1 Surveillance of 16 UK native bat species through conservationist networks

2

uncovers coronaviruses with zoonotic potential

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28 Abstract (max 150 words)

29 There has been limited characterisation of bat-borne coronaviruses in Europe. Here, we 30 screened for coronaviruses 48 faecal samples from 16 of the 17 bat species breeding in 31 the UK and collected through a bat rehabilitation and conservationist network. We 32 recovered nine (two novel) complete genomes across six bat species: four 33 alphacoronaviruses, a MERS-related betacoronavirus, and four closely-related 34 sarbecoviruses. We demonstrate that at least one of these sarbecoviruses can bind and 35 use the human ACE2 receptor for infecting human cells, albeit suboptimally. Additionally, 36 the spike proteins of these sarbecoviruses possess an R-A-K-Q motif, which lies only one 37 nucleotide mutation away from a furin cleavage site (FCS) that enhances infectivity in 38 other coronaviruses, including SARS-CoV-2. However, mutating this motif to an FCS 39 does not enable spike cleavage. Overall, while UK sarbecoviruses would require further 40 molecular adaptations to infect humans, their zoonotic risk is unknown but warrants closer 41 surveillance.

42 Introduction

43 The majority of emerging infectious diseases in humans are zoonotic - arising from animal-to-human transmission of a pathogen¹ - and more than 70% originate in wildlife². 44 45 Coronaviridae is a diverse family of viruses that can infect a broad range of animals and 46 are prone to zoonotic spillovers. There are seven major coronaviruses that can infect 47 humans: SARS-CoV-2 is the agent of the COVID-19 pandemic whose direct ancestor has 48 not vet been identified but its closest relatives have been isolated from horseshoe bats. 49 SARS-CoV-1 caused a major international outbreak in 2002-2004 with around 8,000 50 recorded cases and at least 774 deaths³. MERS-CoV fuels recurrent disease outbreaks 51 in humans through repeated host jumps from its reservoir in camels⁴. Four coronaviruses 52 (HCoV-229, -NL63, -OC43 and -HKU1) circulate endemically in humans and their 53 ancestral reservoirs are believed to be species of bats and rodents, with host jumps into 54 humans likely facilitated by other mammals as bridging hosts^{5–7}. Additionally, multiple cases of host jumps from animals into humans leading to isolated cases or small clusters 55 have been documented for other coronavirus species⁵. Given the current health burden 56 57 exerted by coronaviruses and the risk they pose as possible agents of future epidemics 58 and pandemics, surveillance of animal-borne coronaviruses should be a public health 59 priority. Indeed, the discovery and characterisation of the diversity of coronaviruses 60 harboured by mammals across the world is the first step for designing pre-emptive 61 measures to minimise human or animal exposure. Here, we focus on bats since some 62 human coronaviruses have their ancestral origins in various bat species.

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64 Several studies over the last decade have screened bats across Asia, Africa, the Middle 65 East and Europe for coronaviruses, finding anywhere from 1.5-23% coronavirus prevalence in animals tested⁸⁻¹⁹. A selection of studies representing the diversity of 66 67 previous screening efforts is listed in Supplementary Table 1. These prevalence 68 estimates were primarily obtained via a reverse transcription real-time PCR (RT-PCR) 69 using degenerate primers designed to target most coronaviruses species (i.e., pan-70 coronavirus primers; Supplementary Table 1). Given the vast diversity of coronaviruses, 71 including those yet to be discovered, it is difficult to design primers that can amplify and 72 capture the full diversity of coronaviruses. Our own comparative analysis of published

primer sets show that existing RT-PCR assays²⁰⁻²⁴ underestimate coronavirus 73 74 prevalence (Supplementary Information: Supplementary Fig. 1 and 2). Difficulties with 75 primer design is exacerbated by low RNA concentrations in field samples and RNA 76 degradation, so the large variability in prevalence estimates in these studies may be due 77 to the sensitivity of the primer set used rather than the epidemiology of bat-coronaviruses. 78 While sample RNA quality remains mainly dependent on sample collection and laboratory 79 practices, because untargeted RNA sequencing does not require a priori knowledge of 80 sequence information, it provides a more accurate estimate of viral diversity and 81 prevalence. Hence, we chose this approach over RT-PCR to survey coronaviruses in UK 82 bats.

83

84 Sequencing-based surveillance data can be used to assess the zoonotic potential of 85 novel viruses, that is, the likelihood that these viruses can infect humans in the future. 86 This includes *in silico* assessments that determine the degree of sequence and structural 87 homology to other known and closely-related human-infecting viruses^{11,25}. Even more 88 compelling evidence can be obtained *in vitro*, and one of the most direct assessments is 89 to isolate and test the infectivity of novel viruses in human cells^{25,26}. However, this would 90 increase the risk of exposure to these potentially infectious agents, necessitating stringent 91 biosafety precautions. Additionally, isolation of novel viruses via cell culture without prior 92 knowledge of their cell tropism and receptor usage can be challenging. A lower risk and 93 effective alternative is to measure the binding efficiency of viral entry proteins to host 94 receptors¹⁰, or to assess efficiency of viral entry into human cell lines via a pseudovirus 95 assay²⁵, which expresses only the viral entry protein in a non-infectious reporter system. 96 While observed binding and cellular entry in these low-risk assays do not indicate that a 97 virus can replicate effectively in human cells, they provide an indication of which human 98 cell receptors can be exploited by novel viruses during infection, which are one key 99 determinant of viral infection. Despite the importance of functional validation, many 100 studies to date fall short of providing *in vitro* or even *in silico* assessments of zoonotic risk 101 (Supplementary Table 1).

103 There are 17 bat species that breed in the United Kingdom (UK), most of which can roost 104 in domestic buildings, churches, barns and other man-made structures. The high habitat 105 overlap with humans places bats in close proximity to domesticated and farmed animals, 106 which can serve as potential bridging hosts for transmitting bat-borne viruses to 107 humans²⁷. However, multiple factors have to align for the successful emergence of a 108 zoonotic pathogen in humans, including the frequency of exposure, the ability of the 109 pathogen to infect humans and its capacity for onward human-to-human transmission²⁸. 110 The relative risks of these various factors for zoonotic spillover remains largely unknown 111 and may vary depending on pathogen and geographical context. All UK bat species are 112 protected by law across the UK with licences required for work related to bats. So, 113 although direct contact is rare among the general public, it is far more common for the 114 small proportion of the population comprising bat scientists, ecologists, conservationists 115 and bat rehabilitators that undertake regular research, monitoring, surveillance, and bat 116 rehabilitation work.

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118 Only two coronavirus surveillance studies have been conducted in UK bats to date^{11,19}. 119 The first, published a decade ago, screened seven bat species and detected 120 alphacoronaviruses in Daubenton's bat and Natterer's bat (Myotis daubentonii and M. 121 *nattereri*, *respectively*)¹⁹. The other, from 2021, screened faecal samples from lesser 122 horseshoe bats (*Rhinolophus hipposideros*) and recovered the whole genome sequence 123 of a single sarbecovirus, RhGB01 (MW719567)¹¹. However, neither study provided direct 124 in vitro assessments of zoonotic risk. Accordingly, the viral diversity and zoonotic potential 125 of UK bat viruses remains largely unknown. This is equally true of most other UK 126 mammals. However, given that the evolutionary origins of many coronaviruses of human 127 health concern can be traced back to bats, assessing their zoonotic potential in UK bats 128 is a top priority, before moving on to other animal groups.

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To address this knowledge gap, we used an existing UK network of bat rehabilitators and conservationists to collect faecal samples from UK bats. Faeces from all but one bat species breeding in the UK (the grey long-eared bat, *Plecotus austriacus*, the rarest species in the UK) were collected and subsequently screened using deep RNA 134 sequencing to characterise the genomic diversity of bat-borne coronaviruses in the UK.

135 To assess their zoonotic potential, we then tested the ability of a subset of these

136 coronaviruses to bind human-cell receptors *in vitro*, which is a key requisite for human

137 infection.

138 Results

139 Untargeted RNA sequencing recovers nine complete coronavirus genomes including two140 new species

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142 We performed deep RNA sequencing on 48 faecal samples from 16 of the 17 UK breeding 143 bat species, with wide geographic coverage and over two years (Supplementary Fig. 3). 144 Through taxonomic assignment of sequencing reads using *Kraken2*²⁹, we detected the 145 presence of at least 30 viral families, 53% of which primarily infect non-mammalian hosts 146 such as plants, insects and bacteria (Supplementary Fig. 4a). Additionally, the total 147 relative abundance of viral species that infect non-mammalian hosts was significantly 148 higher than that for mammalian viruses (two-sided Mann-Whitney U test, U=1393, 149 p=0.004; Supplementary Fig. 4b). These findings indicate that the faecal virome in UK 150 bats largely comprises viruses that do not necessarily infect them, nor other mammals, 151 including humans.

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We next focused on coronaviruses due to their relevance to human health and recovered nine complete genomes (96-100% completeness; assessed by CheckV³⁰) and five partial contigs (<3%) across six UK bat species (*M. daubentonii*, *P. pipistrellus*, *P. pygmaeus*, *P. auritus*, *R. ferrumequinum*, and *R. hipposideros*), detecting coronaviruses amongst 29% of the samples. The nine complete genomes were assessed by CheckV to be of high-quality³⁰ and read alignments to these genomes indicated an even coverage of reads with a median coverage of 548-7958 reads per position (Supplementary Fig. 5; Table 1).

161 A global phylogenetic tree based on alignment-free genetic distances³¹ revealed the 162 genus and subgenus membership of these new coronaviruses (Fig. 1a; see Methods). 163 We then followed with local maximum-likelihood phylogenetic analyses to determine their 164 precise placement within each subgenus (Fig. 1b-d). These phylogenetic analyses reveal 165 that the nine novel genomes we recovered comprise four alphacoronaviruses from the 166 Pedacovirus subgenus, five betacoronaviruses including one merbecovirus, and four 167 sarbecoviruses (Fig. 1). Three of the coronaviruses recovered from *M. daubentonii* (which 168 we call MdGB01-03) form a well-supported clade with other pedacoviruses isolated from

the same bat species in Denmark (Fig. 1b). One coronavirus sequenced from *P. pipistrellus* (PpiGB01) falls as a sister lineage to the above clade. Another coronavirus from *P. auritus* (PaGB01) is related to MERS-CoV-like merbecoviruses isolated from *Hypsugo*, *Pipistrellus*, and *Vespertilio* spp. from Western Europe and China (Fig. 1c).
Four coronaviruses isolated from *R. ferrumequinum and R. hipposideros* (RfGB01-02 and RhGB07-08, respectively) are closely related to the previously described UK bat sarbecovirus, RhGB01³² (Fig. 1d).

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177 Of the nine coronaviral genomes recovered here, two represent new species. Indeed, 178 pedacovirus PpiGB01 from *P. pipistrellus* was relatively divergent from its closest match, 179 a pedacovirus previously isolated from *M. daubentonii* (less than 81% nucleotide 180 sequence identity; Table 1). Similarly, merbecovirus PaGB01 shares less than 82% 181 sequence identity to its closest match, a merbecovirus isolated from *P. kuhlii* in Italy 182 (Table 1). Overall, our surveillance efforts have extended our knowledge of the existing 183 diversity of coronaviruses. Looking at their genomic structures, we identified one new gene in each of these new species (Supplementary Information; Supplementary Fig. 6). 184 185

186 Viruses that are able to infect a broad range of hosts have been associated with a higher 187 risk of emerging as infectious diseases that can transmit between humans^{33,34}. Here, the 188 four sarbecovirus genomes, representing one viral species, were recovered from two 189 distinct horseshoe bat species, R. ferrumequinum and R. hipposideros. RhGB07, 190 RhGB08, RfGB01, RfGB02 share 97-100% identity with RhGB01 previously described in 191 *R. hipposideros*¹¹. To better understand how these viruses might be shared among the 192 two hosts, we looked at the habitat distribution of each horseshoe bat species. The two 193 horseshoe bat species share a large proportion of their habitats, with 33% of their 194 occurrence records reported at the same geographical coordinates. Furthermore, species 195 distribution modelling predicted that 45% of the total land area occupied by the two 196 species is shared (Supplementary Fig. 7a). Since the two Rhinolophus species can share 197 roosts³⁵, these results indicate a potentially high frequency of direct contact, which may 198 facilitate viral sharing and thus account for the isolation of RhGB01-like sarbecoviruses 199 that are closely related from these two species.

200 To extend this analysis, we examined both observed and predicted distributions of all 17 201 UK bat species to identify potential viral sharing hotspots for future surveillance work. By 202 analysing 42,953 occurrence records, we identified three regions near Bristol, 203 Birmingham and Brighton with particularly high species diversity (up to 16 species in a 204 single 5x5 km grid; Supplementary Fig. 7b). Additionally, we identified regions within the 205 UK, especially in Wales and the south coast of England where the habitats of the greatest 206 number of different bat species are predicted to coincide (Supplementary Fig. 7c). 207 Alongside an understanding of the ecology of native species, including co-roosting and 208 foraging behaviours, such information is a useful resource for future surveillance studies, 209 and for prioritising focal areas of potential high risk.

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Sarbecoviruses recovered from UK bats can bind the human ACE2 receptor for cellularentry

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214 We tested whether representatives of the newly identified UK coronaviruses (the 215 sarbecoviruses RhGB07 and RfGB02, the merbecovirus PaGB01, and the pedacovirus 216 PpiGB01) could use human cellular receptors for viral entry to assess their zoonotic 217 potential. We successfully incorporated the spike proteins of these UK coronaviruses into 218 lentivirus-based pseudoviruses (see Methods). We then tested the ability of these spike-219 expressing pseudoviruses to infect human cells expressing the human receptors, 220 angiotensin-converting enzyme 2 (hACE2), dipeptidyl peptidase-4 (hDPP4) and 221 aminopeptidase N (hAPN), which are the primary receptors exploited by SARS-CoV-2, 222 MERS-CoV and HCoV-229E, respectively.

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None of the spike pseudoviruses of the UK coronaviruses could enter cells using any of the receptors except RhGB07, which showed significantly higher entry into cells overexpressing hACE2 compared to those not expressing hACE2 (Fig. 2a; p<0.0001). As expected, SARS-CoV-2, MERS-CoV and HCoV-229E showed significantly higher entry into cells overexpressing hACE2, hDPP4 and hAPN, respectively (Fig. 2a; p<0.0001). Additionally, VSV-G pseudoviruses, which can enter cells regardless of their receptor expression, showed comparably high entry across all groups (Fig. 2a; unnormalised data

shown in Supplementary Fig. 8a). Additionally, using biolayer-inteferometry (BLI), we confirmed that the RhGB07 spike is able to bind hACE2 with a dissociation constant, K_d = 253nM (Fig. 2b). However, the binding affinity of RhGB07 spike to hACE2 is approximately 17-fold lower than that to SARS-CoV-2 spike (K_d = 15nM) (Fig. 2b).

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236 Given the lower binding affinity of RhGB07 spike compared to SARS-CoV-2, we then 237 investigated if, like SARS-CoV-2, RhGB07 spike-expressing pseudoviruses can infect 238 human cells expressing lower (HEK293T-hACE2 – HEK293Ts stably transduced with 239 hACE2) or physiological levels of hACE2 (Calu-3 lung, and Caco-2 colorectal cell lines). 240 Alongside this, we tested the entry of RfGB02, PaGB01 and PpiGB01 spike 241 pseudoviruses in the same cell lines in case they use a human receptor not otherwise 242 tested as in Fig. 2a. As positive controls, we included the spike proteins from other 243 coronaviruses, BANAL-20-52/SARS-CoV-2 (wild-type Wuhan-Hu-1 with D614G), MERS-244 CoV, and HCoV-229E, which can efficiently enter these cell lines using hACE2^{36,37}, 245 hDPP4 and hAPN, respectively. We also included the negative control, RaTG13, which 246 can bind hACE2 but cannot enter cells expressing lower or physiological levels of 247 hACE2³⁷. As expected, RaTG13 could not enter any of these cell lines, while all positive 248 controls showed significantly higher entry into these cell lines than 'bald' pseudoviruses 249 not expressing any spike protein (p<0.01; Fig. 2c). In contrast, none of the UK spike 250 pseudoviruses tested, including RhGB07, displayed significant entry into any of these 251 human cell lines (p>0.05; Fig. 2c).

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Separately, we asked if other host proteins are necessary for efficient cellular entry of the UK coronaviruses. In particular, the transmembrane serine protease 2 (TMPRSS2) protease has been shown to greatly enhance the entry efficiency of MERS-CoV³⁸ and HCoV-229E³⁹ spike pseudoviruses into human cells. However, PaGB01 and PpiGB01, which fall in the same subgenus as MERS-CoV and HCoV-229E, respectively, could not enter TMPRSS2-overexpressing cells (Fig. 2d).

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260 RhGB07 can bind and use hACE2 for cellular entry but RfGB02 cannot, despite the high
261 98% sequence identity of their spike proteins. This begs the questions as to how RhGB07

262 might have acquired the ability to use hACE2, and whether this might be associated with 263 the usage of bat ACE2 orthologues. To investigate this, we tested the entry of RhGB07 264 and RfGB02 spike pseudoviruses into human cells expressing the ACE2 orthologues 265 from four bat species (R. ferrumequinum, R. pusillus, Myotis lucifugus, and Rousettus 266 *leschenaultia*). Unfortunately, throughout the course of this study, there was no publicly 267 available ACE2 sequence for *R. hipposideros* (from which RhGB07 was recovered), and 268 no ACE2 transcripts could be identified directly from our metatranscriptomic libraries. 269 Nevertheless, we detected significant cell entry but only through *M. lucifuqus* ACE2 270 (p<0.0001; Fig. 2e; unnormalised data shown in Supplementary Fig. 8b), and neither 271 RhGB07 nor RfGB02 could use *R. ferrumequinum* ACE2 receptors, despite RfGB02 272 being sampled from this species. In contrast, SARS-CoV-2, BANAL-20-52 and RaTG13 273 were all able to efficiently use R. ferrum equinum ACE2 (p < 0.0001; Fig. 2e), indicating that this ACE2 construct could allow sarbecovirus entry. Surprisingly, BLI measurements 274 275 indicate detectable binding of RhGB07 spike to both R. ferrumequinum and M. lucifugus 276 ACE2 (Fig. 2f), which means that RhGB07 can bind *R. ferrumequinum* ACE2 but not 277 enter cells expressing this receptor. This highlights that binding of host ACE2 alone may 278 not be sufficient for efficient viral entry, and that other host cell-virus interactions (e.g., 279 presence of suitable co-receptors) may be required. Taken together, our results indicate 280 that RhGB01-like viruses may not be using ACE2 to infect their native *Rhinolophus* hosts. 281

282 Structural and sequence features of RhGB07 spike explain detectable but inefficient 283 usage of hACE2

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285 To better understand the results of the assays described above, we used the AlphaFold 286 artificial intelligence program⁴⁰ to predict the 3D structure of the receptor-binding domain 287 (RBD) of the RhGB07 spike protein. We then compared it to the resolved RBD structures 288 of SARS-CoV-2⁴¹, BANAL-236 (a close relative of BANAL-20-52)⁸, and RaTG13³⁷ bound 289 to hACE2. Superposition of the RBD structures showed high structural conservation 290 across all four sarbecoviruses (Fig. 3a). Additionally, the 3D structure of the RhGB07 291 RBD near the RBD-hACE2 binding interface was highly similar to that for SARS-CoV-2 (Fig. 3b), which was confirmed by comparing the area of contact surface (894 Å² and 850 292

293 Å², respectively; Supplementary Figure 6). These findings account for the ability of the 294 RhGB07 spike protein to bind hACE2 (Fig. 2a, b). To understand why RhGB07 spike 295 pseudoviruses could not enter cells expressing the ACE2 receptor at physiological levels 296 (Fig. 2c), we compared the level of conservation at key RBD residues of SARS-CoV-2⁴¹ 297 and SARS-CoV⁴² in contact with hACE2. This included sarbecoviruses isolated from Asia, 298 Europe, and Africa, which bind hACE2 with various affinities^{43–45}. All these sarbecoviruses 299 showed conservation at more than half of SARS-CoV-2 (Fig. 3c) or SARS-CoV (Fig. 3d) 300 contact residues, with high levels of conservation at certain positions like Y453, N487, 301 Y489, G502 and Y505 (relative to SARS-CoV-2; Fig. 3c). Previous deep mutational 302 scanning experiments showed that all these positions, except Y453, cannot be mutated 303 without considerable loss of hACE2 binding affinity⁴⁶, indicating that contact residues are 304 important determinants of hACE2 binding. Notably, the novel RhGB01-like 305 sarbecoviruses share only 9/17 SARS-CoV-2 (Fig. 3c) and 7/14 SARS-CoV contact 306 residues (Fig. 3d). This is slightly below RaTG13, which shares 11/17 SARS-CoV-2 and 307 8/14 SARS-CoV contact residues, respectively. In contrast, BANAL-20-52 shares 15/17 308 contact residues with SARS-CoV-2.

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310 These results indicate poorer conservation of these key contact residues in UK 311 sarbecoviruses, which would explain the relatively lower hACE2 usage efficiency, and 312 hence the ability to infect human cells, of RhGB07 and RaTG13 compared to BANAL-20-313 52. Similarly, the poorer conservation of SARS-CoV-2 contact residues in BtKY72 than 314 Khosta-2 (Fig. 3c) may explain the lower binding affinity to hACE2 of the former⁴⁴. 315 Notably, RhGB07 - but not RfGB02 - could enter cells using hACE2 (Fig. 2a), despite 316 their spike proteins sharing the same variants at all SARS-CoV-2 key contact residues 317 (Fig. 3c). The RfGB02 spike has 26 amino acid mutations relative to RhGB07, and only 318 four of these were within the RBD (K337N, H432L, T470A, P487Q; Supplementary Table 319 2). As such, either the residues at these positions are, in addition to the SARS-CoV-2 320 contact residues, important mediators of hACE2-binding, or the remaining 22 non-RBD 321 mutations have caused structural changes that reduce the binding affinity to hACE2, or 322 both. Further experiments delineating these mutational effects could help to shed light on 323 the molecular determinants of sarbecoviral entry into human cells.

324

325 Remarkably, the RhGB01-like sarbecoviruses already possess a R-A-K-Q sequence 326 (spike residues 669-672; Supplementary Fig. 10a), which is one nucleotide away 327 (Gln/CAA to Arg/CGA) from the canonical R-X-K/R-R motif, a furin cleavage site (FCS) 328 that allows cleavage by host furin-like proteases, enhancing the ability of many coronaviruses, including SARS-CoV-2, to infect human cells^{47,48}. This R-A-K-Q motif is 329 330 also found in Khosta-2⁴⁹, a sarbecovirus recovered from *R. hipposideros* in Khosta, 331 Russia, which is at the south-eastern extremes of Europe, but not in BtKY72 from 332 *Rhinolophus* sp. in Kenya⁵⁰ or other sarbecoviruses isolated from Asia. However, western 333 blot analyses indicated that even when we mutated R-A-K-Q to R-A-K-R (i.e., a Q672R 334 mutation), the RhGB07 spike is not cleaved by any human host protease (Supplementary 335 Fig. 10b). Previous studies have shown that the FCS on SARS-CoV-2 (681-RRAR-684) lies on an extended flexible loop that protrudes out of the spike structure, which allows 336 337 access by host furin^{51,52}. Also, it has been shown that deletions that shortened this 338 extended loop prevented efficient cleavage of SARS-CoV-2 spike, which was likely due to reduced accessibility of the FCS⁴⁸. This loop is seven residues shorter in RhGB01-like 339 340 viruses (Supplementary Fig. 10b), which may explain why no cleavage was observed for 341 the RhGB07 R-A-K-R pseudovirus mutant.

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Finally, a recombination analysis of the RhGB01-like and other representative sarbecoviruses indicates a high prevalence of recombination (Supplementary Information; Supplementary Fig. 11), which may accelerate adaptation for infecting novel hosts. Given these findings, the current zoonotic risk of sarbecoviruses in UK bats, while small, cannot be ignored and warrants more extensive surveillance of bats at the national scale.

349 Discussion

The emergence of the COVID-19 pandemic in 2019 is a sobering reminder of the massive impact of zoonotic viruses on global health and economy. Despite this, genomic surveillance in wildlife remains limited. In this study, we used an existing network of bat rehabilitators to obtain geographically and temporally diverse samples from almost all bat species in the UK. We argue that this can be a sustainable and effective surveillance model to identify and characterise novel animal-borne viruses that may or may not yet be able to infect humans but might evolve the ability to do so in the future.

357

358 We provided evidence that at least one sarbecovirus isolated from UK horseshoe bats 359 can bind hACE2 in vitro and discuss these patterns relative to our in silico analyses. Crook 360 et al.¹¹ performed a contact residue analysis, similar to the one we report in Fig. 3c and 361 d, on RhGB01 and suggested that moderate homology in its key contact residues 362 indicates that it is unlikely to bind hACE2. However, our in vitro (Fig. 2) and in silico (Fig. 363 3) results highlight that despite having only moderate conservation of key contact 364 residues, RhGB07 can bind and use hACE2. Additionally, the spike of RhGB07, but not 365 RfGB02, can bind hACE2 for cellular entry, despite identical conservation levels to SARS-366 CoV-2 at key contact residues. These findings indicate that assessing the conservation 367 of key contact residues (Fig. 3c, d) may have limited predictive power for whether a spike 368 protein can bind hACE2, possibly due to multiple structural configurations allowing hACE2 369 binding. This is further evidenced by the different contact residues for SARS-CoV and 370 SARS-CoV-2.

371

372 Our findings indicate that the RhGB01-like viruses likely require further adaptations, 373 particularly in their spike proteins, before they can make a zoonotic jump. Notably, single 374 mutations of some of the SARS-CoV-2 contact residues in sarbecoviral spike proteins 375 have been shown to enable binding of ACE2 from novel host species and improve binding 376 affinity by greater than fivefold⁴⁴. Additionally, a single T403R mutation in the RaTG13 377 spike has been shown to allow the virus to infect human cells⁵³. Given this, we speculate 378 that the genetic barrier precluding effective hACE2 usage for cellular entry into human 379 cells may be small. This may also be the case for the other RhGB01-like sarbecoviruses

sampled previously¹¹. The fact that of the two RhGB01-like viruses we investigated here,
one was capable of infecting hACE2-overexpressing cells and the other not, despite 98%
spike sequence similarity and identical SARS-CoV-2 residues. This further indicates that
minor adaptations in the spike protein may significantly affect its binding affinity with host
receptors, and hence zoonotic potential.

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386 We also identified a R-A-K-Q sequence in all European sarbecoviruses that resembles 387 an FCS precursor, but which is absent in all Asian sarbecoviruses considered 388 (Supplementary Fig. 10a). This supports previous observations that FCSs naturally occur 389 in coronaviruses and have emerged independently at least six times amongst 390 betacoronaviruses⁵⁴. However, even after mutating R-A-K-Q to R-A-K-R, we could not 391 detect any cleavage of RhGB07 spike by human proteases (Supplementary Fig. 10b). These findings indicate that, in addition to acquiring a functional FCS via substitution, 392 393 European sarbecoviruses would likely have to acquire an extended loop structure (like in 394 SARS-CoV-2) via insertion for efficient spike cleavage.

395

396 We found a high prevalence of genetic recombination amongst sarbecoviruses, 397 particularly in the spike gene (Supplementary Fig. 11), which may facilitate viral 398 adaptations to overcome the genetic barrier for a zoonotic jump. This observation is 399 corroborated by other studies that have also suggested an enrichment of recombination 400 signals in or surrounding the sarbecovirus spike gene^{55,56}. Co-infections and subsequent 401 recombination of RhGB01-like viruses with other coronaviruses that already effectively 402 use hACE2 may therefore facilitate zoonotic transmission. As such, the possibility of a 403 future host-jump into humans cannot be ruled out, even if the risk is small. This reiterates 404 the need for individuals that are in frequent contact with bats, such as bat rehabilitators, 405 to adhere to current biosafety practices to reduce their exposure to bat coronaviruses and 406 likewise to reduce the risk of the exposure of bats to human-borne coronaviruses⁵⁷, such 407 as SARS-CoV-2 or the endemic HCoVs. Fortunately in the UK, the risk of zoonotic 408 exposure is minimised for most people through a lack of direct contact (roosting spaces 409 are often well away from human inhabitants) along with the provision of science-based 410 information to roost owners by organisations such as the Bat Conservation Trust
411 (<u>https://www.bats.org.uk</u>).

412

413 Our *in vitro* assays indicate that RhGB01-like sarbecoviruses, including RfGB02 that was 414 directly sampled from this species, do not use *R. ferrumequinum* ACE2 as their primary 415 receptor, which is in line with other studies of bat coronaviruses^{44,58}. Importantly, this 416 raises the question as to what evolutionary mechanisms drive the acquisition of the ability 417 to use hACE2 in bat sarbecoviruses. Given previous associations between pathogen host 418 breadth and their capacity to emerge as zoonotic diseases^{33,34}, we speculate that multi-419 host viruses tend to have 'generalist' cell entry receptors that possess a low genetic 420 barrier to the evolution of zoonotic transmission. More extensive surveillance of the viral 421 sharing dynamics in mammalian hosts, including bats, may provide key insights into the 422 molecular and ecological determinants of zoonotic events. Such studies can leverage 423 both species occurrence data and niche modelling to prioritise regions where a high 424 number of species are likely to be found combined with an understanding of species 425 ecology for quantification of risk.

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The initial spread of SARS-CoV-2 in China, its widely publicised evolutionary origin in *Rhinolophus* bats⁵⁹, and the subsequent identification of other bat-borne sarbecoviruses in Southeast Asia^{8,10}, has focused attentions about the zoonotic risk of coronaviruses in those geographical regions. However, our findings highlight the zoonotic risk of sarbecoviruses may extend beyond Asia, stressing the importance of more extensive surveillance globally.

433

Finally, while it is imperative to better quantify the risk of zoonotic events from bats and design approaches to mitigate risk, bats fulfil important roles in ecosystems globally, including services such as arthropod suppression, pollination and seed dispersal⁶⁰. Some bat species have rapidly declining populations – for example, one third of the most threatened mammalian species in the UK are bats^{61,62}. Recent studies have shown that human-associated stressors such as habitat loss and changes in land-use can be important drivers of zoonotic spillover from wildlife^{63,64}, and that bat culls are ineffective in minimising cross-species transmission⁶⁵. As such, it is vitally important that an
integrated ecological conservation approach is taken that includes maintaining legal
protection, rather than destruction of wildlife and its habitat, in future approaches to
mitigate zoonotic risk.

445 Methods

446 Sample collection

Sampling kits were sent out to various bat rehabilitators in the UK as described previously⁵⁷ for the collection of faeces from bats. These faecal samples (0.02-1g) were immediately stored in 5 ml of RNAlater solution to prevent degradation of RNA. The geographical locations and collection dates for all samples are provided in Supplementary Table 3.

452

453 Murine hepatitis virus (MHV) spike-in control culture

MHV (GenBank AY700211.1) was propagated in an NCTC 1469 clone derivative (NCTC
721) cell line in high glucose DMEM and 10% horse serum. Both the MHV and NCTC cell
line were acquired from the American Type culture Collection (ATCC, Manassas, Virginia,
USA). Cell culture supernatant was isolated for later RNA extraction.

458

459 RNA extraction

460 RNA was extracted from faecal samples using the QIAamp Viral RNA Mini Kit (Qiagen) 461 following the protocol for extracting RNA from stool samples. We used up to 0.5 g of 462 faeces, which was vortexed in 2ml of 0.9% NaCl solution, at 6000rpm for 2 minutes. The 463 supernatant was filtered using a 0.2 µm svringe filter. 280 µl of which was used for RNA 464 extraction. For the MHV spike-in control, we used 140 µl of culture supernatant for RNA 465 extraction. Total RNA was eluted in 80 µl of AVE buffer and stored at -80°C. RNA was 466 quantified using Qubit 2.0 fluorometer (Invitrogen). All faecal extractions were spiked with 467 20 µl of MHV RNA prior to library preparation to act as a sequencing quality control.

468

469 Coronavirus database

To create a database representing the extant global genomic diversity of coronaviruses,
we downloaded all complete *Coronaviridae* (taxid:11118) genomes from *NCBI Virus*,
excluding provirus sequences (accessed 4th July 2022). Additionally, we downloaded all
non-human-associated and non-SARS-CoV-2 betacoronaviruses from GISAID⁶⁶ (n = 29).

To minimise the overrepresentation of certain viral species, we randomly retained 50 isolates for each of the following species: porcine epidemic diarrhoea virus, avian infectious bronchitis virus, MERS-CoV, SARS-CoV and SARS-CoV-2 sequences. This yielded a final dataset comprising 2118 genomes.

478

479 Metagenomic sequencing and assembly

480 All samples were prepared for sequencing using the NEBNext® Ultra[™] Directional RNA 481 Library Prep Kit, with a QIAseg FastSelect rRNA depletion step. Prior to sequencing, we 482 also spiked in MHV RNA (GenBank AY700211.1) as a positive control. Sequencing was 483 carried out using Illumina NovaSeq, paired end 150 bp. Quality control of reads was 484 bbduk.sh v39.01 **BBTools** performed using from the Suite 485 (https://sourceforge.net/projects/bbmap/). Briefly, we trimmed adapter sequences and 486 read ends below Q10, and discarded trimmed reads with average quality below Q10. 487 Reads that mapped to the positive control using Bowtie2 v2.4.5⁶⁷ were removed prior to 488 all downstream analyses. De novo metagenomic assembly was performed on quality-489 controlled or raw reads for each sample using coronaSPAdes v3.15.4⁶⁸. Assembled 490 scaffolds were then gueried using BLASTn against all 2118 genomes in our coronavirus 491 database to determine their most related reference. Scaffolds that could be aligned using 492 BLASTn to coronaviruses in our database and that were already longer than 28kb were 493 considered as complete genomes.

494

495 In some cases, *de novo* assembly yielded multiple scaffolds that were shorter than 28kb 496 but shared the same closest reference. We 'stitched' these scaffolds together using the 497 BLASTn alignment coordinates to the closest coronavirus reference and replaced any 498 gaps with Ns. De novo assembly using raw reads produced better results, producing 499 longer and more complete scaffolds, yielding six >28kb scaffolds (MdGB01, MdGB02, 500 MdGB03, PpiGB01, RfGB01, RfGB02), compared to quality-controlled read assembly 501 which yielded only two (RfGB01, PpiGB01). Further, the two >28kb scaffolds, RfGB01 502 and PpiGB01, generated using either raw or guality-controlled assemblies were identical, 503 suggesting that *de novo* assembly using raw reads were reliable. We hence chose the 504 assemblies generated using raw reads for our downstream analyses. We named the

novel complete genomes following the naming convention for the Sarbecovirus previously
described in a UK bat, RhGB01 – species: 'Rh' (*R. hipposideros*), region the coronavirus
was found in: 'GB' (Great Britain) and the frequency of description: '01' (the first described
in that species and country).

509

510 Genome annotation and characterisation of novel genes

511 Assembled genomes were annotated using Prokka v1.14.6⁶⁹, and annotated genes were 512 inspected to identify and correct erroneous frameshifts that were present in the raw 513 assemblies to produce the final genomes. For the four novel sarbecoviruses (RhGB07, 514 RhGB08, RfGb01, RfGB02) and three of the pedacoviruses (MdGB01, MdGB02, 515 MdG03), we also performed genome alignments to their closest known relative shown in 516 Table 1 to check if erroneous indels were present. The gene annotations were also 517 analysed to determine if these genomes carry any novel genes. We used PSI-BLAST on 518 the online webserver (https://blast.ncbi.nlm.nih.gov/), an iterative search program that is more sensitive than the conventional protein BLAST⁷⁰, to identify distant homologues of 519 520 annotated genes. We additionally used InterProScan^{71,72} to make functional predictions 521 for potentially novel proteins.

522

523 Taxonomic classification of sequencing reads

524 Taxonomic classification of reads was done using Kraken2 v2.1.2²⁹ with the '--paired' flag 525 and using the Viral database maintained by Ben Langmead (7 June 2022 release; 526 https://genome-idx.s3.amazonaws.com/kraken/k2_viral_20220607.tar.gz). This 527 database comprises all genomes available on NCBI RefSeg as of June 2022. We then 528 extracted reads assigned to each viral family (Supplementary Fig. 4a) or viral species 529 (Supplementary Fig. 4b). To minimise the effects of potential read misclassifications, we applied abundance thresholds as described previously⁷³. Briefly, we considered a taxon 530 to be present if greater than 10 read pairs were assigned and if its relative abundance 531 532 was greater than 0.005.

534 Species niche modelling

535 Bat occurrence records data were gathered from the online databases NBN Atlas 536 (https://nbnatlas.org/) and GBIF (www.gbif.org). Records from year 2000-present were 537 included, removing replicate records and those with high coordinate uncertainty. The 538 number of occurrence points used for modelling ranged from 32 (Myotis alcathoe) to 539 16,403 (*Pipistrellus pipistrellus*). An initial 17 environmental variables were identified a 540 priori to be important for predicting bat distributions. Nine were climatic variables 541 averaged across 1980-2010 as described by Karger et al.⁷⁴, and were reduced to five 542 variables using Variance Inflation Factor (VIF), retaining only those with a VIF < 0.5. 543 These were mean annual air temperature, mean diurnal air temperature range, mean 544 daily mean air temperature of the wettest guarter, precipitation seasonality and mean 545 monthly precipitation amount of the warmest guarter. Four variables were derived from 546 the UKCEH Land Cover Map 2019⁷⁵. After merging similar land use classes, distance to 547 woodland, distance to grassland, distance to arable and horticulture, and distance to 548 urban were measured using Euclidean distance tools in ArcMap version 10.8. Two further 549 distance variables were derived from Ordnance Survey polygons (2019, 2021): distance to the nearest road⁷⁶ and distance to the nearest river⁷⁷. Elevation and slope were 550 551 included to describe the topography of Great Britain, and were taken from the LiDAR 552 Composite Digital Terrain Model data at 10m resolution⁷⁸. All spatial data were 553 subsequently reduced to 1000m resolution and projected to British National Grid.

554

555 An ensemble of five supervised binary classifiers was trained to predict the suitability of 556 a land area for each of the 17 UK bat species using the *R* package sdm⁷⁹: random forest 557 (RF), maximum entropy (MaxEnt), multivariate adaptive regression splines (MARS), 558 boosted regression trees (BRT), and support vector machines (SVM). Classifiers were 559 trained to predict whether a particular species was present or 'absent' based on the 13 560 ecological variables described above, using the occurrence data for each species and an 561 equal number of randomly generated pseudo-absence data points across the study area. 562 Training and evaluation were performed using a 5-fold cross-validation protocol, where a 563 random subset comprising 80% of the dataset is used for training and the remaining 20% 564 use for the final evaluation. A final ensemble of all five classifiers that were trained was

565 used to generate the species distribution maps, with the contribution of each individual 566 classifier weighted based on its area under the receiver operating characteristic curve 567 (AUROC) score obtained during training. The resultant species distribution maps indicate 568 habitat suitability as a probability score for each 1 km square grid on the study area, which 569 ranges from 0 (unsuitable habitat) to 1 (suitable habitat). All models across all species 570 performed well, with a median AUROC, sensitivity and specificity of 0.827, 0.854, and 571 0.78, respectively. The individual species distribution maps and model performance 572 metrics are provided in Supplementary Fig. 12 and serve as a useful resource for future 573 studies that seek to understand the geographical range of UK bat species.

574

575 Phylogenetic analyses

576 To place the novel sequences within the global diversity of coronaviruses sequenced to 577 date, we computed alignment-free pairwise Mash distances using Mash v2.3³¹ with a k-578 mer length of 12, and reconstructed neighbour-joining trees⁸⁰ using the *nj* function from 579 the Ape v5.6.2 package in R (Fig. 1a). This alignment-free phylogenetic reconstruction 580 approach circumvents the challenge of aligning highly diverse sequences at the family 581 level, where high frequency of viral recombination may obscure true evolutionary 582 histories⁸¹ and prevent dataset wide alignments. In accordance with previous work⁸², we 583 rooted the neighbour-joining tree to a monophyletic *Deltacoronavirus* clade comprising all 584 10 representative Deltacoronavirus genomes downloaded from NCBI RefSeq.

585

586 From this global phylogeny, we retrieved the pedacovirus (n = 106), merbecovirus (n = 106) 587 113) and sarbecovirus genomes (n = 534) most proximal to the novel assembled 588 genomes. We then aligned genomes from these subgenera separately using the Augur 589 v14.0.0⁸³ wrapper for MAFFT v7.490⁸⁴. Genome positions where more than 20% of 590 sequences were assigned gaps were removed from the alignment. We subsequently 591 reconstructed finer-scale maximum-likelihood trees with IQTree v2.1.4-beta under a GTR+G model, using ultrafast bootstrapping (UFBoot)⁸⁵ and approximate likelihood-ratio 592 593 tests (SH-aLRT)⁸⁶ with 1000 replicates. All phylogenetic trees were visualised either using 594 FigTree v1.4.4 or *agtree* v3.2.1⁸⁷.

596 Recombination analysis

597 We selected 218 sarbecovirus genomes from the local sarbecovirus tree (n = 534) by 598 retaining only one representative each for SARS-CoV (NC 004718) and SARS-CoV-2 599 (MW206198). We subsequently aligned these genomes via the same approach described 600 above but masked all positions with >20% of gaps by replacing the positions with Ns, and 601 removed gaps in the alignment relative to the genome used to root the local sarbecovirus 602 tree, NC 025217. This masked alignment was then analysed using RDP v4.101⁸⁸. Gene 603 annotations for NC_025217 were obtained from GenBank and used to annotate predicted 604 recombinant positions.

605

606 Spike protein homology and conservation of contact residues

607 We extracted the Prokka-annotated spike protein sequences from our novel isolates for 608 further analysis. We peformed multiple sequence alignments of spike proteins from our 609 novel isolates and other sarbecoviruses that have been shown to bind human ACE2^{25,43,44} (BANAL-236, MZ937003.2; SARS-CoV-2, NC_045512.2; SARS-CoV-1, NC_004718.3; 610 611 Rs4084, KY417144.1; RsSHC014, KC881005.1; WIV1, KF367457.1; Rs7327, 612 KY417151.1: Rs4231. KY417146.1; LYRa11, KF569996.1; Pangolin GD-1. 613 EPI_ISL_410721; Pangolin GX-P2V, EPI_ISL_410542; RhGB01, MW719567.1; Khosta-614 2, MZ190138.1; BtKY72, KY352407.1) using Mafft v7.490⁸⁴. Subsequently, pairwise 615 amino acid similarity scores, visualisation of the alignments, and annotatation were 616 performed using the Spike alignments using UGENE v42.0⁸⁹. The accessions of all 617 genome records used in these analyses are provided in Supplementary Table 4.

618

619 Pseudovirus assays

620

To further test the capability of the coronaviruses we identified to infect human cells, we synthesised human codon-optimised, Δ 19-truncated (or equivalent) spike constructs in pcDNA.3.1. The merbecovirus PaGB01 and pedacovirus PpiGB01 were additionally synthesised with GSG-linker Myc tags for detection of spike incorporation into pseudoparticles. Gene synthesis and codon optimisation was performed by GeneArt (Thermo Fisher). Plasmids for human (*Homo sapiens*; BAB40370.1), least horseshoe bat
(*Rhinolophus pusillus*; ADN93477.1), Leschenault's rousette fruit bat (*Rousettus leschenaultia*; BAF50705.1), and little brown bat (*Myotis lucifugusI; XP_023609438.1*) in
pDisplay were used as previously described⁹⁰. Additionally, Greater horseshoe bat
(*Rhinolophus ferrumequinum;* BAH02663.1) ACE2 was synthesised and cloned into
pDISPLAY for this study.

632

633 We maintained human embryonic kidney cells (HEK 293T; ATCC CRL-11268) and 634 human Hepatocyte carcinoma clone 5 (Huh7.5; C. Rice, Rockefeller University, New 635 York, NY) in complete media (DMEM, 10% FBS, 1% non-essential amino acids (NEAA) 636 and 1% penicillin-streptomycin (P/S)). Human lung cancer cells (Calu-3; ATCC HTB-55) 637 and Human epithelial colorectal adenocarcinoma cells (Caco-2; ATCC HTB-37) were 638 maintained in DMEM, 20% FBS, 1% NEAA and 1% P/S. All cells were kept at 5% CO₂, 639 37°C. 293T-hACE2 and Huh7.5-TMPRSS2 cells were generated by transducing HEK 640 293T or Huh7.5 cells with an ACE2 or TMPRSS2-expressing lentiviral vector, MT126⁹¹ 641 and selecting with $2 \mu g m l^{-1}$ puromycin or 4 mg m l⁻¹ G418; after selection, cells were 642 subsequently maintained with $1 \mu g m l^{-1}$ puromycin or 2 mg m l^{-1} G418, respectively. 643

Lentiviral based pseudotyped viruses were generated as previously described⁴⁸. Briefly, 100 mm dishes of 293T cells were transfected using lipofectamine 3000 (Invitrogen) with a mixture of 1 μ g of the HIV packaging plasmid pCAGGs-GAG-POL, 1.5 μ g of the luciferase reporter construct (pCSFLW), and 1 μ g of the plasmid encoding the spike or glycoprotein of interest in pcDNA3.1. After 24 h supernatant was discarded and replaced. PV-containing supernatants were collected at 48 and 72 h post-transfection, passed through a 0.45 μ M filter, and aliquoted and frozen at -80°C.

651

Pseudovirus entry assays were performed as previously described⁴⁸. Briefly, 100 mm dishes of 293T cells were transfected using lipofectamine 3000 (Invitrogen) with 2 μg of the ACE2 encoding plasmid or empty vector. After 24 h, cells were resuspended by scraping and plated into 96 well plates. Cells were overlayed with pseudovirus for 48 h before lysis with reporter lysis buffer (Promega). Caco-2, Calu-3, and 293T-hACE2 cells

were seeded into 96 well plates. Cells were overlayed with pseudovirus for 48 h before lysis with cell culture lysis buffer (Promega). We determined luciferase luminescence on a FLUOstar Omega plate reader (BMF Labtech) using the Luciferase Assay System (Promega). As the RhGB01-like spike proteins appeared to be toxic in HEK 293Ts, the amount of pseudovirus added was standardised by quantifying p24 protein by western blot in a matched concentrated pseudovirus stock.

663

664 We assessed expression of transfected receptors using Western blot assays. Cell 665 suspensions were pelleted by centrifugation at 1000 revolutions per minute (RPM) for 7 666 min at 4°C, then supernatant was removed. Cells were resuspended in 150 µl of cold 667 radioimmunoprecipitation assay (RIPA) buffer (Thermo Fisher) and incubated on ice for 668 30 min. Then, they were spun down at 3750 RPM for 30 min at 4°C. The protein-669 containing supernatants were transferred to sterile Eppendorfs and frozen down at -20°C. 670 Before running a gel, 50 µl of 2-Mercaptoethanol (BME; Sigma) diluted 1:10 in 4X 671 Laemmli Sample Buffer (Bio-Rad, USA) was added to lysates and incubated at 80°C for 672 10 min.

673

To analyse incorporation of spike into the different sarbecovirus pseudoviruses, we concentrated pseudovirus by ultracentrifugation at 100,000 x g for 2 h over a 20% sucrose cushion.

677

678 In all experiments, we confirmed the successful expression of host receptors and spike 679 pseudoviruses using Western blot analyses (Supplementary Fig. 13). For Western 680 blotting, membranes were probed with mouse anti-tubulin (diluted 1/5,000; abcam; 681 ab7291), mouse anti-p24 (diluted 1/2,000; abcam; ab9071), rabbit anti-SARS spike 682 protein (diluted 1/2,000; NOVUS; NB100-56578), rabbit anti-HA tag (diluted 1/2000; 683 abcam; ab9110) or rabbit anti-Myc tag (diluted 1/2000; abcam; ab9106). Near infra-red 684 secondary antibodies, IRDye[®] 680RD Goat anti-mouse (diluted 1/10,000; abcam; ab216776), IRDye[®] 680RD Goat anti-rabbit (diluted 1/10,000; abcam; ab216777), were 685 686 subsequently used. Western blots were visualized using an Odyssey DLx Imaging 687 System (LI-COR Biosciences).

688

689 Alphafold2 (ColabFold) structural analysis

The protein structure model of the RhGB07 RBD was predicted using Alphafold2 as implemented in ColabFold⁹². Default settings were used. The top ranked model was used for all analyses. Structural representations and calculations were done within ChimeraX^{93,94}. RMSD values for structural superpositions were calculated using the matchmaker command. Reported values represent the RMSD of all Calpha carbons. Buried surface area calculations were performed using the measure buriedarea command.

697

698 Biolayer Interferometry (BLI)

699 The RhGB07 spike trimer was designed to mimic the native trimeric conformation of the 700 protein. It consists of a gene synthesized by Genscript of CHO codon-optimized sequence 701 of RhGB07, residues 1-1191, preceded by a u-phosphatase signal peptide⁹⁵, residues 702 969 and 970 mutated to proline (2P) to stabilize the prefusion state of the spike trimer, a 703 putative basic site that may be the site of proteolysis (RAKQ, residues 669-672, was 704 mutated to GASQ), a C-terminal T4 foldon fusion domain to stabilize the trimer complex, 705 followed by C-terminal 8x His and 2x Strep tags for affinity purification. This gene was 706 cloned with the pcDNA3.1(+) vector. The trimeric RhGB07 spike protein was expressed 707 as previously reported as for the SARS-CoV-2 spike transiently expressed in suspension-708 adapted ExpiCHO cells (Thermo Fisher) in ProCHO5 medium (Lonza) at 5 x10⁶ cells/mL 709 using PEI MAX (Polysciences) for DNA delivery⁹⁶. At 1 h post-transfection, dimethyl 710 sulfoxide (DMSO; AppliChem) was added to 2% (v/v). Following a 7-day incubation with 711 agitation at 31 °C and 4.5% CO₂, the cell culture medium was harvested and clarified 712 using a 0.22 µm filter. The conditioned medium was loaded onto Streptactin XT columns 713 (IBA) washed with PBS and eluted with 50 mM biotin in 150 mM NaCl, 100 mM HEPES 714 7.5. Eluted protein was then dialyzed overnight into PBS. The purity of spike trimers was 715 determined to be >99% pure by SDS-PAGE analysis.

Human (residues 19-615), little brown bat (19-629) and greater horseshoe bat (19-615)
ACE2 genes were synthesized by Genscript and cloned in after the human pregnancy
specific glycoprotein 1 signal peptide and is followed by a 3C protease cleavage site, a
mouse IgG2a Fc fragment and a 10x His tag (only for the hACE2 construct). Protein
production was produced exactly as for the RhGB07 spike. The filtered conditioned media
was then subjected to Protein A purification. Eluted protein was dialyzed into PBS.

723

Experiments were performed on a Gator BLI system. Running buffer was 1X PBS. Dimeric mFc-hACE2 and bat ACE2 were diluted to 10 µg/mL and captured with MFc tips (GatorBio). Loaded tips were dipped into 2-fold serial dilution series (highest concentration 3000 nM) of the RhGB07 spike protein. Curves were processed using the Gator software with a 1:1 fit after background subtraction. Plots were generated in Prism v9.

- 730
- 731 Data analysis and visualisation
- All data analyses were performed using R v4.1.0 or Python v3.9.12. Visualizations were
- 733 performed using ggplot v3.3.5⁹⁷.

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- 946

947 Author contributions

VS, CC, ER, GW and TB wrote the grant application that supported this research. VS and
FB supervised the research. Primary analysis on the project was carried out by CCST,
TPP, KYM and CH with contributions from FB, LvD, WDP, DO, and WB. CCST wrote the
initial draft of manuscript, with subsequent rounds of editing from VS, FB and LvD. All
authors provided intellectual contributions to the manuscript.

953 **Declaration of competing interest**

Authors declare that to current knowledge, there are no legal, financial or personalcompeting interests.

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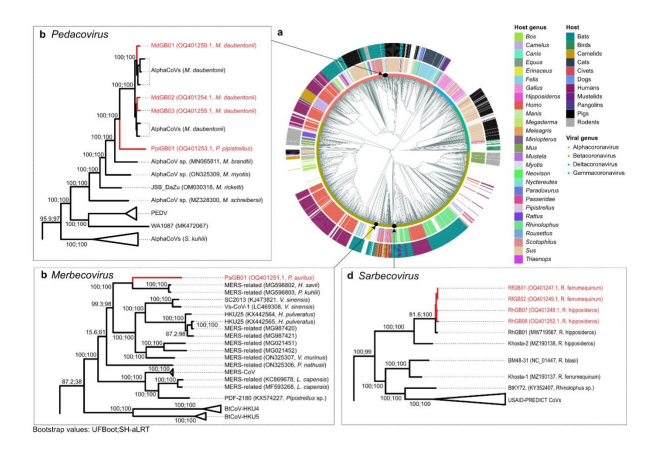
976 Data and code availability

All novel genomes are available in NCBI GenBank under the accessions OQ401247OQ401251, and OQ401253-OQ401255 (BioProject accession PRJNA929706). The raw
sequencing reads generated in this study have also been uploaded to the SRA under the
accessions SRX19257406- SRX19257414. The NBN Atlas datasets used are listed in
Supplementary Table 6. All custom code used to perform the analyses reported here are
hosted on GitHub (https://github.com/cednotsed/bat-CoVs.git).

983 Tables and figures

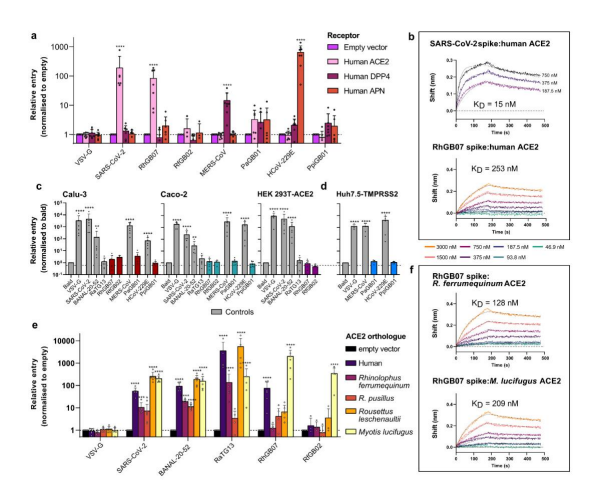
984 Table 1: Summary statistics for novel coronavirus genomes assembled in this 985 study

Host species	Common name	Genome name	Subgenus	Length (bp)	Closest hit			Prop.		
					Accession	Name	BLASTn identity (%)	of query aligned (%)	CheckV completeness	Median coverage
Rhinolophus ferrumequinum	Greater	RfGB01	Sarbecovirus	29308	MW719567	RhGB01	98.1	99.7	97	689
	horseshoe bat	RfGB02	Sarbecovirus	29375	MW719567	RhGB01	98.1	99.4	97	7178
Rhinolophus hipposideros	Lesser	RhGB07	Sarbecovirus	29224	MW719567	RhGB01	97.9	100	97	3809
	horseshoe bat	RhGB08	Sarbecovirus	29217	MW719567	RhGB01	98	100	96	548
Plecotus auritus	Brown long- eared bat	PaGB01	Merbecovirus	30018	MG596803	P. kuhlii MERS- related CoV	81.5	99.7	99	6237.5
Pipistrellus pipistrellus	Common pipistrelle	PpiGB01	Pedacovirus	28247	MN535731	<i>M.</i> daubentonii pedacovirus	80.8	82.9	100	7438
Myotis daubentonii	Daubenton's bat	MdGB01	Pedacovirus	28224	MN535731	<i>M.</i> <i>daubentonii</i> pedacovirus	95.5	99.6	100	7938
		MdGB02	Pedacovirus	28010	MN535733	<i>M.</i> daubentonii pedacovirus	95.4	99.8	100	7874
		MdGB03	Pedacovirus	28227	MN535733	<i>M.</i> <i>daubentonii</i> pedacovirus	95.4	99.8	100	7958



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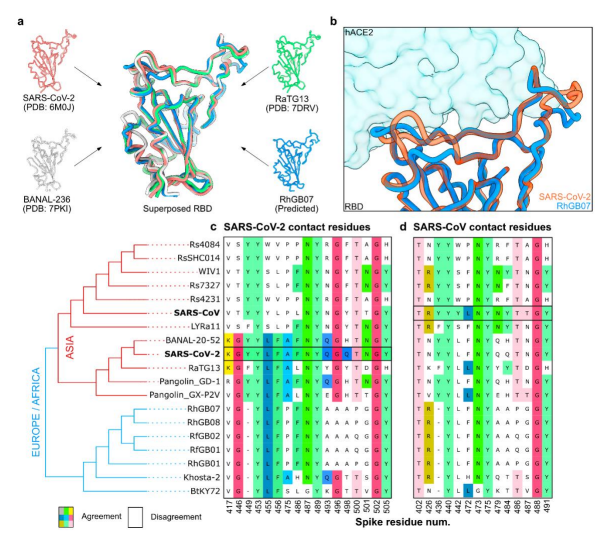
Fig. 1. Phylogenetic placement of novel coronaviruses. (a) Alignment-free phylogeny of the global diversity of coronavirus genomes (n = 2118) and our nine novel genomes. Host genus (inner ring) and their broader host groups (outer ring) are annotated. Local maximum likelihood trees of (b) pedacoviruses (n = 991 106), (c) merbecoviruses (n = 113) and (d) sarbecoviruses (n = 534).



992

993 Fig. 2. RhGB07 can bind and use human ACE2 for cell entry in vitro. (a) Entry of different spike 994 pseudoviruses expressing viral glycoproteins into HEK293T cells transfected with (a) human receptors 995 known to allow entry of human coronaviruses or (e) ACE2 homologues from different species. For (a) and 996 (e), the raw entry values for each pseudoviruses were normalised by their entry into cells transfected with 997 a vector containing no receptor sequence (i.e., 'empty'). The raw entry values of representative repeats are 998 provided in Supplementary Fig. 8 for direct comparisons of absolute entry. Bio-layer interferometry binding 999 curves showing the association and dissociation of SARS-CoV-2 and RhGB07 spike proteins with (b) 1000 hACE2, or (f) with R. ferrumequinum or M. lucifuqus ACE2. (c) Entry of pseudoviruses into different 'normal' 1001 human cell lines that stably express lower or physiological levels of hACE2. All entry measurements are 1002 normalised to those for the 'bald' pseudovirus not expressing any spike protein. (d) Entry of pseudoviruses 1003 into Huh7.5 cells transduced with a human TMPRSS2 vector, normalised to 'bald'. Data from panels (a), 1004 (c) and (d) are compiled from $n \ge 3$ completely independent repeats and plotted as mean + s.d.. Statistical 1005 significance was determined by (a, e) two-way ANOVA or (c, d) one-way ANOVA on log-transformed data 1006 (after determining log normality by the Shapiro-Wilk test and QQ plot) with multiple comparisons against 1007 'empty' vector or 'bald' pseudovirus, respectively. $*0.05 \ge p > 0.01$; $**0.01 \ge p > 0.001$; $***0.001 \ge p > 0.0001$; 1008 *****p*≤0.0001.

1009



1010

1011 Fig. 3. Structural and sequence features of RhGB01-like sarbecoviruses. (a) The solved RBD 1012 structures of SARS-CoV-2, RaTG13, BANAL-236 (close relative of BANAL-20-528) and the AlphaFold-1013 predicted structure of RhGB07 were superposed. (b) The 3D surfaces of the RBD-hACE2 binding interface 1014 for SARS-CoV-2 and RhGB07. Alignment of sarbecovirus spike proteins showing the conservation of key 1015 contact residues involved interactions between (c) SARS-CoV-1 spike and (d) SARS-CoV-2 spike with 1016 hACE2. The sequences shown in the alignments are from Asian, European and African sarbecoviruses 1017 that have been shown to bind hACE243-45. These sequences were ordered based on their genetic 1018 relatedness, as inferred from a consensus maximum likelihood phylogenetic tree reconstructed from their 1019 whole genomes (bottom left).