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CONSTRUCTING THE GENOME OF SARS-COV-2: AN ESSENTIAL STEP TOWARDUNDERSTANDINGTHE COVID-19 OUTBREAK THROUGH PHYLOGENETIC TREES

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Abstract

This paper discusses the emergence of (SARS-CoV-2) is a global concern due to its ability to cause coronavirus disease 2019 (COVID-19). In order better understand this virus and its evolutionary history, a phylogenetic tree was constructed using the Maximum Likelihood Estimate method and a Multiple Sequence Alignment of SARS-CoV-2 and other coronavirus species. This tree will be used to help elucidate the relationship between various coronaviruses and the spread of SARS-CoV-2. This paper made use of 46 SARS-CoV-2 isolates together with two other coronavirus species (Alpha & Beta) retrieved from the GenBank database (National Center for Biotechnology Information) for this exploration. Results indicated that the Maximum Likelihood Estimate approach was successful in producing a reliable phylogenetic tree that displayed the evolutionary history between SARS-CoV-2 and other coronavirus species, as well as delineating the genetic diversity within SARS-CoV-2.

Keywords: SARS-CoV-2 Genome, COVID-19, Maximum-Likelihood, Bioinformatics Tools, Phylogenetic Tree Construction.

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Introduction

An in-depth analysis of the SARS-CoV-2 virus has been conducted with the aim of understanding its evolutionary history. By constructing a phylogenetic tree, scientists have been able to trace the detailed evolution of the virus from its origins to the present day [1]. This study provides an important insight into the spread of the virus, its mutations, and its potential for future mutation [2]. Using genetic sequencing techniques, a clear picture of the virus' genetic makeup has been developed. This data has been used to construct a phylogenetic tree that describes the relationship between different SARS-CoV-2 viruses and provides valuable information about its evolution [3]. The phylogenetic tree has enabled scientists to identify genetic variants and understand the virus' ability to evolve and adapt to its environment [4]. This study provides a comprehensive overview of the virus' evolution and will help researchers in the development of effective treatments and vaccines.

The novel coronavirus pandemic (COVID-19) has changed the world in many ways. Scientists are trying to understand the evolution of the virus that causes this disease, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) [5] [6]. One way to do this is through phylogenetic tree construction [7] [8]. Phylogenetic tree construction is a process used to infer the evolutionary history of a group of related organisms (or viruses) by analyzing their genetic sequences. By constructing a phylogenetic tree, researchers can better understand the evolutionary patterns of SARS-CoV-2 and identify which areas of the world have been most affected by this virus [9]. This information can be used to inform public health strategies and vaccine development [10]. In this paper, we will discuss the basics of phylogenetic tree construction, explain how it is used to study the SARS-CoV-2 virus, and discuss the implications of this paper.

Phylogenetic tree construction involves the analysis of genetic data to create a visual representation of the evolutionary relationships among various organisms. The tree shows how closely related different organisms are and how they have evolved over time [11]. By studying the tree, researchers can gain insights into the transmission and spread of SARS-CoV-2, and identify the genetic changes that have occurred since its emergence.

The tree is then used to estimate how these species are related, this can help us to better understand the virus and its behavior, as well as inform strategies to prevent its spread [12]. Phylogenetic tree construction can be used to follow the development of novel virus strains, and to detect mutations that may be significant for its progression. Maximum likelihood approaches can be used to construct phylogenetic trees, and this approach can be used to examine SARS-CoV-2's evolution. [13].

MATERIAL AND METHODS

Data collection

NCBI's (National Center for Biotechnology Information) database was used to divide the entire genome from gene bank of (46) into (18) Alpha and (28) Beta coronaviruses [14].

Genomic information is a type of data that pertains to the DNA or genetic

material of an organism. It can include information such as accession numbers, definitions, genome size in base pairs (bp), paper titles, host information [15], and country of origin some of them are shown in the Table (1).

Phylogenetic trees were constructed from complete genome sequences by first aligning the sequences and then analyzing them to determine their relatedness [16]. Alignment of the sequences has been made using multiple sequence alignment, such as R language, to find regions of similarity among the sequences [17]. Once the sequences have been aligned, phylogenetic analysis can be performed using a number of methods [18], such as maximum likelihood. This method uses to infer the evolutionary relationships among the sequences [19]. Once the phylogenetic tree has been built, it can then be used to infer the evolutionary history of the species and identify relationships between them as shown in Figure 1.

Table	1:	Provides	details	on	the	genomes	of	SARS-COV-2	and	two	types	of
Coron	avir	ruses (Alph	ia and B	eta).								

ACCESSION	DEFINITION	Genome size (bp)	Host	Country	Lineage
MW924112.1	Alphacoronavirus	28761	zoonotic bat	Korea	Alpha
MH817484.1	Feline alphacoronavirus	29137	Felis catus	Brazil	Alpha
ОК287352.1	Alphacoronavirus	28435	avian species	Yunnan, China	Alpha
OP646598.1	Severe acute respiratory syndrome coronavirus 2	29684	birds and mammals	China, United Kingdom, Germany, Jordan, South Africa, Saudi Arabia, United Arab Emirates	Beta Sarbecovirus
NC_045512	SARS-CoV-2	29903	Homo sapiens	China	Beta Sarbecovirus
MW626421	Severe acute respiratory syndrome coronavirus 2	29802	Bat, Civet, Homo sapiens	Global	Beta Sarbecovirus

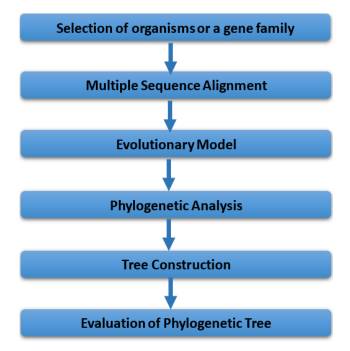


Figure 1: General proceedings for building a Coronavirus phylogenetic tree.

This step involves selecting the organisms or gene family to be included in the phylogenetic analysis [20]. This selection is important as it will determine the type and amount of data that will be used in the analysis, The genomes of alphacoronavirus and betacoronavirus were selected in this paper.

The Coronavirus and SARS-CoV-2 molecular data selection from the NCBI GenBank database provides access to genetic sequence data for the novel coronavirus SARS-CoV-2 and other coronaviruses. It includes the complete genomic sequences for these viruses through:

- 1. NavigatetotheNCBIGenBankhomepage(https://www.ncbi.nlm.nih.gov/genbank/) and select the "Nucleotide" database.
- **2.** Enter the term "coronavirus" in the search box, and select the "All Fields" option.
- **3.** Use the "Organism" and "Gene Family" filters to narrow down your search results to the desired organisms or gene family.
- **4.** Select the "Send to" option for the desired search results, and choose "File", "FASTA" or "GenBank" as the file format.
- **5.** The downloaded file can then be used to analyze the selected organisms or gene family.

The Coronavirus and SARS-CoV-2 molecular data Selection from the NCBI GenBank database is a collection of genetic sequences related to coronaviruses and SARS-CoV-2, the virus responsible for the COVID-19 pandemic. It includes genome sequences and more specialized data such as cDNA sequences, which are transcripts of mRNA sequences that can be used to study gene expression. The data is freely available to the public and is updated regularly [21].

Multiple Sequence Alignment

This step involves aligning the sequences of the selected organisms or gene family to identify similarities and differences in the genetic sequences. This can be done using various software packages such as ClustalW, MUSCLE, and T-Coffee. The goal of MSA is to find the most optimal alignment of the sequences that maximizes the similarity among them. to optimize the alignment of the sequence [22].

And here you will come the process of identifying regions of conserved gene order within the genomes of two different species, it is the "Synteny". It is a term used to describe the arrangement of genes and/or other DNA sequences in the genome that are located on the same chromosome. Before Multiple Sequence Alignment (MSA) used to construct a phylogenetic tree, these sequences must be compared to each other to determine the degree of similarity and evolutionary relatedness [23]. By looking at the synteny of the genes, it is possible to locate conserved blocks of DNA, which can provide evidence of common ancestry. This information can then be used to guide the MSA process, allowing the most accurate phylogenetic tree to be constructed. The FindSynteny() function takes the genomes as input, and it then identifies regions of similarity and rearrangements between the genomes. This is done by comparing and aligning the genomes. The output of the function is a graph showing the synteny blocks, which are regions of conserved gene order between two genomes [24] [25]. The graph also shows any rearrangements or inversions between the two genomes. Generally, synteny matching takes place when two genomes have the same sections. We were able to get the first representation of the synteny of Alpha coronavirus with SARS-CoV-2 Wuhan-Hu-1 (NC_045512) outgroup, which is shown in Figure 2, and Beta coronavirus genomes, shown in Figure 3. We noticed blocks of similar sequences between them. These genomes demonstrate a blend of sequence likeness and divergence, with relatively few recombination events.

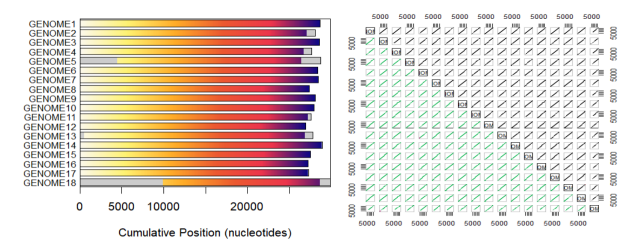


Figure.2. (a) A representation of Alpha coronavirus sequences through adjacent pairs synteny blocks. The blocks are color-coded according to the region they share with the first genome. If the regions are the same, the blocks are the same hue. If they don't share the same region, the block is colored grey. The size of the blocks is proportional to the length of the region they share. This type of representation is useful for visualizing the similarities and differences between different genomes. (b) Dot plot depicting the homologous regions among five Influenza virus A genomes. The dot plots show the similarities between the Alphacoronaviruses. The dot plots also reveal several regions of divergence between the genomes, which likely represent regions of antigenic variability.

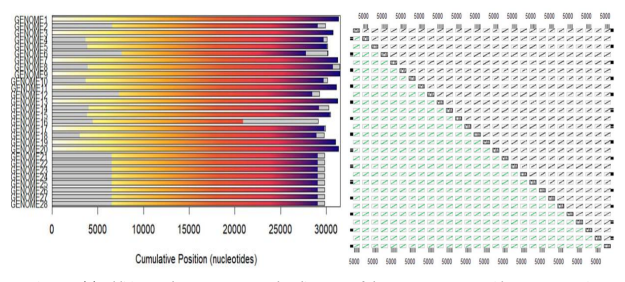


Figure.3. (a) Addition to the synteny maps, the alignment of the two genomes provides a more precise view of their similarity. The alignment can be visualized as a dot plot (b) Dot plot each nucleotide (or amino acid) is represented as a single point in a graph, and the points are connected by black lines if they are in the same location in both genomes. This visualization helps to identify regions of significant similarity, as well as regions of significant difference. The dot plot also allows for the identification of recombinant regions, which are regions of the genome that have been acquired from a different

Evolutionary model

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This step involves selecting a model of evolution that best fits the data. This model will be used to generate a hypothesis of the evolutionary history of the organisms or gene family. Evolutionary tree model selection is an important step in constructing phylogenetic trees for coronaviruses. It involves selecting the appropriate evolutionary tree model to accurately represent the evolutionary relationships between the different coronaviruses. This model selection is based on the alignment of the nucleotide or amino acid sequences of the different coronaviruses. The evolutionary tree model should be chosen based on the characteristics of the sequence data, such as the number of sequence differences, the position of the differences, and the rate of evolution. Once the appropriate evolutionary tree model is chosen which is the GTR-GAMMA model, the phylogenetic tree can be constructed using a variety of software tools that are available. The resulting phylogenetic tree can then be used to identify the evolutionary relationships between the different coronaviruses, which can provide important information about the spread and evolution of the virus [26] [27].

The Maximum Likelihood Estimate (MLE) uses a tree-like structure to represent the evolutionary relationships among the species and calculates the likelihood of the observed data given the model. The model selection is based on the maximization of the likelihood of the observed data given the tree topology. This technique has been used to construct phylogenetic trees for SARS-CoV-2, the virus responsible for the COVID-19 pandemic. MLE is a powerful tool for reconstructing the evolutionary history of any group of organisms, including coronaviruses [28].

Phylogenetic Analysis

This step involves using the evolutionary model and the multiple sequence alignments to infer the phylogenetic relationships among the selected organisms or gene family.

Phylogenetic analysis is a method of analyzing the evolutionary history of organisms. It is used to determine the relationships between different species and to understand the evolutionary processes that have occurred over time. In the case of SARS-CoV-2, phylogenetic analysis is used to construct phylogenetic trees which are diagrams that show the evolutionary relationships between different strains of the virus. Phylogenetic analysis involves collecting data from multiple sources, such as DNA sequences, protein sequences, and gene expression data. This data is then used to create a phylogenetic tree, which is a diagram that shows the evolutionary history of the virus. The phylogenetic tree is then used to compare different strains of the virus and to identify the different lineages of the virus. This information can be used to understand the spread of the virus and to identify which strains may be more dangerous than others. It can also be used to develop treatments and vaccines for SARS-CoV-2 [29].

Tree construction

This step involves constructing a phylogenetic tree based on the inferred relationships from the data obtained from the phylogenetic analysis [7]. The tree is a graphical representation of the evolutionary relationships among the organisms or gene family being studied. Phylogenetic tree construction is a process used to create a visual representation of the evolutionary relationships among different species. This is done by analyzing the genetic sequences of each species and creating a diagram that shows how they are related. The tree is constructed by grouping together species that share more similar genetic sequences, with the most closely related species being grouped together at the tips of the branches. This tree can then be used to study evolutionary processes, such as speciation and adaptation [8] [30].

The tree can also be used to identify the common ancestor of a group of species, and can provide insight into the history and evolution of life on Earth. And then assessing how well the tree matches the original tree. The accuracy of the tree is then assessed using a measure of the tree's robustness, such as bootstrap support. Bootstrapping is a method used to evaluate the accuracy of a phylogenetic tree. It involves randomly sampling the data multiple times to create multiple trees, and then comparing the results. This allows researchers to see how reliable the tree is and determine whether it is accurate enough to make conclusions from. Bootstrapping can also be used to measure the confidence of a particular branch or node in the tree. The higher the bootstrap score, the more reliable the branch or node is considered to be [31]. After the process of Multiple Sequence Aligning as shown in Figure 4:

	and the second	the second s				and the second se
1. NC 048216.1 (NL63-related bat coronavirus strain B9KV/NL63-9b complete genome GT	GGTTGTATTTGTGGTTTGTGA	TAGTOGCTATAATOGA	CTTGTTGCCTAATTTG	TTTAAGTTGAAGGTGTC	TACACACCTITITGATOGTGACAAA!	TTTGTTGGCAC TTTTGAGA
2. NC 046954.1 (Aphacoronavirus Bat-CoV/Pkuhilitaly/3398-19/2015 complete genome GT	GCCTTGTGACTTGGTTTCTTG				GTCACAAC TTTTTGATOGTGATCGG	
3 NC 038861.1 (Transmissible gastroentertils virus complete genome genomic RNA GT	ACGTGATTACCOCATATATCC				AACAAATCTTTTTGAAGGTGATAAA	
4. NC 034972.1 (Coronavirus AcCoV-JC34 complete genome CT	ATOTTOTTOTOGOC TATOTTA	TAATACTCCTTTGTGA	TTGCATGCCTTCTATT.	ACTAAGC TTAAGTTGTC	CACTAATTTGTTGGAGGGGGGGACAAG	TTTGTTGGTAGTTTTGAAC)
5. NC 032730.1 Lucheng Rn rat coronavirus isolate Lucheng-19 complete genome GT	ATGTTATTGTGGGCTATGTAT	TAGTACTOCTATOTO	TTGCATGCCATCTATT	ACCAAGCTTAAGTTGTC	TACCANTITATIGGAAGGCGACAAGI	TTTGTGGGTAGTTTTGAAC
6. NC 032107.1 (NL63-related bat coronavirus strain BBC/NL63-9a complete genome GT	GGC TTTGTGCC TGGTTTTGYI	TTGCCGCACTGTTAGA	GTTGCTACCCAATGTC	TTTAAGTTGAAGATTTC	YACTCAATTGTTTGAAGGTGATAAAT	TTTATTGGWAC TTTTGAGA(
7. NC 030292.1 (Ferret coronavirus isolate FRCoV-NL-2010 complete genome GT	GETTETTTETAGGATATOTEC	TACTOTTTTTCTATO	TTCAATACCATCATTT	TTTAAACTGAAGGTTTC	TACAAACTTATTTGAAGGTGATAAGT	TTCGTAGGTAATTTTGAATC
8 NC 028752.1 Camel alphacoronavirus isolate camel/Ryadh/Ry141/2015 complete genome GT	GECTETETECTTEETACTTT	TTGCTATGTTAACAGO	TCTTTTACCTAGTTTG	TTGAAGC TTAAAGTCTC	GACAAACCTTTTTGAGGGTGACAAA	TTTGTAGGTACATTTGAAA
9 NC 028606 1 (Swine enteric coronavirus strain tas)/213306/2009 complete genome 01	ACGTGATTACCGCATATGTCA	TAGTTTTCCTCTATG	TTCACTCCCTTCACTG	TTTAAAC TTAAAGTTTC	AACAAATCTCTTTGAAGGTGACAAAT	TTTGTGGGTAAC TTTGAAT(
10. NC 028011.1 (BtMr-AlphaCoV/SAX2011 complete genome CT	GECTTATECTCATETATETE	GAGCCTTOGTTCTTGA	ACTTOTTCCCAATTTG	CTAAAAC TTAAAGTTTC	AACTCAATTGTTTGAGGATGGTAAC	TTTATTGGCAAC TTTGAGA(
11. NC 028614 1 (BIRI-AlphaCoV/Mu82013 complete genome 017)	GETTAATCTTTATETATECTT	GTTCTGCTCTTATCGG	TOTTATCCCAAATCTCC	CTCAAACTTAAAGTTAC	TACTCAATTGTTTGATGGTGACAAGT	TTTGTCGGAAATTTTGAAAA
12. NC 028824.1 (BIRT-AlphaCoV//1/2012 complete genome CT	GETTECTTACATEGTATETT	GTGCAGCGCTAATTGC	CCTGATTCCCAATTTG	TTTAAAC TTAAGGTTTC	TACTACITIGITIGAAGGTACIAAA	TTTGTTGGCACGTTTGATG
13. NC 028833.1 (BNv-AlphaCoV/SC2013 complete genome GT	GGCTTCTGAGTTGGTATTTG	TTGCTGGTCTCAGTGG	TGTTATACCATCTTTT	ATCAAGTTGCGTGTCTC	ATCACAGCTTTTTGACGGTGACAAGT	TTTGTCGGTAGTTTTGAAG
14. NC 023760.1 (Mink coronavirus strain WD1127 complete genome GT)	GGATCCTTATTAGTTATTTGT	TAATGTTCTTTTATG	CTETTTACCATETTTE	ITTAAACTAAAAGTCTC	TACACAGTTGTTTGAGGGTGATAAA	ITTGITGGTAATTTTGAGT(
15. LC756668.1 Human coronavirus NL63 HCoVNL63 Fukushima 0230 2018 RNA nearly complete genom CT	GGCTTCTCACATGGTTTAGTT	TTGCTGCATTTTTAGA	GCTTTTACCTAATGTT	TTTAAGTTAAAAATCTC	TACTCAATTGTTTGAAGGTGATAAGT	TTTATAGGTAC TTTTGAGAI
16. ON461751.1 (Human coronavirus 229E isolate 229E 1 partial genome OT	GETTETETECTTEETACTTTE	TTGC TATGTTGACAGO	TTTGTTACCTAGTTTG	TGAAGCTTAAAGTTTC	GACAAATCTTTTCGAAGGTGACAAAT	TTTGTAGGTACATTTGAAA(
17. ON461752.1 (Human coronavirus 229E isolate 229E 2.1 partial genome GT	GETTETETECTTEETACTTTE	TTOCTATOTTGACAGO	TTTGTTACCTAGTTTG	CTGAAGC TTAAAGTTTC	GACAAATCTTTTCGAAGGTGACAAAT	TTTGTAGGTACATTTGAAAL
18. NC 045512.2 Severe acute respiratory syndrome coronavirus 2 isolate Wuhan-Hu-1 complete genomic T	GGATAACAATTGCTTATATCA	TTTGTATTTCCACAAA	GCATTICTATIGGTIC	TTTAGTAATTACC TAAA	GAGACGTGTAGTC TTTAATGGTGTT	TCCTTTAGTACTTTTGAAGA
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		(8	a)			
1. NC 048217.1 (Murine hepatitis virus strain A59 complete genome		GCAGAAGCAAT	TACTOCCCAGATOG		TC TTGGCACAGGGCCCCATGC	TGGAGCCAGTTATGGA
2. NC 045512.2 Severe acute respiratory syndrome coronavirus 2 isolate Wuhan-Hu-1 col		TCAGTCCAAGATGG		CCTAGGAACTGGGCCAGAAGC		
3. NC 006213.1 Human coronavirus OC43 strain ATCC VR-759 complete genome		TOCTOCCACGATGO		TC TGGGAACAGGACCGCATGC		
4. NC 038294.1 (Betacoronavirus England 1 isolate H123990006 complete genome			TGGCTCCCAGGTGG		CACTGGAACTGGACCCGAAGC	
5. NC 039207.1 (Betacoronavirus Erinaceus/VMC/DEU/2012 isolate ErinaceusCoV/2012-17-	4/GER/2012 complete genome	TTAAACAAT	TGGCCCCCAGGTGG			
6. NC 030886.1 (Rousettus bat coronavirus isolate GCCDC1 356 complete genome		TGACCAGTTATG			CACTGTATCTGTG CGAGGG	
7. NC 026011.1 (Betacoronavirus HKU24 strain HKU24-R05005) complete genome			TOCTACCCAGATOG		TCTCGGAACAGGGCCGTATGC	
8. NC 025217.1 (Bat Hp-betacoronavirus/Zhejiang2013 complete genome		TGGACCCGCGCTGG		TETTGGCACAGGCCCTGAG		
9. AC 000192.1 Murine hepatitis virus strain JHM complete genome			TACTOCCCAGATGG		TCTTGGAACAGGGCCCTATGC	TEECOCAGAGTATEEC
10. NC 019843.3 [Middle East respiratory syndrome-related coronavirus isolate HCoV-EMC/	2012 complete genome		TGGCTCCCAGGTGG		CACTOGAACTOGACCCGAAGC	
11. NC 017083.1 Rabbit coronavirus HKU14 complete genome			TOCTOCCACGATOG		TCTGGGAACAGGACCGCATGC	
12. NC 014470.1 (Bat coronavirus BM48-31/BGR/2008 complete genome			TCAGCCCGCGCTGG		TC TAGGAAC TGGACCAGAGGC	the second s
13. NC 012936.1 (Rat coronavirus Parker complete genome		- GCAGAAGCAAT	TACTCCCCAGATGG	T ATTTTTACTA	TCTTGGGACGGGCCCCCATGC	TGGAGCCAGTTTCGGA
14. NC 009019.1 (Tylonycteris bat coronavirus HKU4 complete genome		TTAAACAAT	TGGCTCCCAGATGG		CACTGGGACTGGACCAGAGGC	TAATCTACCTTTTAGA
15. NC 009020.1 (Pipistrellus bat coronavirus HKU5 complete genome		CAAAACCTC	TGGCACCCAGGTGG	T ATTTCTACTA	CACTGGAACAGGACCTGAGGC	AAATCTCCCTTTCAGA
16. NC 009021.1 Rousettus bat coronavirus HKU9 complete genome	TTGATAATAAGG	TTGTTGGTGGTTGC	CAGAACATTAATG.	AGCCCTGCCCAGAACCCCATA	CGGTGTTTCATTTGCA	
17. NC 006577.2 Human coronavirus HKU1 complete genome	- ACAAAAGCAGT	TGTTACCGAGATGG	T - ATTTCTACTA	TCTCGGTACCGGCCCATATGC	CAATGCATCCTATGGT	
18. NC 004718.3 (SARS coronavirus Tor2 complete genome	- AATGAAAGAGC	TCAGCCCCAGATGG	T ACTTCTATTA	C C TAGGAAC TGGCCCCAGAAGC	TTCACTTCCCTACGGC	
19. NC 003045.1 Bovine coronavirus isolate BCoV-ENT complete genome	- CCAGCGTCAAT	TGCTACCACGATGG	T ATTTTTACTA	TC TTGGAACAGGACCGCATGC	CAAAGACCAGTATGGC	
20. NC 001846.1 [Murine hepatitis virus strain MHV-A59 C12 mutant complete genome	- GCAGAAGCAAT	TACTOCCCAGATOG	T ATTTTTACTA	TCTTGGCACAGGGCCCCATGC	TGGAGCCAGTTATGGA	
21. 00504244.1 [Severe acute respiratory syndrome coronavirus 2 isolate SARS-CoV-2/h	- AATGAAAGATC	TCAGTCCAAGATGG	T A T T T C T A C T A	C C TAGGAAC TGGGCCAGAAGC	TGGACTTCCCTATGGT	
22. 00504245.1 (Severe acute respiratory syndrome coronavirus 2 isolate SARS-CoV-2/h	m - AATGAAAGATC	TCAGTCCAAGATGG	T ATTTC TAC TA	CCTAGGAACTGGGCCAGAAGC	TGGACTTCCCTATGGT	
23. 00504246.1 Severe acute respiratory syndrome coronavirus 2 isolate SARS-CoV-2/h	uman/JPN/HiroFH377/2022 complete geno	- AATGAAAGATC	TCAGTCCAAGATGG	T ATTTCTACTA	C C TAGGAAC TGGGC C AGAAGC	TGGACTTCCCTATGGT
24. 00504247.1 (Severe acute respiratory syndrome coronavirus 2 isolate SARS-CoV-2/h	uman/JPN/HiroFH378/2022 complete geno	m - AATGAAAGATC	TCAGTCCAAGATGG	T ATTTC TAC TA	C C TAGGAAC TGGGC CAGAAGC	TGGACTTCCCTATGGT
25. 00504248.1 [Severe acute respiratory syndrome coronavirus 2 isolate SARS-CoV-2/h		- AATCAAACATC	TCAGTCCAAGATGG	TATTTCTACTA	C C TAGGAAC TGGGCCAGAAGC	TGGACTTCCCTATGGT
	uman/JPN/HiroFH380/2022 complete geno	- AAISAASAIV				
26. 00504274.1 Severe acute respiratory syndrome coronavirus 2 isolate SARS-CoV-2/h	the annumentation of the balance of the second strength of the		TCAGTCCAAGATGG		CCTAGGAACTGGGCCAGAAGC	
26. 00504274.1 (Severe acute respiratory syndrome coronavirus 2 isolate SARS-CoV-2/h 27. 00504275.1 (Severe acute respiratory syndrome coronavirus 2 isolate SARS-CoV-2/h	uman/JPN/HiroC429/2022 complete genon	- AATGAAAGATC		T ATTTCTACTA	C C <mark>T A G G A A C T G G G C C A G A A G C</mark> C C <mark>T A G G A A C T G G G C C A G A A G C</mark>	T G G A C T T C C C T A T G G T
the state of the	uman/JPN/HiroC429/2022 complete genom uman/JPN/HiroC430/2022 complete genom	≈ - AA <mark>TGAAAGAT</mark> C ≈ - AA <mark>TGAAAGAT</mark> C	T C A G T C C A A G A T G G T C A G T C C A A G A T G G	T A T T T C T A C T A T A T T T C T A C T A	C C TAGGAAC TGGGCCAGAAGC	TGG <mark>ACTTCCCTATGGT</mark> TGG <mark>ACTTCCCTATGGT</mark>

(b)

Figure.4. (a) Comparing Alpha coronavirus sequences to NC_045512.2 (the SARS-CoV-2 reference sequence) using a multiple sequence alignment (MSA). (b) Multiple sequence alignment of Beta

A phylogenetic tree was built for two families of Coronavirus, alpha and beta, with bootstraps as shown in Figures 5, 6.

Evaluation of phylogenetic tree

This step involves assessing the accuracy of the phylogenetic tree by comparing it to other trees that were constructed using different methods and assessing the support for each branch. This allows for the identification of any potential errors in the tree [32].

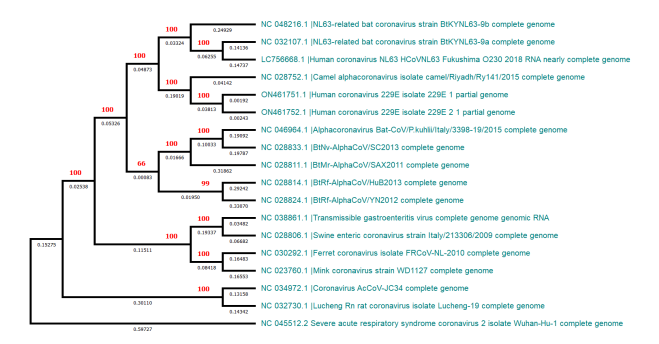


Figure.5. The Maximum Likelihood approach with bootstraps, a phylogenetic tree of (18) Alpha coronaviruses was created, with the SARS-CoV-2 RefSeq.

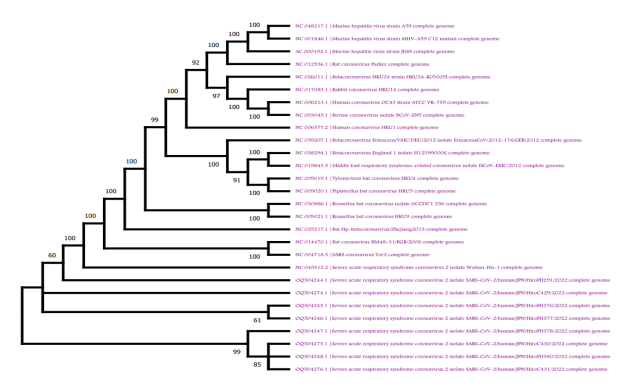


Figure.6. The Maximum Likelihood approach with bootstraps, a phylogenetic tree of (28) Beta coronaviruses was created, with the SARS-CoV-2 RefSeq.

The evaluation of a phylogenetic tree depends on the accuracy of the data used to build the tree. The accuracy of the data is determined by factors such as the quality of the data sources, the reliability of the analysis methods used, and the accuracy of the inference techniques used. If the data used is of high quality, the phylogenetic tree will be more accurate.

The accuracy of the branch lengths can be evaluated by looking at how well the tree reflects the actual evolutionary history of the species. The certainty of the conclusions drawn from the tree can be evaluated by considering the strength of the evidence that supports the tree [33] [34].

Results of Tree construction

The Results of this research paper covers the results of the study and the interpretation of these outcomes. We used the maximum likelihood estimate (MLE) approach to generate three phylogenetic trees based on complete genome sequences. The first tree (Figure 5) was for Alpha coronavirus species only, the second (Figure 6) was for Beta coronavirus species, including SARS-COV-2 Species, and the third (Figure 7) was constructed by merging the Alpha and Beta phylogenetic trees.

Conclusion

In this paper, we used the Maximum Likelihood Estimate (MLE) approach to construct a phylogenetic tree of Alpha and Beta coronavirus to illustrate the evolutionary ties between SARS-CoV-2 and (Alpha and Beta families) of coronavirus. This resulted in accurate and dependable phylogenetic tree. It is likely that different variants of the COVID-19 virus will emerge quickly, so the different strains of SARS-CoV-2 must be kept an eye on. Utilizing the phylogenetic tree and the Maximum Likelihood Estimation (MLE) approach appears to be a successful and advantageous technique for tracking the advancement of SARS-CoV-2's lineage, since the tree generated is accurate.

Future Works

In the future, we plan to build a phylogenetic tree of SARS-CoV-2 to demonstrate the evolutionary links between Alpha, Beta, Gamma, and Delta coronaviruses, which will uncover various SARS-CoV-2 mutations and single nucleotide polymorphisms (SNPs). To contain and treat COVID-19 and stop another outbreak, it is imperative to continually observe the variety and evolution of SARS-CoV-2.

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