

3D Visualizations of Multiple Coronaviruses on Whole Genomes

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Abstract

COVID-19 triggered by SARS-CoV-2 has become a common problem faced by people all over the world. With the development of bioinformatics and the breakthrough progress of gene technologies. It is a challenge topic to use genomic datasets for SARS-CoV-2 research. In this paper, a 3D visualization method is proposed to show A9 module in the Metagenomic Analysis System MAS. Seven coronaviruses of genus were illustrated and briefly analyzed. By comparing the visualization results, various SARS-CoV-2 genomes were represented as 2D and 3D maps under different conditions. Through related specific projections, the characteristics of the coronavirus can be observed intuitively from the projection results to provide an effective viewpoint for studying viral genomes.

Keywords: Coronavirus; Genome; 3D Map; 2D Map; Visualization; Projection

Introduction

In December 2019, a group of people with new coronary infections were discovered. The full genome sequence of the virus was obtained on January 29, 2020, with a total length of 29847 bp. On February 11, the World Health Organization named the new coronary pneumonia disease "COVID-19" and the International Committee of Viral Taxonomy named the virus "SARS-CoV-2". As of April 28, the number of diagnoses worldwide has exceeded 3026981. SARS-CoV-2 [1] has now seriously threatened the health of the global public, and it has attracted widespread attention from people around the world. Research on SARS-CoV-2 at home and abroad is also increasing.

The outer layer of the coronavirus has an envelope, and the shape is spherical or elliptical, with polymorphism. The genome is a linear single-stranded RNA virus, which is a large class of viruses that are ubiquitous in nature [2-6]. The Coronaviruses are divided into four genera by the International Virus Classification Committee: α , β , γ and δ . Among them, HCoV-OC43, HCoV-NL63, HCoV-229E, HCoV-HKU1, SARS-CoV, MERS-CoV and SARS-CoV-2 belongs to β genus [7-10]. Studying the similarities and differences between these seven coronaviruses from the perspective of genome sequence [11-15] visualization plays a vital role in preventing and controlling new coronaviruses and preventing the spread of disease.

Data visualization technology is a technology that displays abstract data in an intuitive graph or image, thereby facilitating people to conduct research and analysis [16,17]. There are many visualization methods for genomic sequences: most of the genomic sequence visualization models are implemented by DNA walking technology. For example, Gates-Nandy model [18], but this model has information degradation problem. In order to solve the problem of degradation and data loss, the researchers proposed a CGR model [19], a three-dimensional visualization model [20] and a worm model based on the Gates-Nandy model [21]. In 2003, Randić [22] proposed a spectral visualization model, which is different from the Gates-Nandy model. It consists of four parallel lines with the same distance and

four bases (adenine A, thymine T, bird the connection of purine G and pyrimidine C) also solves the problem of information degradation. However, the above visualization models for genomic sequences are not suitable for processing long DNA sequence data, and the analysis methods for visualization results are not universal. In 2014, Feng Haiqing, *et al.* proposed a gray-scale image-based DNA sequence visualization model [23]. This model converts one-dimensional DNA sequence information into two-dimensional 256-color gray by encoding four bases the degree of image greatly compresses the length of the DNA sequence visualization, and also has a high spatial tightness. However, the visualization of the model has noise, which is not conducive to researchers to observe more effective information.

According to the properties of genome sequence, a new visualization method of genome sequence is developed based on the research idea of data visualization technology and based on variant logic system [24-28]. By adjusting the parameters, it can adapt to the genomic sequence with a huge amount of data. By processing the seven coronavirus genome sequences, new graphical results can be obtained, and the result graphs clearly converge. From the results, we can find the correlation between the data and clarify the changing rules and phenomena, so that we can observe some characteristics of coronavirus from the perspective of variant.

Materials and Methods

Structurally speaking, variant logic is composed of four primitives. The four primitives represent four different states. There are symmetrical states and complementary states between two pairs. All states are combined to form the entire space. Segmentation is a major feature of variant. Dividing the data into equal segments can quickly process large amounts of data. The state value of each segment can be obtained by calculation. The visual model based on variant system is to illustrate the state set of each segment.

The visualization model processing flow is shown in figure 1.

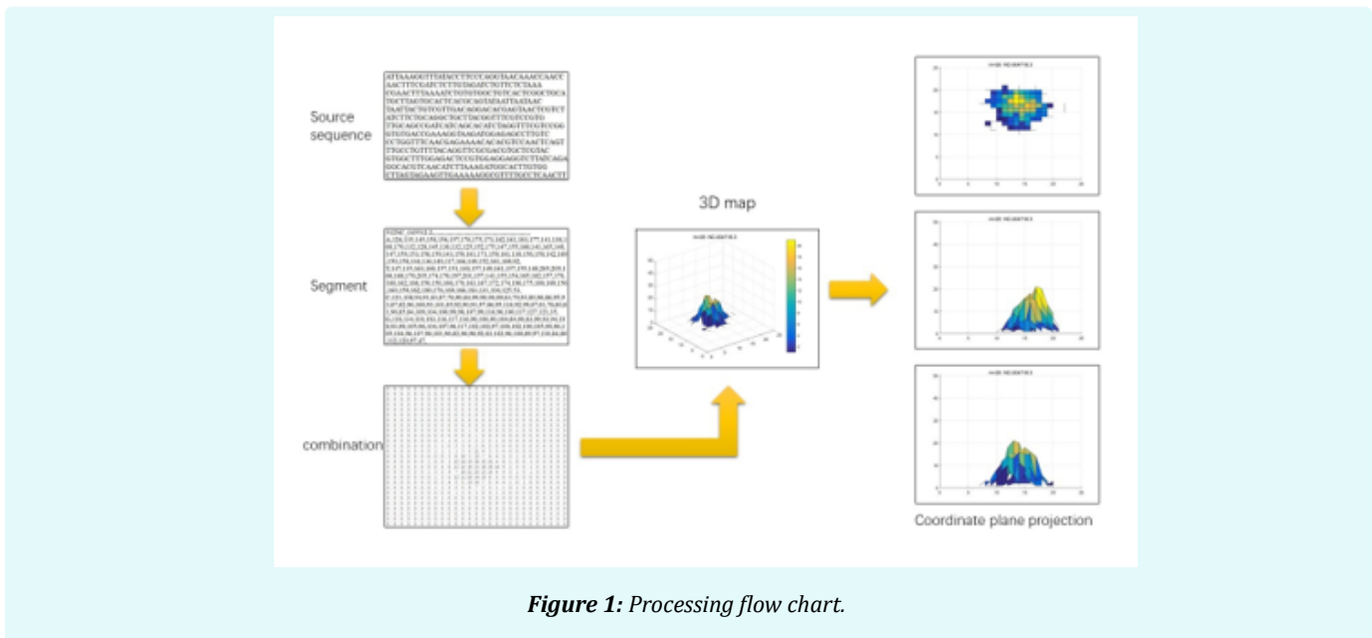


Figure 1: Processing flow chart.

The processing flow is: input the viral genome sequence, and segment the genome sequence according to the segmentation value m to obtain the segmentation result. Calculate the distribution of A, G, C, T in the segmentation results respectively to obtain the elementary states X_A, X_G, X_C, X_T . After the elementary states are obtained, two combinations can be selected between the elementary states to obtain superposition states $X_{(A+T)}, X_{(A+C)}, X_{(A+G)}, X_{(G+C)}, X_{(G+T)}, X_{(C+T)}$. Choosing the values in the primitive state and the superposition state to project can get the graphical results.

Introduction of the main modules in the model:

1. **Segmentation result:** The segmentation result is affected by the segmentation value. The segmentation value is usually represented by m , which represents the number of bases in each segment after the entire sequence is divided equally. The genome sequence is divided into n/m segments by changing the size of the segment value m , each segment has m bases. Controlling the size of m can adjust the resolution and effective area of the result.
2. **Primitive state:** The proportion of four bases in each segment is the primitive state, which represents the proportion of four bases in the genome sequence. The four primitive states in the variable-value system have a substitution and complementarity relationship, which perfectly fits the principle of complementary pairing between bases in the genome. The four symbols X_A , X_G , X_C , and X_T are used to represent the states of the four bases A, G, C, and T in each segment.
3. **Superposition state:** Each primitive state can be combined with each other and can also be combined with three or four primitive states. A state like this is called a superposition state. Each superposition state has a different meaning. There are six superposition states combined by two elementary states, which are $X_{(A+T)}$, $X_{(A+C)}$, $X_{(A+G)}$, $X_{(G+C)}$, $X_{(G+T)}$, $X_{(C+T)}$ said.
4. **Graphical results:** This experiment uses three-dimensional charts and two-dimensional projection charts to analyze the data.

Three-dimensional graph: It can display the features of primitive states and superimposed states. The three-dimensional map has a larger space capacity and can display more features. Two kinds of projections are randomly selected from the four primitive states and superposition states, respectively, as the x-axis and y-axis, and their values are accumulated at the corresponding positions to generate the z-axis, and finally a three-dimensional diagram can be generated. Here select $X_{(A+T)}$ and $X_{(A+G)}$ to generate a three-dimensional map.

Two-dimensional projection map: The three-dimensional image can clearly see the overall structure of the viral genome sequence, and the specific details can be projected onto a two-dimensional plane for observation. Projecting the 3D map onto other planes of the coordinate system generates a 2D projection of the 3D map.

Projecting various genomic sequences into three-dimensional space can observe its overall features from various angles, and the details contained in the overall features can also be displayed in the two-dimensional projection diagram generated by it.

Results and Discussion**Data introduction**

In the variable value visualization model, in order to better analyze the spatial distribution characteristics and cycle characteristics of the seven coronavirus genome sequences, they are expressed in two-dimensional and three-dimensional maps, respectively. In order to ensure the reliability of the data, the whole genome sequences of seven viruses were downloaded from NCBI. The corresponding sequences and sizes of virus names are as follows.

Name	Serial number	Data length
HCoV-OC43	NC-006213.1	30741 bp
HCoV-NL63	NC-005831.2	27553 bp
HCoV-229E	NC-002645.1	27317 bp
HCoV-HKU1	NC-006577.2	29926 bp
SARS-CoV	NC-004718.3	29751 bp
MERS-CoV	NC-019843.3	30119 bp
SARS-CoV-2	NC-045512.2	29924 bp

Table 1: Whole genome sequence information of seven viruses.

From the data information, it can be seen that the lengths of the whole genome sequences of the seven coronaviruses are not much different, in an order of magnitude. The genome sequences of living organisms are all in the millions. Although the genome sequences of coronaviruses are not long, they are about 30,000. Using variable value results to project data into an $n * n$ matrix, useful information can be observed simply and intuitively.

Projection selection

In the process of variable value processing, each link can be adjusted, this feature makes the method adapt to various data. However, the impact of each variable on the results makes it necessary to unify the comparison. Therefore, the appropriate value is calculated by the control variable method, and then the results obtained by the fixed parameters are compared. The two main parameters that affect the result are the m value and the other is the selection and combination.

The three-dimensional space is affected by the value of m in all aspects. The most direct impact is the size of the resulting graph and the speed of processing the data. We use the mean, variance and standard deviation as indicators to determine the effect of m value on the three-dimensional space. The table lists the mean, variance and standard deviation of the seven coronavirus sequences at $m = 26$.

The mean value reflects the amount of information contained in the unit space in the three-dimensional space, and the variance and standard deviation represent the degree of dispersion of the data points in the data set. It can be seen from the table that the indicators of the seven coronaviruses are not much different. Therefore, it has better observation effect when $m = 26$.

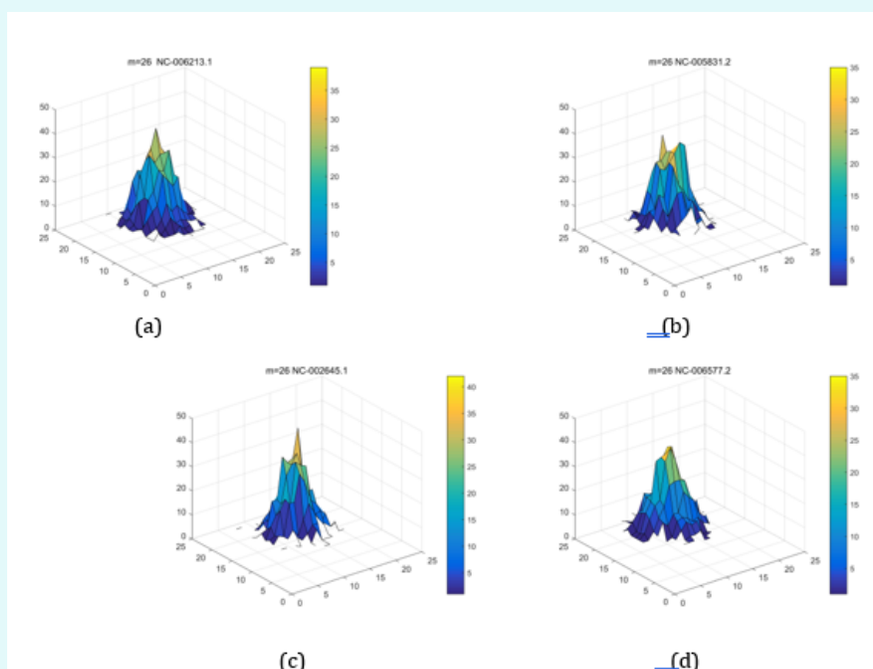
Name	HCoV-OC43	HCoV-NL63	HCoV-229E	HCoV-HKU1	SARS-CoV	MERS-CoV	SARS-CoV-2
Mean	1.62	1.45	1.44	1.58	1.57	1.58	1.57
Variance	26.64	23.61	23.61	24.25	25.45	26.01	25.97
Standard deviation	5.16	4.85	4.85	4.92	5.04	5.10	5.09

Table 2: Mean, variance and standard deviation of seven coronaviruses.

By comparing the display area of the results, select the combination of $X_{(A+T)}$ and $X_{(A+G)}$ to get a three-dimensional graphical result. Therefore, for the three-dimensional graphical result, select the result of the combination of $X_{(A+T)}$ and $X_{(A+G)}$ when $m = 26$.

Results show

Under the fixed parameters $m = 26$, $X_{(A+T)}$ and $X_{(A+G)}$ combination, seven kinds of coronavirus get three-dimensional map (Figure 2).



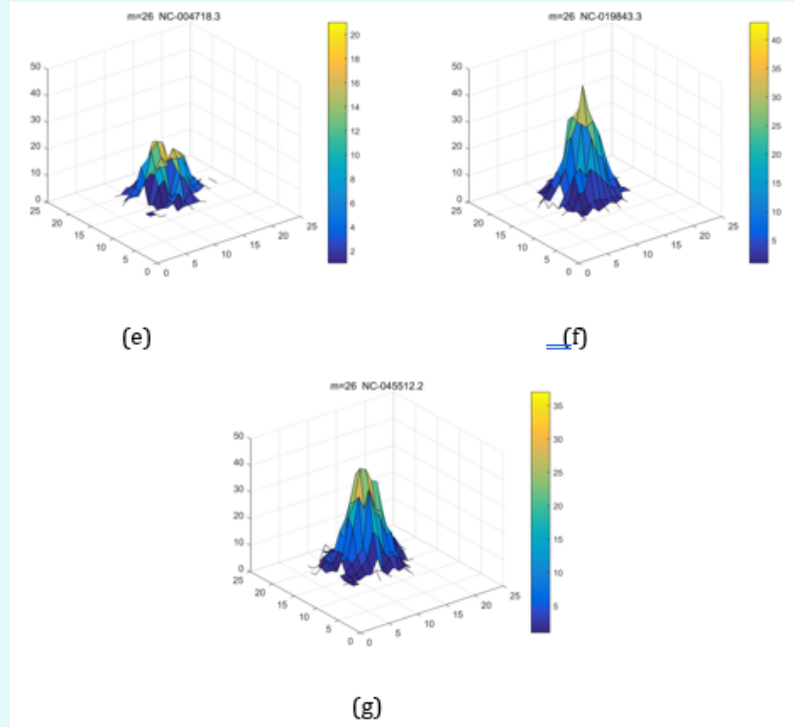


Figure 2: $m=26$, 3D graphical results of HCoV-OC43, HCoV-NL63, HCoV-229E, HCoV- HKU1, SARS-CoV, MERS-CoV and SARS-CoV-2.

Rotate and project the 3D image to obtain the projection results of the 3D image on other planes. The specific value of the three-dimensional projection can be seen by observing the image of the plane projection.

Figure 3 is the result of projecting a three-dimensional image onto the xy plane. Figure 4 is the result of projecting a three-dimensional image onto the xz plane, figure 5 is the result of projecting a three-dimensional image onto the yz plane.

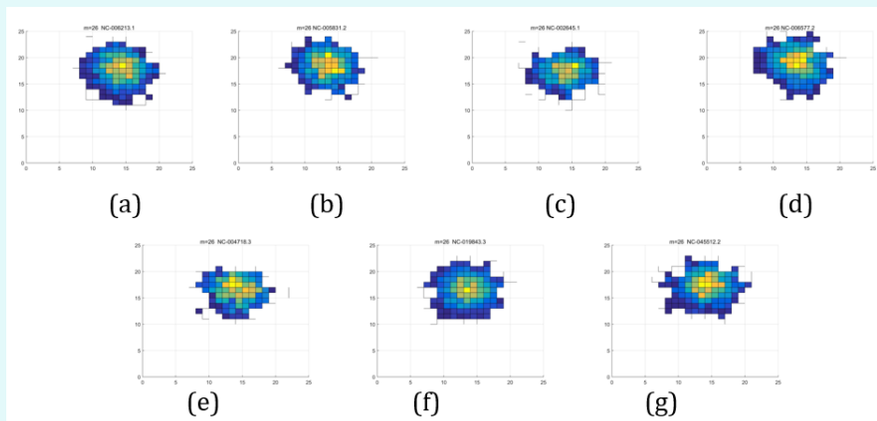


Figure 3: $m = 26$, HCoV-OC43, HCoV-NL63, HCoV-229E, HCoV-HKU1, SARS-CoV, MERS-CoV and SARS-CoV-2 three-dimensional images are projected onto the xy plane.

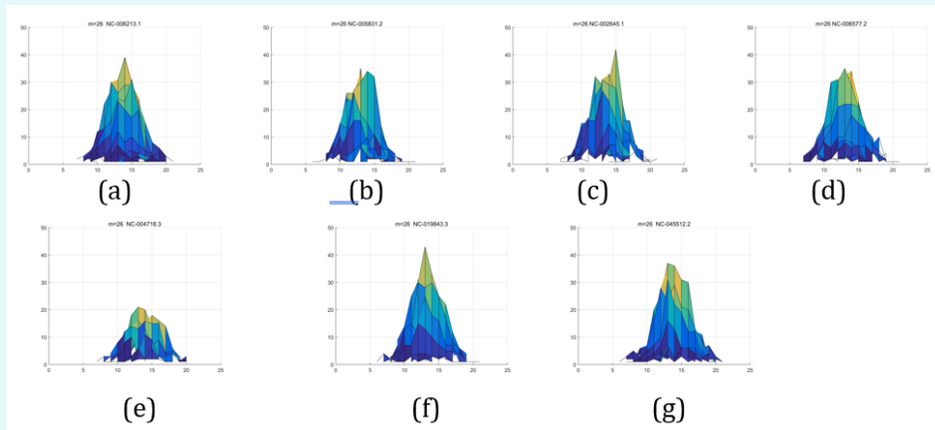


Figure 4: $m = 26$, HCoV-OC43, HCoV-NL63, HCoV-229E, HCoV-HKU1, SARS-CoV, MERS-CoV and SARS-CoV-2 three-dimensional images are projected onto the xz plane.

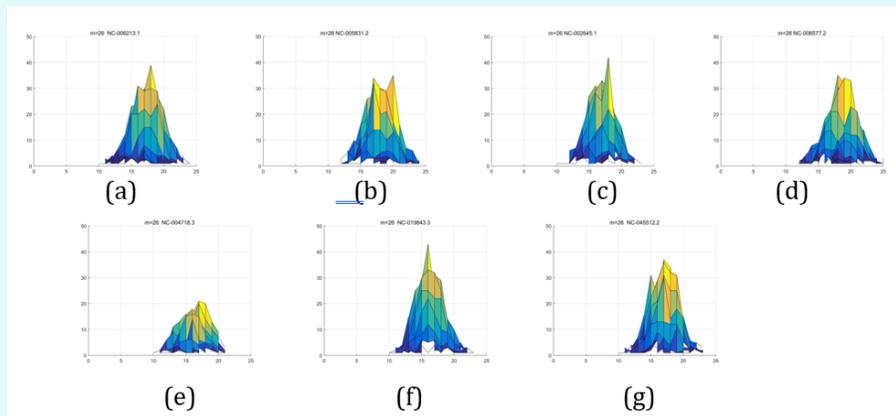


Figure 5: $m = 26$, HCoV-OC43, HCoV-NL63, HCoV-229E, HCoV-HKU1, SARS-CoV, MERS-CoV and SARS-CoV-2 three-dimensional images are projected onto the yz plane.

Result analysis

As can be seen from the three-dimensional graphical results, the graphical results of the seven coronaviruses are clustered in the middle and distributed around. They are all distributed in the upper left corner of the space. The special one is that the peak value of e is obviously lower than that of the other six coronaviruses. Acf has a clear single peak. The result of mapping the three-dimensional graphic results to the xy coordinate plane. From the figure, it is more accurate to observe the single peak position of acf. The distribution of the double peaks of several other coronaviruses is also different.

The result of mapping the three-dimensional graphic results to the xz coordinate surface and the yz coordinate result supplements the deficiencies of other surfaces. It is precisely seen that the result value of e is 21, and the result value of c is 42 is twice that of e.

Conclusion

A variable value visualization model was established for seven kinds of coronaviruses, and the data was converted into easy-to-understand graphs. The two-dimensional and three-dimensional graphs were suitable for different variable value calculation stages. Calculate the m value to adjust the best visual effect. Use three-dimensional projection to observe the distribution of superimposed states. By

projecting the three-dimensional image to two-dimensional to accurately observe the state of the superimposed state at various angles in the three-dimensional image. The results show that the results obtained by the visualization method based on variable value system can clearly see the connection and difference between the seven viruses. The study of new coronaviruses provides a new non-biological method.

Conflict of Interest

No conflict of interest has claimed.

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