Protein Structure and Sequence Reanalysis of 2019-nCoV Genome Refutes Snakes as Its Intermediate Host and the Unique Similarity between Its Spike Protein Insertions and HIV-1

Chengxin Zhang, Wei Zheng, Xiaqiang Huang, Eric W. Bell, Xiaogen Zhou, and Yang Zhang*

ABSTRACT: As the infection of 2019-nCoV coronavirus is quickly developing into a global pneumonia epidemic, the careful analysis of its transmission and cellular mechanisms is sorely needed. In this Communication, we first analyzed two recent studies that concluded that snakes are the intermediate hosts of 2019-nCoV and that the 2019-nCoV spike protein insertions share a unique similarity to HIV-1. However, the reimplementation of the analyses, built on larger scale data sets using state-of-the-art bioinformatics methods and databases, presents clear evidence that rebuts these conclusions. Next, using metagenomic samples from Manis javanica, we assembled a draft genome of the 2019-nCoV-like coronavirus, which shows 73% coverage and 91% sequence identity to the 2019-nCoV genome. In particular, the alignments of the spike surface glycoprotein receptor binding domain revealed four times more variations in the bat coronavirus RaTG13 than in the Manis coronavirus compared with 2019-nCoV, suggesting the pangolin as a missing link in the transmission of 2019-nCoV from bats to human.

KEYWORDS: 2019-nCoV, metagenome assembly, Malayan pangolins, spike protein

INTRODUCTION

The 2019 novel coronavirus (2019-nCoV), also known as SARS-CoV-2 and HCoV-19, is the pathogen behind COVID-19, a new type of pneumonia that initially caused an outbreak in Wuhan, China, and has since spread to most countries in the world. The rapid transmission across country borders and the large number of confirmed cases prompted the World Health Organization (WHO) to declare COVID-19 as a global pandemic on March 11, 2020. As of March 23, there are at least 332,930 and 14,510 patients who have been diagnosed with and have died of COVID-19 worldwide, respectively. Among the affected countries, China has the largest population of confirmed cases (81,610) and the second highest death toll (3,276). Meanwhile, Europe and North America have also been hit hard: 59,138 and 31,573 cases were confirmed in Italy and the United States, which are the nations with the highest number of 2019-nCoV infected patients in their respective continents, with the number of deaths in Italy (5,476) surpassing that of China. Understanding the viral infection mechanisms and animal hosts is of high urgency for the control and treatment of the 2019-nCoV virus. Whereas it is now commonly recognized that bats such as Rhinolophus affinis may serve as the natural reservoir of 2019-nCoV, it is still unclear which animal served as the intermediate host that brought the bat coronavirus to human hosts. Whereas multiple studies suggest the Malayan pangolin (Manis javanica) as another host, some studies have proposed that the pangolin may be a natural host rather than an intermediate host.1-3

During 2019-nCoV’s infection of host cells, a critical virion component is the spike surface glycoprotein, also known as the S protein. Spike proteins constitute the outermost component in a coronavirus virion particle and are responsible for the recognition of angiotensin-converting enzyme 2 (ACE2), a transmembrane receptor on mammalian hosts that is utilized by the coronavirus to enter the host cells.5,6 Therefore, the spike protein largely determines the host specificity and infectivity of a coronavirus.

In this Communication, we first analyzed the results of two recent studies,7,8 which have spurred numerous interests and discussions in the community and society regarding the sequence and structure of the spike protein in 2019-nCoV and the identification of its intermediate hosts. In particular, the study by Pradhan et al. reported the identification of four unique insertions that were shared only with HIV-1 and were "unlikely to be fortuitous in nature."10 Although the work has been questioned by the scientific community, rumors and conspiracy theories based on these studies still widely circulate among the general public.11 We therefore believe that there is an urgent
need to systematically examine the bases and conclusions of these studies in serious scientific reports. To further examine the animal hosts of the 2019-nCoV spread, we next assembled the draft genome of a highly related coronavirus using metagenomic samples from *Manis javanica*. The alignment results of the assembled genome sequences, in particular, on the spike proteins, suggest the importance of pangolins in the evolution of 2019-nCoV and its transmission from bats to humans.

**MATERIALS AND METHODS**

**Protein Sequence Alignment**

Global protein sequence alignment of the full-length coronavirus spike proteins was performed by MUSCLE and visualized by SeaView.

**Structure Prediction of Spike-ACE2 Complex**

We used C-I-TASSER to create structural models of the full-length spike protein. Here C-I-TASSER is an extended pipeline of I-TASSER and utilizes the deep convolutional neural-network-based contact maps to guide the Monte Carlo fragment assembly simulations. Because the RBD domain of the spike exhibits different conformations relative to the remaining portion of the protein, the DEMO pipeline was then used to reassemble the domains and to construct a complex structure consisting of the spike trimer and the extracellular domain of human ACE2 using the ACE2-bound conformation of the SARS-CoV spike glycoprotein (PDB ID: 6ACJ) as a template.

Our complex modeling did not use the template originally used in the Pradhan et al. study (PDB ID: 6ACD) because it did not include the ACE2 receptor.

**Relative Synonymous Codon Usage Analysis**

As per the previous study, the relative synonymous codon usage (RSCU) for codon \(j\) in a species is calculated as

\[
X_j = \frac{k_j}{\sum k_j}
\]

where \(k_j\) is the number of codons synonymous to codon \(j\) (including \(j\) itself) and \(p_j\) is the probability of the respective amino acid being encoded by codon \(j\) among all \(k_j\) synonymous codons in the protein coding sequences (CDSs) of the whole genome. The difference in codon usage in two different species (a virus versus a vertebrate in our case) is defined by the squared Euclidean distance of RSCU, that is

\[
d = \sqrt{\sum_{j=1}^{N} \left( X_j - X'_j \right)^2}
\]

Here \(N = 61\) is the number of codons that encodes amino acids, thereby excluding the three stop codons. \(X_j\) and \(X'_j\) are the RSCUs for codon \(j\) in the virus and in the vertebrate, respectively. In our report, the codon usages of all vertebrates are taken from the CoCoPUTS database, which was last updated in January 2020. This database was therefore much more recent than the Codon Usage Database, which was last
updated in 2007, that was used in the previous research.\textsuperscript{11} To obtain the codon usage of coronaviruses, we imported the GenBank annotations of the three coronavirus genomes to SnapGene (GSL Biotech) to export the codon usage table based on GenBank annotations. \textit{CodonW}\textsuperscript{21} was not used for the codon usage calculation as in the previous study because it cannot account for the -1 frameshift translation of the first open reading frame (ORF) in the coronavirus genome.

\section*{RESULTS AND DISCUSSION}

\subsection*{2019-nCoV Spike Protein Does Not Include Insertions Unique to HIV-1}

In a recent manuscript entitled “Uncanny Similarity of Unique Inserts in the 2019-nCoV Spike Protein to HIV-1 gp120 and Gag”,\textsuperscript{10} Pradhan et al. presented a discovery of four novel inserts unique to 2019-nCoV spike protein (Figure 1). They further concluded that these four inserts are part of the receptor binding site of 2019-nCoV and that these insertions shared “uncanny similarity” to human immunodeficiency virus 1 (HIV-1) proteins but not to other coronaviruses. These claims resulted in considerable public panic and controversy in the community,\textsuperscript{12} even after the manuscript was withdrawn. To investigate whether the conclusions by Pradhan et al. are scientifically precise, we reanalyzed the structural location and sequence homology of the four spike protein inserts discussed therein.

Because the full-length structure of the spike protein in 2019-nCoV was not available at the beginning of this study, we used C-I-TASSER\textsuperscript{15} to model its tertiary structure as part of our

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“If there are multiple redundant hits for the same gene from different strains of the same species removed, then only one hit is shown. The sequence identity is calculated as the number of identical residues divided by the query length. Only the sequence portion aligned to the query is shown. In this table, we also list the closest BLAST hit from bat coronavirus RaTG13, which is known to be closely related to 2019-nCoV."\textsuperscript{3,14}
Because this proteases such as cathepsin L (CTSL) to produce the S1 and S2 interfaces with ACE2 receptor binding, the spike protein can be cleaved by host ACE2 receptor binding. Here it is important to note that following Pradhan et al., which stated that the insertions are located on the spike protein, in contrast with the original conclusion made by DEMO structural models are located outside the RBD of the 2019-nCoV, which are available at [https://zhanglab.ccmb.med.umich.edu/C-I-TASSER/2019-nCoV/](https://zhanglab.ccmb.med.umich.edu/C-I-TASSER/2019-nCoV/). The 2019-nCoV spike model was then assembled with the human ACE2 structure, in contrast with the original conclusion made by DEMO structural models are located outside the RBD of the 2019-nCoV, which are available at [https://zhanglab.ccmb.med.umich.edu/C-I-TASSER/2019-nCoV/](https://zhanglab.ccmb.med.umich.edu/C-I-TASSER/2019-nCoV/) by DEMO18 to form a spike model was then assembled with the human ACE2 structure, where the C-I-TASSER model shares a high structure similarity, with a TM score of 0.95,24,25 to the cryo-EM structure. Because the experimental structure covers only 75% of the residues in the full-length sequence, with several important residues on the receptor binding domain (RBD) of the spike protein missing, our following analysis will mainly be built on the C-I-TASSER reconstrcuted full-length model. We note that C-I-TASSER, also known as “Zhang-Server”, is the top ranked automated server for protein structure prediction in the Critical Assessment of Protein Structure Prediction round 13 (CASP13) challenge (http://www.predictioncenter.org/casp13/zscores_final.cgi?model_type=best&gp_type=server only) among all 39 servers from the community.

C-I-TASSER improves our previously developed I-TASSER structure prediction protocol26 by incorporating a deep-learning-based contact map prediction.17,27 On all 121 CASP13 targets, the average TM score of the C-I-TASSER first model (0.674) is 8.0% higher than that of I-TASSER (0.624) and 0.15% higher than that of C-QUARK (0.673), which is our only other automated CASP13 server and was ranked in second place in CASP13.

As shown in Figure 2B, all four insertions in the C-I-TASSER/DEMO structural models are located outside the RBD of the spike protein, in contrast with the original conclusion made by Pradhan et al., which stated that the insertions are located on the interface with ACE2. Here it is important to note that following ACE2 receptor binding, the spike protein can be cleaved by host proteases such as cathepsin L (CTSL) to produce the S1 and S2 isoforms to facilitate viral entry into host cells.28,29 Because this cutting site immediately follows insertion 4 (IS4) (Figure 1 arrow) along the 2019-nCoV spike protein sequence, there is a possibility that IS4 could affect the cleavage of the spike protein. Regardless, all of the insertions are not directly related to receptor binding.

To investigate the viral homologues of the four insertions, we further performed a BLAST sequence search of these four insertions against the nonredundant (NR) sequence database, restricting the search results to viruses (taxid: 10239) but leaving other search parameters at default values. The top five sequence homologues (including the query itself) identified for each insertion are listed in Table 1. In contrast with the previous claim that the four insertions are unique to 2019-nCoV and HIV-1, all four insertion fragments can be found in other viruses. In fact, an HIV-1 protein is among the top BLAST hits for only one of the four insertion fragments, whereas three of the four insertion fragments are found in bat coronavirus RaTG13. Moreover, partially due to the very short length of these insertions, which range from six to eight amino acids, the E values of the BLAST hits, which is a parameter used by BLAST to assess the statistical significance of the alignments and usually needs to be <0.01 to be considered significant,30 are all >4, except for a bat coronavirus hit for IS2. These high E values suggest that the majority of these similarities are likely to be coincidental.

Given that three out of the four insertion fragments are found in the bat coronavirus RaTG13, it is tempting to assume that these “insertions” may be directly inherited from bat coronaviruses. Currently, there are at least seven known human coronaviruses (2019-nCoV, SARS-CoV, MERS-CoV, HCoV-229E, HCoV-OC43, HCoV-NL63, and HCoV-HKU1), where many of them, including severe acute respiratory syndrome-related coronavirus (SARS-CoV) and Middle East respiratory syndrome-related coronavirus (MERS-CoV), were shown to be transmitted from bats.3,31 To further examine the evolutionary relationship between the 2019-nCoV and the bat coronavirus in comparison with other human coronaviruses, we used MUSCLE to create a multiple sequence alignment (MSA), presented in Figure 3, for all seven human coronaviruses and two bat coronaviruses, RaTG13 and RaSHC014, which have been considered to be the ancestors of 2019-nCoV and SARS-CoV, respectively.5,31,34 Among the four “insertions” (ISs) of the 2019-nCoV, IS1 has only one residue different from the bat coronavirus, and three out of seven residues are identical to MERS-CoV. IS2 and IS3 are both identical to the bat coronavirus, RaTG13, it is tempting to assume that these “insertions” may be directly inherited from bat coronaviruses. Currently, there are at least seven known human coronaviruses (2019-nCoV, SARS-CoV, MERS-CoV, HCoV-229E, HCoV-OC43, HCoV-NL63, and HCoV-HKU1), where many of them, including severe acute respiratory syndrome-related coronavirus (SARS-CoV) and Middle East respiratory syndrome-related coronavirus (MERS-CoV), were shown to be transmitted from bats.3,31 To further examine the evolutionary relationship between the 2019-nCoV and the bat coronavirus in comparison with other human coronaviruses, we used MUSCLE to create a multiple sequence alignment (MSA), presented in Figure 3, for all seven human coronaviruses and two bat coronaviruses, RaTG13 and RaSHC014, which have been considered to be the ancestors of 2019-nCoV and SARS-CoV, respectively.5,31,34 Among the four “insertions” (ISs) of the 2019-nCoV, IS1 has only one residue different from the bat coronavirus, and three out of seven residues are identical to MERS-CoV. IS2 and IS3 are both identical to the bat coronavirus. For IS4, although the local sequence alignment by BLAST did not hit the bat coronavirus in Table 1, it has a
Snakes have the smallest codon usage distance among all the vertebrates, including frogs and birds. This suggests that snakes are not the intermediate hosts of 2019-nCoV, as proposed by Ji et al. 35 The lack of prior biological evidence that zoonotic viruses evolve their codon usage to resemble that of their animal hosts and the results of this study indicate that there is no evolutionary correlate to the bat coronavirus in the 2019-nCoV genome. The results encourage the need for a careful benchmark of RSCU analysis in viruses that evolve their codon usage to resemble that of their animal hosts, including SARS-CoV and MERS-CoV.

**Figure 4.** Inability of RSCU analysis for coronavirus host identification for (A) 2019-nCoV, (B) SARS-CoV, and (C) MERS-CoV. The vertebrate species (frogs) with the lowest squared Euclidean distances of RSCU (x axis) to the coronavirus is colored in dark gray, whereas the vertebrate (frog) with the lowest RSCU distance and sufficient statistics is colored in light gray. The snakes proposed by Ji et al. as intermediate hosts (Naja atra and Bungarus multicinctus snakes) are colored in black. Confirmed hosts (Rhinolopus affinis and Manis javanica for 2019-nCoV, Rhinolopus sinicus and Paguma larvata for SARS-CoV, and Camelus dromedarius for MERS-CoV, as well as Homo sapiens for all three coronaviruses) are colored in white. These data show not only that snakes are not the vertebrates with the lowest RSCU distances to 2019-nCoV but also that unrelated species such as frogs and snakes have smaller RSCU distances to known hosts of all three coronaviruses. These data suggest that the closeness of RSCU is not indicative of a potential pathogen–host relation.
and those of all 102,367 vertebrate species in the CoCoPUTS database. To test whether this kind of analysis can recover known hosts of well-studied coronaviruses, SARS-CoV (NCBI accession NC_004718) and MERS-CoV (NCBI accession NC_019843) were also included. The codon usage frequency is converted to the squared Euclidean distance of RSCU in two separate analyses: one based on all vertebrates (Supplementary Figure S1A–C) and the other based on the subset of vertebrates with enough statistics, that is, >2000 known CDSs (Supplementary Figure S1D–F), roughly corresponding to 10% of all protein coding genes in a typical vertebrate genome.

As shown in Figure 4A, snakes are not the vertebrates with the lowest RSCU distances to 2019-nCoV, suggesting that the implementation of RSCU analysis by Ji et al. was incomplete. More importantly, the data in Figure 4 show that animals unrelated to coronavirus transmission, such as frogs and snakes, consistently have smaller RSCU distances to known hosts of all three coronaviruses. For example, the top-ranking vertebrates with the lowest RSCU distances to the three different coronaviruses are two kinds of frogs (Megophrys feae and Liophryne schlaginhaufeni), whereas another frog (Xenopus laevis) has the smallest RSCU distances among all vertebrates with sufficient sequences. Part of the reason for the failure of RSCU in intermediate host identification, as shown in Supplementary Table S1, is that different coronaviruses, such as SARS-CoV and MERS-CoV, that are known to utilize different intermediate hosts (Paguma larvata and Camelus dromedarius, respectively), have almost no difference in RSCU (squared RSCU distance = 0.12). These data suggest that the RSCU analysis on its own is not specific enough to discriminate coronaviruses from different vertebrate hosts. In this regard, the failure is not merely due to the use of outdated databases or the small number of species included in the original analysis but is, in fact, caused by the incorrect biological assumption that coronaviruses will evolve their RSCU to resemble that of their hosts.

Metagenome Assembly Suggests Pangolins as Potential Hosts of 2019-nCoV

In a recent study,7 Xiao et al. first identified coronavirus sequences in pangolins that are highly similar to 2019-nCoV. In addition, three independent groups also reported the identification of 2019-nCoV-like coronaviruses sequences from metagenomics samples taken from the Malayan pangolin (Manis javanica),8,9 making the pangolin a likely intermediate host of the 2019-nCoV.
To further examine the possibility, we tried to reassemble a draft genome sequence of the coronavirus using the metagenomic samples of *Manis javanica*. To this end, we first collected a set of all publicly available metagenome samples for pangolin, including 11 samples from lung, 8 samples from spleen, 2 samples from lymph (NCBI accession PRJNAS73928),37 and 4 samples from feces (NCBI accession PRJNA476660),38 from the NCBI Sequence Read Archive (SRA) database using the prefetch command of SRA Toolkit version 2.10.3. These samples were converted to paired-end sequencing reads in FASTQ format by fast-dump. A quality check by FastQC version 0.11.9 showed that whereas the 4 samples from PRJNA476660 do not contain adaptor sequences, all 21 samples from PRJNAS73928 contain Illumina universal adaptors. Therefore, for these 21 samples, Trimmomatic version 0.3930 was used to remove adaptor sequences using the flag "ILLUMINACLIP:adapters.fa:2:30:10:2:keepBothReads LEADING:3 TRAILING:3 MINLEN:36". To remove contaminations from the host and from human researchers, only read pairs that could be mapped to *Manis javanica* or *Homo sapiens* genomes by bowtie2 version 2.3.5.1 were retained for further analysis. These sequences were converted from SAM format of bowtie2 back to FASTQ format by SAMtools version 1.10 and bedtools33 version 2.29.2. Following these quality-control processes, we next determined which of the 25 previously mentioned samples include a 2019-nCoV-like sequence by two searches at the protein and nucleotide levels. In the protein-level search, the 2019-nCoV spike protein sequence was searched by BLASTp30 through protein sequences directly assembled from sequencing reads of a metagenome sample by Plass, a protein-level metagenome sequence assembler,14 to identify if there were any close hits with an E value <0.01. Meanwhile, the nucleotide-level search selected samples where more than one pair of sequencing reads could be mapped to the 2019-nCoV genome (NCBI accession: MN908947.3) by bowtie. Both searches consistently reported that only the lung samples (SRA accessions: SRR10168376, SRR10168377, and SRR10168378) contain 2019-nCoV-like sequences. Therefore, the sequences were assembled into nucleotide and protein contigs by MEGAHIT and Plass, respectively. The assembled nucleotide and protein sequences were then aligned by BLASTn and BLASTp to the whole genome and the spike protein of 2019-nCoV, respectively, at an E-value cutoff of <0.01. Finally, we separately merged all nucleotide and protein alignments into a single pairwise alignment between 2019-nCoV and the *Manis* coronavirus (*Manis*-CoV); when multiple *Manis*-CoV hits cover the same 2019-nCoV region, the hit with the highest sequence identity to 2019-nCoV is used in the merged alignment.

Figure 5A presents a sketch of the draft genome for the *Manis*-CoV as compared with the released 2019-nCoV genome.45 Overall, the assembled sequences cover 73% of the 2019-nCoV genome with 91% sequence identity. More importantly, the protein sequences assembled from these *Manis* lung samples include a partial pangolin coronavirus spike protein that is 92% identical to the 2019-nCoV spike protein (Figure 5B). This sequence identity is relatively high, considering that spike proteins are critical for the coronaviruses to invade into host cells and have the largest diversity in coronavirus genomes due to evolutionary pressure to adapt to receptors on different hosts. Notably, there are only 5 residue positions in the *Manis* coronavirus that are different from 2019-nCoV on the spike receptor binding domain compared with 19 different residue positions between 2019-nCoV and bat coronavirus RaTG13 for the same domain (Figure 5B, black box). These data imply that pangolins such as *Manis javanica* can either be the intermediate hosts of 2019-nCoV for the transmission of bat coronaviruses to humans or serve as alternative natural hosts, together with bats, to provide the genetic material for the origin of 2019-nCoV. Nevertheless, considering that *Manis javanica* individuals with coronavirus infections are usually in poor or even critical health condition37 and previously known natural coronavirus hosts (such as bats) are usually asymptotic after infection, thus allowing long-term virus-host coexistence and coevolution, we believe that it is more likely that *Manis javanica* is an intermediate host rather than a natural host.

Approximately one-quarter of nucleotides are missing in our assembled *Manis* coronavirus draft genome, partly because compared with whole-genome sequencing, metagenome sequencing usually has a lower read depth and more assembly errors caused by the mixture of diverse species in the samples. A higher quality genome with better coverage should, in theory, be attainable if the *Manis* coronavirus can be isolated and cultured in vitro using a mammalian cell line and is subjected to whole-genome sequencing.

## CONCLUSIONS

Because of the scarcity of experimental and clinical data as well as the urgency to understand the infectivity of deadly coronaviruses, we have been increasingly relying on computational analyses to study the 2019-nCoV virus in terms of protein structures, functions, phylogeny, and interactions at both molecular and organismal levels. Indeed, within less than 1 month of the publication of the 2019-nCoV genome in January 2020, multiple bioinformatics analyses regarding 2019-nCoV have been either published or posted as preprints. Whereas such expeditious analyses provide much needed insights into the biology of the 2019-nCoV virus, there is a caution to avoid overinterpretation of the data in the absence of comprehensive benchmarks or follow-up experimental validations.

In this Communication, we have investigated two recently published computational analyses regarding intermediate host identification and the analysis of spike protein insertions. In both cases, we found that the conclusions proposed by the original studies do not hold in the face of more comprehensive replications of these analyses. In particular, we found that the unique sequence “inserts” found by Pradhan et al. are, in fact, shared by multiple viruses, especially with the segments from the bat coronavirus RaTG13, revealing the close evolutionary relation to the latter species. In addition, our benchmark results showed that the data based on RSCU are not specific enough to discriminate the relation between coronaviruses and vertebrates, which contradicts the conclusion by Ji et al. regarding snakes as an immediate host of the 2019-nCoV.

Finally, we assembled a draft genome of the 2019-nCoV-like coronavirus using the metagenomic samples from the lung of *Manis javanica*, which shows an overall coverage of 73% of 2019-nCoV with 91% sequence identity. In particular, the spike protein in the assembled genome, which is critical for the virus to recognize host receptors and therefore bears a high speed of variation, shares a high sequence identity with 2019-nCoV, with only 5 residue position differences compared with 19 differences between 2019-nCoV and bat coronavirus RaTG13. These data provide evidence of the possible evolutionary relations among RaTG13, the *Manis* coronavirus, and 2019-nCoV.
Second, the 91% sequence identity between the Manis coronavirus and 2019-nCoV is high enough to confirm an evolutionary relation between the two viruses but not high enough to consider them as the same viral species. To put this into perspective, the viral sequence from intermediate hosts of SARS-CoV and MERS-CoV are 99.8 and 99.9% identical to their human versions, respectively. Therefore, even with the discovery of Manis coronavirus, further searching for other potential intermediate hosts should be continued.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jproteome.0c00129.

Figure S1. Top 20 vertebrate species ranked in ascending order of squared Euclidean distance of RSCU to 2019-nCoV, SARS-CoV, and MERS-CoV. Table S1. Squared Euclidean distances of RSCU among coronaviruses and representative vertebrate species (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**

Yang Zhang — Department of Computational Medicine and Bioinformatics and Department of Biological Chemistry, University of Michigan, Ann Arbor, Michigan 48109-2218, United States; orcid.org/0000-0001-7290-1324

Email: zhng@umich.edu

**Authors**

Chengxin Zhang — Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, Michigan 48109-2218, United States; orcid.org/0000-0002-2984-9003

Wei Zheng — Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, Michigan 48109-2218, United States; orcid.org/0000-0002-2984-9003

Xiaoqiang Huang — Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, Michigan 48109-2218, United States; orcid.org/0000-0002-1005-848X

Eric W. Bell — Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, Michigan 48109-2218, United States; orcid.org/0000-0002-3419-4398

Xiaogen Zhou — Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, Michigan 48109-2218, United States; orcid.org/0000-0001-6839-1923

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.jproteome.0c00129

**Author Contributions**

Y.Z. conceived and designed this study. C.Z. performed the RSCU analysis and structure analysis. C.Z. and W.Z. performed the sequence analysis of the spike protein. X.G. performed the domain assembly of the spike protein. C.Z., W.Z., X.H., E.W.B., and Y.Z. wrote the manuscript.

**Notes**

The authors declare no competing financial interest.

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**ABBREVIATIONS**

2019-nCoV, 2019 novel coronavirus; ACE2, angiotensin-converting enzyme 2; HIV-1, human immunodeficiency virus 1; IS, insertion; SARS-CoV, severe acute respiratory syndrome-related coronavirus; RBD, receptor binding domain; CDS, protein coding sequence; MERS-CoV, Middle East respiratory syndrome-related coronavirus; RSCU, relative synonymous codon usage; Manis-CoV, coronavirus infecting Manis javanica

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