1 2 3	Title: Evolutionary differences in the ACE2 reveals the molecular origins of COVID-19 susceptibility
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14	
15	Abstract:
16	We explore the energetic frustration patterns associated with the binding between the SARS-CoV-
17	2 spike protein and the ACE2 receptor protein in a broad selection of animals. Using energy
18	landscape theory and the concept of energy frustration-theoretical tools originally developed to
19	study protein folding—we are able to identify interactions among residues of the spike protein and
20	ACE2 that result in COVID-19 resistance. This allows us to identify whether or not a particular
21	animal is susceptible to COVID-19 from the protein sequence of ACE2 alone. Our analysis
22	predicts a number of experimental observations regarding COVID-19 susceptibility,
23	demonstrating that this feature can be explained, at least partially, on the basis of theoretical means.
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#### 26 Introduction

27

28 The coronavirus disease 2019 (COVID-19) caused by the severe acute respiratory syndrome

- 29 coronavirus 2 (SARS-CoV-2) has affected the lives of millions of people in a worldwide
- pandemic. The hallmark of COVID-19 is its high degree of contagiousness between individuals.
- 31

32 SARS-CoV-2 is believed to gain entry in to the host cell through its interaction with the

33 Angiotensin-converting enzyme 2 (ACE2) receptor on the host cell surface(Zhou et al., 2020),

similar to SARS-CoV-1(Li et al., 2003). Recently, the structure of the SARS-CoV-2 viral spike

35 glycoprotein bound to the human ACE2 receptor was determined using X-ray crystallography

36 (Wang et al., 2020), providing a crucial starting point for any molecular modeling of the viral

37 interaction with ACE2. The structure of this crucial complex has also been independently

determined using cryo-EM(Yan et al., 2020) and X-ray crystallography(Shang et al., 2020).

39

40 What are the molecular interactions that give rise to interaction specificity between the viral

41 spike and ACE2 receptor? Hints at the molecular origins of COVID-19 susceptibility can be

42 found analyzing the susceptibility of different organisms to the coronavirus. The ACE2 receptors

43 are found in a diverse span of the animal kingdom, including mammals, birds, and aquatic life,

44 which have varying degrees of COVID-19 susceptibility(BS et al., 2021; Gautam, Kaphle,

45 Shrestha, & Phuyal, 2020; Goldstein, 2020; Kumakamba et al., 2020; Muñoz-Fontela et al.,

46 2020; Mykytyn et al., 2020; Oude Munnink et al., 2021; Palmer et al., 2021; Shi et al., 2020; Sia

47 et al., 2020; Sit et al., 2020).

48

For example, it is known that mice are immune to COVID-19 while on the contrary the Bronx
Zoo tiger Nadia(Goldstein, 2020) had tested positive for COVID-19 and also exhibited many of
the symptoms observed in infected humans. To date, a number of animals have been classified as
either being susceptible or immune to COVID-19(BS et al., 2021; Gautam et al., 2020;
Goldstein, 2020; Kumakamba et al., 2020; Muñoz-Fontela et al., 2020; Mykytyn et al., 2020;
Oude Munnink et al., 2021; Palmer et al., 2021; Shi et al., 2020; Sia et al., 2020; Sit et al., 2020).

56 The evolutionary divergence in the sequences of the ACE2 receptor found in different organisms 57 can be related to such susceptibility of infection(Becker et al., 2020; Damas et al., 2020; Frank, Enard, & Boyd, 2020; Lam et al., 2020; Luan, Lu, Jin, & Zhang, 2020; Martínez-Hernández et 58 al., 2020; Melin, Janiak, Marrone, Arora, & Higham, 2020). Here, we explore the molecular 59 mechanisms by which some sequence variants of the ACE2 receptor appear to confer resistance 60 to infection by virtue of their reduced binding affinity to the viral spike protein. We examine the 61 62 sequences for ACE2 receptor across a selection of 63 representative animals and identify the 63 residue interactions that are responsible for the reduced binding affinity, and thus COVID-19 resistance, using the concept of energetic frustration from the theory of protein folding (Onuchic, 64 Luthey-Schulten, & Wolynes, 1997; Onuchic & Wolynes, 2004). Here, energetic frustration 65 66 refers to unfavorable interactions between residues in a given protein structure that cannot be 67 mitigated without structural rearrangement or residue level mutations. In the context of the 68 ACE2/spike complex, frustrated interactions between residues of the ACE2 and the spike 69 glycoprotein can also exist for a given structure of the protein complex.

70

71 The rarity of kinetics traps observed in the folding of proteins indicates that, in general, proteins 72 do not exhibit a high amount of energetic frustration, which would instead create those kinetic 73 traps(Onuchic et al., 1997; Onuchic & Wolynes, 2004). While folding kinetics suggest proteins 74 to be "minimally frustrated", some local frustration may be present; for example, local 75 frustration could be functionally useful for tuning conformational dynamics. In protein 76 complexes, a site frustrated in the monomeric protein may become less frustrated when the 77 protein is bound to its counterparts, thus guiding specific association(Ferreiro, Hegler, Komives, & Wolynes, 2007; Parra et al., 2016). 78 79 In the case of the complex formed between the SARS-CoV-2 viral spike glycoprotein and the

80 ACE2 receptor, we use changes in energy frustration as a proxy for changes in binding affinity.

81 We use the crystal structure of the viral spike bound to the human ACE2 as a template to

82 construct molecular models of the interaction between the viral spike and the ACE2 receptors of

83 these different animals. We then calculate the changes in frustration with respect to the reference

84 point constituted by the human ACE2 sequence (Ferreiro et al., 2007; Parra et al., 2016). This

85 allows us to identify key residues of the ACE2 protein that appear to inhibit the binding of the

spike glycoprotein and to predict whether or not a particular animal will be susceptible to

87	COVID-19.	The novelty	of our a	approach	and the k	ev to o	ur results	resides i	n the fa	ct that	while
07	COVID-17.		y or our o	approach	and the K	Cy 10 0	ui resuits	icsides i	n une na	ci mai,	, winne

- 88 our procedure is based on the only input of the protein sequences of ACE2 receptor, our
- 89 approach does incorporate a great deal of structural information about the protein complex,
- 90 which is extracted from the crystal structure(Wang et al., 2020), and physico-chemical details
- about the energetics of protein folding and docking, which is synthetized in the energy function
- 92 and results from decades of developments(Onuchic & Wolynes, 2004).
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- 95
- 96

# 97 Materials & Methods

### 98 ACE2 protein sequences

99 The majority of ACE2 protein sequences were previously annotated from the genome assembles

and sequencing data from the DNA Zoo Consortium(Dudchenko et al., 2017) (Available for

101 download: <u>https://www.dnazoo.org/post/the-first-million-genes-are-the-hardest-to-make-r</u>). The

102 full length protein sequences of the ACE2 proteins for mouse (*Mus musculus*), ferret (*Mustela* 

103 putorius furo), chicken (Gallus gallus domesticus), pig (Sus), duck (Anas platyrhynchos), Syrian

- 104 golden hamster (*Mesocricetus auratus*), and mink (*Neovison vison*) were obtained from the
- 105 Uniprot database(The UniProt, 2021) to supplement the sequences derived from the DNA Zoo.

106 In total, 63 representative ACE2 sequences were used in our study (Table S1). For comparative

analysis, a multiple sequence alignment was generated for the ACE2 sequences using Clustal

108 Omega(Madeira et al., 2019).

109

110

### 111 Homology Modeling

112 The crystal structure of the SARS-Cov-2 glycoprotein spike bound to the human ACE2 protein

served as our starting template for constructing models of the glycoprotein spike bound to the

- ACE2 protein of other animals. We used the SWISS-MODEL (Waterhouse et al., 2018) to create
- homology models of 63 representative animals (Table S1) for a full list.

116

#### 117

### **118 Frustration Analysis**

- 119 We performed an energy landscape analysis on the predicted ACE2-spike complex for different
- animals using the configurational frustration index(Ferreiro et al., 2007; Parra et al., 2016):

121 
$$F_{ij} = \frac{\left(H_{ij} - \left\langle H_{i'j'}^{decoy} \right\rangle\right)}{\sqrt{\frac{1}{N} \sum_{k=1}^{N} \left(H_{i'j'} - \left\langle H_{i'j'}^{decoy} \right\rangle\right)}}$$
(1)

Here,  $H_{ij}$  represents the pairwise interaction energy between residues *i* and *j* in a given structure using the Associative Memory, Water Mediated, Structure and Energy Model

124 (AWSEM)(Davtyan et al., 2012), a coarse-grained model widely used to study problems of

protein folding and protein-protein association and assembly. The native energies  $H_{ij}$  are

- 126 compared directly to *N* number of different configurational realizations between residues *i* and *j*,
- 127 thereby generating a distribution of decoy energies with a mean of  $\langle H_{i'j'}^{decoy} \rangle$  and a standard

128 deviation of 
$$\sqrt{N^{-1}\sum_{k=1}^{N} (H_{i'j'} - \langle H_{i'j'}^{decoy}\rangle)}$$
. Hence,  $F_{ij}$  is a type of Z-score that measures how  
129 favorable a particular pair of interactions are within a protein or protein complex with respect to  
130 a distribution of decoys. Frustrated (unfavorable interactions) are denoted by  $F_{ij} < 0$  while  $F_{ij} > 0$   
131 are considered favorable; in particular,  $F_{ij} < -1$  is considered highly frustrated while  $F_{ij} > 1$  is  
132 considered minimally frustrated.

133

In our analysis, we found that it was useful to compare the configurational frustration between aninterprotein residue pair with the same pair from the human ACE2-spike complex:

136 
$$\Delta F_{ij}^{(Species)} = F_{ij}^{(Species)} - F_{ij}^{(Human)}$$
(2)

137 We find that  $\Delta F_{ij}^{(Species)} < -1.5$  robustly identifies highly frustrated interactions that result in 138 COVID-19 resistence. On the other hand, if all of the inter-protein residue interactions between 139 the ACE2 receptor and the spike do not exhibit high levels of frustration (i.e.,  $\Delta F_{ij}^{(Species)} > -1$ ) 140 we identify that species as being highly susceptible to COVID-19. For completeness, if the most

- 141 frustrated interprotein interactions fall between  $-1.5 < \Delta F_{ij}^{(Species)} \le -1$  that species is predicted to
- 142 be moderately susceptible.
- 143

### 144 Evolutionary distance between ACE2 proteins

145 The Jukes-Cantor distance is used to quantify the evolutionary distance between aligned ACE2

146 proteins in our study:  $d = -\frac{19}{20} \log \left( 1 - \frac{20}{19} p \right)$ , where p is the p-distance—i.e., the number of

residue sites between two compared sequences that are different divided by the sequence lengthof the multiple sequence alignment.

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### 151 **Results & Discussion**

### 152 Comparative frustration analysis of ACE2-spike complex for different species

153 By examining the comparative differences between the inter-protein interactions with respect to

the human ACE2-spike complex, we are able to identify residue-interactions outliers that

155 represent a significant disruption to the ACE2/spike interaction relative to the human ACE2-

spike interaction.

157

158 Shown in Figure 1 are plots of our frustration analysis for mouse (*Mus musculus*) and tiger

159 (*Panthera tigris*), which are used as representative examples of animals have been

160 experimentally observed to be resistant(Muñoz-Fontela et al., 2020) and susceptible(Goldstein,

161 2020) to COVID-19, respectively.

162 We observe a single frustrated residue pair between 31N of the mouse ACE2 and 484E of the

spike protein on the map of configurational frustration (Figure 2A/2C), which appears as an

outlier in the histogram of  $\Delta F_{ii}^{(Mouse)} \sim -2$ . Similar highly frustrated outliers can be found for the

- 165 other animals that are known to resist COVID-19 (Figure S1), such as chicken (*Gallus gallus*
- 166 *domesticus*)(Shi et al., 2020) and duck (*Anas*)(Shi et al., 2020). We find that a threshold value of
- 167  $\Delta F_{ii}^{(Species)} < -1.5$  robustly identifies residue pairs that appear to confer COVID-19 resistence.
- 168 Likewise, a histogram of  $\Delta F_{ij}^{(Tiger)}$  exhibits comparable levels of frustration to that of the human

169 ACE2 and spike (Figure 1B)—similar findings are obtained for other animals with known

susceptibilities to COVID-19 (Figure S2), such as white-tailed deer (*Odocoileus* 

171 *virginianus*)(Palmer et al., 2021), European rabbit (*Oryctolagus cuniculus*)(Mykytyn et al.,

172 2020), and pig (*Sus scrofa*)(BS et al., 2021).

173

We further apply this analysis for identifying frustrated outliers in our other modeled complexes, 174 175 thereby predicting whether a particular animal is susceptible to COVID-19. A detailed summary 176 of our results is shown in Figure 2, which includes experimental observations that corroborate or 177 are inconsistent with our predictions. Other animals that have been experimentally observed to 178 be susceptible to COVID-19, such as mink (*Neovison vison*) (Oude Munnink et al., 2021), and 179 Syrian golden hamster (*Mesocricetus auratus*)(Sia et al., 2020), are identified as being 180 moderately susceptible by our computational approach. Coronavirus consensus PCR-primer 181 sequences have been detected with high frequency in populations of straw-colored fruit bats 182 (Eidolon helvum)(Kumakamba et al., 2020), which have been predicted to exhibit moderate 183 susceptibility by the computational approach.

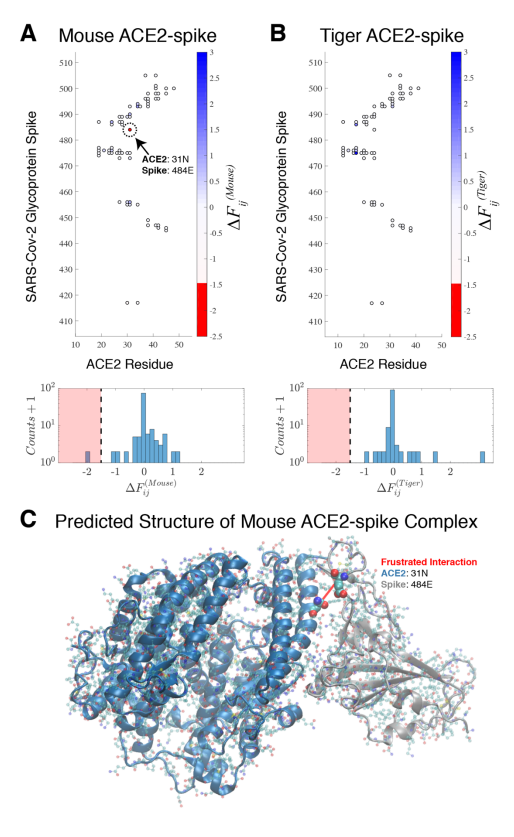
184

Taken together, our findings summarized in Figure 2 show that an energy landscape-based 185 186 approach can identify the molecular origins of COVID-19 susceptibility and resistance. 187 Experimental observations corroborate 10 out of 12 of the computational predictions. One 188 apparent inconsistency between the predictions and experimental observation regards the 189 susceptibility of ferrets (Mustela putorius furo), which have been observed to replicate SARS-190 Cov-2 specifically in their upper respiratory tract(Shi et al., 2020). Our analysis of the ferret 191 ACE2-spike complex reveals a single highly frustrated inter-protein interaction between 34Y of 192 ACE2 and 403R of the spike protein. However, it has been noted(Damas et al., 2020; Shi et al., 193 2020) that ferrets have a unique respiratory biology, which may offer an explanation for this 194 apparent discrepancy. Another apparent inconsistency is observed with our predictions for dogs 195 (*Canis lupus familiaris*). Our frustration analysis predicts two highly frustrated inter-protein contacts within the ACE2/spike complex: 33Y of ACE2 with 417K of the spike and 325E of the 196 197 ACE2 with 502G of the spike. Yet the susceptibility of dogs still remains somewhat 198 controversial—while viral susceptibility and the production of antibody responses have been 199 detected in dogs(Sit et al., 2020), viral replication has been reported to be poor(Shi et al., 2020).

<sup>200</sup> 

201	Our energy landscape-based predictions for COVID-19 susceptibility are closely related to
202	similar approaches that examined sequence differences in ACE2 sequences of different animals
203	in the context of a structural model of the ACE2/spike complex(Damas et al., 2020; Lam et al.,
204	2020; Luan et al., 2020). Frustration analysis yields a benefit to computational estimates of
205	binding affinity because it compares the interaction energies between ACE2 and the spike
206	glycoprotein with respect to alternative configurations (i.e., decoys) to assess how favorable a
207	particular interaction is in the binding interface. In particular, the majority of our predictions are
208	consistent with those of Damas et al(Damas et al., 2020), which makes predictions that are
209	consistent with the same 10 out of 12 experiment observations that are highlighted in Figure 2.
210	However, validation of the different models that exists is limited by the relatively small number
211	of confirmed cases of COVID-19 in animals.
212	
213	By in large, we find that our simple model appears to be consistent with many experimental
214	observations of COVID-19 infections across different animals despite only considering the

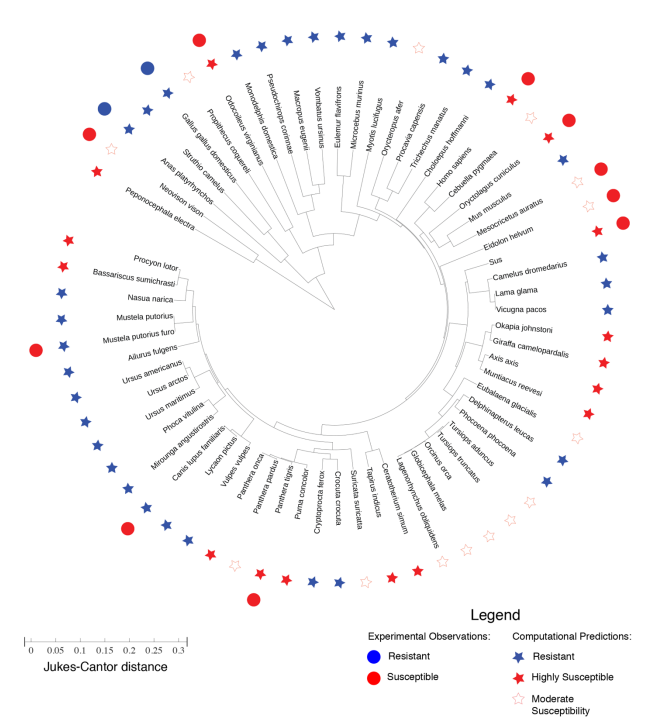
215 ACE2-spike protein interaction.



#### Figure 1. Comparative analysis of the frustration indices with respect to those observed in

### 219 human ACE2-spike complex reveals outliers that indicate COVID-19 resistance. The

- 220 configurational frustration index relative to the frustration in the human ACE2-spike complex is
- shown for (A) mouse (*Mus musculus*) and (B) tiger (*Panthera tigris*) on a contact map
- 222 illustrating select contacts between the SARS-Cov-2 spike and the ACE2 protein. Corresponding
- histograms of the frustration index between all contacts between the ACE2 and spike protein are
- also shown. (A) and (B) are representative examples of animals that are resistant to COVID-19
- and susceptible to COVID-19, respectively. Animals that resist COVID-19 appear to have
- 226 frustrated outliers that represent highly unfavorable residue interactions compared to the human
- ACE2-spike complex (i.e.,  $\Delta F_{ii}^{(Species)} < -1.5$ ). For the mouse, a single frustrated interaction
- between residue 31N of the ACE2 protein and 484E of the spike glycoprotein appears to confer
- 229 COVID-19 resistance. (C) The frustrated interaction is plotted on the modeled 3D structure of
- the spike glycoprotein bound to the mouse ACE2 receptor.



232

233 Figure 2.

### 234 Phylogenic tree representing the evolutionary distance between ACE2 proteins of different

- species. The lengths in the radial direction denote the Jukes-Cantor distance (See Materials &
- 236 Methods) as a measure of evolutionary distance between any two ACE2 proteins. The
- 237 experimental observation of SARS-CoV-2 resistance/susceptibility are plotted alongside the
- 238 computational predictions for resistance/susceptibility based on our frustration analysis of the

ACE2-spike complex. See the Legend for more details. There is a consistency between the

- 240 computational predictions and the experimental observations for mouse (Mus musculus), chicken
- 241 (Gallus gallus domesticus)(Shi et al., 2020), duck (Anas platyrhynchos)(Shi et al., 2020), mink
- 242 (Neovison vison)(Oude Munnink et al., 2021), bat (Eidolon helvum)(Kumakamba et al., 2020),
- 243 Syrian golden hamster (Mesocricetus auratus)(Sia et al., 2020), tiger (Panthera tigris), white-
- tailed deer (*Odocoileus virginianus*)(Palmer et al., 2021), European rabbit (*Oryctolagus*
- 245 *cuniculus*)(Mykytyn et al., 2020), and pig (Sus scrofa)(BS et al., 2021). However, apparent
- inconsistencies are found for ferret (*Mustela putorius furo*)(Shi et al., 2020) and dog (*Canis lupis*
- *familiaris*) (Shi et al., 2020; Sit et al., 2020)—however, SARS-Cov-2 has only been observed to
- replicate in the upper respiratory tract of ferrets(Shi et al., 2020), and viral replication has been
- observed to be low in dogs(Shi et al., 2020).
- 250
- 251

# 252 **Conclusion**

253 The COVID-19 pandemic and the spread of other coronaviruses in recent years requires an

- 254 indirect approach to understanding the molecular determinants behind susceptibility and
- resistance. Here, we constructed structural models of the ACE2-spike glycoprotein complex for a
- wide range of animals with ACE2 receptors. Using an energy landscape theory-based analysis
- 257 we are able to uncover specific inter-protein interactions between the ACE2 and spike that
- appear to confer COVID-19 resistance. Our predictions appear to be consistent with many of the experimental observations regarding animal susceptibility, providing a structural explanation to
- those observations.
- 261

Our analysis reveals that the evolutionary distance between ACE2 proteins is not sufficient to predict COVID-19 susceptibility (Figure 2). Rather, an energy landscape-based analysis appears necessary to assess the interactions between the ACE2 protein and the SARS-Cov-2 spike glycoprotein.

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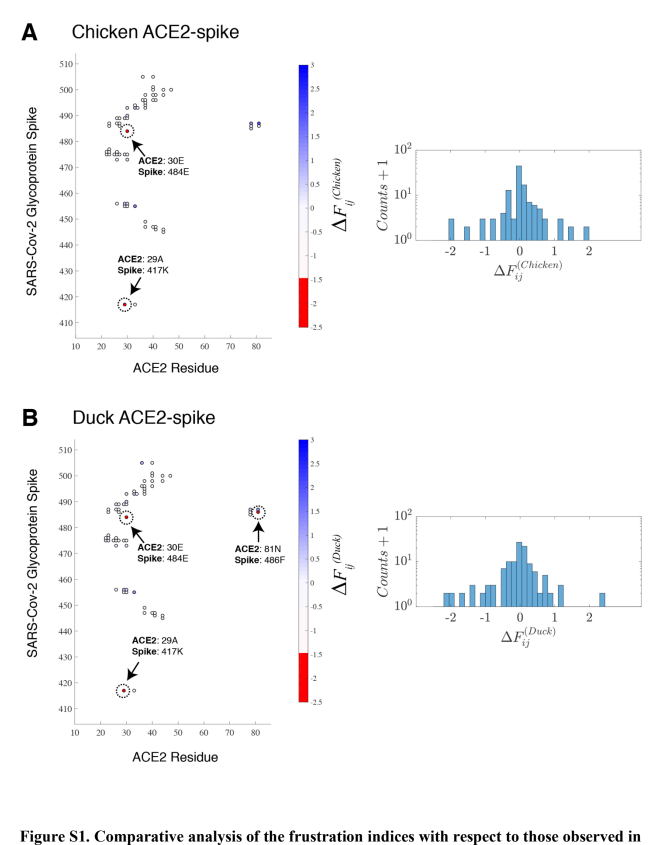
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- 273 Institute of Texas. Scholar in Cancer Research.
- 274



human ACE2-spike complex for additional examples of animals with known COVID-19

resistance. The configurational frustration index relative to the frustration in the human ACE2-

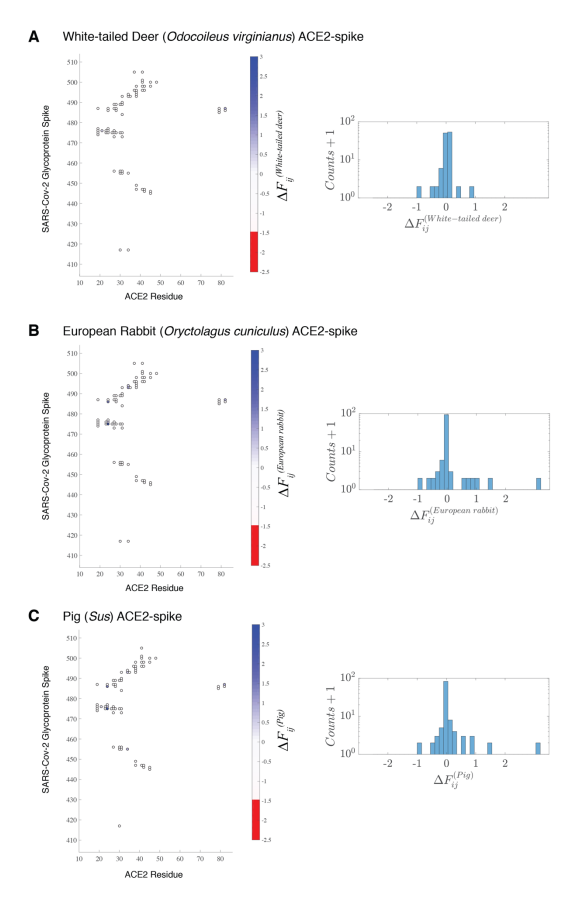
- spike complex is shown for (A) chicken (Gallus gallus domesticus) and (B) duck (Anas
- 284 *platyrhynchos*) on a contact map illustrating select contacts between the SARS-Cov-2 spike and
- the ACE2 protein. Corresponding histograms of the frustration index between all contacts
- between the ACE2 and spike protein are also shown. Animals that resist COVID-19 appear to
- 287 have frustrated outliers that represent highly unfavorable residue interactions compared to the
- human ACE2-spike complex (i.e.,  $\Delta F_{ij}^{(Species)} < -1.5$ ). For the chicken, two highly frustrated
- interactions are identified: between (1) 30E of the ACE2 and 484E of the spike and (2) 29A of
- the ACE2 and 417K of the spike. For the duck, three highly frustrated interactions are identified:
- between (1) 30E of the ACE2 and 484E of the spike, (2) 29A of the ACE2 and 417K of the
- spike, and (3) 81N of the ACE2 and 486F of the spike. These frustrated interactions appear to
- confer COVID-19 resistance.

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295

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297



300	Figure S2. Comparative analysis of the frustration indices with respect to those observed in
301	human ACE2-spike complex for additional examples of animals with known COVID-19
302	susceptibility. The configurational frustration index relative to the frustration in the human
303	ACE2-spike complex is shown for (A) white-tailed deer (Odocoileus virginianus), (B) European
304	rabbit (Oryctolagus cuniculus), and (C) pig (Sus) on a contact map illustrating select contacts
305	between the SARS-Cov-2 spike and the ACE2 protein. Corresponding histograms of the
306	frustration index between all contacts between the ACE2 and spike protein are also shown.
307	Animals that are susceptible to COVID-19 appear to have comparable levels of frustration
308	compared to human ACE2-spike complex.
<ul> <li>309</li> <li>310</li> <li>311</li> <li>312</li> <li>313</li> <li>314</li> <li>315</li> <li>316</li> <li>317</li> <li>318</li> </ul>	Becker, D. J., Albery, G. F., Sjodin, A. R., Poisot, T., Dallas, T. A., Eskew, E. A., Carlson, C. J. (2020). Predicting wildlife hosts of betacoronaviruses for SARS-CoV-2 sampling prioritization: a modeling study. <i>bioRxiv</i> , 2020.2005.2022.111344. doi:10.1101/2020.05.22.111344
319 320 321 322 323 324 325	<ul> <li>BS, P., G, S., MM, P., C, EH., E, M., P, M., &amp; CE, L. (2021). Susceptibility of Domestic Swine to Experimental Infection with Severe Acute Respiratory Syndrome Coronavirus 2. <i>Emerg Infect Dis.</i>, 27(1), 104-112. doi:<u>https://dx.doi.org/10.3201/eid2701.203399</u></li> <li>Damas, J., Hughes, G. M., Keough, K. C., Painter, C. A., Persky, N. S., Corbo, M., Lewin, H. A. (2020). Broad host range of SARS-CoV-2 predicted by comparative and structural analysis of ACE2 in vertebrates. <i>Proceedings of the National Academy of Sciences</i>, 117(36), 22311. doi:10.1073/pnas.2010146117</li> </ul>
326 327 328 329 330 331 332 333 334 335 336 337 338 339	<ul> <li>Davtyan, A., Schafer, N. P., Zheng, W., Clementi, C., Wolynes, P. G., &amp; Papoian, G. A. (2012). AWSEM-MD: Protein Structure Prediction Using Coarse-Grained Physical Potentials and Bioinformatically Based Local Structure Biasing. <i>The Journal of Physical Chemistry</i> <i>B</i>, <i>116</i>(29), 8494-8503. doi:10.1021/jp212541y</li> <li>Dudchenko, O., Batra, S. S., Omer, A. D., Nyquist, S. K., Hoeger, M., Durand, N. C., Aiden, E. L. (2017). De novo assembly of the <em>Aedes aegypti</em> genome using Hi-C yields chromosome-length scaffolds. <i>Science</i>, <i>356</i>(6333), 92. doi:10.1126/science.aal3327</li> <li>Ferreiro, D. U., Hegler, J. A., Komives, E. A., &amp; Wolynes, P. G. (2007). Localizing frustration in native proteins and protein assemblies. <i>Proceedings of the National Academy of Sciences</i>, <i>104</i>(50), 19819. doi:10.1073/pnas.0709915104</li> <li>Frank, H. K., Enard, D., &amp; Boyd, S. D. (2020). Exceptional diversity and selection pressure on SARS-CoV and SARS-CoV-2 host receptor in bats compared to other mammals. <i>bioRxiv</i>, 2020.2004.2020.051656. doi:10.1101/2020.04.20.051656</li> </ul>

- Gautam, A., Kaphle, K., Shrestha, B., & Phuyal, S. (2020). Susceptibility to SARS, MERS, and
  COVID-19 from animal health perspective. *Open Vet J*, 10(2), 164-177.
  doi:10.4314/ovj.v10i2.6
- Goldstein, J. (2020). Bronx Zoo Tiger Is Sick With the Coronavirus. *New York Times*. Retrieved
   from <u>https://www.nytimes.com/2020/04/06/nyregion/bronx-zoo-tiger-coronavirus.html</u>
- Kumakamba, C., Niama, F. R., Muyembe, F., Mombouli, J.-V., Kingebeni, P. M., Nina, R. A., . .
  Lange, C. E. (2020). Coronavirus surveillance in Congo basin wildlife detects RNA of multiple species circulating in bats and rodents. *bioRxiv*, 2020.2007.2020.211664.
  doi:10.1101/2020.07.20.211664
- Lam, S. D., Bordin, N., Waman, V. P., Scholes, H. M., Ashford, P., Sen, N., . . . Orengo, C. A.
  (2020). SARS-CoV-2 spike protein predicted to form complexes with host receptor
  protein orthologues from a broad range of mammals. *Scientific Reports, 10*(1), 16471.
  doi:10.1038/s41598-020-71936-5
- Li, W., Moore, M. J., Vasilieva, N., Sui, J., Wong, S. K., Berne, M. A., . . . Farzan, M. (2003).
   Angiotensin-converting enzyme 2 is a functional receptor for the SARS coronavirus.
   *Nature*, 426(6965), 450-454. doi:10.1038/nature02145
- Luan, J., Lu, Y., Jin, X., & Zhang, L. (2020). Spike protein recognition of mammalian ACE2
   predicts the host range and an optimized ACE2 for SARS-CoV-2 infection. *Biochemical and Biophysical Research Communications*, *526*(1), 165-169.
   doi:https://doi.org/10.1016/j.bbrc.2020.03.047
- Madeira, F., Park, Y. M., Lee, J., Buso, N., Gur, T., Madhusoodanan, N., . . . Lopez, R. (2019).
   The EMBL-EBI search and sequence analysis tools APIs in 2019. *Nucleic Acids Research*, 47(W1), W636-W641. doi:10.1093/nar/gkz268
- Martínez-Hernández, F., Isaak-Delgado, A. B., Alfonso-Toledo, J. A., Muñoz-García, C. I.,
   Villalobos, G., Aréchiga-Ceballos, N., & Rendón-Franco, E. (2020). Assessing the
   SARS-CoV-2 threat to wildlife: Potential risk to a broad range of mammals. *Perspectives in Ecology and Conservation*. doi:https://doi.org/10.1016/j.pecon.2020.09.008
- Melin, A. D., Janiak, M. C., Marrone, F., Arora, P. S., & Higham, J. P. (2020). Comparative
   ACE2 variation and primate COVID-19 risk. *Communications Biology*, 3(1), 641.
   doi:10.1038/s42003-020-01370-w
- Muñoz-Fontela, C., Dowling, W. E., Funnell, S. G. P., Gsell, P.-S., Riveros-Balta, A. X.,
   Albrecht, R. A., . . Barouch, D. H. (2020). Animal models for COVID-19. *Nature*,
   *586*(7830), 509-515. doi:10.1038/s41586-020-2787-6
- 373 Mykytyn, A. Z., Lamers, M. M., Okba, N. M. A., Breugem, T. I., Schipper, D., van den Doel, P.
   374 B., . . . Haagmans, B. L. (2020). Susceptibility of rabbits to SARS-CoV-2. *bioRxiv*,
   375 2020.2008.2027.263988. doi:10.1101/2020.08.27.263988
- Onuchic, J. N., Luthey-Schulten, Z., & Wolynes, P. G. (1997). THEORY OF PROTEIN
   FOLDING: The Energy Landscape Perspective. *Annual Review of Physical Chemistry*,
   48(1), 545-600. doi:10.1146/annurev.physchem.48.1.545
- Onuchic, J. N., & Wolynes, P. G. (2004). Theory of protein folding. *Current Opinion in Structural Biology*, 14(1), 70-75. doi:<u>https://doi.org/10.1016/j.sbi.2004.01.009</u>
- Oude Munnink, B. B., Sikkema, R. S., Nieuwenhuijse, D. F., Molenaar, R. J., Munger, E.,
  Molenkamp, R., . . . Koopmans, M. P. G. (2021). Transmission of SARS-CoV-2 on mink
  farms between humans and mink and back to humans. *Science*, *371*(6525), 172.
- doi:10.1126/science.abe5901

- Palmer, M. V., Martins, M., Falkenberg, S., Buckley, A., Caserta, L. C., Mitchell, P. K., . . . Diel,
  D. G. (2021). Susceptibility of white-tailed deer (<em&gt;Odocoileus
  virginianus&lt;/em&gt;) to SARS-CoV-2. *bioRxiv*, 2021.2001.2013.426628.
  doi:10.1101/2021.01.13.426628
- Parra, R. G., Schafer, N. P., Radusky, L. G., Tsai, M.-Y., Guzovsky, A. B., Wolynes, P. G., &
  Ferreiro, D. U. (2016). Protein Frustratometer 2: a tool to localize energetic frustration in
  protein molecules, now with electrostatics. *Nucleic Acids Research*, 44(W1), W356W360. doi:10.1093/nar/gkw304
- Shang, J., Ye, G., Shi, K., Wan, Y., Luo, C., Aihara, H., . . . Li, F. (2020). Structural basis of
   receptor recognition by SARS-CoV-2. *Nature*, *581*(7807), 221-224. doi:10.1038/s41586 020-2179-y
- Shi, J., Wen, Z., Zhong, G., Yang, H., Wang, C., Huang, B., ... Bu, Z. (2020). Susceptibility of
   ferrets, cats, dogs, and other domesticated animals to SARS–coronavirus 2. *Science*,
   *368*(6494), 1016. doi:10.1126/science.abb7015
- Sia, S. F., Yan, L.-M., Chin, A. W. H., Fung, K., Choy, K.-T., Wong, A. Y. L., ... Yen, H.-L.
   (2020). Pathogenesis and transmission of SARS-CoV-2 in golden hamsters. *Nature*,
   583(7818), 834-838. doi:10.1038/s41586-020-2342-5
- 402 Sit, T. H. C., Brackman, C. J., Ip, S. M., Tam, K. W. S., Law, P. Y. T., To, E. M. W., ... Peiris,
   403 M. (2020). Infection of dogs with SARS-CoV-2. *Nature*, *586*(7831), 776-778.
   404 doi:10.1038/s41586-020-2334-5
- The UniProt, C. (2021). UniProt: the universal protein knowledgebase in 2021. *Nucleic Acids Research, 49*(D1), D480-D489. doi:10.1093/nar/gkaa1100
- Wang, Q., Zhang, Y., Wu, L., Niu, S., Song, C., Zhang, Z., ... Qi, J. (2020). Structural and
  Functional Basis of SARS-CoV-2 Entry by Using Human ACE2. *Cell*, 181(4), 894904.e899. doi:<u>https://doi.org/10.1016/j.cell.2020.03.045</u>
- Waterhouse, A., Bertoni, M., Bienert, S., Studer, G., Tauriello, G., Gumienny, R., . . . Schwede,
   T. (2018). SWISS-MODEL: homology modelling of protein structures and complexes.
   *Nucleic Acids Research*, 46(W1), W296-W303. doi:10.1093/nar/gky427
- Yan, R., Zhang, Y., Li, Y., Xia, L., Guo, Y., & Zhou, Q. (2020). Structural basis for the
  recognition of SARS-CoV-2 by full-length human ACE2. *Science*, *367*(6485), 1444.
  doi:10.1126/science.abb2762
- Zhou, P., Yang, X.-L., Wang, X.-G., Hu, B., Zhang, L., Zhang, W., . . . Shi, Z.-L. (2020). A
  pneumonia outbreak associated with a new coronavirus of probable bat origin. *Nature*,
  579(7798), 270-273. doi:10.1038/s41586-020-2012-7
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