An analysis of k-mer frequency features with SVM and CNN for viral subtyping classification
Un análisis de atributos de frecuencia de k-mer con SVM y CNN para la clasificación de subtipos de virus

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Abstract

Viral subtyping classification is very relevant for the appropriate diagnosis and treatment of illnesses. The most used tools are based on alignment-based methods, nevertheless, they are becoming too slow due to the increase of genomic data; for that reason, alignment-free methods have emerged as an alternative. In this work, we analyzed four alignment-free algorithms: two methods use k-mer frequencies (Kameris and Castor-KRFE); the third method used a frequency chaos game representation of a DNA with CNNs; and the last one processes DNA sequences as a digital signal (ML-DSP). From the comparison, Kameris and Castor-KRFE outperformed the rest, followed by the method based on CNNs.

Keywords: CNN, genome, viral subtyping, k-mer, Kameris, Castor, ML-DSP.

1 Introduction

In Virology, viral subtyping refers to a fundamental unit of virus nomenclature within a defined specie [1]. It is very important for taxonomic studies and disease treatment.

Each virus, could be divided into different subtypes, for instance, Human Immunodeficiency Viruses (HIV) are classified into two types: HIV-1 and HIV-2. HIV-1 is related to viruses in chimpanzees and gorillas, while HIV-2 is related to viruses of the sooty mangabey primate. Furthermore, HIV-1 is divided into groups: M, N, O, and P. Inside M group, there are subtypes A, B, C, D, E, F, G, H, I, K, K, and L that are related to the location where they emerged. Also, there are Circulating Recombinant Forms (CRFs) that refers to the different combination between HIV-1 subtypes [2].

The appropriate virus subtype classification is relevant for disease treatment, for example, one of the obstacles to the HIV treatment is its high genetic variability [2]. The HIV-1’s genome is diverged by about 15% [3]. Furthermore, some authors claim that a correct classification of HIV-1 subtypes is required for clinical management [4].

There are several methods for viral classification, they could be grouped into alignment-based, organism-specifics, and alignment-free methods. [5]. One of the most popular method used for genome similarity is the alignment-based method BLAST [6]. Furthermore, alignment-based is not suitable for viral genome classification because of their genetic variation (recombination, shuffling, and horizontal gene transfer events) [7]. Also, we need prior knowledge to adjust some parameters in order to get good results [8]. In addition, the majority of the alignment-based and organism-specifics are proprietary and they require laboratories and servers to transmit the sequence, this expose sensitive information [1].
In this study, we analyzed the performance of k-mer frequency like feature vectors with Support Vector Machines (SVM) and Convolutional Neural Networks (CNN) for viral subtyping classification. We compared Kameris [1], Castor-KRFE [5], and an own implementation, using Frequency Chaos Game Representation (FCGR) with CNN. Additionally, we include a method based on digital signal processing [9].

The work is structured as follows: in Section 2, we present the most relevant related works, Section 3 describes the methods, parameters and data used in order to replicate the experiment, Section 4 describes the results, Section 5 comments the results, Section 6 shows the conclusions and finally in Section 7, we show the future work.

2 Viral subtyping classification

The most popular tools for viral subtyping classification are REGA [10], SCUAEUL [11] and USEARCH [12], all of them are based on alignment-based methods and phylogenetics analysis. Furthermore, there are pair-wise alignment-based methods like BLAST [6], TurboBLAST [13], ScalaBLAST [14] and multiple sequence alignment (CLUSTAL [15]).

On the other hand, there are alignment-free methods. They compute a feature vector and then, they use a distance algorithm or a machine learning model in order to classify a genome [16, 17]. The viral (subtype) classification problem in alignment-free methods takes as input, a set of sequences \( S = \{ s_1, s_2, ..., s_n \} \), where \( s_i \) is a string of characters \( \omega = \{ A, C, G, T \} \) and \( y \) representing the class of each sequence. Some alignment-free methods are Comet [18], Castor [19], VirFinder [20], a data warehouse with machine learning algorithms [21], an open source framework Kameris [1] and Castor-KRFE [5].

Blaisdell [22] proposed one of the first methods using k-mer frequencies. These k-mer frequencies have been used for phylogenetic studies and machine learning models during the past decades [23, 24, 25, 26]. Other researches used k-mer for sequence prediction [27] or sequence assembly [28]. The k-mers can be viewed as sub-strings of a sequence, for example, for the sequence \( s = \{ A, C, T, G, A, C \} \), the 3-mers set is: \( \{ ACT, CTG, TGA, GAC \} \), and the 2-mers set: \( \{ AC, CT, TG, GA, AC \} \). The alignment-free methods base on k-mer frequencies compute the frequency of each element in a k-mer set, they usually experiment with different values of \( k \) in order to get the best \( k \) for a given dataset.

Moreover, some researches have used Chaos Game Representations (CGR) as a genomics signature along with machine learning models; For instance, CGR genomic signatures for HIV-1 sub-typing [29], CGR for the analysis of Circulating Recombinant Forms (CRFs) of HIV-1 [30], classification of the genotypes of Human Papilloma Virus (HPV) [31], CGR for classification of viral pathogens [32], and more recently, CGR for rapid classification of COVID-19 [33].

2.1 Kameris

Proposed by Solis-Reyes [1] in 2018, they computed the k-mer frequencies using a Frequency Chaos Game Representation (FCGR), this FCGR is derived from the Chaos Game Representation (CGR) [34]. A FCGR could be represented in Fig. 1, each cell in the matrix represents a specific k-mer, the first cell, for example, represents the 2-mer AA, and it appears 5 times in a sequence. Besides, in Fig. 2, we present how each cell of a FCGR matrix represents a specific k-mer, the \( C \) quadrant sub-divided into CA, CC, CG, CT and the \( T \) quadrant sub-divided into ACT, CCT, GCT, TCT, the division depends on the number of levels or the \( k \) value in k-mer, in this way each k-mer has a unique position in a FCGR matrix. Another example is shown in Fig. 3, here we present the FCGR matrix for one (1-mer), two (2-mer) and three (3-mer) levels, in the case of \( L = 2 \) we could see all the possible 2-mers of a sequence. Also, in Fig. 1, we represent the number of times each k-mer appears in a sequence. In this way, it is easy to store the k-mer frequencies in the FCGR.

![Figure 1: A FCGR k-mer example, each k-mer is represented as a cell in the matrix, and the frequency of each k-mer is represented as the pixel value.](image)

After processing the FCGR as a feature vector, it is applied a Single Value Decomposition (SVD) for dimensionality reduction. Then, they used a Support Vector Machine (SVM). The authors evaluated their work for the classification of HIV-1 virus subtypes.

2.2 Castor-KRFE

Proposed by Lebatteux et al. [5] in 2019, they computed the k-mer frequencies in a different form than Kameris. For example, from a set of sequences \( S = \{ s_1, s_2, ..., s_n \} \), the authors obtain the k-mers frequencies presented only in the dataset. For example
Figure 2: A FCGR matrix. The $G$ quadrant subdivided into the corresponding $G$-endings and the $TG$ quadrant subdivided into the corresponding $TG$-endings.

Figure 3: A FCGR example for one, two and three levels. Each cell represents a specific k-mer.

If $S = \{ACTGC, TTGCG\}$, the 3-mers will be: $\{ACT, CTG, TGC, TTG, GCG\}$ avoiding repetition, then the feature vector will be: $\{1, 1, 2, 1, 1\}$, we noticed that the 3-mer $TGC$ appears twice in the dataset. In contrast to Kameris, Castor-KRFE obtains a feature vector smaller than Kameris. Moreover, Castor-KRFE uses Recursive Feature Elimination (RFE) based on SVM to eliminate features. After that, they used SVM as a classifier. Also, they evaluated their proposal in different viral datasets obtaining the best $k$ and the number of features for each dataset.

2.3 FCGR and CNN

In this method, we used Frequency Chaos Game Representation (FCGR) of a DNA sequence along with Convolutional Neural Network (CNN) to classify virus subtyping.

The problem of represent DNA sequences as images, was first proposed by H.J. Jeffrey et al. [34]. A formal definition is denote by Eq. (1), for a DNA sequence $s_1, s_2, ..., s_n$, the corresponding CGR sequence $(X_n)$ is given by:

$$X_0 = \left(\frac{1}{2}, \frac{1}{2}\right), X_n = \frac{X_{n-1} + W}{2}$$  (1)

where $W$ represents the coordinates of the corners of the unit square $A = (0, 0); C = (0, 1); G = (1, 1); T = (1, 0)$ if $s_n$ is $a, c, g, t$ respectively. In this form, we plotted a point for each value of $X_n$. For example, in Fig. 4 (left), we plotted the points according to the sequence $s = T, G, C, A$. Moreover, in Fig. 4 (right), we plotted the complete CGR of a HIV-1 whole genome. Jeffrey et al. [34] demonstrated that, each specimen have different fractal patterns in its CGR.

Figure 4: Chaos game representation of a DNA sequence. Left: CGR point of sequence: $s = T, G, C, A$. Right: CGR of a HIV-1 whole genome

Then, Wang et al.[35] proved that CGR’s patterns are completely determined by frequencies of oligonucleotides of all lengths (a deep study of CGR is presented by Joseph et al. [36]). These frequencies of oligonucleotides are commonly known as k-mer frequencies and they could be represented as a FCGR (Fig. 1). Moreover, these frequencies could be interpreted as pixel intensities to represent an image (Fig. 5).

Figure 5: A FCGR (k=5) of a HIV-1 genome.

Finally, we used a Convolutional Neural Network (CNN) to classify a sequence. We used the CNN’s architecture proposed by Fabijańska et al. [37] (Fig. 7) and other two small models detailed in Fig. 6. We did not use Max Pooling, because the images from a FCGR with $k = 5$ were small (32x32).

Figure 6: CNN’s architecture proposed.
2.4 ML-DSP

Machine Learning with Digital Signal Processing (ML-DSP), is proposed by Randhawa et al. [9]. In this case, a DNA sequence is mapped to numbers, where each base $T/C = 1$ and $A/G = -1$. Then, the authors applied a Discrete Fourier Transform (DFT) and its magnitude spectrum. Finally, a Pearson Correlation Coefficient is computed. Then, they used a Support Vector Machine (SVM) to train the model.

2.5 Summary

We compared the performance of Kameris [1], Castor-KRFE [5], CNNs and ML-DSP. The first two methods, use k-mer frequencies, but they differ in the way they computed it. Kameris computed all the $4^k$ possible k-mers using a FCGR matrix, it has an efficient way of getting the frequencies they but got a big feature vector. In contrast, Castor-KRFE computed just the k-mers frequencies presented in the training set, they computed it inefficiently but, they got a feature vector smaller than Kameris. Moreover, Kameris used SVD for dimensionality reduction meanwhile Castor-KRFE use RFE for feature elimination. The third method, uses a FCGR images and CNNs. Finally, the last method (ML-DSP), consider a DNA sequence as a digital signal.

In Table 1, we presented the methods described before, from now on we are going to use the nomenclature of this table for each method.

3 Materials and Method

We used the datasets from Castor\(^1\), they are the first 10 rows of Table 2. HBVGENCG is a group of sequences of Hepatitis-B virus; HIVRPGC, HIVSUBCG and HIVSUBPOL are sequences of HIV-1; INFSUBHA, INFSUBMP and INSUBFNA are sequences from Influenza virus; EBOSPECG, RHISPECG and HPVGENCG are sequences of Ebola, Rhinovirus and Human papillomavirus respectively.

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\(^1\)http://castor.bioinfo.uqam.ca/

| Method name | Description |
|-------------|-------------|
| Kameris-SVD | Kameris with dimensionality reduction SVD. The feature vector was reduced to 10%. |
| Kameris     | Kameris without dimensionality reduction. |
| Castor-KRFE | Castor with feature elimination RFE. |
| Castor      | Castor without feature elimination. |
| CNN-1, CNN-2 and CNN-3 | Convolutional Neural Networks (CNN) with FCGR as images. We used three architectures: CNN-1, CNN-2 and CNN-3 (Fig. 6 and 7). |
| ML-DSP      | Machine Learning Digital Signal Processing (ML-DSP) uses Discrete Fourier transform and Pearson Correlation Coefficient. |

Also, we used a group of dataset proposed by Randhawa et al. [9], these datasets are the last 9 rows of Table 2.

For CNN models, we used Adam optimizer, mini-batch size of 128 and 100 epochs. Moreover, We used seed of 1 for random numbers in weights initialization.

For processing time analysis (Table 6), we used an Intel i5 processor of 1.9 GHz, and 8 GBytes of memory RAM.

4 Results

In this section, we show a comparison between the methods proposed (Kameris, Castor, ML-DSP, CNN-1, CNN-2 and CNN-3). Additionally, we compared the processing time of these methods against BLAST. In the experiments, $k$ refers to k-mer.

4.1 F-score and feature vector size

Kameris-SVD/Kameris and Castor-KRFE/Castor are very similar so, we compared them exhaustively. We evaluated the f-score metric with and without dimensionality reduction and feature elimination. A comparison of f-score is presented in Fig. 8, we used k-mers, ranging from $k=1$ to $k=9$. We concluded that Kameris, and Castor without applying dimensionality reduction and feature elimination have the same results. In contrast, when we apply SVD to Kameris, the f-score decreases slightly, but when we apply RFE
Table 2: The datasets used in the experiments.

| Data sets     | Average seq. length | No. of classes | No. of instances |
|---------------|---------------------|----------------|------------------|
| HIVGRPCG      | 9164                | 4              | 76               |
| HIVSUGCG      | 8922                | 18             | 597              |
| HIVSUBPOL     | 1211                | 28             | 1352             |
| INFSUBHMA     | 1719                | 2              | 10825            |
| INFSUBMP      | 759                 | 2              | 21421            |
| INSUBFMA      | 1416                | 2              | 10715            |
| EOBSCP       | 18917               | 5              | 751              |
| HBVGENCG      | 3189                | 8              | 230              |
| RHISPEG      | 369                 | 3              | 1316             |
| HPVGENCG      | 7610                | 3              | 125              |
| Primates      | 16626               | 2              | 148              |
| Dengue        | 10595               | 4              | 4721             |
| Protists      | 31712               | 3              | 159              |
| Fungi         | 49178               | 3              | 224              |
| Plants        | 277931              | 2              | 174              |
| Amphibians    | 17530               | 3              | 290              |
| Insects       | 15689               | 7              | 898              |
| 3classes      | 16292               | 3              | 2170             |
| Vertebrates   | 16806               | 5              | 4322             |

to Castor, the f-score decreases more than Kameris.

Moreover, we evaluated how the feature vector size of Kameris and Castor increases according to k (Fig. 9). As we can see, since $k = 6$ Kameris’s feature vector is too big, it is bigger than Castor’s feature vector, and that is because Kameris’s feature vector size is $4^k$ (it considers all possible k-mers) meanwhile Castor just considers the k-mers presented in the dataset. Also, we evaluated how the feature vector size of Kameris-SVD and Castor-KRFE increases according to $k$ (Fig. 10), Kameris-SVD’s feature vector size is bigger than Castor-KRFE’s feature vector since 5-mers.

Figure 8: F-score (y-axis) comparison between Kameris-SVD/Kameris and Castor-KRFE/Castor for k-mers ranging from k=1 to k=9 (x-axis).

Figure 9: Feature vector size of Kameris and Castor according to $k$. We used HIVSUBPOL dataset for this experiment.

Figure 10: Feature vector size of Kameris-SVD and Castor-KRFE according to $k$. We used HIVSUBPOL dataset for this experiment.

In Table 3, we show the best f-score that we got with the smaller value of $k$, for Kameris-SVD and Castor-KRFE. Kameris-SVD slightly outperform Castor-KRFE, meanwhile, Castor-KRFE’s feature vector is smaller than Kameris’ feature vector. Furthermore, in Table 4, we show the f-score of Kameris-SVD and Castor-KRFE with 5-mers and 10-fold validation.

In Table 5, we show the f-score of the methods proposed. In this case, we used Kameris and Castor without dimensionality reduction and feature elimination because, they got better results in this way. We noticed that ML-DSP got poor results meanwhile, Kameris and Castor got the best results.

4.2 Processing time

Also, we compared the processing time of each algorithm against BLAST. We choose BLAST because
Table 3: The best f-score that we got with the smaller value of $k$. Also, the number of features is presented.

| Dataset      | (k-mer) | f-score | nfeatures |
|--------------|---------|---------|-----------|
| HIVGRPCG     | 4       | 1.0000  | 12        |
| HIVSUBCG     | 5       | 0.9983  | 47        |
| HIVSUBPOL    | 7       | 0.9761  | 99        |
| Castor-KRFE  |         |         |           |

Table 4: F-score and standard deviation of Kameris-SVD and Castor-KRFE (5-mer) with 10-fold validation.

| Dataset      | Kameris-SVD | Castor-KRFE |
|--------------|-------------|-------------|
|              | f-score     | sd          |
| HIVGRPCG     | 1.000       | 0.000       |
| HIVSUBCG     | 0.999       | 0.003       |
| HIVSUBPOL    | 0.965       | 0.022       |
|              | f-score     | sd          |
| HIVGRPCG     | 1.000       | 0.000       |
| HIVSUBCG     | 0.9937      | 0.008       |
| HIVSUBPOL    | 0.9629      | 0.023       |

this is the most used method for viral subtyping classification and similarity analysis. For example, REGA compares a DNA sequence (query) against 87 other sequences (HIV-1 samples), then according to the phylogenetic analysis, the method classifies a sequence. A simple query on these tools could take some minutes but, we can not measure exactly this time because this is web service.

For Kameris, Castor, ML-DSP, and CNN (CNN-1, CNN-2 and CNN-3), we measured the processing time to get the feature vector; for BLAST, we measured the processing time to align the query sequence against 76 other sequences (the smallest dataset have 76 instances); we used the bioblast library. For all the experiments, we run each method 10 times and then, we took the average time.

In Table 6, we show a comparison of the processing time. Castor, ML-DSP and CNN (CNN-1, CNN-2 and CNN-3) got the best results, meanwhile BLAST got poor results against the others.

4.3 Limitations

In this study, we noticed that the processing time to compute k-mer frequency could take too much time since 7-mer. Additionally, bigger genomes need bigger k-mers [5]. In this context, we need efficient algorithms in order to compute k-mer frequency. Moreover, there are datasets with a small number of samples that are not appropriated to use in machine learning.

5 Discussion

Kameris and Castor without their dimensional reduction and feature elimination are almost the same, despite Castor got a small feature vector (the extra attributes of Kameris are zero values). Meanwhile, if we applied SVD and RFE to Kameris and Castor respectively, the f-score slightly is reduced. Moreover, according to the results in Table 3 and 4, kameris-SVD outperformed Castor-KRFE.

In Table 5, ML-DSP got poor results, meanwhile the authors presented good results in their paper [9].

In Table 6, we compared the processing time against BLAST. When we use BLAST for viral classification, we need to align a query sequence against others to find the most similar, and then we could infer the class of a sequence. Nevertheless, this method depends strongly on the amount of instances in a dataset. Currently, the datasets in Bioinformatics are increasing every day, so BLAST will be slower.

6 Conclusions

In this work, we evaluated four methods for viral subtyping classification, based on alignment-free algorithms. Two methods use k-mer frequencies (Kameris and Castor), the third method uses CNNs, and finally the last method treats a DNA sequence as a signal.

Kameris-SVD and Castor-KRFE, compute k-mer frequencies, then they used SVD and RFE for dimensionality reduction and feature elimination. Finally, they used a SVM as a classifier. We noticed that Kameris-SVD outperformed slightly Castor-KRFE. Moreover, if we did not use SVD and RFE, they got the same f-score. Additionally, Castor-KRFE got a smaller feature vector than Kameris-SVD, it is very important considering the huge amount of data in Bioinformatics.

ML-DSP processed a DNA sequence as a digital signal, but this method got poor results against CNNs, Kameris, and Castor. Kameris and Castor without SVD and RFE got the best accuracy, but they are
Table 5: F-score of the methods proposed (5-mer). We put a ‘-’ for the methods that couldn’t converge.

| Dataset     | CNN-1 | CNN-2 | CNN-3 | ML-DSP | Kameris | Castor |
|-------------|-------|-------|-------|--------|---------|--------|
| Primate     | 1     | 1     | 0.964 | 1      | 1       | 1      |
| Dengue      | 1     | 1     | 1     | 0.998  | 1       | 1      |
| Protists    | 1     | 1     | 0.882 | 0.805  | 1       | 1      |
| Fungi       | 0.978 | 1     | 0.923 | 0.859  | 1       | 1      |
| Plants      | 0.882 | 0.85  | 0.917 | 0.78   | 0.882   | 0.972  |
| Amphibians  | 0.983 | 1     | 0.982 | 0.967  | 1       | 1      |
| Insects     | 0.961 | 0.949 | 0.966 | 0.893  | 0.994   | 0.994  |
| 3classes    | 1     | 1     | 1     | 1      | 1       | 1      |
| Vertebrates | 0.995 | 0.998 | 0.995 | 0.985  | 0.998   | 0.998  |
| HIVGRPCG    | 0.909 | 0.913 |       | 0.838  | 1       |        |
| HIVSUBCG    | 0.962 | 0.975 | 0.978 | 0.686  | 1       |        |
| HIVSUBPOL   | 0.981 | 0.981 | 0.981 |       | 0.993   | 0.993  |
| INFSUBHA    | 1     | 1     | 1     | 1      | 1       | 1      |
| INFSUBMP    | 0.986 | 0.989 | 0.989 | 0.935  | 0.989   | 0.988  |
| EBOSEPSCG   | 1     | 1     | 1     | 0.982  | 1       | 1      |
| HBVGENCG    | 1     | 1     | 1     | 0.773  | 1       |        |
| RHISPECG    | 1     | 1     | 1     | 1      | 1       | 1      |
| HPVGENCG    | 1     | 1     | 1     | 1      | 1       | 1      |

Table 6: Time processing in milliseconds, for each method to process a DNA sequence, we used 5-mer for each method. CNN stands for the time processing to compute FCGR images in CNN-1, CNN2 and CNN-3.

| Dataset     | BLAST  | Kameris | Castor | ML-DSP | CNN   | Sequence length |
|-------------|--------|---------|--------|--------|-------|-----------------|
| Primate     | 0.609  | 0.052   | 0.006  | 0.005  | 0.005 | 16499           |
| Dengue      | 0.175  | 0.033   | 0.003  | 0.004  | 0.004 | 10313           |
| Protists    | 0.227  | 0.066   | 0.007  | 0.006  | 0.006 | 24932           |
| Fungi       | 0.402  | 0.240   | 0.026  | 0.026  | 0.017 | 190834          |
| Plants      | 2.160  | 0.299   | 0.034  | 0.043  | 0.029 | 103830          |
| Amphibians  | 0.397  | 0.057   | 0.006  | 0.006  | 0.005 | 16101           |
| Insects     | 0.135  | 0.045   | 0.005  | 0.005  | 0.004 | 14711           |
| 3classes    | 0.110  | 0.044   | 0.005  | 0.004  | 0.004 | 14496           |
| Vertebrates | 0.417  | 0.049   | 0.006  | 0.005  | 0.005 | 16442           |
| HIVGRPCG    | 0.229  | 0.027   | 0.003  | 0.003  | 0.003 | 8654            |
| HIVSUBCG    | 0.351  | 0.026   | 0.003  | 0.003  | 0.003 | 8589            |
| HIVSUBPOL   | 0.130  | 0.004   | 0.001  | 0.001  | 0.001 | 1017            |
| INFSUBHA    | 0.095  | 0.006   | 0.001  | 0.001  | 0.002 | 1704            |
| INFSUBMP    | 0.086  | 0.003   | 0.001  | 0.001  | 0.001 | 759             |
| INSUBFNA    | 0.081  | 0.005   | 0.001  | 0.001  | 0.001 | 1413            |
| EBOSEPSCG   | 0.209  | 0.058   | 0.006  | 0.006  | 0.005 | 18828           |
| HBVGENCG    | 0.146  | 0.011   | 0.001  | 0.001  | 0.002 | 3182            |
| RHISPECG    | 0.073  | 0.001   | 0.000  | 0.000  | 0.001 | 369             |
| HPVGENCG    | 0.115  | 0.022   | 0.002  | 0.002  | 0.003 | 7100            |

followed by CNNs.

Also, we compared the processing time against BLAST. In this case, Castor, ML-DSP and CNNs got the best results. Moreover, there is a big difference against BLAST.

7 Future work

K-mer frequencies are good descriptors for viral subtyping classification. Nevertheless, despite the good performance, they have a high cost for $k > 5$; large genomes require higher values of $k$. For that reason, we are going to evaluate another algorithms like the inverted-index technique to compute k-mer frequencies. Moreover, we are going to evaluate the use of another deep learning technique for viral classification.

Competing interests

The authors have declared that no competing interests exist.
Authors’ contribution
V.E. Machaca conceived the idea, wrote the program, developed the experiments, analyzed the results, wrote and revised the manuscript.

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Citation: V.E. Machaca. An analysis of k-mer frequency features with SVM and CNN for viral subtyping classification. Journal of Computer Science & Technology, vol. 20, no. 2, pp. 99–107, 2020.

DOI: 10.24215/16666038.20.e11.

Received: March 21, 2020 Accepted: October 5, 2020.

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