Evolutionary relationships and sequence-structure determinants in human SARS coronavirus-2 spike proteins for host receptor recognition

Lalitha Guruprasad*, School of Chemistry, University of Hyderabad, Hyderabad 500046, India.

*Email: lalitha.guruprasad@uohyd.ac.in

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Abstract:

Coronavirus disease 2019 (COVID-19) is a pandemic infectious disease caused by novel Severe Acute Respiratory Syndrome coronavirus-2 (SARS CoV-2). The SARS CoV-2 is transmitted more rapidly and readily than SARS CoV. Both, SARS CoV and SARS CoV-2 via their glycosylated spike proteins recognize the human angiotensin converting enzyme-2 (ACE-2) receptor. We generated multiple sequence alignments and phylogenetic trees for representative spike proteins of CoV and CoV-2 from various host sources in order to analyze the specificity in SARS CoV-2 spike proteins required for causing infection in humans. Our results show that two sequence motifs in the N-terminal domain; "MESEFR" and "SYLTPG" are specific to human SARS CoV-2. In the receptor binding domain (RBD), two sequence motifs; "VGGNY" and "EIYQAGSTPCNGV" and a disulfide bridge connecting 480C and 488C in the extended loop are structural determinants for the recognition of human ACE-2 receptor. The complete genome analysis of representative SARS CoVs from bat, civet, human host sources and human SARS CoV-2 identified the bat genome (GenBank code: MN996532.1) as closest to the recent novel human SARS CoV-2 genomes. The bat CoV genomes (GenBank codes: MG772933 and MG772934) are evolutionary intermediates in the mutagenesis progression towards becoming human SARS CoV-2.

Introduction:

In the last two decades, zoonotic coronaviruses, Severe Acute Respiratory Syndrome coronavirus SARS CoV (2002) (Dorsten et al., 2003) and Middle East Respiratory Syndrome coronavirus (MERS CoV) (2012) (Azhar et al., 2014) have caused acute respiratory diseases in humans that have resulted in several deaths. The present coronavirus disease 2019 (COVID-19) is a pandemic respiratory disease caused by the novel SARS CoV-2. The initial infection started in Wuhan, Hubei province, China in December 2019 and very soon became a global outbreak, infecting population in almost every country in the world causing a total of 184,248 deaths and 2,638,852 infections as of 23rd April 2020 (https://www.worldometers.info/coronavirus/). Within a short span of time this pandemic has caused major social and economic disruptions. Compared to other coronaviruses, the novel SARS CoV-2 appears to be spreading more rapidly and readily, posing a challenging task before the administrative and scientific communities. The SARS CoV-2 is transmitted from
person to person contact via respiratory secretions during coughing and sneezing. Infection of this highly pathogenic virus can cause acute respiratory distress syndrome which impacts the lung and heart functions. The prominent symptoms of this viral infection are flu, severe respiratory, enteric and neurological disorders, resulting in increased white blood cells and kidney failure. There are no vaccines or drugs available to combat this deadly infectious disease and there is no strategic plan to treat the infected patients. Hence, there is a need to develop specific anti-CoV-2 vaccines and drugs to treat infected patients, in order to reduce viral shedding and further transmission in populations.

The SARS CoV-2 comprises positive-sense single-stranded RNA genome of size 29-30 kb and belongs to the coronaviridae family and betacoronavirus sub-family. Mammals such as bats are the main reservoir of betacoronaviruses, but due to the zoonotic contacts and viral genomic mutations, SARS CoV-2 has recently crossed species and caused infections in humans (Wu et al., 2020). Research findings have pointed that previous zoonotic CoV infections such as, SARS CoV, that first infected humans in the Guangdong province of southern China in 2002 was transmitted from bats and civets (Xu et al., 2004, Marra et al., 2003, Rota et al., 2003, Ksiazek et al., 2003, Holmes & Enjuanes 2003), the MERS CoV that originated in bats was first identified from camel to human transmission in Saudi Arabia in 2012 (Azhar et al., 2014, Chan et al., 2015, Sabir et al., 2016, Azhar et al., 2014a, Omrani et al., 2015). These coronaviruses have crossed species and resulted in causing human infections leading to mortality. It is reported that civet SARS CoV can also infect humans (Wang et al., 2005, Li et al., 2006). The SARS-like CoVs from some bats and civets are predicted to result in human infections (Menachery et al., 2015, Wang et al., 2018) due to their changing genomic RNA sequences, importantly in the spike protein regions (Song et al., 2005, Menachery et al., 2016). Since January 2020, several complete genome sequences of viral CoVs-2 isolated from infected patients belonging to various geographical locations, such as, Australia, China, Denmark, Finland, Hungary, India, Italy, Japan, South Korea, USA and Vietnam have been deposited in the GenBank (https://www.ncbi.nlm.nih.gov/genbank/).

At the genomic level, the nucleotide sequences of SARS CoV and SARS CoV-2 share 79.6% sequence identity (Zhou et al., 2020). The viral RNA stores the genetic information and also serves to translate into structural and non-structural proteins (NSPs) of SARS CoV-2. The SARS CoV uses angiotensin converting enzyme-2 (ACE-2) as receptor for entry into human epithelial cells (Li et al., 2003) to cause the infection. Zhou et al., 2020 have carried out virus
infectivity studies on HeLa cells and have shown that SARS CoV-2 also uses ACE-2 as receptor for cellular entry.

It has been reported that the first SARS CoV-2 infection in Wuhan, China is caused from the original host, bats (Zhou et al, 2020), and in less than 4 months transmitted among the human populations in the entire world. The initial contact between the SARS CoV/CoV-2 and human host is via recognition between the heavily glycosylated cell envelope spike protein of the virus and the ACE-2 receptor of the human host resulting in an infection. In order to understand the specificity, estimate the extent of similarities and variations in the SARS CoV-2 spike proteins required for binding to the host receptor, we analyzed the representative spike protein sequences. Further, in order to estimate the evolutionary progression of the bat SARS CoV genome, such that it is able to adapt to a human host as a novel coronavirus causing COVID-19, we have analyzed the complete genomes of the bat, civet, human SARS CoV and human SARS CoV-2. We have carried out computational analyses on the nucleotide and protein sequences by generating multiple sequence alignments (MSAs), constructing phylogenetic trees and analyzing the three-dimensional structure of the spike proteins to address the above.

Materials and Methods:

The spike proteins were retrieved from the NCBI database, using the sequence similarity search BLAST program (Schäffer et al., 2001) with human SARS CoV-2 spike protein as the query sequence (NCBI code: QHD43416.1) from the genome (GenBank code: MN908947.3) (Wu et al., 2020). The complete genome nucleotide sequences of SARS CoV from bats, civets, and SARS CoV and CoV-2 from human host were obtained from NCBI virus database (https://www.ncbi.nlm.nih.gov/labs/virus/vssi/#/virus?SeqType_s=Nucleotide) in FASTA format. Only complete genome sequences without any ambiguity in nucleotide composition were considered for analyses. The redundancy in each dataset was removed using the CD-HIT program (Li & Godzik 2006).

The nucleotide and protein sequence homology analyses based on MSA reveals the substitutions, deletions and insertions at each position. To understand the evolutionary relationships between the members, the MSAs were further processed to generate phylogenetic trees - a pictorial representation of the evolutionary relationships between
related members of various sequences analyzed. The MSAs and phylogenetic trees of the spike proteins and complete genomes were generated using the Next Generation Phylogeny.fr web service available at https://NGPhylogeny.fr (Lemoine et al., 2019). The protocol takes all sequences (nucleotide or protein) as input in FASTA file format and generates the MSA and phylogenetic tree. In the NGPhylogeny server, we have selected FastME 2.0 program that infers phylogenies using a distance approach since it is capable of handling large datasets (Lefort et al., 2015). Based on the input FASTA sequences, a MSA is generated that adopts Multiple Alignment using Fast Fourier Transform (MAFFT) (Katoh & Standley 2013) with gap extension penalty; 0.123 and gap opening penalty; 1.53. The MSA generated is parsed through Block Mapping and Gathering with Entropy (BGME) software for selecting regions suitable for phylogenetic inference (Criscuolo & Gribaldo 2010). This method uses a sliding window size; 3, Maximum entropy threshold; 0.5, gap rate cutoff; 0.5, minimum block size; 5, matrix: PAM250 for DNA and BLOSUM62 for proteins. FastME estimates phylogenies that employ distance-based methods from MSAs using TN93 and LG as substitution models for DNA and proteins, respectively. In the distance-based methods, pairwise distances between all pairs of sequences are generated as a square matrix. The sequence pair with shorter pairwise distances are clustered together more closely in the phylogenetic tree. Tree refinement was performed using Subtree Pruning and Regrafting (SPR) with Balanced version of Minimum Evolution (BalME), with a decimal precision for branch length set to 6. Finally, the phylogenetic trees were generated using Interactive Tree Of Life program (iTOL) v4 (Letunic and Bork, 2019).

Results and discussion:

Analyses of the spike proteins of SARS Cov and SARS CoV-2

The SARS CoV and SARS CoV-2 spike proteins retrieved from various host sources have a sequence length ranging between 1240 to 1273 amino acids. Structurally, a spike protein is characterized by three regions; 1) the N-terminal extracellular domain, 2) a transmembrane anchor domain and 3) an intracellular segment. The N-terminal extracellular domain comprises a receptor binding subunit (S1) and a membrane-fusion subunit (S2). The S1 subunit comprises two domains, an N-terminal domain (NTD) and RBD. The sequence analyses of spike proteins from the various host sources is shown in the MSA in supplementary Figure S1 and the phylogenetic tree in Figure 1. From Figure 1, it is observed
that the paralogous proteins from individual host sources are associated with a distinct clade, the spike proteins from human SARS CoV-2 share highest sequence similarity according to the least pairwise distances. Also, the orthologous spike proteins from other host species are highly similar according to the low pairwise distances. Therefore, it is intriguing to see that despite high sequence identity between the spike proteins from various host sources, only some SARS CoVs and SARS CoV-2 are able to bind the human host ACE-2 receptor. In order to understand this, we have analyzed the MSA of the spike proteins.

From the supplementary Figure S1, it is observed that the N-terminal ~500 amino acids, comprising the S1 subunit vary to a moderate extent among all host sources, relative to the later region that shares higher sequence identity. The region between ~300-500 amino acid residues is crucial in spike proteins, as it forms the RBD that recognizes the ACE-2 receptor which allows entry of the virus into the host cells. A sequence motif "PRRA", from 681P to 684A (amino acid numbering is according to NCBI code: QHD43416) that is gained only in the human SARS CoV-2 spike proteins is referred to as a furin cleavage site (Wang et al., 2020, Ou et al., 2020). In this work, we identify the sequence motifs specific to human SARS CoV-2; a six residues insertion "MESEFR" from 153M to 158R and another six residues insertion sequence motif "SYLTPG" from 247S to 252G. The bat SARS CoV spike protein QHR63300 (GenBank code:MN996532.1) also comprises the identical sequence motifs as above. In two bat spike proteins, AVP78042 (GenBank code:MG772934) and AVP78031 (MG772933), a six residues sequence motif "SIREFA" and a three residue sequence motif "GDP" are present at equivalent positions, respectively. The insertion sequence motifs specific to human SARS CoV-2, 153MESEFR158 and 247SYLTPG252 are associated with the NTD and distant from the ACE-2 binding site. From the three-dimensional structure of human SARS CoV-2, we infer that this region is likely to be exposed towards the surface of the spike protein in NTD. In human SARS CoV-2, there are two insertion regions; a five residues insertion "VGGNY" from 445V to 449Y, and a thirteen residues insertion "EIYQAGSTPCNGV" from 471E to 483V. The bat SARS CoV spike protein QHR63300 has also gained equivalent insertions with the sequences; “EGGNF” and “EIYQAGSKPCNGQ”. It is interesting to note that the bat SARS CoV QHR63300 has already acquired a thirteen residues sequence motif, whereas all other spike proteins that recognize the ACE-2 receptor in SARS CoVs comprise a twelve residues sequence motif. The Figure 2A was generated by editing the MSA in Figure S1 to depict the sequence motifs discussed above in representative
spike proteins from the four host sources (bat, civet, human SARS CoV and human SARS CoV-2).

The two sequence motifs, "VGGNY" and "EIYQAGSTPCNGV", are part of the RBD and involved in recognition of the ACE-2 receptor in human host. Their absence in the bat SARS CoV at equivalent positions may be responsible for their inability to bind human ACE-2. To study this, we have analyzed the three-dimensional structures of human SARS CoV (PDB code: 6ACG) (Song et al., 2018) and the RBD domain of human SARS CoV-2 (6M17) complexed with ACE-2 (Yan et al., 2020). The structures were superimposed and amino acid residues within 5Å distance from the ACE-2 receptor were identified and highlighted in the pairwise sequence alignment of RBD (Figure 2B). Despite significant amino acid mutations in this region in both proteins, the structures are highly superimposable (Figure 2C). The location corresponding to the deletion regions of the two sequence motifs in bat SARS CoV with respect to human SARS CoV-2 match with the ACE-2 binding region (Figure 2D). The second insertion sequence motif, i.e., 471EIYQAGSTPCNGV483 in the RBD domain forms an extended loop in human SARS CoV-2 and is stabilized by a disulfide bond between 480C and 488C. This disulfide bond is conserved in all spike proteins that comprise the two insertion sequence motifs in RBD. In bat SARS CoV spike proteins, the position equivalent to 488C is replaced by a conserved glycine residue and therefore this disulfide bond would be absent. Interestingly, the bat SARS CoV spike protein QHR63300 which has acquired the insertion sequences in RBD also possesses the disulfide bond. We believe that the presence of the two sequence motifs in RBD and the disulfide bond that stabilizes the conformation of the extended loop are required for the recognition of human ACE-2.

The sequence motifs identified in this work serve as potential candidate epitopes for the design of antibodies specific for human SARS CoV-2 recognition. Our analyses suggest that the bat spike protein QHR63300 has undergone significant evolutionary changes such that it resembles the human CoV-2 spike protein more than the bat CoV which may have lead to the transmission of CoV from bat to human as the novel SARS CoV-2. Our results also suggest that the bat SARS CoV spike proteins; AVP78042 and AVP78031, are in progression of acquiring mutations towards becoming SARS CoV-2-like proteins. The phylogenetic tree in Figure 1 showing the proteins; QHR63300, AVP78042 and AVP78031 close to the human SARS CoV-2 are in support of our hypothesis.
Complete genome analyses of SARS CoV and SARS CoV-2

The representative complete genomes of nucleotide sequences from bat, civet, human CoV and human CoV-2 genomes were analyzed. The MSA is shown in Figure S2 and the phylogenetic tree in Figure 3. From Figure 3, it is observed that the human SARS CoV-2 genomes cluster into one clade (pairwise distance is lower than 0.002) revealing high identity that suggest their recent evolution. The bat SARS CoV genome (GenBank code: MN996532.1) is also member of this clade (pairwise distance between 0.042-0.043) indicating that it is the closest homolog to the human SARS CoV-2 among the bat genomes. The two bat SARS CoV genomes (GenBank codes: MG772933.1 and MG772934.1) are also close to the human SARS CoV-2 clade. The human and civet SARS CoV genomes cluster into another distinct clade. The members of bat SARS CoV clade have undergone maximum evolutionary changes as observed in Figure 3. Based on these results, we propose that the bat SARS CoV genomes have diverged the most during the last 18 years (since its detection in 2002) and have evolved closer to civet and human SARS CoV genomes. The bat SARS CoV genome (GenBank code: MN996532.1) has diverged significantly into the recent novel human SARS CoV-2 genomes, whereas, the bat CoV genomes (GenBank codes: MG772933 and MG772934) are intermediates during the evolution of bat SARS CoV into human novel SARS CoV-2. We believe genomes such as these are likely to undergo further evolutionary mutations and become adaptable to infecting human populations at the opportunistic moment and thus pose a potential threat.

Conclusions:

Two sequence motifs: "MESEFR" and "SYLTPG" in the NTD of the spike protein specific to human SARS CoV-2 and two sequence motifs in the RBD; "VGGNY" and "EIYQAGSTPCNGV" that interact with ACE-2 may be exploited as potential candidates for antibody design. The phylogenetic analyses of the bat, civet, human SARS CoV and human SARS CoV-2 genomes show that the bat SARS CoV genome (GenBank code: MN996532.1) is closest homolog of human SARS CoV-2. The two other bat SARS CoV genomes (GenBank codes: MG772933 and MG772934) that have gained insertion sequence motifs in NTD are intermediates in the evolution of bat genomes into human SARS CoV-2.
Conflict of interest:

The author declares no potential conflict of interest.

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Figure 1. Phylogenetic tree of SARS CoV and SARS CoV-2 spike proteins from human CoV-2 (green), human CoV (dark green), bat CoV (red), civet CoV (violet), cat CoV (orange), swine CoV (blue), mouse CoV (bright yellow), SARS CoV recombinant spike proteins and from lab adapted cells (black).
Figure 2A. Portions of the alignment of spike proteins extracted from the multiple sequence alignment (Figure S1) showing the insertion sequence motifs and their locations within the NTD, RBD and furin cleavage sites for human SARS CoV-2 (*), bat SARS CoV (@), civet SARS CoV (#), human SARS CoV ($).
**Figure 2B.** Pairwise sequence alignment corresponding to the RBD from human SARS CoV (6ACG) and human SARS CoV-2 (6M17). The residues that lie within 5Å in RBD from ACE-2 are highlighted in 6ACG (green) and 6M17 (magenta). The start and end amino acid residues numbers in RBD are shown. ‘*’ indicates identical residues, ‘.’ indicates conservative amino acid residue substitutions, ‘.’ indicates weakly conserved amino acid residue substitutions in the alignment.
Figure 2C. Structural superposition of human SARS CoV (PDB code: 6ACG, green) and human SARS CoV-2 (6M17, majenta) and the long H1 helix in ACE-2 (blue) are shown along with the side chains of amino acid residues in RBD that lie within 5Å from ACE-2.
Figure 2D. Structural superposition of human SARS CoV (6ACG, green) and human SARS CoV-2 (6M17, majenta) and the long H1 helix in ACE-2 (blue) are shown. The equivalent regions corresponding to deletions associated with the RBD sequence motifs in bat SARS CoV are indicated in the structure of 6M17 (red). The C480-C488 disulfide bond connecting the extended loop in 6M17 is shown (yellow).
Figure 3. Phylogenetic tree of SARS CoV and SARS CoV-2 complete genomes. Human SARS CoV-2 (green), human SARS CoV (dark green), bat SARS CoV (red), civet SARS CoV (violet).
Legend to supplementary Figures:

**Figure S1.** Multiple sequence alignment of spike proteins from SARS CoV and SARS CoV-2 from different host sources.

**Figure S2.** Multiple sequence alignment of complete genomes from human SARS CoV-2, human SARS CoV, bat SARS CoV and civet SARS CoV.