Comparative sequence analysis of SARS-CoV-2 suggests its high transmissibility and pathogenicity

Kartika Padhan1, Mohammad K Parvez*,2 & Mohammed S Al-Dosari2

1Center for Advanced Tissue Imaging, National Institute of Allergy & Infectious Diseases, National Institutes of Health, Bethesda, MD 20892, USA
2Department of Pharmacognosy, College of Pharmacy, King Saud University, Riyadh 11451, Saudi Arabia

*Author for correspondence: Tel.: +96 6 614 675 132; mohkhalid@ksu.edu.sa

Aim: Because the highly pathogenic SARS-CoV-2 is newly introduced to humans, we aimed to understand the unique features of its genome and proteins, crucial for high transmissibility and disease severity.

Materials & methods: The available genome and protein sequences of SARS-CoV-2 with known human and nonhuman CoV were analyzed using multiple sequence alignment programs.

Results: Our analysis revealed some unique mutations in SARS-CoV-2 spike, ORF1a/b, ORF3a/3b and ORF8. The most interesting ones were in the spike angiotensin-converting enzyme 2 receptor binding-motif and generation of a furin-like cleavage site as well as deletions of ORF3a ‘diacidic motif’ and the entire ORF3b.

Conclusion: Our data suggest that SARS-CoV-2 has diverged from SARS-CoV-1 but is most close to bat-SL-CoV. Unique mutations in spike and ORF3a/b proteins strongly endorse its adaptive evolution, enhanced infectivity and severe pathogenesis in humans.

First draft submitted: 20 June 2020; Accepted for publication: 23 February 2021; Published online: 8 March 2021

Keywords: bat-SL-CoV • COVID-19 • pathogenesis • SARS-CoV-1 • SARS-CoV-2 • spike protein

Following the 1918 ‘Spanish Flu’ pandemic, outbreaks of emerging or re-emerging novel viral diseases, such as 1956 ‘Asian Flu’, 1968 ‘Hong Kong Flu’ and 2009 ‘Swine Flu’ have exerted great toll on human health [1]. For centuries, coronaviruses (CoV), the causative agents of common cold have been known to infect humans. Of the known six human CoV (HCoV), the severe acute respiratory syndrome CoV (SARS-CoV-1) originating in China caused the first CoV pandemic during 2002–2003 [2]. Later, a second such pandemic caused by the Middle-East respiratory syndrome CoV (MERS-CoV) originated in Arabian Peninsula during 2012–2013 [2]. Fortunately, most of the countries remained unaffected because both SARS-CoV-1 and MERS-CoV were quickly contained within months.

The ongoing pandemic caused by the novel SARS-CoV-2 disease (COVID-19) that also originated in China is the seventh HCoV infection [3,4]. At the time of writing (February 2021), SARS-CoV-2 has infected over 42 million of the world’s population, causing >1 million deaths (https://coronavirus.jhu.edu/map.html). The phylogenetic analysis of SARS-CoV-2 genome has shown its close similarity (96%) with bat-SARS-like CoV (SL-CoV), indicating its zoonosis in horseshoe bats or civet cats [5]. In the absence of specific therapeutics, several repurposed drugs are currently under Phase II/III clinical trials or approved for emergency use [6–8]. Fortunately, of the leading vaccine candidates under final stages of trials, five have been recently granted approval in some countries.

SARS-CoV-2 belongs to the genus Betacoronavirus (Beta-CoV), and has a plus-sense single-strand genomic RNA (~30 kb) defined into 13 open reading frames (ORF), which code for its structural, nonstructural and accessory proteins [5]. The four structural proteins are crown-like spike (S), envelope (E), membrane (M) and nucleocapsid (N). The S protein has two structural subunits, wherein the ‘S1’ subunit contains the cellular angiotensin-converting enzyme 2 (ACE2) receptor-binding domain (RBD) and the ‘S2’ subunit possesses structural elements required for cell membrane fusion. The M protein is a trans-membrane glycoprotein crucial for membrane fusion, whereas the E protein is required for virion assembly and morphogenesis [5]. In case of SARS-CoV-1, the N protein is highly...
antigenic and is used as a serological marker [9]. The nonstructural replicase proteins (pp1a: nsp1–nsp11 and pp1b: nsp12–nsp16) are involved in mRNA synthesis and replication, and the accessory proteins (3a, 3b, 6, 7a, 7b, 8 and 9b) participate in modulating host innate immunity. Upon infection, SARS-CoV-2 first gets attached to the naso-/oro-pharyngeal inner linings and then moves down to the lungs, which are even richer in ACE2, and triggers cell damage. The bat-SL-CoV S protein is known to bind to civet and horseshoe bat ACE2 receptors [10]. Similarly, SARS-CoV-2 S protein also binds to the ACE2 of airway epithelium, alveolar type-2 pneumocytes that produce pulmonary surfactant [11]. In SARS-CoV-1 patients, the role of apoptosis in lung epithelial cells damage as well as hematological changes including lymphopenia, thrombocytopenia and occasionally leucopenia has been observed, suggesting it role in disease severity [12]. Moreover, the SARS-CoV-1 N, 3a, 3b and 7a proteins are reported to induce apoptosis in cultured cells [13,14].

Theoretical framework
SARS-CoV-2 has faster ‘human-to-human’ transmission rates and higher pathogenicity than SARS-CoV-1 and MERS-CoV [15]. Notably, the success behind containment of SARS-CoV-1 was due to the fact that the majority of infections happened in hospital setting where spread occurred during late and symptomatic phase [16,17]. Unlike SARS-CoV-1, most of the spread of SARS-CoV-2 is occurring through asymptomatic infection [18], which is a bottleneck for its quick containment. In addition, SARS-CoV-2 has been reported to survive longer than SARS-CoV-1 on certain surfaces including cardboard, plastic and stainless steel [19], which raises the higher chances of its fomite transmission. In general, HCoV do not cause life-threatening disease. However, owing to zoonotic origin of SARS-CoV-1 and SARS-CoV-2, humans lack natural immunity making them aggressively pathogenic [20]. Lacking of this pre-existing immunity, called ‘herd immunity’ is an important reason why naive humans have a much delayed time to develop adaptive immune responses against SARS-CoV-2.

Moreover, SARS-CoV-2 has an incubation period of 2–14 days, which is higher than that of MERS-CoV (2–7 days) and SARS-CoV-1 (2–7 days) [21,22]. The symptoms include fever, cough and breathlessness, which may manifest from mild pneumonia to severe illness [22,23]. Nearly 80% of COVID-19 cases remain asymptomatic or show very mild and self-recovering symptoms and about 15% cases show high fever, pneumonia and breathlessness, whereas up to 5% develop respiratory or multiorgan failure and death [22]. Also, COVID-19 patients with diabetes or hypertension have significantly increased expression of cellular ACE2 receptors, putting them on high risk of mortality [24]. Clinical studies have shown that COVID-19 patients with severe pneumonia may rapidly progress to acute respiratory stress syndrome, septic shock or multiorgan failure and deaths [24]. Nonetheless, unlike SARS-CoV-1 and MERS-CoV, the precise mechanism of modulation of host innate immune responses and severe pathogenesis by SARS-CoV still remains elusive [21]. Moreover, digestive symptom and liver inflammations are also reported in hospitalized COVID-19 patients, which are attributed to cytotoxic T cells and Kupffer cells activities [25–30]. Unlike SARS and MERS cases, cardiac disease, arrhythmia and hypertension have been observed twice as much among COVID-19 critical patients [31,32].

Although, the clinical manifestations of COVID-19 are well understood now, the mechanism(s) underlying its high infection rate and pathogenicity is hitherto not clearly established. Several recent studies have reported comparative sequence analysis revealing some important aspects of specific mutations among SARS-CoV-2 isolates from different geographical regions [33–38]. In this report, we, therefore, analyzed the available human and nonhuman CoV genome and protein sequences to have an insight into its high transmissibility and disease severity.

Materials & methods
Data collection
A structured online literature search for peer-reviewed preprint and published articles was conducted on the PubMed, Europe PMC, Medline and Google Scholar portals, using phrases: ‘coronavirus’, ‘SARS-CoV’, ‘MERS-CoV’, ‘SARS-CoV-2’ and ‘bat SARS-like CoV’ as well as ‘COVID-19 infection’, ‘transmission’, ‘pathogenesis’, ‘clinical manifestations’, ‘CoV RNA’ and ‘protein sequences’ etc. The representative genome and protein sequences of various CoV isolates from humans, bats, civets and pangolin were retrieved from NCBI GenBank (www.ncbi.nlm.nih.gov/sars-cov-2) as well as GISAID (www.gisaid.org) database (Table 1).

Multiple sequence analysis
Translation of protein sequence from cDNA sequence was performed using the sequence analysis application MacVector. Genome and proteins/ORFs sequence alignment was analyzed using multiple sequence alignment
Table 1. List of representative human and nonhuman coronavirus isolates used for sequence alignments.

| Species | Coronavirus | Isolates          | Year   | GenBank accession n |
|---------|-------------|-------------------|--------|---------------------|
| Human   | SARS-CoV-2  | Wuhan Hu-1        | 2020   | NC_045512.2         |
|         | SARS-CoV-2  | USA-WA-1          | 2020   | MN985325.1          |
|         | SARS-CoV-1  | Urbani            | 2002–2003 | AY278741.1         |
|         | SARS-CoV-1  | Tor2              | 2002–2003 | AY274119.3         |
|         | SARS-CoV-1  | G003T10013        | 2003–2004 | AYS25636.1         |
|         | H-CoV-OC43  | UK/London/2011    | 2011   | KU131570.1          |
|         | MERS-CoV    | EMC-2012          | 2011   | NC_019843.3         |
| Civet   | SARS-CoV    | SZ3               | 2003   | AY304486.1          |
|         | SARS-CoV    | Civet007          | 2004   | AY572034.1          |
| Bat     | SL-CoV      | ZXC21             | 2015   | MG772934.1          |
|         | SL-CoV      | WIV16             | 2013   | KT444582.1          |
|         | SL-CoV      | RaTG13            | 2013   | MN996532.1          |
| Pangolin| SL-CoV      | MP789             | 2020   | MT084071.1          |

Table 2. Comparative sequence analysis of different coronavirus genomes (A) and proteins (B, C and D), showing percent identity between sequences.

A. RNA genome

| SARS-CoV-2 (Hu-1) | SARS-CoV-1 (Urbani) | SL-CoV (Civet-SZ3) | SL-CoV (Bat-RaTG13) | H-CoV-OC43 | MERS-CoV |
|-------------------|---------------------|-------------------|---------------------|------------|----------|
| SARS-CoV-2 Wuhan Hu-1 | –                   | 79%               | 79%                 | 96%        | 51%      |
| SARS-CoV-1 Urbani | 79%                 | –                 | 99%                 | 94%        | 50%      |

B. Proteins

| Orf1a/1b | Spike (S) | 3a | Envelop (E) | Membrane (M) | Nucleocapsid (N) |
|----------|-----------|----|-------------|---------------|------------------|
| SARS-CoV-2 vs CoV-1 | 86%       | 76% | 72%         | 94%            | 90%              |

C. ‘S’ protein

| Extracellular domain | Cytoplasmic domain | TM | RBD | RBM | Overall |
|----------------------|--------------------|----|-----|-----|---------|
| SARS-CoV-2 vs CoV-1  | 75%                 | 97% (mut: n = 1) | 90% (mut: n = 1) | 73% | 50%     | 76%     |

D. ‘S’ protein

| SARS-CoV-2 (2002–2003) | SARS-CoV-1 (2003–2004) | Civet-SL-CoV (SZ3) | Bat-SL-CoV-RaTG13 | MERS-CoV | H-CoV-OC43 |
|------------------------|------------------------|--------------------|--------------------|----------|------------|
| SARS-CoV-2             | 76%                    | 76%                | 75%                | 97%      | 26%        |
| SARS-CoV-1             | –                      | 98%                | 98%                | 96%      | 27%        |

FP: Fusion peptide; mut: Mutation; RBD: Receptor-binding domain; RBM: Receptor-binding motif; TM: Transmembrane domain.

Results

Comparison of genome sequences of SARS-CoV-2 with other CoV

In the present study we compared the genome and protein sequences of SARS-CoV-2, SL-CoV, MERS CoV and other HCoV. Our comparative analysis showed no similarity of SARS-CoV-2 with MERS-CoV and the common cold causing HCoV-OC43. The SARS-CoV sequence from horseshoe bats, civets and humans had strong similarity in RNA sequences (Table 2A). Interestingly, genome analysis of the reference Wuhan Hu-1 and USA WA-1 sequences revealed up to 87 and 96% identity with bat-SL-CoV-ZXC21 and bat-SL-CoV-RaTG13 isolates, respectively. This indicated that SARS-CoV-2 had a separate line of evolution from bat-SL-CoV or bat-like mammals as compared with SARS-CoV-1.

Mutational analysis of SARS-CoV-2 S protein

Amino acid sequence alignment of SARS-CoV-2 with SARS-CoV-1 showed most mutations acquired in the S protein (Table 2B). Interestingly, the RBM residues located within RBD extracellular domain showed the most
variability, suggesting it a ‘hotspot’ of rigorous mutations (Table 2C). Similar to genome sequences, the SARS-CoV-2 S protein had the most identical sequences with those of bat-SL-CoV (Table 2D). One of the unique features of SARS-CoV-2 S protein is the presence of a furin-like cleavage site (Figure 1) with insertions of proline–arginine–arginine–alanine (PRRA) residues, not reported in HCoV or bat-SL-CoV isolates. In line with this, we also did not observe this furin-like cleavage site in any of the known CoV sequence (Figure 1). However, whether the ‘PRRA’ insertion would have evolved in bats or another intermediate mammal, like civet or pangolin remains another area of further investigation.

Mutational analysis of ORF3a/3b

Unlike other Beta-CoVs, SARS-CoVs express unique ORFs where ORF3a is the largest. Our sequence analysis revealed SARS-CoV-2 encoded 274 residues long 3a as a major mutational ‘hotspot’ with only 72% similarity between SARS CoV-1 (Table 2B). We also observed mutations leading to deletion of the 3a diacidic motif (EXD) in SARS-CoV-2. Notably, a significant truncation in ORF3b (also called orf4 elsewhere) due to introduction of multiple stop codons was also observed. Therefore, the truncated 3b significantly differentiated SARS-CoV-2 from other Beta-CoVs (Figure 2). Taken together, our analysis has highlighted some of the interesting features of SARS-CoV-2 that may be crucial for its pathogenicity and evolution.

Discussion

Approximately 80% of viruses that infect humans are zoonotic, which are initially ill adapted in a new host, slowly replicated and inefficiently transmitted [39]. Therefore, their ‘animal-to-human’ and ‘human-to-human’ transmission greatly depend on their evolution to virulent strains that can well adapt to human hosts. RNA viruses, due to the high replication-fidelity rate (∼10^{-4} error/site/cycle) of their RNA polymerase, are more genetically diversified than DNA viruses [1]. In addition to this, post-transcriptional nucleotide modifications, genetic reassortment or virus–host recombination may further lead to the establishment of stable strains or lineages in human populations [1]. In a recent analysis, the most prevalent A→G mutation observed in SARS-CoV-2 RNA are suggested to be caused by the host RNA-deamination mechanism [40]. In view of this, both viral RNA replication errors and host RNA-modification system have significant impact on mutation rates toward host adaptation and pathogenicity.

Previously, the phylogenetic analysis has shown very close similarity of SARS-CoV-2 genome (∼96% identity) and S protein (∼80% identity) with bat-SL-CoV (ZXC21 and ZC45) [41–44]. Notably therein, while the ‘S1’
Comparative sequence analysis of SARS-CoV-2 suggests its high transmissibility & pathogenicity

Research Article

SARS-CoV-2 RNA genome

Figure 2. Genome organization of the SARS-CoV-2. The open reading frames (ORF1a and 1b) encoding two large polyproteins, pp1a and pp1b, structural proteins that includes spike (S), envelope (E), membrane (M) and nucleocapsid (N) and accessory proteins (not drawn to scale). Compare with the accessory protein ‘3b’, which is truncated in SARS-CoV-2.

subunit is highly variable (~70% similarity), ‘S2’ sequences are conserved and shares ~99% identity with both bat-SL-CoV and SARS-CoV-1 [44,45]. Notably, five critical amino acids in the RBD differ between SARS-CoV-2 and SARS-CoV-1, suggesting strong binding of SARS-CoV-2 S protein with ACE2 receptor and high infectivity [46,47]. Our data showed insignificant similarity of SARS-CoV-2 with MERS-CoV and HCoV-OC43. Interestingly, the two SARS-CoV2 isolates (Wuhan Hu-1 and USA WA-1) revealed up to 87 and 96% sequence identity, respectively with bat-SL-CoV. This indicated that SARS-CoV-2 could have a separate line of evolution from bat-SL-CoV or bat-like mammals (e.g., civet and pangolin) as compared with SARS-CoV-1. Notably, SARS-CoV-1 has been already detected in masked-palm or gem-faced civet cats, which are commonly sold at Chinese wildlife/wet markets [14,48]. In line with this, during the early phase of the SARS-CoV-1 outbreak in China over 40% of the infected individuals were associated with wildlife market or restaurant workers [10]. Likewise, some of the initial cases of SARS-CoV-2 infection were also suggested to be linked to a Wuhan wet market, indicating its possible transmission from bats, civets or pangolins to humans [49]. Our data showed most mutations were acquired in the SARS-CoV-2 S proteins, especially in the RBM, suggesting its enhanced binding to ACE2 as compared with SARS-CoV-1. In addition, we did not observe furin-like cleavage site in any of the known CoV, except in SARS-COV-2, as reported elsewhere [50,51]. Cleavage of SARS-CoV-2 S protein by furin leads to its open conformation that gives advantage of enhanced binding to host ACE2 receptors [51]. Moreover, neuropilin-1 that binds to furin and cleaves substrates has been reported to significantly improve the ACE2-mediated entry of SARS-CoV-2 [52]. Because the role of furin-like proteases in virus entry is a characteristic of pathogenic flu viruses [53], its presence in SARS-CoV-2 strongly supports its high transmissibility and pathogenicity in humans.

The recent comparative sequence analysis of S, E, M, N and ORF8 proteins of several SARS-CoV-2 isolates of 15 countries with the reference Wuhan Hu-1 has revealed substitutions and/or deletion in S protein at 13 sites, substitutions at three sites in N protein and one substitution in M protein [54]. Another sequence analysis has identified frequent mutations in S, N, ORF1ab and ORF8 proteins in 20 SARS-CoV-2 isolates and evaluated their potential in protein stability and possible functional consequences [34]. Of these, co-occurrences of some mutations across different proteins have suggested their structural and/or functional interactions among other viral proteins, and their involvement in virus adaptability and enhanced transmission. Notably, analysis of structural stability of S protein mutants has further indicated the viability of specific variants that could be more prone to their global distribution, temporally and spatially [54]. Sequence analysis of Russian SARS-CoV-2 isolates, including those from other countries, has revealed a set of seven common mutations in S and N proteins, suggesting their multiple
import to Russia, local circulation and varying patterns of spread [36]. Meta-analysis of SARS-CoV-2 isolates within the USA has also identified over 900 unique variants in at least three samples [59]. These included 487 missense and 348 synonymous mutations, four in-frame deletions, five stop codon insertions/deletions and 66 intergenic recombinations. In another study, though diversity of SARS-CoV-2 strains seemed to emerge globally, there was no geographical clustering observed, which suggested their multiple introductions [37]. Interestingly however, their 5’ terminal sequences were more variable as compared with 3’ termini, indicating S, E, ORF1ab and ORF3a as key drivers of diversity, notably RBD as mutational hotspot.

Previous study on SARS-CoV-1 has revealed the presence of a ‘EXD’ motif within the internalization motif of ORF3a that regulates the surface expressions, interaction and internalization of 3a and S proteins [56]. Our sequence analysis has shown mutations leading to deletion of ‘EXD’ in SARS-CoV-2 3a. Notably, the SARS-CoV-2 ORF3b was found to be truncated due to introduction of multiple stop codons, which significantly differentiated it from other Beta-CoVs that code for eight accessory proteins. In line with the reported SARS-CoV-1 truncated 3a activity [57], a recent study has shown the interference of SARS-CoV-2 3b with host interferon system [58]. Because 3a has been reported to modulate IL-2 promoter and interfere with interferon signaling [59], it would be interesting to study the consequence of such observed mutations in clinical settings.

Moreover, SARS-CoV-2 ORF3b and ORF8 has been found to induce a strong antibody response in the early and late phase of infection [60]. In SARS-CoV-2 ORF8, a substitution (Leu→Ser) in 23 isolates has been observed, suggesting their high impact on protein functionality and pathogenesis [37]. Moreover, a proposed phylogenetic tree with representative HCoV and bat-CoV has identified at least two hypervariable hotspots in ORF8 protein, one of which showing a Leu→Ser substitution [33]. Notably, analysis of new SARS-CoV-2 isolates in Italy has reported no evidence for the putative 382-nucleotide deletion in ORF8 as reported in Singapore [58,61]. Taken together, our analysis has endorsed some of the interesting features of SARS-CoV-2 that may be crucial for its pathogenicity and high transmission in humans.

In addition to acquired mutations, differential host factors, such as age, health, physiology, nutritional status, past-exposure, travel history, co-infections, immune-competence, comorbidities and genetics significantly determine the susceptibility to a novel virus [62]. A recent phylogenetic analysis of SARS-CoV-2 sequences from Hong Kong has shown their linkage to European isolates [63]. Interestingly therein, despite insignificant variations between their genomes, they had different clinical presentations, suggesting a more important role of host factors in pathogenicity than mutations. New viruses introduced to humans may further evolve into more aggressive strains as seen for SARS-CoV-2. Herein, the intricate ‘host–pathogen–environment’ interplay is very important in the understanding of the evolution and adaptation of such novel viruses [1]. In view of this, while the emergence of SARS-CoV-2 in naive regions is caused primarily by human movement, local emergence is driven by a combination of environmental and socio-traditional changes [62]. Notably, viral transmission rates are often higher in dense than in sparse populations and social contacts greatly enhance their human-to-human spread. Very interestingly, a recent analysis of SARS-CoV-2 genomic data has revealed its accelerated and high transmission in Italy because of ‘air pollution’ measured with days exceeding the limits set for PM10 (particulate matter 10 μm) [64]. In particular, hinterland cities with average set limits along with low wind speed had a very high infection rates compared with coastal cities with high wind speed. This study suggested the accelerated ‘polluted air-to-human’ transmission dynamics of SARS-CoV-2 [64]. Moreover, inhalations of aerosolized or splattered infectious virus particles have been implicated in a community spread. Previously, a study on hundreds of residents of a housing society in Hong Kong showed that the building’s faulty drainage significantly contributed to the aerosolization and respiratory spread of fecal SARS-CoV-1 [65]. In view of this, shedding of infectious SARS-CoV-2 in stool and urine of COVID-19 patients warrants the risk of its ‘waterborne or fecal–oral’ as well as ‘airborne or respiratory’ transmission [66]. Further, the respiratory, flu or pneumonia viruses, including certain HCoV survive in cold seasons and gradually wane with a rise in temperature. Nonetheless, although the emergence of SARS-CoV-2 in colder weather indicated for its plausible seasonality, arrival of summer did not affect the infection rate [62].

The current surveillance of transmission dynamics of infectious pathogens is mainly based on reproduction number ($R_0$) and fatality rates, which has been adopted as a real-time monitoring of COVID-19 pandemic [62]. Few mathematical–computational models that combined a framework for host, epidemiological and molecular data for SARS-CoV-2 have demonstrated understanding of patterns of evolution, global spread and country-by-country distribution [67,68]. However, due to the rapid increase in RNA sequencing data, mutation rates differ from viral protein-to-protein and study-to-study. A recently proposed mathematical model has highlighted the emergence phenomena of SARS-CoV-2 and the effects of evolutionary adaptations on spreading processes [69].
Another such model offers quantification (Index $c$: contagions) of the environmental risk of exposure to future COVID-19 epidemics in a given region [70]. These theoretical models could be helpful in formulating a proactive epidemiological and environmental strategy in the prevention of such pandemics.

**Conclusion**

Mutations acquired by SARS-CoV-2 during ‘human-adaptation’ and ‘human-to-human’ spread could provide insights into its transmission dynamics that together with clinical and epidemiological data can predict disease prognosis. In view of this, our genome and protein sequence analysis of SARS-CoV-2 has revealed several novel mutations, the most important ones in the ACE2 receptor binding-motif and generation of a furin-like cleavage site in the spike protein. This suggests high infectivity of SARS-CoV-2 in humans through enhanced cell attachment and facilitated entry. Observed mutations within the replicase protein may be crucial for the enhanced replication of the viral genome. In addition, mutations within the accessory proteins (3a, 3b etc.) could have significant roles in evading or modulating host innate immune system and sustaining virus replication. Nonetheless, the consequence of such mutations on virus infectivity and tissue-tropism remain to be studied in animal models. Since COVID-19 is spread even during the asymptomatic phase of the disease, it will be interesting to study the replication of the virus in early phase and how the innate and adaptive immune system responds to the life cycle of SARS-CoV-2, associated with high transmission and pathogenesis. Nonetheless, a larger sample size, including the recently emerged SARS-CoV-2 mutant strains and a rigorous analysis using more advanced tools would further enhance our knowledge on the subject.

**Summary points**

- The underlying mechanisms of high transmission rate and pathogenicity of SARS-CoV-2 remain poorly understood.
- Our comparative sequence analysis of SARS-CoV-2 and other CoVs identifies unique mutations in spike, ORF1a/b, ORF3a/3b and ORF8.
- Its most conserved E, M, and N protein sequences have, however, undergone fewer mutations.
- The most crucial mutations are in the spike ACE2 binding domain and creation of a furin-like cleavage site as well as deletion of ORF3b.
- Sequence analysis reveals that SARS-CoV-2 has diverged from SARS-CoV-1 but is most close to bat-SL-CoV.
- The roles of unique mutations are, therefore, envisaged in the high transmission and pathogenicity SARS-CoV-2.

**Author contributions**

K Padhan and MK Parvez contributed in conceptualization, methodology, software, data analysis and manuscript writing. MS Al-Dosari contributed in data analysis, manuscript writing and editing.

**Acknowledgments**

The authors thank the Deanship of Scientific Research, King Saud University for funding.

**Financial & competing interests disclosure**

The authors have received a grant from the Deanship of Scientific Research, King Saud University (Grant no: RG-1435-053). The authors have no other relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript apart from those disclosed.

No funded writing assistance was utilized in the production of this manuscript.

**References**

1. Parvez MK, Parveen S. Evolution and emergence of pathogenic viruses: past, present, and future. *Intervirology* 60(1-2), 1–7 (2017).
2. WHO. Summary of probable SARS cases with onset of illness. 1 November 2002 to 31 July 2003. www.who.int/csr/sars/country/table2004_04_21/en/
3. Ren LL, Wang YM, Wu ZQ et al. Identification of a novel coronavirus causing severe pneumonia in human: a descriptive study. *Chin. Med. J. (Engl).* 133(9), 1015–1024 (2020).
4. Wu F, Zhao S, Yu B et al. A new coronavirus associated with human respiratory disease in China. *Nature* 579, 265–269 (2020).
5. Chen Y, Liu Q, Guo D. Emerging coronaviruses: genome structure, replication, and pathogenesis. *J. Med. Virol.* 92(4), 418–423 (2020).
6. Grein J, Ohmagari N, Shin D et al. Compassionate use of remdesivir for patients with severe Covid-19. N. Engl. J. Med. 382(24), 2327–2336 (2020).
7. Jeyanathan M, Alkhami S, Small F, Miller MS, Litchy BD, Xing Z. Immunological considerations for COVID-19 vaccine strategies. Nat. Rev. Immunol. 20(10), 615–632 (2020).
8. Parvez MK, Padhan K. Current advances in novel SARS-CoV-2 disease (COVID-19) treatment and intervention strategies. Coronaviruses 2, 1–6 (2021).
9. Leung DT, Tam FC, Ma CH et al. Antibody response of patients with severe acute respiratory syndrome (SARS) targets the viral nucleocapsid. J. Infect. Dis. 190(2), 379–386 (2004).
10. Ge XY, Li JL, Yang XL et al. Are patients with hypertension and diabetes mellitus at increased risk for COVID-19 infection? Lancet 395(10223), 497–506 (2020).
11. Grein J, Ohmagari N, Shin D et al. Compassionate use of remdesivir for patients with severe Covid-19. N. Engl. J. Med. 382(24), 2327–2336 (2020).
12. Parvez MK. Gastrointestinal and hepatobiliary manifestations of COVID-19: potential implications for healthcare resource-deficient countries. Gastroenterol. Hepatol. Lett. 2, 7–11 (2020).
13. Gu J, Han B, Wang J. COVID-19: gastrointestinal manifestations and potential fecal–oral transmission. Gastroenterology 158(6), 1518–1519 (2020).
14. Zhao J, Zhu Y, Luo MY et al. The genome and proteome of SARS-CoV-2. Science 367(6483), 1986–1993 (2020).
15. Zhang C, Shi L, Wang FS. Liver injury in COVID-19: management and challenges. Lancet Gastroenterol. Hepatol. 5(5), 428–430 (2020).
16. Adams DH, Hubscher SG. Systemic viral infections and collateral damage in the liver. Am. J. Pathol. 168(4), 1057–1059 (2006).
17. Parvez MK. Gastrointestinal and hepatobiliary manifestations of COVID-19: potential implications for healthcare resource-deficient countries. Gastroenterol. Hepatol. Lett. 2, 7–11 (2020).
18. Liu Y, Yang Y, Zhang C et al. Clinical and biochemical indexes from 2019-nCoV infected patients linked to viral loads and lung injury. Sci. China Life Sci. 63(3), 364–374 (2020).
19. Fang L, Karakiulakis G, Roth M. Are patients with hypertension and diabetes mellitus at increased risk for COVID-19 infection? Lancet Respir. Med. 8(4), e21 (2020).
20. Higny J, Feye F, Foret F. COVID-19 pandemic: overview of protective-ventilation strategy in ARDS patients. Acta Clin. Belg. 2020, 1–3 (2020).
21. Huang C, Wang Y, Li X et al. Clinical features of patients infected with 2019 novel coronavirus in Wuhan, China. Lancet 395(10223), 497–506 (2020).
22. Kakodkar P, Kaka N, Baig MN. A comprehensive literature review on the clinical presentation, and management of the pandemic coronavirus disease 2019 (COVID-19). Cureus 8(4), e21 (2020).
23. van Doremalen N, Bushmaker T, Morris DH et al. Aerosol and surface stability of SARS-CoV-2 as compared with SARS-CoV-1. N. Engl. J. Med. 383(12), 1564–1567 (2020).
24. Wang HYF, Lam HYS, Fong AH et al. Frequency and distribution of chest radiographic findings in COVID-19 positive patients. Radiology 296(2), E72–E78 (2019).
25. Laha S, Chakraborty J, Das S, Manna SK, Biwas S, Charterjee R. Characterizations of SARS-CoV-2 mutational profile, spike protein stability and viral transmission. Infect. Genet. Evol. 85, 104445 (2020).
26. Higny J, Feye F, Foret F. COVID-19 pandemic: overview of protective-ventilation strategy in ARDS patients. Acta Clin. Belg. 2020, 1–3 (2020).
27. Ceraolo C, Giorgi FM. Genomic variance of the 2019-nCoV coronavirus. J. Med. Virol. 92(5), 522–528 (2020).
28. Parvez MK. COVID-19 and coronaviral hepatitis: evidence of collateral damage. Acta Clin. Belg. 2020, 1–3 (2020).
29. Parvez MK. Gastrointestinal and hepatobiliary manifestations of COVID-19: potential implications for healthcare resource-deficient countries. Gastroenterol. Hepatol. Lett. 2, 7–11 (2020).
30. Liu Y, Yang Y, Zhang C et al. Clinical and biochemical indexes from 2019-nCoV infected patients linked to viral loads and lung injury. Sci. China Life Sci. 63(3), 364–374 (2020).
31. Fang L, Karakiulakis G, Roth M. Are patients with hypertension and diabetes mellitus at increased risk for COVID-19 infection? Lancet Respir. Med. 8(4), e21 (2020).
32. Higny J, Feye F, Foret F. COVID-19 pandemic: overview of protective-ventilation strategy in ARDS patients. Acta Clin. Belg. 2020, 1–3 (2020).
33. Ceraolo C, Giorgi FM. Genomic variance of the 2019-nCoV coronavirus. J. Med. Virol. 92(5), 522–528 (2020).
34. Laha S, Chakraborty J, Das S, Manna SK, Biwas S, Charterjee R. Characterizations of SARS-CoV-2 mutational profile, spike protein stability and viral transmission. Infect. Genet. Evol. 85, 104445 (2020).
Benvenuto D, Demir AB, Giovanetti M, Ciccozzi M, Cassone A. Evidence for mutations in SARS-CoV-2 Italian isolates potentially affecting virus transmission. J. Med. Virol. 92(10), 2232–2237 (2020).

Kozlovskaya L, Piniaeva A, Ignatyev G et al. Isolation and phylogenetic analysis of SARS-CoV-2 variants collected in Russia during the COVID-19 outbreak. Int. J. Infect. Dis. 99, 40–46 (2020).

Sheikh JA, Singh J, Singh H et al. Emerging genetic diversity among clinical isolates of SARS-CoV-2: lessons for today. Infect. Genet. Evol. 84, 104330 (2020).

Licastro D, Rajasekharan S, Dal Monego S, Curcio F, de Rosa R. Isolation and full-length genome characterization of Sarscov-2 from covid-19 cases in northern Italy. J. Virol. 94, e00543–20 (2020).

Cleaveland S, Laurenson MK, Taylor LH. Diseases of humans and their domestic mammals: pathogen characteristics, host range and the risk of emergence. PLoS. Trans. R. Soc. Lond. B Biol. Sci. 356, 991–999 (2011).

Li Y, Yang X, Wang N et al. SNPs or RNA modifications? Concerns on mutation-based evolutionary studies of SARS-CoV-2. PLoS ONE 15, e0238490 (2020).

Gorbunova AE, Baker SC, Baric RS et al. The species Severe acute respiratory syndrome-related coronavirus: classifying 2019-nCoV and naming it SARS-CoV-2. Nat. Microbiol. 5, 536–544 (2020).

Kumar S, Maurya VK, Prasad AK, Bhart MLB, Saxena SK. Structural, glycosylation and antigenic variation between 2019 novel coronavirus (2019-nCoV) and SARS coronavirus (SARS-CoV). ViruDis. 31, 13–21 (2020).

Chan JF, Kok KH, Zhu Z et al. Genomic characterization of the 2019 novel human-pathogenic coronavirus isolated from a patient with atypical pneumonia after visiting Wuhan. Emerg. Microbes Infect. 9, 221–236 (2020).

Walls AC, Park YJ, Tortorici MA, Wall A, McGuire AT, Veesler D. Structure, function, and antigenicity of the SARS-CoV-2 spike glycoprotein. Cell 181, 281–292 (2020).

Coutard B, Vallee C, de Lambarlere X, Canard B, Seidah NG, Decroly E. The spike glycoprotein of the new coronavirus 2019-nCoV contains a furin-like cleavage site absent in CoV of the same clade. Antiviral Res. 176, 104’742 (2020).

Wan Y, Shang J, Graham R, Baric RS, Li F. Receptor recognition by the novel coronavirus from Wuhan: an analysis based on decade-long structural studies of SARS coronavirus. J. Virol. 94, e00127–20 (2020).

Wrapp D, Wang N, Corbett KS et al. Cryo-EM structure of the 2019-nCoV spike in the prefusion conformation. Science 367, 1260–1263 (2020).

Guan Y, Zheng BJ, He YQ et al. Isolation and characterization of viruses related to the SARS coronavirus from animals in southern China. Science 302, 276–278 (2003).

Li Q, Guan X, Wu P et al. Early transmission dynamics in Wuhan, China, of novel coronavirus-infected pneumonia. N. Engl. J. Med. 382, 1199–1207 (2020).

Tai W, He L, Zhang X et al. Characterization of the receptor-binding domain (RBD) of 2019 novel coronavirus: implication for development of RBD protein as a viral attachment inhibitor and vaccine. Cell Mol. Immunol. 17, 613–620 (2020).

Walls AC, Park YJ, Tortorici MA, Wall A, McGuire AT, Veesler D. Structure, function, and antigenicity of the SARS-CoV-2 spike glycoprotein. Cell 181, 281–292 (2020).

Cantuti-Castelvetri L, Ojha R, Pedro LD et al. Neuropilin-1 facilitates SARS-CoV-2 cell entry and infectivity. Science 370, 856–860 (2020).

Klenk HD, Garten W. Host cell proteases controlling virus pathogenicity. Trends Microbiol. 2, 39–4 (1994).

Shah A, Rashid F, Aziz A, Jan AU, Suleman M. Genetic characterization of structural and open reading Fram-8 proteins of SARS-CoV-2 isolates from different countries. Gene Rep. 21, 100886 (2020).

Shen L, Dien Bard J, Biegel JA, Judkins AR, Gai X. Comprehensive genome analysis of 6,000 USA SARS-CoV-2 isolates reveals haplotype signatures and localized transmission patterns by state and by country. Front. Microbiol. 11, 573430 (2020).

Tan YJ. The Severe Acute Respiratory Syndrome (SARS)-coronavirus 3a protein may function as a modulator of the trafficking properties of the spike protein. Virol. J. 2, 5 (2020).

Varshney B, Agnihotram S, Tan YJ, Baric R, Lal SK. SARS coronavirus 3b accessory protein modulates transcriptional activity of RUNX1b. PLoS ONE 7, e29542 (2012).

Mimakshi R, Padhan K, Rani M, Khan N, Ahmad F, Jameel S. The SARS Coronavirus 3a protein causes endoplasmic reticulum stress and induces ligand-independent downregulation of the type 1 interferon receptor. PLoS ONE 4, e8342 (2009).

Konno Y, Kimura I, Uriu K et al. SARS-CoV-2 ORF3b is a potent interferon antagonist whose activity is increased by a naturally occurring elongation variant. Cell Rep. 32, 108185 (2020).

Hachim A, Kavian N, Cohen CA et al. ORF8 and ORF3b antibodies are accurate serological markers of early and late SARS-CoV-2 infection. Nat. Immunol. 21, 1293–1301 (2020).

Su YCF, Anderson DE, Young BE. Discovery and genomic characterization of a 382-nucleotide deletion in ORF7b and ORF8 during the early evolution of SARS-CoV-2. mBio 11, e01610–01620 (2020).
62. Parvez MK, Jagirdar RM, Purty RS et al. COVID-19 pandemic: understanding the emergence, pathogenesis and containment. World Acad. Sci. J. 2, 18 (2020).

63. Au CH, Chan WS, Lam HY et al. Genome sequences of SARS-CoV-2 strains detected in Hong Kong. Microbiol. Resour. Announc. 9, e00697–20 (2020).

64. Coccia M. Factors determining the diffusion of COVID-19 and suggested strategy to prevent future accelerated viral infectivity similar to COVID. Sci. Total Environ. 729, 138474 (2020).

65. Yu IT, Li Y, Wong TW et al. Evidence of airborne transmission of the severe acute respiratory syndrome virus. N. Engl. J. Med. 350, 1731–1739 (2004).

66. Parvez MK. Gut feeling: the plausible fecal-oral transmission route of Covid19. J. Infect. Dis. Epidemiol. 6, 141–143 (2020).

67. Anderson RM, Fraser C, Ghani AC et al. Epidemiology, transmission dynamics and control of SARS: the 2002–2003 epidemic. Philos. Trans. R. Soc. Lond. B Biol. Sci. 359, 1091–1105 (2004).

68. Hufnagel L, Brockmann D, Geisel T. Forecast and control of epidemics in a globalized world. Proc. Natl Acad. Sci. USA 101, 15124–15129 (2004).

69. Eletreby R, Zhuang Y, Carley KM, Yagan O, Poor HV. The effects of evolutionary adaptations on spreading processes in complex networks. Proc. Natl Acad. Sci. USA 117, 5664–5670 (2020).

70. Coccia M. An index to quantify environmental risk of exposure to future epidemics of the COVID-19 and similar viral agents: theory and practice. Environ. Res. 191, 110155 (2020).