constant population size coalescent model. Each analysis was run for 2.5 x 10⁸ generations, with sampling every 2 x 10⁴ steps. All BEAST computations were performed on the CIPRES Science Getaway Portal⁹⁰. Convergence of the chain was assessed in Tracer so that the effective sample size (ESS) of all parameters was > 200 after removing at least 10% of the chain as burn-in.

470 Ancestral state reconstruction and transition rates

A Bayesian discrete phylogeographic approach implemented in BEAST 1.8.4 was used to reconstruct the
ancestral state of each node in the phylogenetic tree for three discrete traits: host family, host genus

and zoogeographic region. An asymmetric trait substitution model was applied. These analyses were
performed for each trait on the complete dataset and random subsets. Maximum clade credibility (MCC)
tree annotated with discrete traits were generated in TreeAnnotator and visualized using the software
SpreaD3⁹¹.

477	For each analysis, a Bayesian stochastic search variable selection (BSSVS) was applied to estimate the
478	significance of pairwise switches between trait states using Bayesian Factor (BF) as a measure of
479	statistical significance ⁴⁷ . BF were computed in SpreaD3. BF support was interpreted according to Jeffreys
480	1961 ⁹² (BF > 3: substantial support, BF > 10: strong support, BF > 30: very strong support, BF > 100:
481	decisive support) and only strongly supported transitions were presented in most figures, following a
482	strategy used in other studies ^{93,94} . We also estimated the count of state switching events (Markov
483	jumps) ^{48,49} along the branches of the phylogenetic tree globally (for the three discrete traits) and for
484	each strongly supported (BF > 10) transition between character states (for bat families and ecoregions
485	only). Convergence of the MCMC runs was confirmed using Tracer. The rate of state switching events
486	per unit of time was estimated for each CoV genus by dividing the total estimated number of state
487	switching events by the total branch length of the MCC tree.
488	To assess the phylogenetic relationships among SARS-CoV-2 and other CoVs from the Sarbecovirus
489	subgenus, we also reconstructed a MCC tree in BEAST 1.8.4 and median-joining network in Network
490	10.0 ⁹⁵ including all <i>Sarbecovirus</i> sequences, two sequences of SARS-CoV-2 isolated in humans (GenBank
491	accession numbers: MN908947 and MN975262), one sequence of SARS-CoV (GenBank accession
492	number: NC_004718), eight sequences from Malayan pangolins (<i>Manis javanica</i>) (GISAID accession
493	numbers: EPI_ISL_410538-410544, EPI_ISL_410721) and one from <i>Rhinolophus malayanus</i> (GISAID

494 accession number: EPI_ISL_412977).

495 **Phylogenetic diversity**

The Mean Phylogenetic Distance (MPD) and the Mean Nearest Taxon Distance (MNTD) statistics⁵⁰ and 496 497 their standardized effect size (SES) were calculated for each zoogeographic region, bat family and genus using the R package picante⁹⁶. MPD measures the mean phylogenetic distance among all pairs of CoVs 498 499 within a host or a region. It reflects phylogenetic structuring across the whole phylogenetic tree and 500 assesses the overall divergence of CoV lineages in a community. MNTD is the mean distance between 501 each CoV and its nearest phylogenetic neighbor in a host or region, and therefore it reflects the 502 phylogenetic structuring closer to the tips and shows how locally clustered taxa are. SES MPD and SES 503 MNTD values correspond to the difference between the phylogenetic distances in the observed 504 communities versus null communities. Low and negative SES values denote phylogenetic clustering, high 505 and positive values indicate phylogenetic over-dispersion while values close to 0 show random 506 dispersion. The SES values were calculated by building null communities by randomly reshuffling tip 507 labels 1000 times along the entire phylogeny. Phylogenetic diversity computations were performed on 508 both the complete dataset and random subset for each trait. A linear regression analysis was performed 509 in R to assess the correlation between CoV phylogenetic diversity (MPD) and bat species richness in 510 China. Total species richness per province or region was estimated using data from the IUCN spatial 511 database while sampled species richness corresponds to the number of bat species sampled and tested 512 for CoV per province or region in our datasets.

The inter-region and inter-host values of MPD (equivalent to phylogenetic β diversity), corresponding to the mean phylogenetic distance among all pairs of CoVs from two distinct hosts or regions, and their SES were estimated using the function *comdist* of the R package phylocomr⁹⁷. The matrices of inter-region and inter-host MPD were used to cluster zoogeographic regions and bat hosts in a dendrogram according to their evolutionary similarity (phylo-ordination) using the function *hclust* with complete linkage method of the R package stats (R core team). These computations were performed on both the complete dataset and random subset.

520 Mantel tests and isolation by distance

521	Mantel tests performed in ARLEQUIN 3.5 ⁹⁸ were used to compare the matrix of viral genetic					
522	differentiation (F_{st}) to matrices of host phylogenetic distance and geographic distance in order to					
523	evaluate the role of geographic isolation and host phylogeny in shaping CoV population structure. The					
524	correlation between these matrices was assessed using 10,000 permutations. To gain more resolution					
525	into the process of evolutionary diversification, these analyses were also performed at the host genus					
526	and province levels. To calculate phylogenetic distances among bat genera, we reconstructed a					
527	phylogenetic tree including a single sequence for all bat species included in our dataset. Pairwise					
528	patristic distances among tips were computed using the function $distTips$ in the R package adephylo ⁹⁹ .					
529	We then averaged all distances across genera to create a matrix of pairwise distances among bat					
530	genera. Pairwise Euclidian distances were measured between province centroids and log transformed.					
531	Mantel tests were performed with and without genera and provinces including less than four viral					
532	sequences to assess the impact of low sample size on our results.					
533	Data availability					
534	GenBank accession numbers of sequences generated in this study and previously published sequences					
535	included in our analysis are available in the Supplementary Note 1 and Supplementary Tables 34 and 35.					
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768 Author contributions

- 769 K.J.O., H.E.F, J.H.E., L-F.W., Z.S. and P.D. created the study design, initiated field work and set up sample
- collection and testing protocols. B.H., G.Z., L.Z., H.L., A.A.C and Z.L. collected samples or provided
- data. B.H., B.L., and W.Z. performed laboratory work. A.L. carried out the analyses and drafted the
- 772 manuscript with K.J.O, C.Z.-T. and P.D. All authors reviewed and edited the manuscript
- 773 **Competing interests**: The authors declare no competing interests.
- 774 Figure legends

Fig. 1 Geographic sampling. Pie chart (A) showing the number of sequences of each CoV genus (alphaCoVs and beta-CoVs) available for each zoogeographic region and map of China provinces (B) showing
the number of *RdRp* sequences available for each province, in bold grey for alpha-CoVs and black for
beta-CoVs. Province colors correspond to the zoogeographic region to which they belong: NO, Northern
region; CN, Central northern region; SW, South western region; CE, Central region; SO, Southern region;
HI, Hainan island. The three beta-CoV sequences from HI were included in the SO region. Provinces
colored in grey are those where CoV sequences are not available.

782 Fig. 2 Phylogenetic trees and ancestral host reconstructions. Alpha-CoV (A) and beta-CoV (B) maximum 783 clade credibility annotated trees using complete datasets of RdRp sequences and bat host family as 784 discrete character state. Pie charts located at the root and close to the deepest nodes show the state 785 posterior probabilities for each bat family. Branch colors correspond to the inferred ancestral family 786 with the highest probability. Branch lengths are scaled according to relative time units (clock rate = 1.0). 787 Well-supported nodes (posterior probability > 0.95) are indicated with a black dot. The ICTV approved 788 CoV subgenera were highlighted: Rhinacovirus (L1), Decacovirus (L2), Myotacovirus (L3), Pedacovirus 789 (L5), Nyctacovirus (L6), Minunacovirus (L7) and an unidentified lineage (L4) for alpha-CoVs; and 790 Merbecovirus (Lineage C), Nobecovirus (lineage D), Hibecovirus (lineage E) and Sarbecovirus (Lineage B) 791 for beta-CoVs.

Fig. 3 Phylogenetic relationships within the *Sarbecovirus* subgenus (beta-CoVs). Maximum clade
credibility tree (A) including 202 *RdRp* sequences from the *Sarbecovirus* subgenus isolated in bats, two
sequences of SARS-CoV-2 and one sequence of SARS-CoV isolated in humans and eight sequences
isolated in Malayan pangolins (*Manis javanica*). Well-supported nodes (posterior probability > 0.95) are
indicated with a black dot. Tip colors correspond to the host genus, SARS-CoV-2 sequences and SARSCoV sequence are highlighted in grey and black, respectively. Median-joining network (B) including 202 *RdRp* sequences from the *Sarbecovirus* lineage isolated in bats, two sequences of SARS-CoV-2 and one

799 sequence of SARS-CoV isolated in humans and eight sequences isolated in Malayan pangolins (Manis 800 *javanica*). Colored circles correspond to distinct CoV sequences, circle size is proportional to the number 801 of identical sequences in the data set. Small black circles represent median vectors (ancestral or 802 unsampled intermediate sequences). Branch length is proportional to the number of mutational steps 803 between haplotypes. 804 Fig. 4 Inter-family host switches. Strongly supported host switches between bat families for alpha- (A) 805 and beta-CoVs (B). Arrows indicate the direction of the switch; arrow thickness is proportional to the 806 switch significance level, only host switches supported by strong Bayes factor (BF) > 10 are shown. 807 Histograms of total number of host switching events (state changes counts using Markov jumps) from/to 808 each bat family along the significant inter-family switches for alpha- (C) and beta-CoVs (D). 809 Fig. 5 Inter-genus host switches. Strongly supported host switches between bat genera for alpha- (A) 810 and beta-CoVs (B) and their significance level (Bayes factor, BF). Only host switches supported by strong 811 BF values > 10 are shown. Line thickness is proportional to the switch significance level. Red lines 812 correspond to host switches among bat genera belonging to different families, black lines correspond to 813 host switches among bat genera from the same family. Arrows indicate the direction of the switch. 814 Genus names are colored according to the family they belong to using the same colors as in Fig. 2 and 3. 815 Fig. 6 CoV spatiotemporal dispersal in China. Strongly supported dispersal routes (Bayes factor, BF > 10) 816 over recent evolutionary history among China zoogeographic regions for alpha- (A) and beta-CoVs (B). 817 Arrows indicate the direction of the dispersal route; arrow thickness is proportional to the dispersal 818 route significance level. Darker arrow colors indicate older dispersal events. Histograms of total number 819 of dispersal events (Markov jumps) from/to each region along the significant dispersal routes for alpha-820 (C) and beta-CoVs (D). NO, Northern region; CN, Central northern region; SW, South western region; CE, 821 Central region; SO, Southern region; HI, Hainan island.

822	Fig. 7 Phylogenetic diversity. Metrics of CoV phylogenetic diversity within each bat family (A), genus (B)
823	and zoogeographic regions (C): standardized effect size of Mean Phylogenetic Distance (SES MPD), on
824	the left panels; and standardized effect size of Mean Nearest Taxon Distance (SES MNTD), on the right
825	panels. One-tailed p-values (quantiles) were calculated after randomly reshuffling tip labels 1000 times
826	along the entire phylogeny. Values departing significantly from the null model (p-value < 0.05) are
827	indicated with an asterisk, all exact p-values are available in Supplementary Tables 14-27. NO, Northern
828	region; CN, Central northern region; SW, South western region; CE, Central region; SO, Southern region;
829	HI, Hainan island.
830	Fig. 8 Phylogenetic diversity. Standardized effect size of Mean Phylogenetic Distance (SES MPD) and
831	phylogenetic ordination among bat host families (A, B) and genera (C, D) for alpha- and beta-CoVs.
832	Boxplots for each host family and genus show the mean (cross), median (dark line within the box),
833	interquartile range (box), 95% confidence interval (whisker bars), and outliers (dots), calculated from all
834	pairwise comparisons between bat families (n=10 for alpha-CoVs and n=6 for beta-CoVs) and genera
835	(n=91 for alpha-CoVs and n=105 for beta-CoVs).
836	Fig. 9 Phylogenetic diversity. Standardized effect size of Mean Phylogenetic Distance, SES MPD) and
837	phylogenetic ordination among zoogeographic regions for alpha- (A) and beta-CoVs (B). Boxplots for
838	each region show the mean (cross), median (dark line within the box), interquartile range (box), 95%
839	confidence interval (whisker bars), and outliers (dots), calculated from all pairwise comparisons between
840	regions (n=15 for alpha-CoVs and n=10 for beta-CoVs). NO, Northern region; CN, Central northern
841	region; SW, South western region; CE, Central region; SO, Southern region; HI, Hainan island.

843 Figure 1



846 Figure 2



851 Figure 3



853 Figure 4



854

856 Figure 5



857

859 Figure 6



861 Figure 7



864 Figure 8



866 Figure 9





867