Variants Within over a Hundred Complete COVID-19 Genomes and the Impact on Health Security

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Abstract: 102 complete COVID-19 genomes have been collected from the viral genomes database to track and characterize novel variants. The data were treated bioinformatically so that 172 variants, with 127 unique and 45 polymorphic variants were found. The 127 unique variants consist of 76 missense, 39 synonymous, 6 non-coding, 5 deletions and 1 insertion. The 45 polymorphic variants consist of 25 missense, 15 synonymous, 4 non-coding and 1 in-frame-deletion. Most common variants are 28144T>C (33 missense), 8782C>T (31 synonymous), followed by missense 11083G>T (11 samples), 18060C>T (9 samples) and 26144G>T (7 samples). L3606F, S5932F and L84S are the amino acid changes in the last three common variants. Most variants were found in ORF1ab gene within the region encoded for domains (nsp4 and nsp6) and in the coding ORF8 gene. The variant 28144T>C could be among the main enhancers of viral transmission. There is a tendency for a national specificity of the most recorded variants. The virus outbreak could be between countries or dependent on the place of origin. Reasonable evidence of Chinese origin of the virus could be possible and thus more genomes should be collected and analyzed to understand the origin and the reason for its outbreak. This could support human health security by either finding out suitable vaccines or managing health precautionary measures.

Keywords: Coronavirus, Genome, Variants, Mutation, Health Security

Introduction

Since the virus is like SARS-CoVs, the International Committee on Taxonomy of Viruses (ICTV) termed the newly discovered Coronavirus (2019-nCoV) SARS-CoV-2 (Cascella et al., 2020). It was first reported in Wuhan, the largest metropolitan area in China on December 31st, 2019. Therefore, the (WHO, 2020) acronymized this 2019 disease as “COVID-19”. CoVs can infect different species including birds, mice, bats, livestock and humans (Wang et al., 2006; Ge et al., 2013; Chen and Guo, 2016), with a serious infection in humans, causing pneumonia.

Systematically, CoVs belong to the subfamily Orthocoronavirinae of the family Coronaviridae and the order Nidovirales. Coronavirinae contains four genera: Alphacoronavirus, Betacoronavirus (4 subgenera) (Chan et al., 2013), Gammacoronavirus and Deltacoronavirus (Chen et al., 2020). Bats and rodents are the probable source of alpha and Betacoronaviruses, whereas birds could be the source of Gamma and delta viruses (Cascella et al., 2020). Over 210,000 COVID-19 complete genomes have so far been sequenced with an approximate genome length of 30000 bases. As fast as the virus spreads, more variants accumulate with the possibility of further emerging virulent strains. The virus genome is found in a single-stranded positive-sense RNA (+ssRNA) (Perlman and Netland, 2009) acquiring 5′-cap structure and 3′-poly-A tail (Chen et al., 2020). As a severe global health threat, CoVs outbreaks are, most probably, unavoidable in the future. There is an urgent need, thus, to identify the possible variants in the 102 sequenced genomes that might support in understanding the reasons for the virus outbreak and in producing an effective therapy and vaccine against the virus.

Materials and Methods

The COVID-19 page (https://bigd.big.ac.cn/ncov) available in China’s National Genomics Data Center (NGDC) was accessed on the 27th of September 2020.
From 2019-nCoV genomes available on this page and in three Genbank databases, 102 publicly complete genomes have been collected and are listed in Table 1. As has been previously used (Koyama et al., 2020; Matsuda et al., 2020), the two longest and identical NC_045512 and MN908947 sequences (29903 bp) were used as reference genomes to coordinate the differences in the ribosomal slippage of the 102 used genomes. The genome data were checked for the accuracy of the alignments and the aligned data could be obtained from the author upon request. Since genomes acquire differences in their start and end points, their lengths have been adjusted to the lengths of NC_045512 and MN908947 and their variants were numbered according to the positions of these two genomes. All genomes were first aligned to each other using BioEdit Sequence Alignment Editor (Hall et al., 2011) and the FASTA file of the aligned data was executed to MacClade v.4.10 program (Maddison and Maddison, 2002) and manually aligned to the reference genomes. Using MacClade v.4.10, mutations were recorded and data were transferred to the Paup v. 4.0b10 (Swofford, 2002) program for phylogenetic analysis.

After deleting ambiguous, gap-containing sites and N or mixed bases sites, the remaining 29532 sites were analyzed by Maximum-Parsimony (MP), Neighbor-Joining (NJ) and Maximum-Likelihood (ML) methods in Paup v. 4.0b10. For MP, heuristic searches of 10 random stepwise additions were conducted by Tree Bisection Reconnection (TBR) branch swapping and 1000 bootstrap replications. For NJ analysis (Saitou and Nei, 1987; Tamura and Nei, 1993) distance option and 1000 bootstrap replications were used. As the MT226610 sequence was rapidly evolved, it was used as an outgroup. For ML, heuristic searches by axis additions and Nearest-Neighbor Interchange (NNI) branch-swapping were adjusted. Other conditions for the ML analysis like gamma shape parameter of 1.5214 and 4 rate categories were also adjusted. The substitution rate matrix of the data was as follow: R(a) = 0.1571, R(b) = 0.7500, R(c) = 0.2121, R(d) = 0.6049, R(e) = 2.6308 and R(f) = 1.00. Likelihood settings from best-fit model (GTR+I+G) were selected by AIC in Modeltest Version 3.06.

Table 1: Organization of the reference + ssRNA genome used in this study. Nsp = non structural protein; ORF = Open Reading Frame

| Gene   | Position | Product                      | Notes                                      |
|--------|----------|------------------------------|--------------------------------------------|
| 5'UTR  | 1-265    | 5'UTR                        | cap                                        |
| Orf1ab | 266-805  | nsp1                         | Leader protein                             |
|        | 806-2719 | nsp2                         | Transmembrane domain 2                     |
|        | 2720-8554| nsp3                         | 3C-like protease                           |
|        | 8555-10054| nsp4                        | Putative transmembrane domain              |
|        | 10055-10972| nsp5                       | ssRNA-binding protein                      |
|        | 10973-11842| nsp6                       | Growth-factor-like protein                 |
|        | 11843-12901| nsp7                       |                                            |
|        | 12092-12685| nsp8                       |                                            |
|        | 12686-13042| nsp9                       |                                            |
|        | 13025-13441| nsp10                      |                                            |
|        | 13442-13480| nsp11                      |                                            |
|        | 13476-13503| Stem_loop                  | Frameshifting stimulation element         |
|        | 13488-13542| Stem_loop                  | Frameshifting stimulation element         |
|        | 13442-13468, 13468-16235| nsp12                  | RNA-dependent RNA polymerase              |
|        | 16237-18093| nsp13                    | Helicase                                   |
|        | 18040-19620| nsp14                    | 3'-to-5' exonuclease                       |
|        | 19621-20658| nsp15                    | endoRNAse                                  |
|        | 20659-21552| nsp16                    | O-ribose methyltransferase                 |
| S      | 21563-25384| spike glycoprotein        | Surface glycoprotein                       |
| ORF3a  | 25393-26220| ORF3a protein             |                                            |
| E      | 26245-26472| envelope protein           |                                            |
| M      | 26523-27191| membrane glycoprotein     |                                            |
| ORF6   | 27202-27387| ORF6 protein              |                                            |
| ORF7a  | 27394-27759| ORF7a protein             |                                            |
| ORF7b  | 27756-27887| ORF7b                     |                                            |
| ORF8   | 27894-28259| ORF8 protein              |                                            |
| N      | 28274-29533| nucleocapsid phosphoprotein| ORF9 protein                             |
| ORF10  | 29558-29674| ORF10 protein             |                                            |
| ORF10  | 29609-29644| --                         | 3'UTR pseudoknot stem-loop 1              |
| ORF10  | 29629-29657| --                         | 3' UT pseudoknot stem-loop 2              |
| 3'UTR  | 29675-29903| 3'UTR                      |                                            |
Results and Discussion

The COVID-19 genome is approximately 30,000 bp. It contains within its 5′-proximal two-thirds, nonstructural protein coding regions (nsp1-nsp16; replicase genes) encoded on orf1ab gene. Within the 3′-proximal one-third of the genome, structural proteins and nonessential accessory protein coding regions are encoded (ORF3-ORF10, S, E, M, N) (Paul, 2006). The 5′ UTR (265 b) comprises the genome cap and the 3′ UTR (~300 b) forms the tail (Table 1). The recorded variants were counted between positions 33 and 29830 to avoid base bias. The mixed bases were not considered in the analysis. 172 total variants are found, 127 unique and 45 polymorphic (Table 2). The unique variants (Table 3) are 121 substitutions (76 missense, 39 synonymous and 6 non-coding), 5 deletions (2 in-frame-deletions, 2 out-of-frame-deletions and 1 non-coding) and 1 insertion. The polymorphic variants (Table 4) are 45 substitutions (25 missense, 15 synonymous, 4 non-coding and 1 in-frame-deletion) (Fig. 1).

In agreement with previous studies (Koyama et al., 2020; Matsuda et al., 2020), most common variants were 8782C>T (ORF1ab) and 28144T>C (ORF8) in 31 samples followed by 11083G>T (nsp6) in 11 samples and 18060C>T (nsp14) in 9 samples. The occurrences of 8782C>T and 28144T>C concur and most of the other common variants are subsets of these most common ones. 8782C>T is synonymous; however, 11083G>T (L3606F) and 18060C>T (S5932F) and 28144T>C (L84S) are missense causing amino acid changes in nsp6, nsp14 and ORF8, respectively. The variant 28144 T>C which exhibited an amino acid change in ORF8 protein from Leucine to Serine, was indicated in the polypeptide involved in enhancing virus transition from bat to human (Chen et al., 2020; Nguyen et al., 2020). As this variant was recorded in thirteen North Americans, ten central Asians (Chinese including Wuhan residents), three Japanese, one Taiwanese, one Indian and one Spanish, it could be among the main enhancers of viral transmission.

Table 2: Statistics of the 172 recorded mutations in the 102 COVID-19 genomes

| Variants     | Substitution | Deletion |
|--------------|--------------|----------|
|              | Missense     | Synonymous| Non-coding | In-frame-deletion | Out-of-frame deletion | Non-coding | Insertion |
| Unique       | 76           | 39        | 6         | 2               | 2                   | 1         | 1         |
| Polymorphic  | 25           | 15        | 4         | 1               | -                   | -         | -         |

Table 3: Unique monomorphic variants (substitution, deletion and insertion), their positions, number of cases and country distribution found in the 102 CoVs genomes

| #   | Variant | Type       | Amino acid change | Accession number, country |
|-----|---------|------------|-------------------|---------------------------|
| 1   | 35A>T   | Non-coding |                   | MT163716, USA             |
| 2   | 36C>T   | Non-coding |                   | MT163716, USA             |
| 3   | 75C>A   | Non-coding |                   | MT049951, CHN             |
| 4   | 186C>T  | Non-coding |                   | MT093631, CHN             |
| 5   | 382A>T  | Synonymous |                   | LC521925, JP              |
| 6   | 359-382 | In-frame-deletion | G32_V39   | LC521925, JP              |
| 7   | 490T>A  | Synonymous |                   | MT044257, USA             |
| 8   | 514T>C  | Synonymous |                   | MT188340, USA             |
| 9   | 654G>A  | Synonymous |                   | MT123293, CHN             |
| 10  | 686-694 | In-frame-deletion | K141_F143 | MT044258, USA             |
| 11  | 1102C>T | Synonymous |                   | MT188339, CHN             |
| 12  | 1348C>T | Synonymous |                   | GWHACDD01000001, PAK      |
| 13  | 1385C>T | Missense   | H374Y             | MT159720, USA             |
| 14  | 1397G>A | Synonymous |                   | GWHACDD01000001, PAK      |
| 15  | 1548G>A | Missense   | S428N             | MN994467, USA             |
| 16  | 1912C>T | Synonymous |                   | LC521925, JP              |
| 17  | 2091C>T | Missense   | T609I             | MT027064, USA             |
| 18  | 2269A>T | Synonymous |                   | MT066156, ITA             |
| 19  | 2277T>C | Missense   | I671T             | MT012098, IND             |
| 20  | 2446T>C | Synonymous |                   | MT163716, USA             |
| 21  | 2717G>A | Missense   | G818S             | MT093571, SWE             |
| 22  | 2971G>T | Missense   | M902I             | MT049951, CHN             |
| 23  | 3037C>T | Synonymous |                   | MT192765, USA             |
| 24  | 3177C>T | Missense   | P971L             | MT044257, USA             |
| 25  | 3259G>T | Missense   | T998G             | MT159707, USA             |
| 26  | 3411C>T | Missense   | E1049V            | MT163716, USA             |
| 27  | 3738C>T | Missense   | P1158L            | MT159705, USA             |
| 28  | 3792C>T | Missense   | E1176V            | LC522973, JP              |
| No. | Mutation   | Type       | Reference ID | Location |
|-----|------------|------------|--------------|----------|
| 29  | 4288G>T    | Synonymous | MT226610, CHN |          |
| 30  | 4307A>C    | Missense   | MT226610, CHN |          |
| 31  | 5572G>T    | Synonymous | MT163716, USA |          |
| 32  | 5784C>T    | Missense   | MT152824, USA |          |
| 33  | 5845A>T    | Missense   | MT159715, USA |          |
| 34  | 6031C>T    | Synonymous | MT039890, SKorea |       |
| 35  | 6035A>G    | Missense   | MT188341, USA |          |
| 36  | 6363C>T    | Missense   | MT159712, USA |          |
| 37  | 6636C>T    | Missense   | MT184910, USA |          |
| 38  | 6968C>A    | Missense   | MT123293, CHN |          |
| 39  | 6996T>C    | Missense   | MT123293, CHN |          |
| 40  | 7479A>G    | Missense   | MT118835, USA |          |
| 41  | 9034A>G    | Synonymous | MT066176, TWN |          |
| 42  | 9157T>C    | Missense   | MT184910, USA |          |
| 43  | 9274A>G    | Synonymous | MT093571, SWE |          |
| 44  | 9474C>T    | Missense   | MT159706, USA |          |
| 45  | 9491C>T    | Missense   | MT066176, TWN |          |
| 46  | 9561C>T    | Missense   | MT012098, IND |          |
| 47  | 9924C>T    | Missense   | MT012098, IND |          |
| 48  | 10036C>T   | Synonymous | MT184911, USA |          |
| 49  | 10507C>T   | Synonymous | MT159708, USA |          |
| 50  | 11207G>C   | Missense   | MT184911, USA |          |
| 51  | 11237T>G   | Synonymous | MT226610, CHN |          |
| 52  | 11557G>T   | Missense   | MT159712, USA |          |
| 53  | 11750C>T   | Missense   | MT184910, USA |          |
| 54  | 11956C>T   | Synonymous | MT159712, USA |          |
| 55  | 12041G>C   | Missense   | MT123293, CHN |          |
| 56  | 12115C>T   | Synonymous | MT159708, USA |          |
| 57  | 12160G>C   | Synonymous | MT159708, USA |          |
| 58  | 12202G>C   | Synonymous | MT159708, USA |          |
| 59  | 12208G>T   | Synonymous | MT159708, USA |          |
| 60  | 12355G>C   | Missense   | MT159708, USA |          |
| 61  | 12378G>A   | Missense   | MT184911, USA |          |
| 62  | 12464G>T   | Missense   | MT159712, USA |          |
| 63  | 12467G>T   | Missense   | MT159712, USA |          |
| 64  | 12491C>T   | Synonymous | MT159712, USA |          |
| 65  | 12514G>C   | Synonymous | MT159712, USA |          |
| 66  | 12534C>T   | Missense   | MT159712, USA |          |
| 67  | 12572G>T   | Missense   | MT159712, USA |          |
| 68  | 12578G>T   | Missense   | MT159712, USA |          |
| 69  | 12582G>T   | Missense   | MT159712, USA |          |
| 70  | 12608G>T   | Missense   | MT159712, USA |          |
| 71  | 12660G>C   | Missense   | MT159712, USA |          |
| 72  | 12685G>C   | Missense   | MT159712, USA |          |
| 73  | 12773G>T   | Missense   | MT159712, USA |          |
| 74  | 12793G>T   | Missense   | MT159712, USA |          |
| 75  | 13072C>T   | Synonymous | MT159712, USA |          |
| 76  | 13225C>G   | Synonymous | MT159712, USA |          |
| 77  | 13226T>C   | Missense   | MT093571, SWE |          |
| 78  | 14657C>T   | Missense   | MT012098, IND |          |
| 79  | 15597T>C   | Missense   | MT039890, SKorea |       |
| 80  | 15607T>C   | Synonymous | MT093571, SWE |          |
| 81  | 16467A>G   | Missense   | MT188341, USA |          |
| 82  | 17000C>T   | Missense   | MT159715, USA |          |
| 83  | 17247T>C   | Missense   | MT126808, BRA |          |
| 84  | 17376A>G   | Missense   | MT093571, SWE |          |
| 85  | 18512C>T   | Missense   | LC521925, JP |          |
| 86  | 18603T>C   | Missense   | MT106054, USA |          |
| 87  | 18814C>T   | Synonymous | MT192765, USA |          |
### Table 3: Continue

| # | variant | Type   | aa change | gene          | Accession number (country)          |
|---|---------|--------|-----------|---------------|-------------------------------------|
| 89| 18975T>A| Missense| F6237Y    | MT106054, USA |                                     |
| 90| 19065T>C| Missense| L6267P    | MT007544, AUSTR|                                     |
| 91| 19175A>C| Missense| M6304L    | MT106054, USA |                                     |
| 92| 19610C>T| Missense| T6449R    | MT123291, CHN |                                     |
| 93| 20281T>C| Synonymous|           | MT163719, USA |                                     |
| 94| 20299-30301| Out-of-frame deletion| N6678N (nsp15) | MT039887, USA |                                     |
| 95| 20670G>A| Missense| R6802H    | NMDC60013002-10, Wuhan |                                     |
| 96| 20679G>A| Missense| R6805Q    | NMDC60013002-10, Wuhan |                                     |
| 97| 20936C>T| Missense| R6891C    | MT039890, SKorea|                                     |
| 98| 20980G>C| Missense| Q6905M    | MT226610, CHN |                                     |
| 99| 21147T>C| Missense| L6961S    | MT188339, CHN |                                     |
|100| 21647T>A| Missense| Y23N      | MT049951, CHN |                                     |
|101| 21707C>T| Missense| H44Y      | MT027064, USA |                                     |
|102| 21784T>A| Missense| F152L     | MT226610, CHN |                                     |
|103| 21386-21388| Insertion| S7041     | MT188341, USA |                                     |
|104| 21997-21999| Out-of-frame deletion| Y144Y (S gene) | MT012098, IND |                                     |
|105| 22033C>A| Missense| G176V     | MT159716, USA |                                     |
|106| 22104G>T| Synonymous|           | MT184910, USA |                                     |
|107| 22224C>G| Missense| S216F     | MT039890, SKorea|                                     |
|108| 22303T>G| Missense| S242R     | MT007544, AUSTRIA|                                     |
|109| 22432C>T| Synonymous|           | MT049951, CHN |                                     |
|110| 22785G>T| Missense| R403M     | MT012098, IND |                                     |
|111| 23185C>T| Synonymous|           | MT188341, USA |                                     |
|112| 23403A>G| Synonymous|           | MT192765, USA |                                     |
|113| 23955T>G| Missense| F792Y     | MT093571, SWE |                                     |
|114| 25775G>T| Missense| C120L     | MT039890, SKorea|                                     |
|115| 26354T>A| Missense| L22G      | MT039890, SKorea|                                     |
|116| 27493C>T| Missense| P61S      | NMDC60013002-09, Wuhan |                                     |
|117| 27925C>T| Missense| T18M      | MT106054, USA |                                     |
|118| 28253C>T| Synonymous|           | NMDC60013002-09, Wuhan |                                     |
|119| 28409C>T| Missense| P47S      | MT159718, USA |                                     |
|120| 28792A>T| Synonymous|           | MN994467, USA |                                     |
|121| 28878G>A| Missense| S203N     | MT106052, USA |                                     |
|122| 28916G>A| Missense| G216S     | MT188339, USA |                                     |
|123| 29230C>T| Synonymous|           | MT159720, USA |                                     |
|124| 29301A>T| Missense| D344V     | MT135043, CHN |                                     |
|125| 29705G>T| Non-coding|           | LC522973, JP |                                     |
|126| 29742G>A| Non-coding|           | MT106052, USA |                                     |
|127| 29750-29759| Non-coding del|          | MT007544 (AUSTRIA) |                                     |

### Table 4: Polymorphic variants (substitutions), their positions, number of cases and country distribution found in the 102 CoV's genomes

| # | variant | Type         | aa change | gene          | Accession number (country)          |
|---|---------|--------------|-----------|---------------|-------------------------------------|
| 1 | 241C>T  | Non-coding   | 5' UTR    | GWHACDD010000001(PAK), MT192765(USA) |                                     |
| 2 | 254C>T  | Non-coding   | 5' UTR    | MT184910, MT184908 (USA) |                                     |
| 3 | 508-522 | In-frame-deletion G82_V86 orf1ab | MT044258, MT159716 (USA) |                                     |
| 4 | 614G>A   | Missense     | A117T     | MT027062, MT027063 (USA) |                                     |
| 5 | 1691A>G  | Missense     | I476V     | MT027063 (USA), MT050493 (IND) |                                     |
| 6 | 2662C>T  | Synonymous   |           | LC522973-LC522975 (JP) |                                     |
| 7 | 3099C>T  | Missense     | D954I     | MT159717, MT184912 (USA) |                                     |
| 8 | 4402T>C  | Synonymous   |           | MT135041-MT135044 (CHN) |                                     |
| 9 | 5062G>T  | Missense     | L1599A    | MT135041-MT135044 (CHN) |                                     |
|10 | 5084A>G  | Missense     | I1607V    | MT027062, MT027063 (USA) |                                     |
|11 | 6501C>T  | Missense     | P2079L    | MT027063 (USA), MT050493 (IND) |                                     |
|12 | 6819G>T  | Missense     | S2185I    | MT123293, MT123291(China) |                                     |
|13 | 8782C>T  | Synonymous   |           | MN938384, MN975262, MT049951, MT123292, MT226610, MT135041-4, GWHACDD0100000014(CHN), MN985325, MN997409, MT020880, MT020881, MT044257, MT106052, MT106054, MT152824, MT163717-19, MT188339, MT188341 (USA), MT066175 (TAW), LC522973-LC522975 (JP), MT050493, MT050493 (IND), MT198651, MT198652 (ESP) |                                     |
|14 | 9477T>A  | Missense     | F3071Y    | MT198651, MT198652 (ESP) |                                     |
|15 | 10232C>T | Missense     | R3323C    | MT192772, MT192773 (Vietnam) |                                     |
|16 | 11083G>T | Missense     | L3606F    | LC528252-33 (JP), MT126808 (BRA), MT163716, MN997409, MT184910-13 (USA), MT226610 (CHN), GWHACDD010000001 (PAK) |                                     |
Table 5: Length, country and accession number of COVID-19 genome sequences used in this study

| #  | Country       | Length | Database | Accession number            |
|----|---------------|--------|----------|------------------------------|
| 1  | Wuhan         | 29903  | Genbank  | MN908947                     |
| 2  | China         | 29838  | Genbank  | MN938384                     |
| 3  | China         | 29891  | Genbank  | MN975262                     |
| 4  | USA           | 29882  | Genbank  | MN985325                     |
| 5  | USA           | 29882  | Genbank  | MN988713                     |
| 6  | USA           | 29882  | Genbank  | MN994467                     |
| 7  | USA           | 29882  | Genbank  | MN994468                     |
| 8  | USA           | 29882  | Genbank  | MN997409                     |
| 9  | Finland       | 29806  | Genbank  | MT020781                     |
| 10 | USA           | 29882  | Genbank  | MT020880                     |
| 11 | USA           | 29882  | Genbank  | MT020881                     |
| 12 | Japan         | 29848  | Genbank  | LCS21925                     |
| 13 | USA           | 29882  | Genbank  | MT020706                     |
| 14 | USA           | 29882  | Genbank  | MT020703                     |
| 15 | USA           | 29882  | Genbank  | MT020704                     |
| 16 | Taiwan        | 29870  | Genbank  | MT066175                     |
| 17 | Japan         | 29878  | Genbank  | LCS22973                     |
| 18 | Japan         | 29878  | Genbank  | LCS22974                     |
| 19 | Japan         | 29878  | Genbank  | LCS22975                     |
| 20 | Japan         | 29878  | Genbank  | LCS22972                     |
| 21 | USA           | 29879  | Genbank  | MT039887                     |
| 22 | USA           | 29858  | Genbank  | MT044258                     |
| 23 | USA           | 29882  | Genbank  | MT044257                     |
| 24 | South Korea   | 29903  | Genbank  | MT039890                     |
| 25 | China         | 29903  | Genbank  | MT049951                     |
| 26 | Taiwan        | 29870  | Genbank  | MT066176                     |
| 27 | Nepal         | 29811  | Genbank  | MT072688                     |
| 28 | China         | 29860  | Genbank  | MT093631                     |
| 29 | Sweden | 29886 | Genbank | MT093571 |
| 30 | USA    | 29882 | Genbank | MT106052 |
| 31 | USA    | 29882 | Genbank | MT106053 |
| 32 | USA    | 29882 | Genbank | MT106054 |
| 33 | USA    | 29882 | Genbank | MT118835 |
| 34 | China  | 29882 | Genbank | MT123291 |
| 35 | China  | 29891 | Genbank | MT123290 |
| 36 | Japan  | 29902 | Genbank | MT126808 |
| 37 | Japan  | 29902 | Genbank | MT126823 |
| 38 | USA    | 29878 | Genbank | MT152824 |
| 39 | Brazil | 29876 | Genbank | MT126808 |
| 40 | USA    | 29903 | Genbank | MT163716 |
| 41 | China  | 29903 | Genbank | MT135041 |
| 42 | China  | 29903 | Genbank | MT135042 |
| 43 | China  | 29903 | Genbank | MT135043 |
| 44 | China  | 29903 | Genbank | MT135044 |
| 45 | USA    | 29897 | Genbank | MT163717 |
| 46 | USA    | 29903 | Genbank | MT163718 |
| 47 | USA    | 29903 | Genbank | MT163719 |
| 48 | India  | 29851 | Genbank | MT050493 |
| 49 | India  | 29854 | Genbank | MT012098 |
| 50 | USA    | 29882 | Genbank | MT159717 |
| 51 | USA    | 29882 | Genbank | MT159718 |
| 52 | USA    | 29882 | Genbank | MT159719 |
| 53 | USA    | 29882 | Genbank | MT159720 |
| 54 | USA    | 29882 | Genbank | MT159721 |
| 55 | USA    | 29882 | Genbank | MT159722 |
| 56 | USA    | 29882 | Genbank | MT159705 |
| 57 | USA    | 29882 | Genbank | MT159706 |
| 58 | USA    | 29882 | Genbank | MT159710 |
| 59 | USA    | 29882 | Genbank | MT159707 |
| 60 | USA    | 29882 | Genbank | MT159708 |
| 61 | USA    | 29882 | Genbank | MT159709 |
| 62 | USA    | 29882 | Genbank | MT159711 |
| 63 | USA    | 29882 | Genbank | MT159712 |
| 64 | USA    | 29882 | Genbank | MT159713 |
| 65 | USA    | 29882 | Genbank | MT159714 |
| 66 | USA    | 29882 | Genbank | MT159715 |
| 67 | USA    | 29882 | Genbank | MT159716 |
| 68 | China  | 29923 | Genbank | MT123292 |
| 69 | China  | 29871 | Genbank | MT123293 |
| 70 | Italy  | 29867 | Genbank | MT066156 |
| 71 | Japan  | 29903 | Genbank | MT159705 |
| 72 | USA    | 29882 | Genbank | MT184907 |
| 73 | USA    | 29880 | Genbank | MT184908 |
| 74 | USA    | 29882 | Genbank | MT184909 |
| 75 | USA    | 29882 | Genbank | MT184910 |
| 76 | USA    | 29882 | Genbank | MT184911 |
| 77 | USA    | 29882 | Genbank | MT184912 |
| 78 | USA    | 29882 | Genbank | MT184913 |
| 79 | USA    | 29783 | Genbank | MT188339 |
| 80 | USA    | 29845 | Genbank | MT188340 |
| 81 | USA    | 29835 | Genbank | MT188341 |
| 82 | USA    | 29829 | Genbank | MT192765 |
| 83 | Taiwan | 29862 | Genbank | MT192759 |
| 84 | Vietnam| 29891 | Genbank | MT192772 |
| 85 | Vietnam| 29891 | Genbank | MT192773 |
| 86 | Spain  | 29611 | Genbank | MT198651 |
| 87 | Spain  | 29782 | Genbank | MT198652 |
| 88 | China  | 29899 | Genbank | MT226610 |
| 89 | Pakistan| 29836 | Genome Warehouse | GWHACDD01000001 |
| 90 | Wuhan  | 29899 | Genome Warehouse | GWHABKF0000000001 |
Table 5: Continue

| 91  | Wuhan  | 29889 | Genome Warehouse | MT044258-USA CGTGCTTTTG GAGACTCCGT GGAGGAGGTC TTCATCAGAGG CACGTCACA |
|-----|--------|-------|-----------------|----------------------|
| 92  | Wuhan  | 29890 | Genome Warehouse | MT188341-USA CGTGCTTTTG GAGACTCCGT GGAGGAGGTC TTCATCAGAGG CACGTCACA |
| 93  | Wuhan  | 29891 | NMDC            | MT159716-USA CGTGCTTTTG GAGACTCCGT GGAGGAGGTC TTCATCAGAGG CACGTCACA |
| 94  | Wuhan  | 29890 | NMDC            | LC521925-JP CGTGCTTTTG GAGACTCCGT GGAGGAGGTC TTCATCAGAGG CACGTCACA |
| 95  | Wuhan  | 29891 | NMDC            |                        |
| 96  | Wuhan  | 29896 | NMDC            |                        |
| 97  | Wuhan  | 29891 | NMDC            |                        |
| 98  | Australia | 29893 | Genbank        | MT007544 |
| 99  | Wuhan  | 29903 | Genbank        | NC_045512 |
| 100 | China  | 29871 | Genbank        | MN996530 |
| 101 | China  | 29894 | Genbank        | MN996528 |
| 102 | India  | 29874 | Genbank        | MT050943 |

354 403
MT044258-USA CGACCTCA--------------------------TGAGCTG GTACGCAAGAC TCGAGGCCAT
MT188341-USA GCACCTCATG GTCACTTTA ATGCATTGCAG TCGAGGCCAT
MT159716-USA GCACCTCA--------------------------TGAGCTG GTACGCAAGAC TCGAGGCCAT
LC521925-JP GCACCTCATG GTCACTTTA ATGCATTGCAG TCGAGGCCAT

500 549
MT044258-USA GCACCTCA--------------------------TGAGCTG GTACGCAAGAC TCGAGGCCAT
MT188341-USA GCACCTCATG GTCACTTTA ATGCATTGCAG TCGAGGCCAT
MT159716-USA GCACCTCA--------------------------TGAGCTG GTACGCAAGAC TCGAGGCCAT
LC521925-JP GCACCTCATG GTCACTTTA ATGCATTGCAG TCGAGGCCAT

600 699
MT044258-USA AAAAGAGGCTG TTGCCTTAGT TTACGGCGCC GATCTA--------------------------GACTT
MT188341-USA AAAAGAGGCTG TTGCCTTAGT TTACGGCGCC GATCTAAAGT CATTGACTT
MT159716-USA AAAAGAGGCTG TTGCCTTAGT TTACGGCGCC GATCTAAAGT CATTGACTT
LC521925-JP AAAAGAGGCTG TTGCCTTAGT TTACGGCGCC GATCTAAAGT CATTGACTT

(A) 3 In-frame deletions at positions: C359 - T382, T508 - T532 & A686 - T694

2061 20310
MT050943-IND TAGAATTAGC TATGGATGAA TTCACTTGAAG GTATAAACAT AGAAGGCTAT
MT012098-IND TAGAATTAGC TATGGATGAA TTCACTTGAAG GTATAAACAT AGAAGGCTAT
MT044258-USA TAGAATTAGC TATGGATGAA TTCACTTGAAG GTATAAACAT AGAAGGCTAT
MT039887-USA TAGAATTAGC TATGGATGAA TTCACTTGAAG GTATAAACAT AGAAGGCTAT

19251 22000
MT050943-IND AAGTCTGTGA ATTTCAATTT TGAATATGGCG CATTGGGGG TGGTTATTAC
MT012098-IND AAGTCTGTGA ATTTCAATTT TGAATATGGCG CATTGGGGG TGGTTATTAC
MT044258-USA AAGTCTGTGA ATTTCAATTT TGAATATGGCG CATTGGGGG TGGTTATTAC
MT039887-USA AAGTCTGTGA ATTTCAATTT TGAATATGGCG CATTGGGGG TGGTTATTAC

(B) 2 Out-of-frame deletions in MT039887-USA & MT012098-IND

21350 21399
MT044258-USA GAGGGAATAC AATCCCAATT CAGTTGCTTCT CATTT---C TTTTTATGAC
MT188341-USA GAGGGAATAC AATCCCAATT CAGTTGCTTCT CATTT---C TTTTTATGAC
MT159716-USA GAGGGAATAC AATCCCAATT CAGTTGCTTCT CATTT---C TTTTTATGAC
LC521925-JP GAGGGAATAC AATCCCAATT CAGTTGCTTCT CATTT---C TTTTTATGAC

(C) 1 Insertion in MT188341-USA

Fig. 1: In-frame, out-of-frame deletions and an insertion in USA, Japanese and Indian variants
Fig. 2: NJ tree constructed by using 29532 sites of the collected genomes. Bootstrap values are shown at nodes whenever they are above 50%
Fig. 3: ML phylogenetic tree constructed by the dataset using the modeltest GTR + I + G
For the 101 missense variants, 80 variants are found in the longest ORF1ab gene distributed in the cleaved nonstructural proteins (NSP1-NSP16). However, more variants are found in the structural protein genes (S, ORF3a and N). MT226610-CHN was the fastest evolving strain as it exhibited 29 substitutions. One of the out-of-frame-deletions is found close to 3’end of nsp15 protein of an Indian strain and the other one is found close to 5’end of S protein of the USA strain (Fig. 1). The first mutation probably did not alter the O-ribose methyltransferase (nsp16) since it is located at the end of the gene, while the second could alter the post-translational spike, glycoprotein. This mutation may increase disease susceptibility (Zimmerman et al., 1997) or stop protein function indicating that it is not necessary for efficient viral transmission. It is not known that S deletion enhances virulence or transmission rates of the virus and it is not known whether the strain acquiring this deletion could successfully transmit to a new host (Assiri et al., 2016).

Fortunately, this study collected various COVID-19 genomes from the same place of origin as shown in Table 5 (51 genomes from USA, 25 from China (including 10 from Wuhan) and 8 from Japan). This supports that the novel mutations found herein could reflect the diversity of the place of origin rather being acquired during spreading of the infection (Matsuda et al., 2020). It is therefore an indication that stopping virus outbreak is possible in the short-term future. However, the constructed tree (Fig. 2) indicated that viruses from the same country did not form a single group, which suggests that CoVs-19 were introduced to each country several times (Koyama et al., 2020; Matsuda et al., 2020) and it, thus, may be difficult to follow the virus origin. The phylogeny of the maximum-likelihood analysis (Fig. 3) indicated a possible transmission scenario of the virus. The tree referred to Chinese origin of COVID-19 and showed its transmissions to USA, Spain, Japan and India.

Researchers sequenced a lot of SARS-CoV-2 genomes and shared results during the pandemic. The sequenced data allowed public health officials to evaluate the relevant epidemiological parameters such as the reproductive number and virus introduction into new regions. Knowing the possibilities for the outbreak is still managing health precautionary measures which could be conducted in daily life (Hopkins, 2020). Understanding genetic framework of COVID-19 genome enhances WHO’s ability to analyze the risk of the virus introduction into countries and define the response actions and prioritization of resources, as well as the possible capacity to manage the virus outbreak. The implementation of action plans for health security is occurring globally with varied progress rates (Samhouri et al., 2018) and is actively supported by WHO to enhance operational readiness for the virus in countries (Al-Mandhari et al., 2020).

Conclusion

In conclusion, the virus is still considered a threat to human health security as there is lack of knowledge about the origin and the reasons for its outbreak. Chinese origin could be possible. Two debates about the virus outbreak are either the diversity of the place of origin or spreading the infection through individuals’ movements between countries. Emergence of new variants by releasing more genomes could help in clarifying the virus origin, the reasons of its outbreak and the development of vaccines or effective precautions.

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Ethics

This article is original and contains unpublished material.

Conflict of Interest

The author declares that he has no conflict of interest. No ethical approval for this study is needed since it depended on the data deposited in the Genbank database.

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