# SARS-CoV-2 spike protein predicted to form stable complexes with host receptor protein orthologues from mammals, but not fish, birds or reptiles

Lam SD<sup>1,2,†</sup>, Bordin N<sup>2,†</sup>, Waman VP<sup>2</sup>, Scholes HM<sup>2</sup>, Ashford P<sup>2</sup>, Sen N<sup>2,3</sup>, van Dorp L<sup>4</sup>, Rauer C<sup>2</sup>, Dawson NL<sup>2</sup>, Pang CSM<sup>2</sup>, Abbasian M<sup>2</sup>, Sillitoe I<sup>2</sup>, Edwards SJL<sup>5</sup>, Fraternali F<sup>6</sup>, Lees JG<sup>7</sup>, Santini JM<sup>2</sup>, Orengo CA<sup>2,\*</sup>

1 Department of Applied Physics, Faculty of Science and Technology, Universiti Kebangsaan Malaysia, Bangi, Selangor, 43600, Malaysia

2 Institute of Structural and Molecular Biology, University College London, London, WC1E 6BT, UK 3 Indian Institute of Science Education and Research, Pune, 411008, India

4 UCL Genetics Institute, University College London, London, WC1E 6BT, UK

5 Department of Science and Technology Studies, University College London, London, WC1E 6BT, UK 6 Randall Division of Cell and Molecular Biophysics, Guy's Campus, New Hunt's House, King's College London, London, SE1 1UL, UK

7 Department of Biological and Medical Sciences, Faculty of Health and Life Sciences, Oxford Brookes University, Oxford, OX3 OBP, UK

+ These authors contributed equally

\* To whom correspondence should be addressed: c.orengo@ucl.ac.uk

#### Abstract

The coronavirus disease 2019 (COVID-19) global pandemic is caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). SARS-CoV-2 has a zoonotic origin and was transmitted to humans via an undetermined intermediate host, leading to infections in humans and other mammals. To enter host cells, the viral spike protein binds to its receptor, angiotensin-converting enzyme 2 (ACE2), and is then processed by transmembrane protease serine 2 (TMPRSS2). Whilst receptor binding contributes to the viral host range, changes in energy of spike protein:ACE2 complexes in other animals have not been widely explored. Here, we analyse interactions between the spike protein and orthologues of ACE2 and TMPRSS2 from a broad range of 215 vertebrate species. Using models of spike protein:ACE2 orthologue complexes, we calculated their changes in energy, correlated to mammalian COVID-19 infection data. Across vertebrates, mutations are predicted to have more impact on the function of ACE2 than TMPRSS2. Finally, we provide phylogenetic analysis demonstrating that human SARS-CoV-2 strains have been isolated in animals. Our results suggest that SARS-CoV-2 can infect a broad range of mammals, but not fish, birds or reptiles. Susceptible animals could serve as reservoirs of the virus, necessitating careful ongoing animal management and surveillance.

**Keywords:** SARS-CoV-2, COVID-19, spike protein, ACE2, TMPRSS2, residue mutations, structural bioinformatics

### Introduction

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is a novel coronavirus that emerged towards the end of 2019 and is responsible for the coronavirus disease 2019 (COVID-19) global pandemic. Available data suggests that SARS-CoV-2 has a zoonotic source(1), with the closest sequence currently available deriving from the horseshoe bat(2). As yet, the transmission route to humans, including the intermediate host, is unknown. So far, little work has been done to assess the animal reservoirs of SARS-CoV-2, or the potential for the virus to spread to other species living with, or in close proximity to, humans in domestic, rural, agricultural or zoological settings.

Coronaviruses, including SARS-CoV-2, are major multi-host pathogens and can infect a wide range of non-human animals(3–5). SARS-CoV-2 is in the Betacoronavirus genus, which includes viruses that infect economically important livestock, including cows(6), pigs(7), mice(8), rats(9), rabbits(10), and wildlife, such as antelope and giraffe(11). Severe acute respiratory syndrome coronavirus (SARS-CoV), the betacoronavirus that caused the 2002-2004 SARS outbreak(12), likely jumped to humans from its original bat host via civets. Viruses genetically similar to human SARS-CoV have been isolated from animals as diverse as civets, racoon dogs, ferret-badgers(3) and pigs(13), suggesting the existence of a large host reservoir. It is therefore probable that SARS-CoV-2 can also infect a wide range of species.

Real-world SARS-CoV-2 infections have been reported in cats(14), tigers(15), dogs(16,17) and minks(16,17). Animal infection studies have also identified cats(18) and dogs(18) as hosts, as well as ferrets(18), macaques(19) and marmosets(19). Recent *in vitro* studies have also suggested an even broader set of animals may be infected(20–22). As with humans, age is also a predictive risk factor for rhesus macaques, with SARS-CoV-2 causing worse pneumonia in older animals(23). To understand the potential host range of SARS-CoV-2, the plausible extent of zoonotic and anthroponotic transmission, and to guide surveillance efforts, it is vital to know which species are susceptible to SARS-CoV-2 infection.

The receptor binding domain (RBD) of the SARS-CoV-2 spike protein (S-protein) binds to the extracellular peptidase domain of angiotensin I converting enzyme 2 (ACE2) mediating cell entry(24). ACE2 is highly expressed in the endothelium, kidney, lungs and heart(25,26), where it plays a role in regulating blood pressure(27), with loss of function increasing the risk of heart failure(28). The sequence of ACE2 is highly conserved across vertebrates, suggesting that SARS-CoV-2 could use orthologues of ACE2 for cell entry. The structure of the SARS-CoV-2 S-protein RBD has been solved in complex with human ACE2(29) and identification of critical binding residues in this structure have provided valuable insights into viral recognition of the host receptor(29–33). Compared with SARS-CoV, the SARS-CoV-2 S-protein has a 10-22-fold higher affinity for human ACE2(29,30,34) thought to be mediated by three classes of mutations in the SARS-CoV-2 S-protein(35). Similarly, variations in human ACE2 have also been found to increase affinity for S-protein(36) receptor binding. These factors may contribute to the host range and infectivity of SARS-CoV-2.

Both SARS-CoV-2 and SARS-CoV additionally require the transmembrane serine protease (TMPRSS2) to mediate cell entry. Together, ACE2 and TMPRSS2 confer specificity of host cell types that the virus can enter, with the S-protein preferentially binding to cells that co-express both ACE2 and TMPRSS2(26,37). Upon binding to ACE2, the S-protein is cleaved by TMPRSS2 at two cleavage sites on separate loops, which primes the S-protein for cell entry(38). TMPRSS2 has been docked against the SARS-CoV-2 S-protein, which revealed its binding site to be adjacent to these two cleavage sites(37).

An approved TMPRSS2 protease inhibitor drug is able to block SARS-CoV-2 cell entry(39), which demonstrates the key role of TMPRSS2 alongside ACE2. Furthermore, variations in TMPRSS2, as opposed to ACE2, have been suggested as associated with disease severity in COVID-19 patients(40). TMPRSS2 is frequently mutated in prostate cancer, which may contribute to the higher rate of COVID-19 mortality observed in men compared to women of the same age(41–43). As such, both ACE2 and TMPRSS2 represent attractive therapeutic targets against SARS-CoV-2(44).

The structural interplay between ACE2 and the SARS-CoV-2 spike protein has been used to predict possible hosts for SARS-CoV-2. A broad range of hosts have already been proposed, covering hundreds of mammalian species, including tens of bat species(32); primates(45) and a recent more comprehensive study analysed all classes of vertebrates(45,46), including agricultural species of cow, sheep, goat, bison and water buffalo. In addition, sites in ACE2 have been identified as under positive selection in bats, particularly in regions involved in binding the S-protein(32). The impacts of mutations in ACE2 orthologues have also been tested, for example structural modelling of ACE2 from 27 primate species(45) demonstrated that apes and African and Asian monkeys may also be susceptible to SARS-CoV-2. However, whilst cell entry is necessary for viral infection, it may not be sufficient alone to cause disease. For example, variations in other proteins may prevent downstream events that are required for viral replication in a new host. Hence, examples of real-world infections(14–17) and experimental data from animal infection studies(18–22) are required to validate hosts that are predicted to be susceptible.

Here, we analysed the effect of known mutations in orthologues of ACE2 and TMPRSS2 from a broad range of 215 vertebrate species, including primates, rodents and other placental mammals; birds; reptiles; and fish. For each species, we generated a 3-dimensional model of the ACE2 protein structure from its protein sequence and calculated the impacts of known mutations in ACE2 on the stability of the S-protein:ACE2 complex. We correlated changes in the energy of the complex with changes in the structure of ACE2, chemical properties of residues in the binding interface, and experimental infection phenotypes from *in vivo* and *in vitro* animal studies. To further test our predictions, we performed detailed manual structural analyses, presented as a variety of case studies for different species. Unlike other studies that analyse interactions that the S-protein makes with the host, we also analyse the impact of mutations in vertebrate orthologues of TMPRSS2. Our results suggest that SARS-CoV-2 can infect a broad range of vertebrates, which could serve as reservoirs of the virus, supporting future anthroponotic and zoonotic transmission.

# **Material and Methods**

#### Sequence Data

ACE2 protein sequences for 239 vertebrates, including humans, were obtained from ENSEMBL(47) version 99 and eight sequences from UniProt release 2020\_1 (<u>Supplementary Table S1</u>). TMPRSS2 protein sequences for 278 vertebrate sequences, including the human sequence, were obtained from ENSEMBL (<u>Supplementary Table S2</u>).

A phylogenetic tree of species, to indicate the evolutionary relationships between animals, was downloaded from ENSEMBL(47).

#### **Structural Data**

The structure(29) of the SARS-CoV-2 S-protein bound to human ACE2 at 2.45Å was used throughout (PDB ID 6M0J).

# Methods

#### Analyses of sequence similarity between human ACE2 and other vertebrate species

Vertebrate protein sequences were aligned to each other pairwise, and to the human ACE2 sequence, using BLASTp(48).

# Domain family detection, alignment construction and conservation of residues in ACE2 and TMPRSS2

In order to detect highly conserved residues likely to be implicated in the domain function (i.e. ACE2 binding residues, TMPRSS2 binding residues in the active site) we mapped the domain sequence to the appropriate functional family (FunFam) in the CATH classification(49). FunFams are clusters of evolutionary related domain sequences, predicted to have highly similar structures and functions. They have been previously used to analyse the impact of genetic variation on protein function in the context of antibiotic resistance phenotypes(50) and cancer progression(51).

The ACE2 (or TMPRSS2) sequences were scanned against the seed FunFam hidden Markov model library for the next release of CATH-Gene3D(52) v4.3 (unpublished data) with the best matches resolved by cath-resolve-hits(53) using a bit score cut-off of 25 and a minimum query coverage of 80%. Each hit was subsequently re-aligned to the matching FunFam using Clustal(54) in Jalview(55).

The conservation scores for residues in the ACE2 and TMPRSS2 domains were obtained using ScoreCons(56). Information contents of FunFam multiple sequence alignments were calculated as the diversity of positions(56) (DOPs) score. Residues are considered highly conserved for DOPs  $\geq$  70 and ScoreCons  $\geq$  0.7.

#### Structure analyses

#### Identifying residues in ACE2

In addition to residues in ACE2 that contact the S-protein directly, various other studies have also considered residues that are in the second shell, or are buried, and could influence binding(57). Therefore, in our analyses we built on these approaches and extended them to compile the following sets for our study :

- 1. *Direct contact (DC)* residues. This includes a total of 20 residues that are involved in direct contact with the S-protein(29) identified by PDBe(58) and PDBSum(59).
- 2. Direct Contact Extended (DCEX) residues. This dataset includes residues within 8Å of DC residues, that are likely to be important for binding. These were selected by detailed manual inspection of the complex, and also considering the following criteria: (i) reported evidence from deep mutagenesis(57), (ii) in silico alanine scanning (using mCSM-PPI(60)), (iii) residues with high evolutionary conservation patterns identified by the FunFams-based protocol described above, i.e. residues identified with DOPS ≥ 70 and ScoreCons score ≥ 0.7, (iv)

allosteric site prediction, and (v) sites under positive selection. Selected residues are shown in Supplementary Figure S1 and residues very close to DC residues (i.e. within 5Å) are annotated.

To identify residues involved in the S-protein:ACE2 interface, we extracted information from a range of sources including PDBe(58), PDBsum(59), examined structural evidence (such as from crystallography, cryo-EM and homology modelling) in the literature(21,29,37,45,57,61), and performed manual inspection of the human complex (PDB ID 6MOJ) using Chimera(62) (<u>Supplementary Figure S1</u>). We also examined residues identified by other studies including alanine scanning mutagenesis, deep mutagenesis experiments(29,45,61,63,64), and methods identifying sites under positive selection(32,46).

In addition, in order to identify potential allosteric residues within the ACE2 DCEX set, three different allosteric prediction methods were used: AlloSitePro(65), ENM(66) and PARS(67). The protein structure PDB ID 6MOJ(29) was used as input, which contains information on ligand binding sites for zinc, chlorine, and N-Acetylglucosamine. PARS is a normal mode analysis method which calculates normal modes in the presence and absence of a simulated allosteric modulator. A site is predicted as allosteric if the motions are significantly different. ENM identifies hinge residues as potential functional residues, and relies exclusively on inter-residue contact topology. It uses features of spatial clustering properties and relative solvent accessibility to identify functionally important hinge residues. AlloSitePro uses both pocket-based structure features with normal mode analyses. The performance of AllositePro has been endorsed by recent studies(68).

We also identified DC and DCEX sites under positive selection using codon-based methods, including mixed effect model of evolution(69), available at the Datamonkey Adaptive Evolution web-server(70). These methods estimate dN/dS ratio for every codon in an alignment. We analysed evidence of positive selection using a codon alignment of all ACE2 orthogue sequences. Potential recombinant sequences were identified using RDP(71) version 5 and were excluded prior to selection pressure analyses.

#### **Generating 3-dimensional structure models**

#### ACE2

Using the ACE2 protein sequence from each species, structural models were generated for the S-protein:ACE2 complex for 247 animals using the FunMod modelling pipeline(72,73), based on MODELLER(74). Given that the S-protein has the same sequence in PDB ID 6M0J, and ACE2 is conserved in vertebrates, models are expected to be high-quality. Models were refined by MODELLER to optimise the geometry of the complex and the interface. Only high-quality models were used in this analysis, with nDOPE(75) score < -1 and with < 10 DCEX residues missing. This gave a final dataset of 215 animals for further analysis.

The modelled structures of ACE2 were compared against the human structure (PDB ID 6M0J) and pairwise, against each other, using SSAP(76). SSAP returns a score in the range 0-100, with identical structures scoring 100.

#### TMPRSS2

There is currently no solved structure for TMPRSS2. To build a structural model of TMPRSS2, we selected the best TMPRSS2 model template by performing a HMMER(77) search of the human TMPRSS2 sequence (UniProt accession O15393) against the FunFams, identifying four potential structural templates (CATH domain IDs 2f83A05, 5eodA05, 5eokA05 and 5i25A05) within FunFam 2.40.10.10-FF-2 (best E-value = 3e-96). After modelling with FunMod, the structure with the best score (nDOPE = -0.773) was selected, based on PDB ID 5I25.

We identified the relevant residues in TMPRSS2 by extracting the active site and cleavage site residue information available on the human TMPRSS sequence (UniProt accession O15393). After we modelled the human TMPRSS2, two sets of residues were extracted:

Set 1: Direct Contact (ASCS) residues comprising the active site (three residues) and the cleavage site (2 residues).

Set 2: Direct Contact Extended (ASCSEX) residues: This dataset includes the residues from Set 1 and additional residues within 8Å of ASCS residues. We only analysed highly conserved residues (i.e. high ScoreCons  $\geq$  0.7) from Set 2, which are likely to have a functional role (21 residues).

#### Measuring changes in the energy of the S-protein:ACE2 complex in SARS-CoV-2 and SARS-CoV

Since we only have a protein structure for human S-protein:ACE2, we modelled the structure of all the other animal species using established protocols(72,73). We observed very high conservation of ACE2 amongst animals, with most species sharing 60% or more identical residues with human ACE2. We were, therefore, confident of obtaining high quality models in which the binding interface would be well constructed. This was assessed using standard measures and we removed any animal species with poor models from the analysis.

We calculated the changes in binding energy of the SARS-CoV-2 S-protein:ACE2 complex and the SARS-CoV S-protein:ACE2 complex of different species, compared to human, following three different protocols:

- 1. Protocol 1: Using the human complex and mutating the residues for the ACE2 interface to those found in the given animal sequence and then calculating the  $\Delta\Delta G$  of the complex using both mCSM-PPI1(60) and mCSM-PPI2(78). This gave a measure of the destabilisation of the complex in the given **animal relative to the human** complex.  $\Delta\Delta G$  values < 0 are associated with destabilising mutations, whilst values  $\geq$  0 are associated with stabilising mutations.
- 2. Protocol 2: We repeated the analysis with both mCSM-PPI1 and mCSM-PPI2 as in protocol 1, but using the animal 3-dimensional models, instead of the human ACE2 structure, and calculating the  $\Delta\Delta G$  of the complex by mutating the animal ACE2 interface residue to the appropriate residue in the human ACE2 structure. This gave a measure of the destabilisation of the complex in the human complex relative to the given animal. Values  $\leq$  0 are associated with destabilisation of the human complex (i.e. animal complexes more stable), whilst values > 0 are associated with stabilisation of the human complex (i.e. animal complexes less stable).
- 3. *Protocol 3:* We used the PRODIGY server(79) to calculate the binding energy for the human complex and for the 3-dimensional models of the other 215 animal complexes. We then calculated the change in binding energy from the human complex to the animal complex.

mCSM-PPI1 assigns a graph-based signature vector to each mutation, which is then used within machine learning models to predict the binding energy. The signature vector is based upon atomdistance patterns in the wild-type protein, pharmacophore information and available experimental information. The more recent mCSM-PPI2 is a development of the former with an improved description of the signature vector including, among other features, evolutionary information and energetic terms. For both mCSM-PPI1 and mCSM-PPI2 we used the mCSM server for the simulations.

As a third method, we used the PRODIGY web-server which calculates the binding energy as a function of inter-residue contacts and the non-interacting surface of the residues. As this method delivers an absolute binding energy we calculate the relative binding energy as  $\Delta\Delta G = \Delta G_{human} - \Delta G_{animal}$ . The calculations, using the PRODIGY web-server, were performed at 298 and 310 Kelvin.

By exploring multiple protocols and tools we hoped to gain some insights into the dependence of our analyses, and insights derived from them, on the methodology used. mCSM-PPI1 was much faster to run than mCSM-PPI2, so we considered this method as well, since it could allow such analyses to be extended to more animals in the future.

We subsequently correlated  $\Delta\Delta G$  values with available *in vivo* and *in vitro* experimental data on COVID-19 infection data for mammals. Protocol 2, mCSM-PPI2, correlated best with these data. This allowed us to assign thresholds for risk of infection by SARS-CoV-2 on the  $\Delta\Delta G$  values with  $\Delta\Delta G \leq 1$  high risk, 1 <  $\Delta\Delta G < 2$  medium risk and  $\Delta\Delta G \geq 2$  low risk.

#### Change in residue chemistry for mutations

To measure the degree of chemical change associated with mutations occurring in the key interface residues, we computed the Grantham score(80) for each vertebrate compared to the human sequence. For the ACE2-TMPRSS2 comparison we used the same set of organisms as for ACE2, but only those that had at least one TMPRSS2 protein available from ENSEMBL, thus reducing the number of available organisms to 156.

The sums and averages of Grantham scores were obtained for DC and DCEX residues in ACE2. For TMPRSS2 the sum and average Grantham values were obtained for the active site, cleavage site residues (ASCS) and the residues 8Å from the active site (ASCSEX), having ScoreCons  $\geq$  0.7. Data and plots for the various combinations are available in <u>Supplementary Table S3</u>.

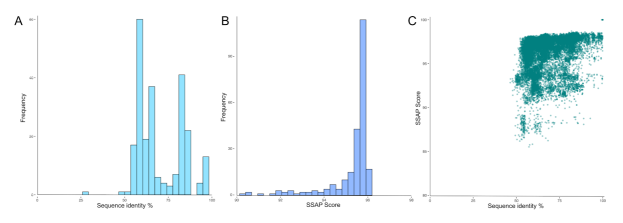
#### Phylogeny of SARS-like betacoronaviruses

Genome assemblies were downloaded for a subset of SARS-CoV (n = 10), SARS-like (n = 27) and SARS-CoV-2 (n = 13) viruses from publicly available data on NCBI(81–88) and GISAID(16,17)H For a full list of metadata, as compiled by NextStrain(89), including contributing and submitting laboratories, see <u>Supplementary Table S4</u>. All assemblies were downloaded and aligned against the SARS-CoV-2 reference genome Wuhan-Hu-1 (NCBI NC\_045512.2) using MAFFT(90). The alignment was manually inspected and the first 130 bp and last 50 bp were masked. A maximum likelihood phylogenetic tree was built on the alignment using RaXML-NG(91) under the GTR+G substitution model with 100 bootstraps to assess branch support. The resulting phylogeny was plotted using ggtree(92) v1.16.6. The spike ORF (21563:25384 nt relative to Wuhan-Hu-1) was extracted and translated to amino acid sequence using Ape(93) v5.3. The alignment of spike protein and receptor binding domain sequences were assessed and visualised using ggtree.

#### Results

#### **Conservation of ACE2 in vertebrates**

We aligned protein sequences of 247 vertebrate orthologues of ACE2. Most orthologues have more than 60% sequence identity with human ACE2 (Figure 1A). For each orthologue, we generated a 3-dimensional model of the protein structure from its protein sequence using FunMod(72,73). We were able to build high-quality models for 236 vertebrate orthologues, with nDOPE scores > -1 (Supplementary Table S5), because 11 low-quality models were removed from the analysis. After this, we removed a further 21 models from the analysis that were missing > 10 DCEX residues, leaving 215 models to take forward for further analysis. The structural models were compared to the human ACE2 protein structure using SSAP(76). Most models had high structural similarity to human ACE2 (SSAP > 90) (Figure 1B and Figure 1C). However, despite sharing > 60% sequence identity to the human ACE2, some models had lower structural similarity (Figure 1C). These changes probably reflect mutations slightly altering the relative orientations of the S-protein and ACE2 across the interface. (Supplementary Figure S2).

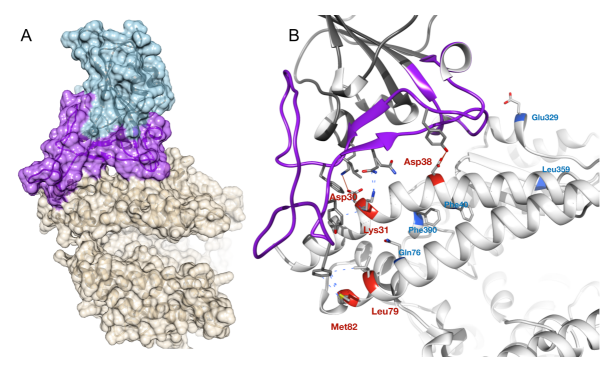


<u>Figure 1.</u> Conservation of ACE2. (a) ACE2 sequence conservation in vertebrates. Distribution of pairwise sequence identities of ACE2 orthologues compared to human ACE2. (b) ACE2 structure conservation in vertebrates. Distribution of pairwise SSAP scores of ACE2 orthologues compared to human ACE2. (c) Pairwise sequence identities and SSAP scores for ACE2 orthologues.

#### Identification of critical S-protein:ACE2 interface residues

Residues in the binding interface between the S-protein and ACE2 were identified in a structure of the complex (PDB ID 6M0J; Figure 2a, Supplementary Figure S1). Residues within 5Å of the interacting protein are reported in PDBe and PDBsum(58,59), which we refer to as 'Direct Contact' (DC) residues. Other important residues likely to influence binding have been identified using a variety of approaches, including structural analysis by us and other groups, alanine scanning, and mutagenesis(21,29,37,45,57,61,94). In addition, we identified highly conserved residues, sites under positive selection and predicted allosteric sites. This larger set includes 42 residues that are not involved in direct contact with the interacting protein but could still influence the binding affinity. 15 of these residues are within 5Å and 27 residues are further away but within 8Å of one, or more, DC residues. We, therefore, compiled a set of residues that includes these residues, together with the DC residues. We refer to this as the direct contact plus 'extended set' of residues ('DCEX' residues) (Figure

<u>2b</u>). Some of the ACE2 DCEX residues are highly-conserved positions in the ACE2 FunFam (see <u>Supplementary Figure S1</u> for summary of evidence for each residue in the DCEX set).



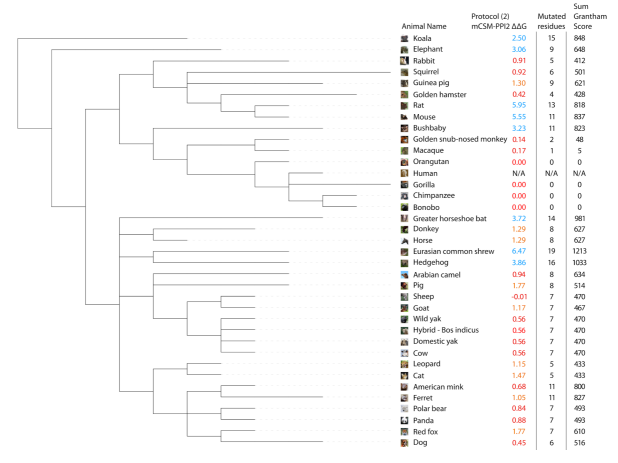
<u>Figure 2</u>. Overview of SARS-CoV-2 S-protein:human ACE2 complex and interface. (a) SARS-CoV-2 Sprotein RBD (light blue, purple) showing the receptor binding motif (purple) at the interface with ACE2 (tan). (b) DC residues in the SARS-CoV-2 S-protein:human ACE2 complex interface. A subset of contact residues that are not conserved in vertebrates are highlighted for DC (red) and DCEX residues (blue) (PDB ID: 6M0J).

#### Changes in the energy of the S-protein:ACE2 complex in vertebrates

We used two protocols, protocols 1 and 2, to assess the relative change in binding energy ( $\Delta\Delta$ G) of the SARS-CoV-2 S-protein:ACE2 complex following mutations in the DC and DCEX residues. Since these residues lie directly in the binding interface (DC residues) or in the secondary shell and are likely to be influencing binding, we hypothesise that the changes in energy of the complex caused by mutations in these residues can be used to gauge the relative risk of infection for each animal. Our third protocol directly measured the binding energies for SARS-CoV-2 S-protein:ACE2 complex for the human structure and the modelled structures of each animal.

We examined how well these approaches correlated with each other (Supplementary Figure S3-S4-S5). We also correlated the  $\Delta\Delta$ G values (protocols 1 and 2) and binding energies (protocol 3) with SARS-CoV-2 infection phenotypes in different animals(2,18–22,95,96) (Supplementary Table S6; phenotype data are sourced from *in vivo*(18,19,95,97,98) and *in vitro*(20–22) infection studies, some of which have not been peer reviewed). We found that protocol 2 employing mCSM-PPI2 (henceforth referred to as P(2)-PPI2), calculated over the DCEX residues, correlated best with the phenotype data (Supplementary Figure S6), justifying the use of animal models to calculate  $\Delta\Delta$ G values in this context. Since this protocol considers mutations from animal to human, lower  $\Delta\Delta$ G values correspond to stabilisation of the animal complex relative to the human complex, and therefore higher risk of infection. To consider these relative  $\Delta\Delta$ G values in an evolutionary context, we annotated phylogenetic trees for all the 215 vertebrate species analysed (Supplementary Figure S7). Figure 3

shows the phylogenetic tree for a subset of these 215 species that humans come into close contact with in domestic, agricultural or zoological settings together with the  $\Delta\Delta G$  values, the number of residue changes compared to human ACE2 sequence and the total chemical shift across for the DCEX residues, for each animal. We show the residues that P(2) reports as stabilising or destabilising for the SARS-CoV-2 S-protein:ACE2 animal complex for DC (Supplementary Figure S8) and DCEX (Supplementary Figure S9) residues.



<u>Figure 3</u> Phylogenetic tree of species that humans come into close contact with in domestic, agricultural or zoological settings. Leaves are annotated by the change in energy of the complex ( $\Delta\Delta G$ ), as measured by protocol 2 - PPI2. Animals are categorised according to risk of infection by SARS-CoV-2, with  $\Delta\Delta G \leq 1$  high risk (red),  $1 < \Delta\Delta G < 2$  medium risk (orange) and  $\Delta\Delta G \geq 2$  low risk (blue). These thresholds were chosen as they agree well with the available experimental data. This tree contains a subset of animals from the larger ENSEMBL tree (<u>Supplementary Figure S7</u>).

The  $\Delta\Delta G$  values measured by P(2)-PPI2 correlate well with the infection phenotypes (<u>Supplementary</u> <u>Table S6</u>). As shown in previous studies, many primates are at high risk(19,45,46). Exceptions include New World monkeys, for which the capuchin and the squirrel monkey all show no infection risk in experimental studies, in agreement with our predicted energies for the complex(20).

According to our relative  $\Delta\Delta G$  values, zoological animals that come into contact with humans, such as pandas, leopards and bears, are also at risk of infection. In the agricultural context, camels, cows, sheep, goats and horses also have relatively low  $\Delta\Delta G$  values suggesting comparable binding affinities to humans, as validated by the experimental data(20) (Supplementary Table S6). Whilst in domestic settings, dogs, cats, hamsters, and rabbits are also at risk from infection.

In general we see a high infection risk for most mammals, with a notable exception for all nonplacental mammals. Importantly, mice and rats are not susceptible, so hamsters and ferrets are being used as models of human COVID-19, instead of mice.

Of the 35 birds tested only the blue tit shows an infection risk. Similarly, the Nile tilapia is the only fish out of the 72 in this study which shows a low change in energy of the complex, suggesting susceptibility to infection. Also, all 14 reptiles and amphibians we investigated do not show any risk.

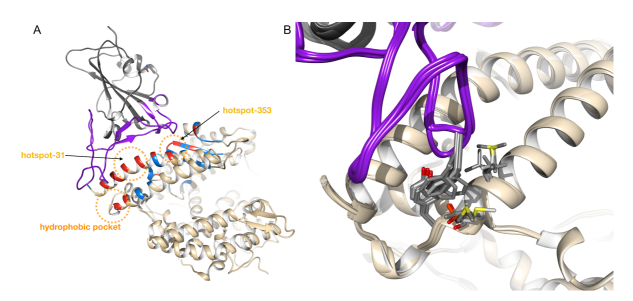
The analysis shows that, with few exceptions, placental mammals are most susceptible to infection by SARS-CoV-2. Our predictions do not always agree with the experimental data. For some cases, we predict that some animals are at medium risk of infection, in conflict with experimental data. For example, we predict that guinea pigs and donkeys have a medium risk of infection, but no infections were observed *in vitro* for these animals(21). However, infection has been observed *in vitro* for horse(21) and horse and donkey have identical DCEX residues and the same  $\Delta\Delta G(21)$  On the other hand, we predict that some animals are at low risk of infection, despite experimental evidence to the contrary. For example, *in vivo* studies have shown that horseshoe bats(2) and marmosets(19) can be infected by SARS-CoV-2, but we predict that both animals have a low risk of infection, in agreement with *in vitro* data on marmosets(20). We consider these discrepancies further using detailed structural analyses below.

#### Structural analyses of S-protein:ACE2 interactions

#### Cross-species comparison

We selected nine animals for structural analyses because they are likely to come into frequent contact with humans in domestic, zoological or agricultural settings (cat, dog, macaque, cow, sheep and camel) or because there were discrepancies between our energy calculations and currently available COVID-19 infection data (guinea pig, marmoset and horseshoe bat).

Structural variations in the S-protein:ACE2 interface amongst models of these nine species are highlighted with reference to the human PDB structure (PDB ID 6MOJ) and in the context of three sites on the interface: hydrophobic pocket, hotspot-353 and hotspot-31(30) (Figure 4a). These sites have previously been identified(30) as key to understanding why the SARS-CoV-2 S-protein binds to human ACE2 with high affinity and how the viral S-protein has evolved to bind with much higher affinity to human ACE2 than SARS-CoV(29,30,34).



<u>Figure 4</u>: Structural features of the S-protein from SARS-CoV-2 and SARS-CoV bound to ACE2. (a) Three key interaction sites for the SARS-CoV-2 and SARS-CoV S-proteins with ACE2. (b) Variations in hydrophobic pocket residues at the S-protein:ACE2 interface and the range of conformations adopted by RBD residue Phe486 in 6 animal species.

#### Hydrophobic pocket

The S-protein in SARS-CoV-2 has increased flexibility of the ACE2 interface loop due to a four residue motif (Gly-Val-Glu-Gly) in place of three in SARS-CoV (Pro-Pro-Ala), allowing for a more compact interface and insertion of SARS-CoV-2 RBD's Phe486 into a hydrophobic pocket in human ACE2 comprising Leu79, Met82 and Tyr83(30). We assessed conformations of S-protein Phe486 and these three pocket residues for each of nine species compared to human. Phe486 binds in a similar conformation with respect to the hydrophobic pocket in humans, marmoset, macaques, guinea pigs, camels, cats and cows (Figure 4b).

#### Hotspot-353

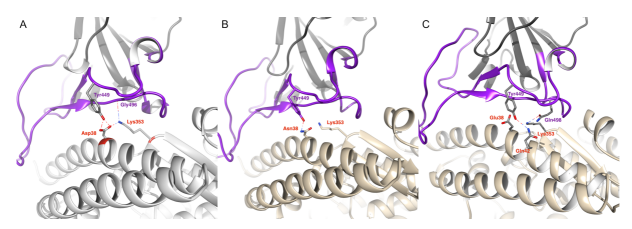
ACE2 Lys353 is highly conserved and a key mediator of the SARS-CoV and SARS-CoV-2 S-protein interfaces(30). Human Lys353 forms a salt bridge with Asp38 and these residues form H-bonds with the S-protein's RBD (Figure 5a). These interface interactions are significantly disrupted in horseshoe bat which has uncharged residue Asn38 in place of aspartate. This leads to loss of the salt bridge with Lys353 and a change in its conformation that no longer has an H-bond with S-protein Gly496 due to an increase in distance to 4.49Å (Figure 5b).

In cat and dog the physicochemically similar but bulkier glutamate replaces aspartate, leading to loss of both the salt bridge and H-bond interactions at this position. However, we observed a reconfiguration of ACE2 residues permitting alternative H-bonds to the S-protein RBD residues (<u>Figure 5c</u>).

#### Hotspot-31

In this hotspot, ACE2 residues Lys31 and Glu35 are highly conserved. However, residue 31 is mutated to Glu in guinea pigs and to Asp in horseshoe bat and camel. Residues on the S-protein in contact with position 31 in ACE2 are hydrophobic. Somewhat diverse conformations were observed at this site in

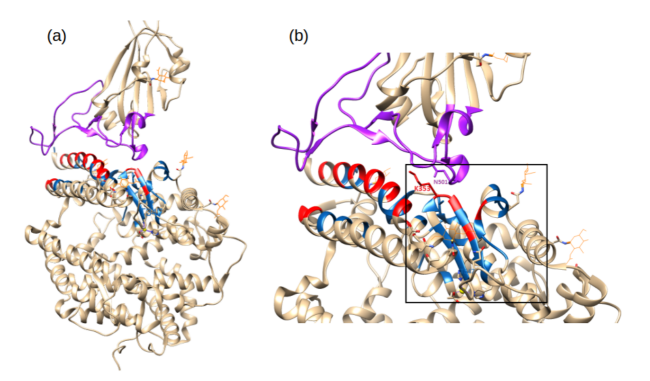
guinea pig, horseshoe bat and camel (<u>Supplementary Figure S10</u>). In guinea pig, mutation of Glu35 to Lys appears to no longer support H-bonds due to a pronounced change in conformation. Therefore, changes at this position are harder to interpret than for the other key sites.



<u>Figure 5:</u> S-protein-ACE2 Interface interactions near the Lys353 hotspot are altered in species with Asp38 variants. (a) Human hotspot with salt-bridge Lys353-Asp38 and H-bonds to S-protein Tyr449 and Gly496. (b) Horseshoe bat Asn38 forms only H-bond with Tyr449. (c) In cat and dog bulkier Glu38 no longer forms salt bridge or H-bonds, but alternative interactions are predicted as H-bonds Lys353-Gln498 and Gln42-Tyr449.

#### Potential allosteric region formed by non-interface residues

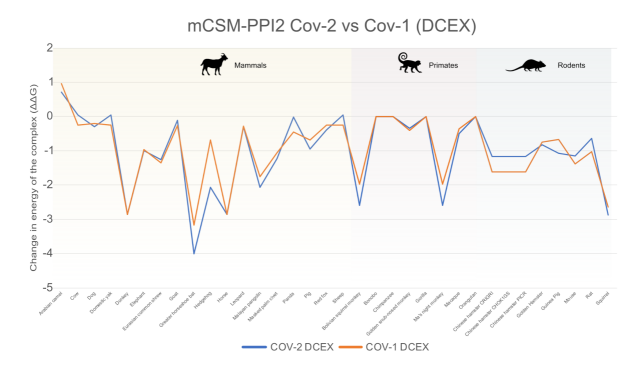
We identified a cluster of residues in our DCEX set (Figure 6) in close proximity to the wellcharacterized hotspot residue K353 of ACE2. These residues lie in the  $\beta$ 3 and  $\beta$ 4 strands that are close to the loop containing K353. Most of these residues are highly conserved (ScoreCons scores  $\geq$  0.7 and 10 residues (59%) have ScoreCons scores  $\geq$  0.90). The proximity of the residues to the interface suggests a possible allosteric role for these very highly conserved residues. Furthermore, various methods for predicting allosteric sites in protein structures confirmed that this region is likely to be involved in allosteric regulation of the binding (see <u>Supplementary Text</u> for more details). Although most of these residues are highly conserved across animals, 168 animals have  $\geq$  one mutation and more than 100 animals have  $\geq$  three mutations in this region. However most of these changes are quite conservative. The only significant change is the L359K mutation, which moderately stabilises the complex in many animals.



<u>Figure 6</u>: Potential allosteric region, identified in this study: The potential allosteric cluster (shown in blue) formed by non-interface residues is indicated in the box. The cluster is formed near the known hotspot residue K353 in ACE2. The cluster is formed by residues (83: 347-350, 352), (84: F356, I358, L359 and M360), which lie close to the loop containing the hotspot residue K353. Additionally residues, M323, T324, Q325, S331, M332, L333, W328 and E329 also form part of the cluster.

#### Changes in energy of the S-protein:ACE2 complex in SARS-CoV-2 and SARS-CoV

Farmed civets, infected by SARS-CoV in the 2002-2004 SARS epidemic(99), are thought to have been intermediate hosts of SARS-CoV(100), and thousands were culled in China to help to control the epidemic(101). Using the SARS-CoV S-protein sequence, we analysed changes in energy of the S-protein:ACE2 complex for the 215 animals in our study. We used a structure of the SARS-CoV S-protein:ACE2 complex (PDB ID 2AJF) to model structures for the other animal species. Changes in energy of the S-protein:ACE2 complex are highly correlated in SARS-CoV-2 and SARS-CoV (Figure 7 and Supplementary Figure S11), suggesting that the range of animals susceptible to the virus is likely to be similar for SARS-CoV-2 and SARS-CoV.



# <u>Figure 7:</u> Changes in the energy of the SARS-CoV:ACE2 and SARS-CoV-2:ACE2 complexes across selected animals. The change in energy of the complex ( $\Delta\Delta G$ ), as measured by protocol (2) - PPI2 over the DCEX residues, is shown on the y-axis.

It should be noted that the binding of the SARS-CoV-2 S-protein:ACE2 complex is 10-22-fold stronger than for SARS-CoV(29,30,34), however both energies are sufficient for an interaction and clearly did enable infection with human hosts.

#### Conservation of TMPRSS2 and its role in SARS-CoV-2 infection

ACE2 and TMPRSS2 are key factors in the SARS-CoV-2 infection process. Both are highly co-expressed in the susceptible cell types, such as type II pneumocytes in the lungs, ileal absorptive enterocytes in the gut, and nasal goblet secretory cells(25). Since both proteins are required for infection of host cells and since our analyses clearly support suggestions of conserved binding of S-protein:ACE2 across animal species, we decided to analyse whether the TMPRSS2 was similarly conserved. There is no known structure of TMPRSS2, but we built a high-quality model from a template structure (PDB ID 5125). Since TMPRSS2 is a serine protease, and the key catalytic residues are known, this allowed us to use FunFams to identify highly conserved residues in the active site and the cleavage site that are likely to be involved in substrate binding. We, therefore, used two sets of residues for our analysis: the active site and cleavage site residues (ASCS) and the active site and cleavage site residues plus residues within 8Å of catalytic residues that are highly conserved in the functional family (ASCSEX).

The sum of Grantham scores for mutations in the active site and cleavage site for TMPRSS2 is zero or consistently lower than ACE2 in all organisms under consideration, for both ASCS and ASCSEX residues (ASCSEX shown in Figure 8). This means that the mutations in TMPRSS2 involve more conservative changes.

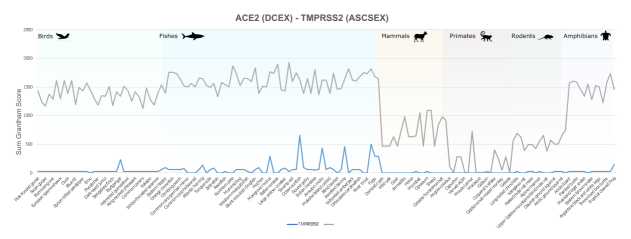
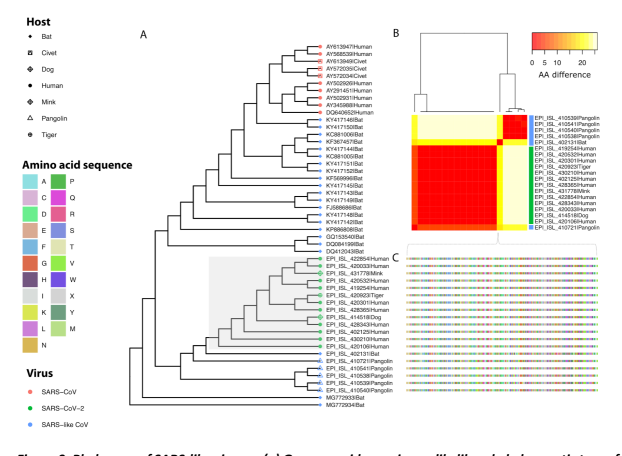


Figure 8: Comparison of Grantham Scores sums for ACE2 and TMPRSS2 for ASCSEX residues

Mutations in ACE2 DCEX residues seem to have a more disruptive effect. Whilst we expect orthologues from organisms that are close to humans to be conserved and have lower Grantham scores, we observed some residue substitutions that have high Grantham scores for primates, such as capuchin, marmoset and mouse lemur. In addition, primates, such as the coquerel sifaka, greater bamboo lemur and Bolivian squirrel monkey, have mutations in DCEX residues with high Grantham scores. These animals may therefore be less susceptible to infection because of these mutations in the protease.

#### Phylogenetic Analysis of SARS-like strains in different animal species

A small-scale phylogenetic analysis was performed on a subset of SARS-CoV-2 assemblies in conjunction with a broader range of SARS-like betacoronaviruses (Supplementary Table S3), including SARS-CoV isolated from humans and civets. The phylogeny is consistent with previous work(2) which identified the virus sampled from horseshoe bats (RaTG3, EPI\_ISL\_402131) as the closest genome sequence to SARS-CoV-2 strains currently available (Figure 9). Aided by a large community effort, thousands of human-associated SARS-CoV-2 genome assemblies are now accessible on GISAID(16,17). To date, these also include one assembly generated from a virus infecting a domestic dog (EPI ISL 414518), one obtained from a zoo tiger (EPI ISL 420923) and one obtained from a mink (EPI\_ISL\_431778). SARS-CoV-2 strains from animal infections all fall within the phylogenetic diversity observed in human lineages (Figure 9a). The receptor binding domain is completely conserved (Figure 9b-c) across both human and animal SARS-CoV-2, with replacements in the spike protein of dog (Sprotein V8L), tiger (S-protein D614G) and mink (S-protein D614G) strains relative to Wuhan-Hu-1 also observed in human-associated lineages(102), consistent with circulation in non-human hosts. Of note, while genome-wide data indicates a closer phylogenetic relationship between SARS-CoV-2 strains and RatG13, the receptor binding domain alignment instead supports a closer relationship with a virus isolated from pangolins (EPI\_ISL\_410721; Figure 9c), in line with previous reports(103). This highlights the importance of considering variation in putative structural determinants of host range, which may support a different topology to that observed based solely on genome-wide analyses.



<u>Figure 9</u>: Phylogeny of SARS-like viruses. (a) Genome-wide maximum likelihood phylogenetic tree of SARS-like betacoronavirus strains sampled from diverse hosts (coloured tip symbols; <u>Supplementary</u> <u>Table S3</u>). NB genome EPI\_ISL\_402131 is sample RaTG13 from a horseshoe bat. (b) Pairwise amino acid differences at the S-protein RBD between human and animal strains of SARS-CoV-2, relative to closely related-SARS-like viruses in animal hosts. (c) Sequence alignment for the spike protein receptor binding domain.

#### Discussion

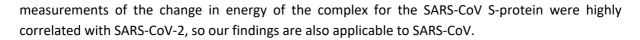
The ongoing COVID-19 global pandemic has a zoonotic origin, necessitating investigations into how SARS-CoV-2 infects animals, and how the virus can be transmitted across species. Given the role that the stability of the complex, formed between the S-protein and its receptors, could contribute to the viral host range, zoonosis and anthroponosis, there is a clear need to study these interactions. However, relative changes in the energies of the S-protein:ACE2 complex have not been explored experimentally or *in silico*. A number of recent studies(20,21,46) have suggested that, due to high conservation of ACE2, some animals are vulnerable to infection by SARS-CoV-2. Concerningly, these animals could, in theory, serve as reservoirs of the virus, increasing the risk of future transmission across species. However, transmission rates across species are not known. Therefore, it is important to try to predict which other animals could potentially be infected by SARS-CoV-2, so that the plausible extent of zoonotic transmission can be estimated, and surveillance efforts can be guided appropriately.

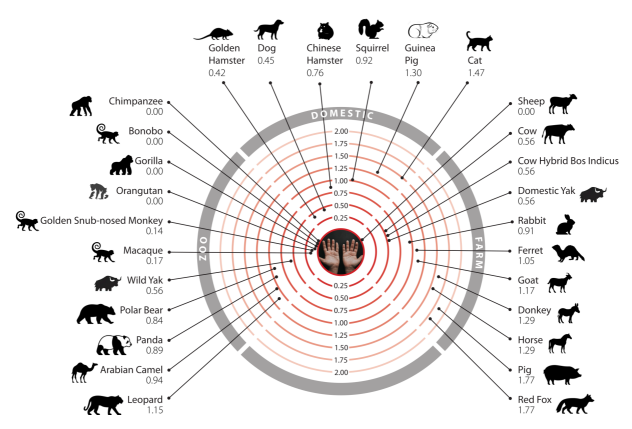
Animal susceptibility to infection by SARS-CoV-2 has been studied *in vivo*(18,19,95,97,98) and *in vitro*(20–22) during the course of the pandemic. Parallel *in silico* work has made use of the protein structure of the S-protein:ACE2 complex to computationally predict the breadth of possible viral hosts. Most studies simply considered the number of residues mutated relative to human ACE2(33,104,105), although some also analyse the effect that these mutations have on the interface stability(36,45,106). The most comprehensive of these studies analysed the number, and locations, of mutated residues in ACE2 orthologues from 410 species(46), but did not perform detailed energy calculations as we have done. Also, no assessment of TMPRSS2 was made. Furthermore, our work is the only study that has so far explored changes in the energy of the S-protein:ACE2 complex on a large scale.

In this study, we performed a comprehensive analysis of the major proteins that SARS-CoV-2 uses for cell entry. We predicted structures of ACE2 and TMPRSS2 orthologues from 215 vertebrate species and modelled S-protein:ACE2 complex and analysed changes in energy of the S-protein:ACE2 complex  $(\Delta\Delta G)$ , in silico, following mutations from animal residues to those in human. Our predictions suggest that, whilst many mammals are susceptible to infection by SARS-CoV-2, birds, fish and reptiles are not likely to be. We manually analysed residues in the S-protein:ACE2 interface, including DC residues that directly contacted the other protein, and DCEX residues that also included residues within 8Å of the binding residues, that may affect binding. We identified these important residues by detailed manual inspection of the complex, evolutionary conservation patterns, in silico alanine scanning, allosteric predictions and detection of sites under positive selection, some of which had been previously reported. We clearly showed the advantage of performing more sophisticated studies of the changes in energy of the complex, over more simple measures (e.g. number and nature of mutated residues) used in other studies. Furthermore, the wider set of DCEX residues that we identified, near the binding interface, had a higher correlation to the phenotype data than the DC residues. In addition to ACE2, we also analysed how mutations in TMPRSS2 impact binding to the S-protein. We found that mutations in TMPRSS2 are less disruptive than mutations in ACE2, indicating that binding interactions in the S-protein:TMPRSS2 complex in different species will not be affected.

To increase our confidence in assessing changes in the energy of the complex, we developed multiple protocols using different, established methods. We correlated these stability measures with experimental infection phenotypes in the literature, from *in vivo*(18,19,95,97,98) and *in vitro*(20–22) studies of animals. Protocol 2, mCSM-PPI2 correlates best with the number of mutations, chemical changes induced by mutations, and infection phenotypes, so we chose to focus our analysis employing this protocol.

Our measurement of changes in energy of the complex, together with high sequence and structural identity of ACE2 and TMPRSS2 orthologues, suggests that a broad range of mammals—but not fish, birds or reptiles—are susceptible to infection by SARS-CoV-2. Our predictions are supported by the findings of experimental studies assessing species that are at risk of infection by SARS-CoV-2(2,18–22,95,96). Humans are likely to come into contact with 26 of these species in domestic, agricultural or zoological settings (Figure 10). Of particular concern are sheep, that have no change in energy of the S-protein:ACE2 complex, as these animals are farmed and come into close contact with humans. We also provide phylogenetic evidence that human SARS-CoV-2 is also present in tigers(15), dogs(16,17) and minks(16,17) (Figure 9), consistent with reports of human-to-animal transmission. Our





<u>Figure 10</u> Mammals that humans come into contact with that are at risk of infection by SARS-CoV-2. Twenty-six mammals are categorised into domestic, agricultural or zoological settings. Numbers represent the change in binding energy ( $\Delta\Delta G$ ) of the S-protein:ACE2. Animals close to the centre are more at risk of infection.

For a subset of species, we performed more detailed structural analyses to gain a better understanding of the nature of the S-protein: ACE2 interface. In some cases, we had found discrepancies between our energy calculations and experimental phenotypes. To test our predictions, we manually analysed how the shape or chemistry of residues may impact complex stability for all DC residues and a selection of DCEX residues. In agreement with other studies(30), we identified a number of locations in ACE2 that are important for binding the S-protein. These locations, namely the hydrophobic cluster near the Nterminus and two hotspot locations near residues 31 and 353, stabilise the binding interface. Five DC residues have species-specific variants and influence how well S-protein can bind utilising these key interface regions. In agreement with our calculations in changes in energy of the S-protein:ACE2 complex, our structural studies do not support (in vivo) infection of horseshoe bat, which has variants at three out of five variant DC residues, one of which (D38N) causes the loss of a salt bridge and Hbonding interactions between ACE2 and S-protein at hotspot 353. These detailed structural analyses are supported by the high Grantham score and calculated total  $\Delta\Delta G$  for the change in energy of the complex. Both dog and cat have a physico-chemically similar variant at this hotspot (D38E), which although disrupting the salt bridge still permits alternative H-bonding interactions between the spike RBD and ACE2.

SARS-CoV-2 is better able to exploit the hydrophobic pocket than SARS-CoV by increased flexibility of its RBD loop and by mutation of L486F(30). Our structures show how SARS-CoV-2 can utilise this pocket for binding at the interface in a wide range of species. Of species with DCEX variants at this pocket, only guinea pig maintains an entirely hydrophobic environment (M82A), whilst also conserving three out of the five variant DC residues. This helps explain why we predict moderate risk of infection, in contrast to the *in vitro* experimental data that reports no infection. For the marmoset, there is contradictory *in vivo* and *in vitro* experimental data. Our energy calculations suggest no risk and the structural analyses support this by identifying a large structural difference caused by a 39 residue insert. This alters the overall structural superposition of the marmoset and human structures, which could affect the energy of the complex. Finally, some DCEX residues were predicted to be allosteric sites, which may be promising drug targets(94).

We applied protocols that enabled a comprehensive study of host range, within a reasonable time, for identifying species at risk from COVID-19 or of becoming reservoirs of the virus. Whilst we felt that these faster methods were justified by the need for timely answers to these questions, there are clearly caveats to our work that should be taken into account. Whilst we use a state of the art modelling tool(74) and an endorsed method for calculating changes in energy of the complex(78), molecular dynamics would possibly give a more accurate picture of energy changes by sampling rotamer space more comprehensively. However, such an approach would be prohibitively expensive at a time when it is clearly important to identify animals at risk as quickly as possible. Furthermore, although the animals we highlight at risk from our changes in binding energy calculations correlate well with the experimental data, there is only a small amount of such data currently available, and many of the experimental papers reporting these data are yet to be peer reviewed. Finally, we restricted our analyses to one strain of SARS-CoV-2. However, other strains may have evolved with mutations that give more complementary interfaces. Recent work reporting a new SARS-CoV-2 strain that can infect mice(107) suggests that this could be the case. Our model of the mouse complex supports a likely risk to mouse with this alternative SARS-CoV-2 strain.

The ability of SARS-CoV-2 to infect host cells to cause COVID-19, which sometimes results in severe disease, ultimately depends on a multitude of other host-virus protein interactions(44). While we do not investigate them all in this study, our results suggest that SARS-CoV-2 can indeed infect a broad range of mammals. As there is a possibility of creating new reservoirs of the virus, we should now consider how to identify such transmission early and to mitigate against such risks. Animals living in close contact with humans should be monitored and farm animals should be protected where possible and managed accordingly(108).

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# Author contributions

SL conceived the idea of analysing structures and effects of mutations in the S-protein:human ACE2 complex. JL conceived the idea of extending the analyses to animal complexes, for animals reported to be infected. JS conceived the idea of extending to a larger set of animals to explore host range. CO conceived the idea of contrasting multiple protocols to validate predictions. SL, CO, JL, JS, LvD designed the experiments. SL, NB, VW, PA, LvD performed the experiments. SL, NB, VW, PA, NS, JS, LvD, CR, IS, JL, CO analysed data. SL, NB, VW, HS, PA, NS, JS, LvD, CR, ND, CSMP, MA, IS, JL, CO contributed to the manuscript and figures. HS, VW, PA, SL, CO wrote the manuscript. JL, SE, FF, JS, CO revised the manuscript.

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