

ACE2 receptor usage reveals variation in susceptibility to SARS-CoV and SARS-CoV-2 infection among bat species

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Bats are the suggested natural hosts for severe acute respiratory syndrome coronavirus (SARS-CoV) and the causal agent of the coronavirus disease 2019 (COVID-19) pandemic, SARS-CoV-2. The interaction of viral spike proteins with their host receptor angiotensin-converting enzyme 2 (ACE2) is a critical determinant of potential hosts and cross-species transmission. Here we use virus-host receptor binding and infection assays to examine 46 ACE2 orthologues from phylogenetically diverse bat species, including those in close and distant contact with humans. We found that 24, 21 and 16 of them failed to support infection by SARS-CoV, SARS-CoV-2 or both viruses, respectively. Furthermore, we confirmed that infection assays in human cells were consistent with those in two bat cell lines. Additionally, we used genetic and functional analyses to identify critical residues in bat ACE2 receptors associated with viral entry restrictions. Our results suggest that many bat species may not be the potential hosts of one or both viruses and that no correlation was identified between proximity to humans and probability of being natural hosts of SARS-CoV or SARS-CoV-2. This study demonstrates dramatic variation in susceptibility to SARS-CoV and SARS-CoV-2 infection among bat species and adds knowledge towards a better understanding of coronavirus-bat interaction.

he unprecedented pandemic of coronavirus disease 2019 (COVID-19), caused by the new coronavirus SARS-CoV-2, has led to major threats to public health and economic development. It is therefore critically important to identify natural or intermediate hosts of SARS-CoV-2 to prevent further spread of COVID-19 and future emergence of similar diseases. Inferred from sequence similarity of human and bat virus genomes, it was suggested that horseshoe bats (Rhinolophus spp.) might be natural hosts of SARS-CoV and SARS-CoV-2 (refs. 1-3). These suggestions have resulted in misguided fears and unwarranted attacks on many bats-including species other than Rhinolophus-thereby seriously impacting efforts towards bat conservation4. Given the remarkable diversity of bats, which includes more than 1,400 species across the globe⁵, assessing the possibility that diverse bat species act as potential hosts of SARS-CoV and SARS-CoV-2 is urgent and crucial for both controlling outbreaks and protecting populations of wildlife.

Angiotensin-converting enzyme 2 (ACE2) is the main host cell receptor of SARS-CoV and SARS-CoV-2 and plays a vital role in mediating viral entry to cause infection 1.6. The interaction of a virus with its host receptor has been repeatedly demonstrated to serve as a primary determinant of host range^{7,8}. In this study, we tested ACE2 orthologues from 46 bat species across the phylogeny, including species occurring in urban and in rural areas, for their ability to support the entry of SARS-CoV and SARS-CoV-2 using virus-host receptor binding and infection assays. Hence, this study assessed whether diverse bat species are potential hosts of SARS-CoV or SARS-CoV-2. Moreover, by determining the correlation between proximity to humans and probability of being natural hosts of the

two viruses, these results provide important insights into pandemic control and wildlife conservation.

Results

Evolution of ACE2 in bats inhabiting urban or rural areas. We collected ACE2 orthologues from 46 bat species across the phylogeny (Fig. 1 and Supplementary Table 1). These species contained 28 species that roost or forage in urban areas near humans and 18 species more restricted to rural areas and hence likely to have minimal contact with humans (Supplementary Table 2). In total, we examined 46 species representing 11 bat families (Supplementary Table 3). After aligning the protein sequences of bat ACE2 orthologues, we examined 25 critical residues involved in the binding of the surface spike glycoprotein (S protein) of SARS-CoV-2 (ref. 9) (Extended Data Fig. 1). Genetic variations were observed in nearly all these 25 sites, which may have led to different abilities to support entry of SARS-CoV and SARS-CoV-2 (ref. 9). Furthermore, we detected at least 22 amino acid sites that are putatively under positive selection (Supplementary Table 4), which is indicative of heterogeneous selection pressure across sites. Notably, four of these positively selected sites are in the binding region of ACE2 to the SARS-CoV-2 S protein (Supplementary Table 4).

Interaction between bat ACE2 orthologues and SARS-CoV or SARS-CoV-2 receptor binding domain. Efficient binding between the S protein and the ACE2 receptor is essential for SARS-CoV and SARS-CoV-2 entry. This binding is mainly mediated by the interaction between the critical residues on the receptor-binding domain

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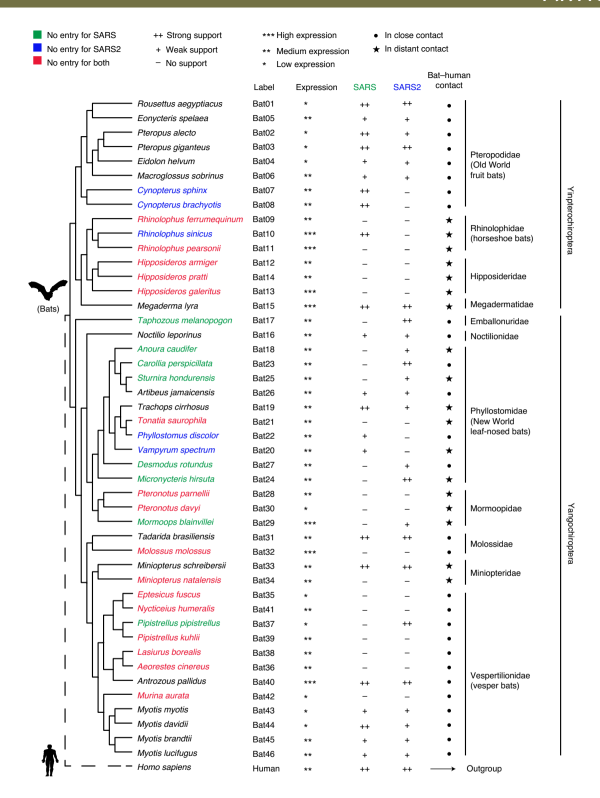


Fig. 1 | Phylogenetic tree of 46 bat species in this study. The labels of bat species in our experiments are indicated. Expression levels determined by western blot (Fig. 2a) are shown with asterisk symbols compared with human ACE2: the triple asterisk indicates high expression, the double asterisk indicates medium expression and the single asterisk indicates low but detectable expression. The ability of bat ACE2 to support SARS-CoV and SARS-CoV-2 pseudovirus entry is shown with different signs (Fig. 3a,b): infection data are presented as percentage mean values of bat ACE2 supporting infection compared with the infection supported by human ACE2. Infection efficiency <5% is indicated with a minus sign, between 5 and 50% with a plus sign and >50% with a double plus sign. Bat phylogeny was taken from previous studies²⁸⁻³⁰.

(RBD) and ACE2. To characterize the receptor function of ACE2 orthologues in a range of diverse bat species, we generated a stable cell library consisting of cell lines expressing the respective 46 bat

ACE2 orthologues through lentiviral transduction of 293T cells lacking ACE2 expression¹⁰. All bat ACE2 orthologues were exogenously expressed at a comparable level after puromycin selection,



Fig. 2 | Expression of bat ACE2 orthologues and their interaction with the SARS-CoV and SARS-CoV-2 RBD. a, Western blot detected the expression levels of ACE2 orthologues on 293T stable cells by targeting the C-terminal 3xFLAG-tag. Glyceraldehyde 3-phosphate dehydrogenase was employed as a loading control. **b**, Visualization of the intracellular bat ACE2 expression level by immunofluorescence assay detecting the C-terminal 3xFLAG-tag. Scale bar, 100 μm. **c,d**, Assessment of the interaction between different ACE2 orthologues and SARS-CoV-RBD-hFc (**c**) or SARS-CoV-2-RBD-hFc (**d**) proteins. Species that do not support efficient binding are underlined. 293T cells stably expressing the different bat ACE2 orthologues were incubated with 5 μg ml⁻¹ of the recombinant proteins at 37 °C for 1h; binding efficiency was examined by Alexa Fluor 488 goat anti-human IgG via fluorescence assay. Scale bar, 200 μm.

as indicated by western blot and immunofluorescence assays detecting the C-terminal 3×FLAG-tag (Fig. 2a,b).

To analyse the interaction, we produced recombinant SARS-CoV or SARS-CoV-2 RBD human immunoglobulin G (IgG) Fc fusion proteins (RBD-hFc), previously reported to be sufficient to bind human ACE2 efficiently^{11,12}. Protein binding efficiency was tested on the bat ACE2 cell library by means of immunofluorescence or flow cytometry targeting the human Fc. As expected, binding was almost undetectable on mock 293T cells but a strong binding signal was detected in the 293T cells expressing human ACE2 (Fig. 2c,d). Consistent with previous reports^{13,14}, SARS-CoV-2 RBD showed

higher binding to human ACE2 than SARS-CoV, which can also be observed on many bat ACE2 orthologues (Fig. 2c,d). Previous reports have shown that only a small fraction of ACE2 orthologues from tested mammalian species could not bind with SARS-CoV-2 S protein (n=6 of 49 species⁷; n=5 of 17 species¹⁵). However, our study revealed that many bat species (n=32 and n=28 of 46 species) do not support efficient binding with SARS-CoV-RBD and SARS-CoV-2-RBD, respectively (Fig. 2c,d). The overall profiles of bat ACE2 to bind to SARS-CoV and SARS-CoV-2 RBD are generally comparable; a few showed contrasting modes of binding preferences (Fig. 2c,d). For instance, Bat22 could bind to SARS-CoV

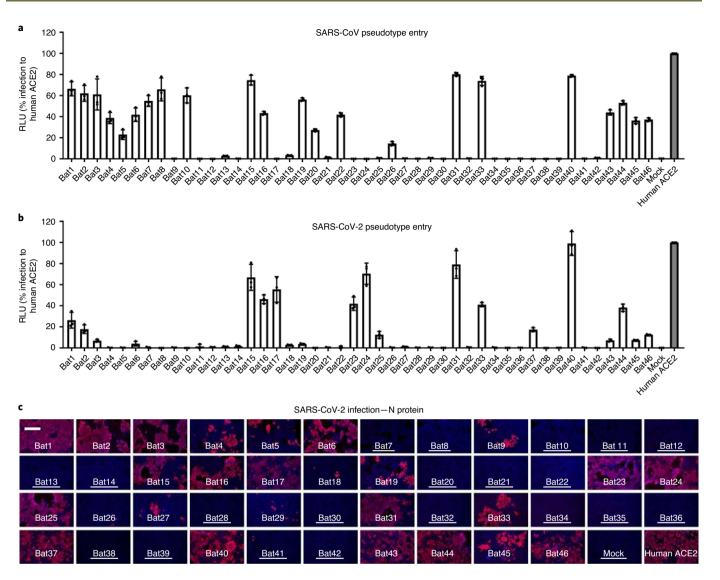


Fig. 3 | Characterization of bat ACE2 orthologues mediating entry of SARS-CoV and SARS-CoV-2 viruses. a,b, Ability of bat ACE2 orthologues to support the entry of SARS-CoV and SARS-CoV-2 pseudovirus. 293T cells expressing bat ACE2 orthologues in a 96-well plate were infected with VSV-dG-Luc pseudotyped with SARS-CoV (a) and SARS-CoV-2 (b) spike proteins, respectively. Intracellular luciferase activity was determined at 20 h post-infection. RLU, relative light unit. c, 293T cells expressing bat ACE2 orthologues were inoculated with the SARS-CoV-2 live virus at an MOI = 0.01. N protein (red) in the infected cells was detected through immunofluorescence assay at 48 h post-infection. Scale bar, 200 μ m. Samples expressing the indicated ACE2 orthologues that showed almost no entry for SARS-CoV-2 live virus are underlined. Data shown are representative results from 3 independent experiments and are presented as the mean \pm s.d. (n=3 for a and n=2 for b).

but not SARS-CoV-2, whereas Bat14, 21 and 40 could bind to SARS-CoV-2 but not SARS-CoV (Fig. 2c,d). Flow cytometry analysis showed consistent results (Extended Data Fig. 2).

Overall, the RBD-hFc binding assays demonstrated that bat ACE2 orthologues showed different affinity and selectivity levels to SARS-CoV and SARS-CoV-2, indicating that the ACE2 receptors of many bat species may not support efficient SARS-CoV and SARS-CoV-2 infection.

Receptor function of bat ACE2 orthologues to support the entry of SARS-CoV and SARS-CoV-2 using pseudotyped and live viruses. To further evaluate the receptor function of different bat ACE2 orthologues, we employed a vesicular stomatitis virus (VSV)-based rhabdoviral pseudotyping system to mimic the coronavirus spike protein-mediated single-round entry¹⁵. SARS-CoV and SARS-CoV-2 pseudotypes were generated by assembling the coronavirus spike proteins and replication-deficient VSV with

the VSV glycoprotein gene replaced with a fluorescence protein (VSV-dG-GFP) or a firefly luciferase (VSV-dG-Luc) reporter¹⁵. Both viruses showed minimal background infection on 293T cells, but efficient infection on 293T-human ACE2 cells (Extended Data Fig. 3). The susceptibility of the 293T cells expressing bat ACE2 orthologues was then examined with SARS-CoV and SARS-CoV-2 pseudotypes. The results showed that bat ACE2 orthologues have varying abilities to support coronavirus entry and different preferences for SARS-CoV and SARS-CoV-2. (Fig. 3a,b and Extended Data Fig. 4). Pseudotypes with green fluorescent protein (GFP) reporter showed similar results (Extended Data Fig. 5). Notably, we found that 24, 21 and 16 of the 46 bat species showed almost no entry for SARS-CoV, SARS-CoV-2 and both viruses, respectively (Figs. 1 and 3a,b and Supplementary Table 5), suggesting that these species are not likely to be potential hosts of either or both coronaviruses. The bat species showing no viral entry include those that occur in urban areas and those more restricted to rural areas (Fig. 1), suggesting that there is no correlation between proximity to humans and probability of being natural hosts of SARS-CoV or SARS-CoV-2. Although horseshoe bats were suggested as potential natural hosts of SARS-CoV and SARS-CoV-2 (refs. ¹⁻³), only one of the three species examined (*Rhinolophus sinicus*) supported SARS-CoV entry; this species was suggested as the potential host of SARS-CoV-2 (Figs. 1 and 3). These results unambiguously indicate that ACE2 receptor usage is species-dependent.

The SARS-CoV-2 S protein used in this study for pseudotyping contains a D614G mutation, which is currently a dominant variation¹⁷. The D614G mutation remarkably improved the in vitro infectivity of SARS-CoV-2 but may not significantly affect the receptor interaction since it is not in the RBD18. Indeed, we identified a very similar susceptibility profile using an original strain without D614G (Extended Data Fig. 4). We further demonstrated that the pseudotyped entry assay mimics the entry of live viruses through a SARS-CoV-2 infection assay (Fig. 3c). As expected, the profile of SARS-CoV-2 N protein expression is highly consistent with the results from the VSV-dG-based pseudotyped virus entry assay, except for some ACE2 that showed relatively higher infection efficiency (for example, Bat43-46) compared with the pseudovirus infection assay, which may be attributed to the different virus strains used (Fig. 3c). In addition, the live virus infection resulted in the phenotype of plaque formation, while the pseudotypes showed evenly distributed, single-round infection (Extended Data Fig. 5), which also partially explains why some bat ACE2 showed higher infection in the live virus infection assay.

When comparing the RBD-hFc binding and pseudotyped entry profiles, we found that binding and susceptibility are not always consistent, although the phenotypes were reproducible. For instance, some species (Bat12, 13, 14) were able to bind to SARS-CoV-2 RBD-hFc efficiently but could not support infection of the same virus, indicating that high binding affinity does not guarantee efficient viral entry (Figs. 2 and 3). In contrast, some species (Bat3-8) were defective or less efficient in SARS-CoV RBD-hFc binding but supported the entry of the same virus to some degree (Figs. 2 and 3). We hypothesize that such minimal binding may be sufficient for viral entry mediated by those ACE2 orthologues; alternatively, additional residues outside the traditional RBD region might be required for efficient interaction. These hypotheses should be tested in the future. Together, our results demonstrated dramatic variation of susceptibility to SARS-CoV and SARS-CoV-2 infection among bat species, suggesting that SARS-CoV and SARS-CoV-2 can selectively use some bat ACE2 as functional receptors for viral entry and many—if not most—bat ACE2 are not favoured by one or both viruses.

Evaluation of critical residues in bat ACE2 orthologues affecting viral binding and entry efficiency or specificity. We comprehensively analysed the relationship between critical RBD binding

sites in bat ACE2 sequences and their ability to support SARS-CoV and SARS-CoV-2 RBD binding and viral entry. Several critical residues were identified that may play critical roles in the determination of species specificity (Extended Data Fig. 1). According to the sequence alignment, two species pairs (Bat33 and Bat34 and Bat38 and Bat40) were selected to demonstrate the role of critical residues in RBD binding and viral entry because they were phylogenetically close but showed contrasting phenotypes for supporting RBD binding and viral entry. Specifically, Bat34 and Bat38 do not support SARS-CoV and SARS-CoV-2 RBD binding and infection, while Bat33 supports efficient binding and infection of both viruses and Bat40 supports infection of both viruses and to a lesser degree SARS-RBD binding (Figs. 2 and 3). We compared their protein sequences and highlighted the residues that may affect RBD interaction. For example, substitutions I27K, N31G and K42E were observed when comparing Bat33 with Bat34, while Q24L, E30K, K35Q and G354N were present between Bat38 and Bat40 (Fig. 4a). We hypothesized that the discrepancy in binding and infection phenotype is determined by their differences in critical residues for RBD interaction. To test this hypothesis, we designed a residue swap mutagenesis assay to investigate the role of critical residues on RBD binding and virus entry (Fig. 4a). We generated four swap mutations and corresponding 293T stable cell lines to test whether these substitutions could achieve gain-of-function and loss-of-function. All bat ACE2 orthologues and related mutants were expressed at a comparable level after lentiviral transduction, as indicated by the immunofluorescence of the C-terminal 3×FLAG-tag (Fig. 4b). Recombinant SARS-CoV and SARS-CoV-2 RBD-hFc proteins were applied to the cells expressing different ACE2 and binding efficiency was evaluated by immunofluorescence (Fig. 4c) and flow cytometry assays (Fig. 4d). As expected, the swap of critical residues on the selected four bat ACE2 changed their receptor function to the opposite, except for Bat38 mutant, which remained unable to bind SARS-CoV RBD-hFc (Fig. 4c,d). GFP (Fig. 4e) and luciferase levels (Fig. 4f) from the pseudotyped virus entry assay and the N protein staining from the live SARS-CoV-2 infection assay (Fig. 4g) further confirmed our hypothesis at the viral entry level. Structure modelling of bat ACE2/SARS-CoV-2-RBD complexes showed that the substitutions of I27K and N31G between Bat33 and Bat34 lead to a reduced packing interaction and the substitution of K42E disrupts the hydrogen bond with Y449, which may be related to the difference of susceptibility between Bat33 and Bat34 (Fig. 4h,i and Extended Data Fig. 6). In comparison, the substitutions of Q24L and E30K between Bat38 and Bat40 destroyed the favourable hydrophilic interactions with N487 and K417, respectively (Extended Data Fig. 6).

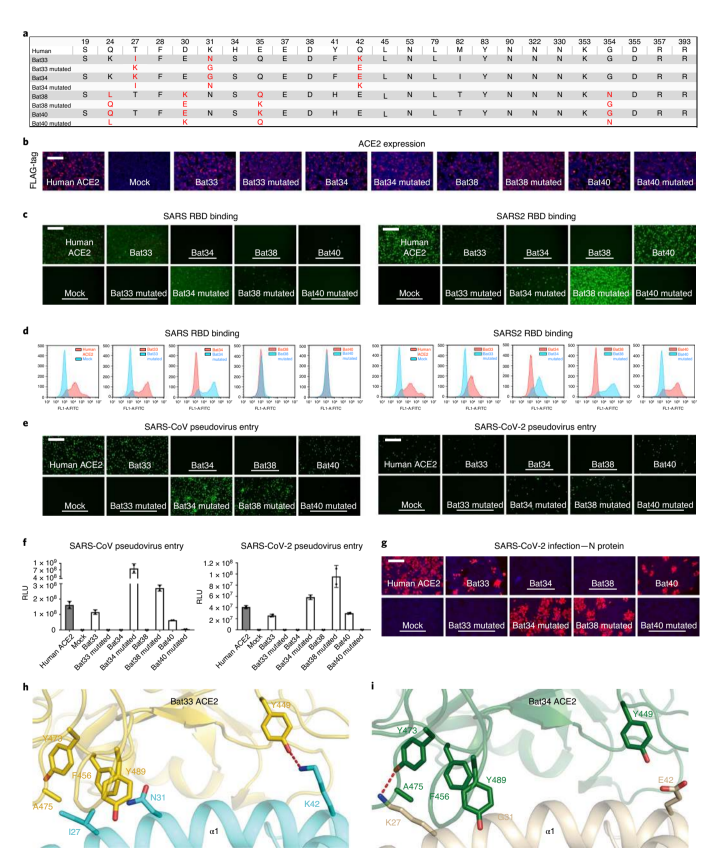
In addition, two bat cell lines, the lung epithelial cell line Tb 1 Lu of *Tadarida brasiliensis* (Bat31) and the kidney epithelial cell line of *Pteropus alecto* (Bat2), were used to validate our findings derived from human HEK293T cells. Endogenous ACE2 expression

Fig. 4 | Evaluation of the critical binding sites determining the species-specific restriction of SARS-CoV and SARS-CoV-2 binding and entry. a, Swap mutagenesis assay to investigate the role of critical residues on bat ACE2 orthologues for tropism determination. Residues involved in RBD (according to the structure between SARS2-RBD and human ACE2, Protein Data Bank 6MOJ) interaction are shown in the table. Residues that changed in the mutagenesis assay are marked in red. b, The expression level of the bat ACE2 orthologues and related mutants in transduced 293T cells was determined by an immunofluorescence assay recognizing the 3xFLAG-tag. Scale bar, 200 μm. c,d, Binding efficiency of SARS2-RBD-hFc and SARS2-RBD-hFc on 293T cells expressing bat ACE2 and related mutants. Cells were incubated with 5 μg ml⁻¹ of recombinant proteins at 37 °C for 1h and then washed and incubated with a secondary antibody recognizing human Fc. Immunostaining (c) and flow cytometry (d) were conducted to show binding efficiency. Scale bar, 200 μm. e,f, Ability of the indicated ACE2 and related mutants to support the entry of coronavirus pseudotypes. The 293T cells expressing the indicated ACE2 and their mutants were infected with SARS-CoV and SARS-CoV-2 pseudotypes expressing GFP (e) and luciferase (f). Infection was analysed at 20 h post-infection. Scale bar, 200 μm. Data are presented as the mean with s.d. (n = 2). g, 293T cells infected by the SARS-CoV-2 live virus at an MOI = 0.01; the infection was examined at 48 h post-infection through N protein (red) immunostaining. Nuclei were stained with Hoechst 33342 (blue). Scale bar, 200 μm. h,i, Comparison of the interface between Bat33/SARS-CoV-2-RBD and Bat34/SARS-CoV-2-RBD. Bat33 and its complexed RBD are coloured cyan and gold, respectively (h); Bat34 and its complexed RBD are coloured wheat and green, respectively (i). The mutated residues in ACE2 and the corresponding residues in SARS-CoV-2-RBD are shown and labelled. The red dotted lines between residues indicate hydrogen or ion

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was almost undetectable in these two cell lines, accounting for at least 1,000 folds lower than the susceptible Vero-E6 cells (Extended Data Fig. 7a). Therefore, these cells cannot support the entry of SARS-CoV and SARS-CoV-2. We successfully generated Tb 1 Lu stable cell lines expressing human ACE2 and bat ACE2 (Bat2, 3,

31, 32) since the transduction efficiency of Tb 1 Lu is much higher than that of PakiT03 cells (Extended Data Fig. 7b). As expected, Tu 1 Lu were susceptible to both SARS-CoV and SARS-CoV-2 when human ACE2 or some bat ACE2 orthologues (Bat2, 3 and 31) were expressed, yet remained non-susceptible when an ACE2 of a closely



related species (Bat32) was expressed (Extended Data Fig. 7c–e). Furthermore, we conducted SARS-CoV and SARS-CoV-2 pseudovirus entry assays on the two bat cell lines transiently transfected with various bat ACE2 (Bat2, 3, 31, 32, 33, 34, 38, 40) and their mutants (mutant Bat33, 34, 38 and 40m). The results were consistent with those derived from human cells, further confirming that ACE2 is the main receptor for the species-specific entry of SARS-CoV and SARS-CoV-2 in these bat cells (Extended Data Fig. 7f,g).

Discussion

Our study provides genetic and functional evidence from ACE2 receptor usage to show dramatic variation of susceptibility to SARS-CoV and SARS-CoV-2 infection among bat species. Our sampling covers representative species from 11 bat families, hence providing a broad picture of bat phylogeny. Moreover, our study included 28 species inhabiting urban areas and 18 species that are not common in cities or do not roost in buildings. Our functional assays demonstrated that there is no correlation between proximity to humans and probability of being natural hosts of SARS-CoV or SARS-CoV-2. Our results are only partially consistent with a recently published prediction based on sequence similarity, which estimated a binding score between ACE2 and the SARS-CoV-2 S protein for each vertebrate species9. The predicted binding scores for all 37 bat species fell into low (n=8) and very low (n=29) categories9, suggesting that all examined bat species are at low risk for SARS-CoV-2 infection. Our study included 36 of the 37 previously examined bat species (Fig. 1 and Extended Data Fig. 1); 21 of these appeared to support SARS-CoV-2 entry by their ACE2 receptors (Figs. 1 and 3), strongly suggesting that these bats are at risk for SARS-CoV-2 infection. These disparities between in silico analyses and functional experiments unambiguously suggest the importance of experimental data for confirmation of in silico analyses since our understanding of ACE2 sequences and structures is incomplete thus far. We note that our functional experiments also have limitations. For instance, it would be interesting to test bat ACE2 function in their respective species-specific cells if all bat cell lines were available. It is also important to note that the lack of viral infection in bat cell lines without ACE2 expression cannot rule out the possibility that alternative receptors are expressed in other cell types. Moreover, there were also some inconsistencies for some bat ACE2 between binding and infection. These inconsistencies may have resulted from some binding events that are defective in triggering the correct conformational change required for downstream entry^{13,19}. In addition, unlike binding, which is a dynamic process²⁰, entry is irreversible and can accumulate over time. Thus, we should be cautious about predicting the ability of ACE2 orthologues to support entry only based on their binding affinities to viral RBDs. Our genetic and functional evidence revealed critical residues of bat ACE2 that are involved in supporting SARS-CoV-2 entry (Fig. 4). However, these residues are not the genetic determinant of New World monkey ACE2 orthologues mediating SARS-CoV-2 entry⁷ and many bat ACE2 orthologues carrying residues that were considered unfavourable in the same study (H41 and E42)7 were fully functional in our study (Fig. 4), further confirming the complexity of ACE2 functionality.

We found that closely related species can show strikingly different ACE2 receptor usage. For example, *Rhinolophus sinicus* can support SARS-CoV entry, whereas its congeneric relatives *Rhinolophus ferrumequinum* and *Rhinolophus pearsonii* cannot (Figs. 1 and 3), even though some polymorphic sites of ACE2 may have occurred in *Rhinolophus sinicus* populations²¹. These findings clearly show that ACE2 receptor usage is species-dependent. Accordingly, although some bats might be potential hosts of ancestral SARS-CoV and SARS-CoV-2 (refs. ¹⁻³), one cannot assume that all bat species or individuals can carry these viruses. Although certain bat species are frequently observed to carry coronaviruses closely related to human

viruses in terms of sequence similarity22, there is no solid and direct evidence showing the initial spillover from bats to humans and other animals. Nevertheless, humans infected with coronavirus should maintain distance from bats that can use ACE2 as a viral receptor because many bat species are endangered and may be susceptible to human coronaviruses²³, as suggested for many other mammals^{9,24}. Indeed, the International Union for Conservation of Nature has assessed that over one-third of bat species are threatened or data-deficient and over half of all bat species have unknown or decreasing population trends²⁵. Thus, bats need protection more than ever. Our study supports the calls that public education on bat biology will reduce the threat to bats^{4,25}. In fact, all bats are potentially safe as long as they are treated with care and respect. We should work collaboratively to combat the pandemic and identify which species are potential hosts and not fear those species that are not hosts of the virus. Instead, we must respect and care for those species that are potential hosts and learn about the impact of human activities on their natural habitats, which may lead to zoonotic spillover events.

Methods

ACE2 sequence acquisition and selective pressure analysis. We obtained 46 full-length coding sequences of bat *ACE2* in this study, of which 32 were taken from a recent study⁹ and 14 were newly extracted from published or recently sequenced genome assemblies (see Supplementary Table 1 for the sources and accession numbers for the sequences and assemblies). Next, we aligned the deduced ACE2 protein sequences using the MUSCLE program (v.3.8.31)²⁶ (see Extended Data Fig. 1 for the resulting alignment). The sequence logo was generated with WebLogo (https://weblogo.berkeley.edu/logo.cgi). We performed selective pressure analysis on bat ACE2 using CodeML implemented in PAML (v.4.9a)²⁷. Two comparisons of site models (M1a and M2a, M8a and M8) were used to predict positively selected sites²⁷. The input tree was the species tree (Fig. 1) taken from previous studies²⁸⁻³⁰.

Cell culture. HEK293T cells (293T, catalogue no. CRL-3216; ATCC) and VERO E6 cells (catalogue no. CRL-1586; ATCC) were cultured in DMEM (Gibco) supplemented with 10% fetal bovine serum (FBS), 2.0 mM of L-glutamine, 110 mg l $^{-1}$ of sodium pyruvate and 4.5 g l $^{-1}$ of p-glucose. The lung epithelial bat cell line Tb 1 Lu (catalogue no. CCL-88; ATCC) and an I1-Hybridoma (catalogue no. CRL-2700; ATCC) cell line secreting a monoclonal antibody targeted against the VSV glycoprotein were cultured in MEM with Earle's balanced salts and 2.0 mM of L-glutamine and 10% FBS. The kidney epithelial bat cell line PakiT03 was a generous gift from L.-F. Wang and was cultured in DMEM:F-12 (Gibco) supplemented with 10% FBS. All cells were cultured at 37 °C in 5% CO $_2$ with the regular passage of every 2–3 d. 293T stable cell lines overexpressing ACE2 orthologues were maintained in growth medium supplemented with 1 μ g ml $^{-1}$ of puromycin.

Plasmids. Human codon-optimized complementary DNA sequences encoding various ACE2 orthologues and their mutants fused with a C-terminal 3×FLAG-tag (DYKDHD-G-DYKDHD-I-DYKDDDDK) were commercially synthesized and subcloned into a lentiviral transfer vector (pLVX-IRES-puro) through the EcoRI and NotI restriction sites. The DNA sequences of human codon-optimized SARS-CoV S protein (CUHK-W1, GenBank AY278554.2) and SARS-CoV-2 S protein (Wuhan-Hu-1, GenBank MN908947) were amplified from the plasmids pCMV/hygro-SARS-CoV-S (VG40150-G-N; Sino Biological) and pCAGGS-SARS-CoV-2-S-c9 (gifted by W. Li, National Institute of Biological Science, Beijing, China) into the pCAGGS vector with a C-terminal 18-amino acid deletion to improve VSV pseudotyping efficiency^{31,32}. The D614G mutation was introduced into the SARS-CoV-2-S coding sequence for higher in vitro infection efficiency. The plasmids for the expression of coronavirus RBD-IgG Fc fusion proteins were generated by inserting the coding sequences of SARS-CoV RBD (amino acids 318-516) and SARS-CoV-2 RBD (amino acids 331-530) into the pCAGGS vector to express fusion proteins with C-terminal human Fc (IgG1) and an N-terminal CD5 secretion leading sequence (MPMGSLQPLATLYLLGMLVASVL).

Generation of ACE2 stable expression cell lines. 293T and Tb 1 Lu cells overexpressing ACE2 orthologues were generated by lentiviral transduction. Specifically, the lentivirus was produced by cotransfection of lentiviral transfer vector carrying ACE2 coding sequences (pLVX-EF1a-Puro; from GENEWIZ) and packaging plasmids pMD2G (plasmid no. 12259; Addgene) and psPAX2 (plasmid no. 12260; Addgene) into 293T cells through Lipofectamine 3000 (Thermo Fisher Scientific). The lentivirus-containing supernatant was collected and pooled at 24 and 48 h post-transfection. 293T and Tb 1 Lu cells were transduced by the

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lentivirus (1:1 dilution of culture medium and lentivirus-containing supernatant) after 16 h in the presence of $8\,\mu g\,ml^{-1}$ polybrene. Stable cells expressing various ACE2 orthologues were selected and maintained in growth medium with puromycin (1 $\mu g\,ml^{-1}$). Cells selected for at least 10 d were considered as stable cell lines and used in different experiments.

Immunofluorescence assay to evaluate the expression levels of ACE2 orthologues. The expression levels of ACE2 orthologues were evaluated by an immunofluorescence assay used to detect the C-terminal 3×FLAG-tags. The 293T or Tb 1 Lu cells used for the analysis were seeded in the poly-lysine-pretreated 96-well plate at a cell density of 5×10^5 per ml (100 µl per well) and cultured for 24 h. Cells were fixed with 4% paraformaldehyde at room temperature for 10 min, permeabilized with 0.2% Triton X-100/PBS at room temperature for 10 min and blocked with 1% BSA at 37 °C for 30 min. Next, they were incubated with a mouse monoclonal antibody targeting the FLAG-tag (9A3, catalogue no. 8146S; Cell Signaling Technology) diluted in 1% BSA/PBS at 37 °C for 1 h. After three rounds of PBS washing, cells were incubated with 2 µg ml $^{-1}$ of the Alexa Fluor 594-conjugated goat anti-rabbit IgG (catalogue no. A11032; Thermo Fisher Scientific) diluted in 1% BSA /PBS at room temperature for 30 min. The nucleus was stained blue with Hoechst 33342 (1:5,000 dilution in PBS). Images were captured with a fluorescence microscope (MI52-N; Mshot).

Production of VSV reporter viruses pseudotyped with coronavirus spike proteins. Coronavirus spike protein pseudotyped viruses (CoV-psV) were packaged according to a previously described protocol using a replicate-deficient VSV-based rhabdoviral pseudotyping system (VSV-dG)33. The VSV-G glycoprotein-deficient VSV exogenously expressing enhanced GFP (VSV-dG-GFP) or firefly luciferase (VSV-dG-Luc) were rescued by a reverse genetics system from Kerafast. To produce CoV-psV, Vero E6 cells were transfected with the plasmids overexpressing SARS2-CoV (pCAGGS-SARS-S-dc) and SARS2-CoV-2 spike proteins (pCAGGS-SARS2-S-dc) through the Lipofectamine 3000 reagent. After 36 h, the transfected cells were transduced with VSV-dG reporter viruses diluted in serum-free Opti-MEM for 1 h at 37 °C (at a genome equivalent = 100). Transduced cells were washed with culture medium once and then replenished with fresh culture medium with I1-hybridoma-cultured supernatant containing anti-VSV monoclonal antibody (1:100 dilution) to neutralize the infectivity of the residual input viruses. The CoV-psV-containing supernatants were collected at 24h after transduction, clarified at 12,000 r.p.m. for 2 min at 4°C and immediately transferred to $-80\,^{\circ}\mathrm{C}$ for storage. The viral titre (genome equivalents) was determined by quantitative PCR with reverse transcription (RT-qPCR). The RNA copies in the virus-containing supernatant were detected using the VSV-L gene sequences (Supplementary Table 6).

Pseudotype entry assay. The 293T or bat lung epithelial (Tb 1 Lu) stable cell lines overexpressing various ACE2 orthologues were trypsinized and resuspended with SARS-CoV or SARS-CoV-2 pseudotyped viruses (at a genome equivalent = 100) in DMEM with 10% FBS. Next, they were seeded at 5×10^4 in a well of a 96-well plate to allow attachment and viral infection simultaneously. At 16-24 h after infection, images of infected cells with GFP expression were acquired with a fluorescence microscope (MI52-N). Intracellular luciferase activity was determined by a Bright-Glo Luciferase Assay Kit (Promega Corporation) and measured with a SpectraMax iD3 Multi-well Luminometer (Molecular Devices) or a GloMax 20/20 Luminometer (Promega Corporation). Alternatively, the PakiT03 bat cell line or Tb 1 Lu cells were transfected with plasmids overexpressing various ACE2 orthologues (based on the pLVX-IRES-puro vector) by Lipofectamine 3000 reagent and infected with the SARS-CoV or SARS-CoV-2 pseudotyped viruses carrying luciferase (at a genome equivalent = 100) at 24 h post-transfection. Cells were lysed using 1× Bright-Glo luciferase assay reagent and determined using a GloMax 20/20 Luminometer.

Coronavirus RBD-hFc binding assay. Recombinant SARS-CoV-RBD-hFc and SARS-CoV-2-RBD-hFc proteins were produced by transient transfection of 293T cells with Lipofectamine 3000. Transfected cells were cultured in FreeStyle 293 Expression Medium (Thermo Fisher Scientific) and the supernatants containing the recombinant proteins were collected at 2 and $4\,\hat{d}$ post-transfection. The RBD-hFc protein concentration was determined by comparing the target protein band with BSA standard dilutions through Coomassie Brilliant Blue staining. The RBD-hFc protein-containing supernatant $(5\,\mu g\,ml^{-1})$ was incubated with the 293T stable cell lines overexpressing different ACE2 orthologues for 1h at 37 °C. Cells were washed twice with DMEM and then incubated with 2 µg ml⁻¹ of Alexa Fluor 488-conjugated goat anti-human IgG (catalogue no. A11013; Thermo Fisher Scientific) diluted in DMEM with 2% FBS for 30 min at 37 °C. For immunostaining, cells were washed twice with PBS. Images were captured with a fluorescence microscope (MI52-N). For the flow cytometry analysis, cells were detached by 5 mM of EDTA/PBS and analysed with a CytoFLEX Flow Cytometer (Beckman Coulter Life Sciences).

SARS-CoV-2 live virus infection assay. The SARS-CoV-2 (strain IVCAS 6.7512) was provided by the National Virus Resource, Wuhan Institute of

Virology, Chinese Academy of Sciences. All SARS-CoV-2 live virus-related experiments were approved by the Level 3 Biosafety Committee (ABSL-3) of Wuhan University. All experiments involving SARS-CoV-2 were performed in the BSL-3 facility. SARS-CoV-2 was amplified on Vero E6 cells and stored at -150 °C; the titre was determined on Vero E6 cells through a plaque assay. 293T cells expressing ACE2 orthologues were seeded on a poly-lysine-coated 96-well plate for 24 h before inoculation. Cells were infected with SARS-CoV-2 at an MOI = 0.01 and then incubated in DMEM with 2% FBS for 48 h before testing. Cells were fixed with 4% paraformaldehyde in PBS at room temperature for 1 h, permeabilized with 0.2% Triton X-100 for 10 min and then blocked with 1% BSA/PBS at 37 °C for 1 h. Cells were subsequently incubated with a mouse monoclonal antibody, SARS-CoV/SARS-CoV-2 Nucleocapsid Antibody (catalogue no. 40143-MM05; Sino Biological), at 1:500 dilution at 37 °C for 1 h and then incubated with 2 µg ml⁻¹ of Alexa Fluor 594-conjugated goat anti-mouse IgG antibody (catalogue no. A-11032; Thermo Fisher Scientific) at 37 °C for 1 h. The nucleus was stained with Hoechst 33342. Images were acquired with a fluorescence microscope (MI52-N).

Quantification of the endogenous expression of ACE2 orthologues by RT–qPCR. Endogenous ACE2 expression levels of Vero E6 (from *Chlorocebus sabaeus*), PakiT03 (from *P. alecto*, Bat2) and Tb 1 Lu (from *T. Brasiliensis*, Bat31) cells were determined by RT–qPCR. In general, total RNA from cells cultured for 24 h was extracted by RNA-Easy Isolation Reagent (catalogue no. R701-02; Vazyme). cDNA was reverse-transcribed from 1 μg of total RNA by the HiScript II 1st Strand cDNA Synthesis Kit (catalogue no. R212-02; Vazyme); 1/20 volume of the cDNA was used as the template for the qPCR assay using the ChamQ Universal SYBR qPCR Master Mix (catalogue no. Q311-02; Vazyme) and a CFX96 Touch Real-Time PCR instrument (Bio-Rad Laboratories). Small nuclear ribonucleoprotein D3 polypeptide was used as an internal control for the normalization of the ACE2 relative expression level, which is presented as percentage expression compared with Vero E6 cell. Data are presented as the mean ± s.e.m. (*n* = 3).

Homology-based structural modelling. Molecular models of different bat ACE2 were predicted by iterative threading assembly refinement (I-TASSER) v.5.1 (ref. ³⁴). Starting from the amino acid sequences, the I-TASSER algorithm constructed the full-length three-dimensional atomic models by structural template identification, followed by template-based fragment assembly simulations. The structure of SARS-CoV-2-RBD was obtained from the previously reported crystal structure of the human ACE2/SARS-CoV-2-RBD complex (Protein Data Bank 6LZG)³⁵. The docking of ACE2 and SARS-CoV-2-RBD was performed using RosettaDock (v.4.0)³⁶. The binding energy of the ACE2/RBD complex was calculated using the InterfaceAnalyzer from the RosettaDock package and is represented in Rosetta energy units³⁷. All types of non-covalent interactions were analysed using RING v.2.0 (ref. ³⁸) and LigPlot+ v.2.2 (ref. ³⁹). Structural alignment and visualization were implemented in PyMOL (v.2.5.0a0)⁴⁰.

Statistical analysis. Data are expressed as mean values with s.d. unless otherwise stated. All experiments were repeated 3–5 times, each yielding similar results.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

All data are available in the manuscript or supporting materials.

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Author contributions

H.Z., H.Y., Y.C. and K.L. designed study. H.Z., H.Y. and H.J. wrote the manuscript. H.Y., H.J., Q.L., Z.Z., Q.X., X.W. and M.G. performed the experiments. H.Y., H.Z., H.J., B.-J.W., L.-F. W. and Y.C. analysed the data.

Competing interests

The authors declare no competing interests.

Additional information

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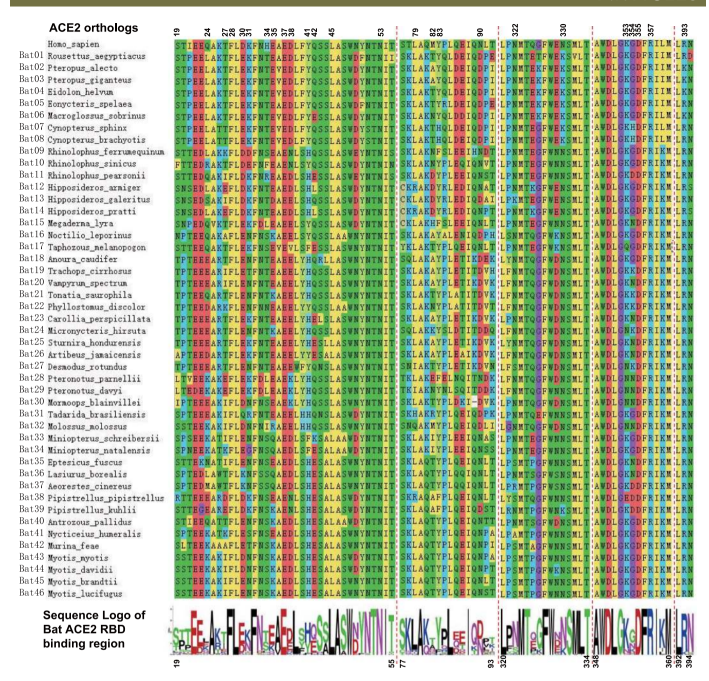
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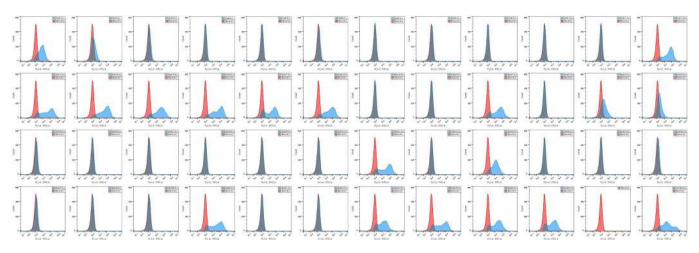
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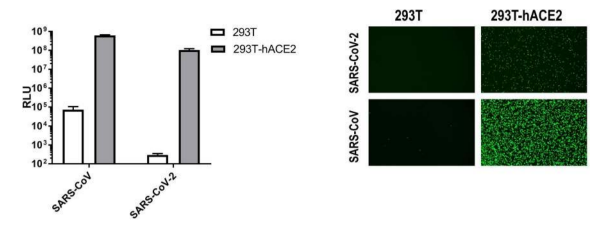


Extended Data Fig. 1 | Sequence alignment of RBD interacting region on 46 bat ACE2 orthologs. The alignment of RBD interaction regions were generated by MEGA-X software. Amino acids were coloured based on their residue features. The residues involved in the interaction between human ACE2 and SARS-CoV-2 RBD were indicated on the top. Sequence logo of the corresponding regions was generated by WebLogo (bottom).



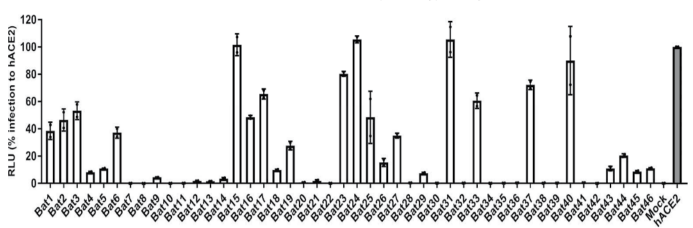
Extended Data Fig. 2 | Flow cytometry analysis of the SARS-CoV-2 RBD-hFc binding to 293T cells expressing different bat ACE2 orthologs. 293T cells expressing bat ACE2 orthologs were incubated with 5 μ g/ml of recombinant SARS-CoV-2 RBD-hFc protein at 37 °C for 1 hour, and then washed and incubated with a Alexa Fluor 488 conjugated secondary antibody recognizing human lgG Fc. Histogram charts were generated by FlowJo with mock cells (sample 47) as negative control.

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Extended Data Fig. 3 | Verification of SARS-CoV and SARS-CoV-2 pseudotypes on 293T cells expressing human ACE2. A high signal/background ratio of viral entry can be achieved on 293T-hACE2 cells for SARS-CoV and SARS-CoV-2 spike protein pseudotyped VSV-dG viruses expressing Firefly Luciferase (left) and GFP (right).

SARS-CoV-2 pseudotype entry



Extended Data Fig. 4 | Infection profile of virus pseudotyped with SARS-CoV-2 S protein without D614G variation. 293T cells expressing the indicated bats ACE2 orthologs were infected with SARS-CoV-2 (Wuhan-Hu-1 strain, without D614G variation) S protein pseudotyped VSV-dG-luc viruses. Luciferase units were determined at 20 hpi. Scale bar = 200 µm. The data are presented as the Mean±SD (n=3).

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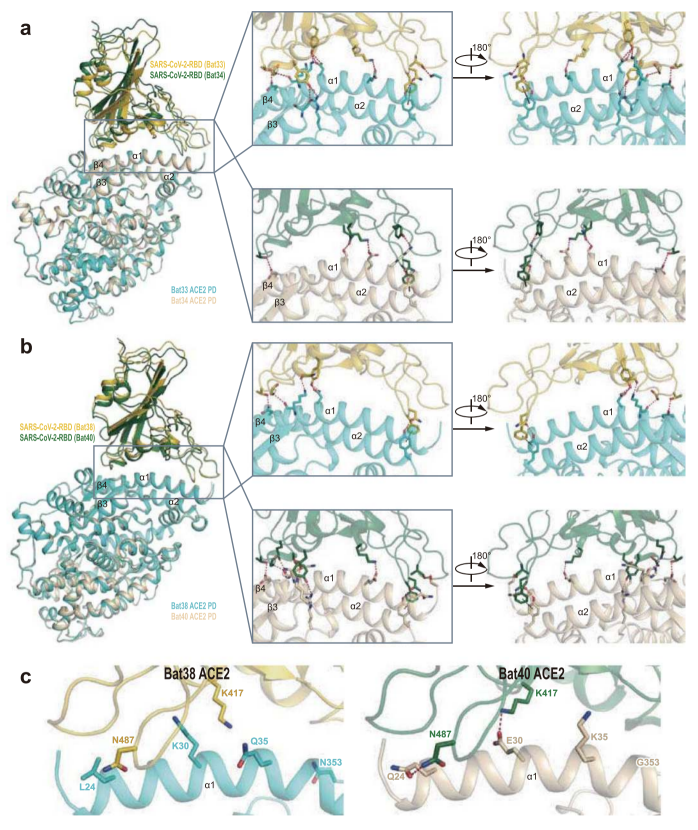
SARS-CoV-GFP infection

Bat1	Bat2	Bat3	Bat4	Bat5	Bat6	Bat7	Bat8	Bat9	Bat10	<u>Bat11</u>	<u>Bat12</u>
<u>Bat13</u>	Bat14	Bat15	Bat16	<u>Bat17</u>	Bat18	Bat19	Bat20	<u>Bat21</u>	Bat22	<u>Bat23</u>	<u>Bat24</u>
Bat25	Bat26	Bat27	<u>Bat28</u>	Bat29	<u>Bat30</u>	Bat31	<u>Bat32</u>	Bat33	<u>Bat34</u>	Bat35	<u>Bat36</u>
<u>Bat37</u>	Bat38	<u>Bat39</u>	Bat40	Bat41	Bat42	Bat43	Bat44	Bat45	Bat46	<u>Mock</u>	hACE2

SARS-CoV-2-GFP infection

Bat1	Bat2	Bat3	Bat4	Bat5	Bat6	Bat7	Bat8	Bat9	<u>Bat10</u>	<u>Bat11</u>	<u>Bat12</u>
<u>Bat13</u>	Bat14	Bat15	Bat16	Bat17	Bat18	Bat19	<u>Bat20</u>	<u>Bat21</u>	Bat22	Bat23	Bat24
Bat25	Bat26	Bat27	<u>Bat28</u>	Bat29	<u>Bat30</u>	Bat31	Bat32	Bat33	Bat34	<u>Bat35</u>	<u>Bat36</u>
Bat37	<u>Bat38</u>	<u>Bat39</u>	Bat40	<u>Bat41</u>	<u>Bat42</u>	Bat43	Bat44	Bat45	Bat46	<u>Mock</u>	hACE2

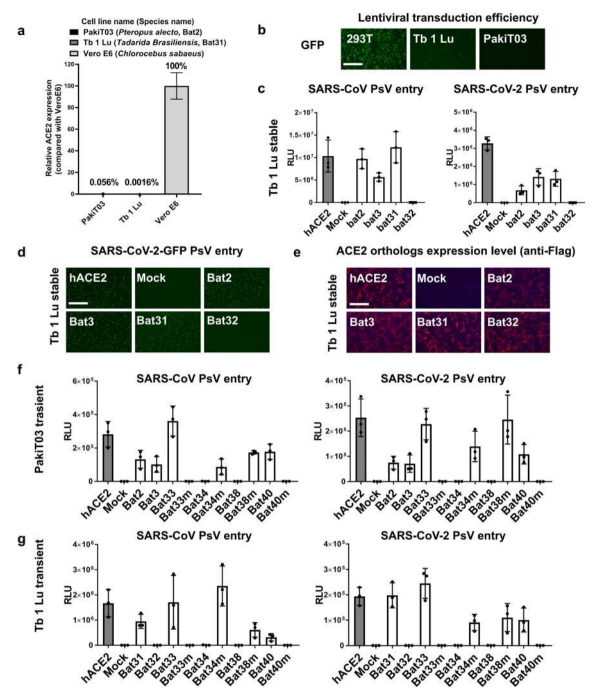
Extended Data Fig. 5 | The ability of bat ACE2 orthologs to support the entry of SARS-CoV and SARS-CoV-2 pseudotyped VSV-dG-GFP viruses. 293T cells expressing the ACE2 orthologs of the indicated bats were infected with SARS-CoV and SARS-CoV-2 pseudotyped VSV-dG-GFP viruses. Images were captured at 20 hpi. Scale bar = $200 \, \mu m$.



Extended Data Fig. 6 | See next page for caption.

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Extended Data Fig. 6 | Interface comparison between bat ACE2 with SARS-CoV-2-RBD. a, Structure alignment for the Bat33/SARS-CoV-2-RBD and Bat34/SARS-CoV-2-RBD. Bat33 and its complexed RBD are coloured cyan and gold, respectively; Bat34 and its complexed RBD are coloured wheat and green, respectively. The inset shows the interface of SARS-CoV-2-RBD with different ACE2. The residues which form hydrophilic interactions and aromatic interactions are shown as sticks. Here and subsequently, dotted lines between residues indicate interaction types: red for hydrogen bonds or ionic bonds and black for π - π interactions. The right panel was obtained by clockwise rotation of the middle panel along a longitudinal axis. **b**, Structure alignment for the Bat38/SARS-CoV-2-RBD and Bat40/SARS-CoV-2-RBD. Bat38 and its complexed RBD are coloured cyan and gold, respectively; Bat40 and its complexed RBD are coloured wheat and green, respectively. The inset shows the interface of SARS-CoV-2-RBD with different ACE2. The residues which form hydrophilic interactions and aromatic interactions are shown as sticks. The right panel was obtained by clockwise rotation of the middle panel along a longitudinal axis. **c**, Comparison of the interface between Bat38/SARS-CoV-2-RBD and Bat40/SARS-CoV-2-RBD. The mutated residues in ACE2 and the corresponding residues in RBD are shown and labelled.



Extended Data Fig. 7 | ACE2 is the primary receptor for the species-specific entry of SARS-CoV and SARS-CoV-2 in bat cell lines. a, The relative expression level of ACE2 orthologs in Vero-E6 and bat cell lines determined by Quantitative reverse transcription PCR(RT-qPCR). The data are presented as the Mean±SEM (n=3). b, Lentiviral transduction efficiency of 293T (high), Tb 1 Lu (medium) and PakiTO3 (very low). Cells were transduced with lentivirus carrying EGFP, and the images were captured at 72h post-transduction. Tb 1 Lu cell were selected for generating stable cell lines because of its higher transduction efficiency. c-e, The entry of SARS-CoV and SARS-CoV-2 pseudovirus into Tb 1 Lu cells stably expressing different ACE2 orthologs. The CoV entry efficiency was examined on Tb1-Lu stable-cells two weeks post lentiviral transduction (c,d). The data in c are presented as the Mean±SD (n=3). The expression level of ACE2 orthologs was detected by immunofluorescence detecting the C terminal fused Flag tags (e). f-g, The entry of SARS-CoV and SARS-CoV-2 pseudovirus into bat cell lines transiently transfected with human and bat ACE2 orthologs. Scale bar = 200 μm. The data are presented as the Mean±SD (n=3).

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		Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.					
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Da	ata analysis	GraphPad Prism 7 was used in this study. FlowJo version 10 was used for analyzing flow cytometry data					
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Randomization	In this study, v	we included as many bat ACE2 sequences as possible. No grouping is applied				
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Antibodies used Flag tag (9A3, #8146S, Cell signaling technology, United States); Secondary goat anti-mouse antibody conjugated with Alexa (A11032, Thermo Fisher Scientific, United States); Alexa Fluor 488 conjugated Goat anti-Human IgG (A11013, Thermo Fishe Scientific, United States)						
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Eukaryotic co	ell lines					
Policy information a	about <u>cell line</u>	<u> </u>				
Cell line source(s)		HEK293T cells (293T, ATCC, CRL-3216); Tb 1 Lu cells (ATCC, CCL-88); PakiT03 cells (a gift from professor Peng Zhou); VERO-E6 cells (ATCC, CRL-1586); I1-Hybridoma (ATCC, CRL-2700)				

Cell lines used in this study were not tested for mycoplasma contamination

Authentication

(See <u>ICLAC</u> register)

Mycoplasma contamination

Commonly misidentified lines

Not authenticated

No commonly misidentified line was used

Flow Cytometry

Plots

Confirm that:					
The axis labels state the ma	arker and fluorochrome used (e.g. CD4-FITC).				
The axis scales are clearly v	visible. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).				
All plots are contour plots	with outliers or pseudocolor plots.				
A numerical value for num	ber of cells or percentage (with statistics) is provided.				
Methodology					
Sample preparation	The 293T cells stably overexpressing different ACE2 orthologs were incubated with RBDs and antibody in two consecutive staining steps. The first was an incubation with SARS-CoV-RBD-hFc or SARS-CoV-2-RBD-hFc (5-10 μ g/ml) in DMEM with 2% FBS for 1 hour at 37 $^{\circ}$ C. The second utilized an Alexa Fluor 488 conjugated Goat anti-Human IgG (A11013, Thermo Fisher Scientific, United States) in 2 μ g/ml for additional 30min at 37 $^{\circ}$ C. The stained cells were detached by 5mM EDTA/PBS before strained through a 70 μ m cell mesh, and then analyzed with a CytoFLEX Flow Cytometer (Beckman Coulter, United States). More information are available on Methods sections				
Instrument	CytoFLEX Flow Cytometer (Beckman Coulter, United States)				
Software	FlowJo version 10				
Cell population abundance	We did not count the cell population percentage				
Gating strategy	The RBD-hFc binding 293T cells were gated as Alexa Fluor 488 conjugated Goat anti-Human IgG+ single cells. More information are available on Methods sections				
Tick this box to confirm that	at a figure exemplifying the gating strategy is provided in the Supplementary Information.				