# Clustering the Normalized Compression Distance for Influenza Virus Data

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Abstract. The present paper analyzes the usefulness of the normalized compression distance for the problem to cluster the hemagglutinin (HA) sequences of influenza virus data for the HA gene in dependence on the available compressors. Using the CompLearn Toolkit, the built-in compressors zlib and bzip2 are compared.

Moreover, a comparison is made with respect to hierarchical and spectral clustering. For the hierarchical clustering, hclust from the R package is used, and the spectral clustering is done via the kLine algorithm proposed by Fischer and Poland (2004).

Our results are very promising and show that one can obtain an (almost) perfect clustering. It turned out that the **zlib** compressor allowed for better results than the **bzip2** compressor and, if all data are concerned, then hierarchical clustering is a bit better than spectral clustering via kLines.

### 1 Introduction

The similarity between objects is a fundamental notion in everyday life. It is also fundamental to many data mining and machine learning algorithms, and, in particular to clustering algorithms. Often the similarity between objects is measured by a domain-specific distance measure based on features of the objects. For defining the right domain-specific distance measure one needs special knowledge about the application domain for extracting the relevant features beforehand. Such an approach does not only cause difficulties, but includes a certain danger or risk of being biased.

If one is pursuing the approach to design data mining algorithms based on domain knowledge, then the resulting algorithms tend to have many parameters. By using these parameters, one can then control the algorithms' sensitivity to certain features. Determining how relevant particular features are is often difficult and may require a certain amount of guessing. Expressing this differently,

<sup>\*</sup> Supported by MEXT Grant-in-Aid for Scientific Research on Priority Areas under Grant No. 21013001.

one has to tune the algorithms which is requiring domain knowledge and a larger amount of experience. Furthermore, it may be expensive, error prune and time consuming to arrive at a suitable tuning.

However, as a radically different approach, the paradigm of parameter-free data mining has emerged (cf. Keogh *et al.* [11]). The main idea of parameter-free data mining is the design of algorithms that have no parameters and that are universally applicable in all areas.

The problem is whether or not such an approach can be realized at all. It is only natural to ask how an algorithm can perform well if it is not based on extracting the important features of the data and if we are not allowed to adjust its parameters until it is doing the right thing. As expressed by Vitányi *et al.* [21], if we a priori know the features, how to extract them, and how to combine them into exactly the distance measure we want, we should do just that. For example, if we have a list of cars with their color, motor rating, etc. and want to cluster them by color, we can easily do that in a straightforward way.

So the approach of parameter-free data mining is aiming at scenarios where we are not interested in a certain similarity measure but in *the* similarity between the objects themselves.

The main goal of the present paper is to test the usefulness of this approach in the domain of influenza viruses. Our data are gene sequences for the hemagglutinin of influenza viruses. The hemagglutinin of influenza viruses is important, since it is responsible for binding the virus to the cell it infects. So far, 16 subtypes of influenza hemagglutinin are known. More details are given in Subsection 3.1.

The definite method used by biologists to determine the subtype of the influenza hemagglutinin is based on the antiserum that prevent the docking of the virus. So intuitively, the similarity between the gene sequences for the hemagglutinin of influenza viruses should be large if they have the same subtype and small if the have a different subtype. Therefore, it seems justified to test the paradigm of parameter-free data mining in this domain.

The most promising approach to this paradigm uses Kolmogorov complexity theory [13] as its basis. The key ingredient to this approach is the so-called *normalized information distance* (NID) which was developed by various researchers during the past decade in a series of steps (cf., e.g., [4, 12, 9]). The idea behind it is quite intuitive. If two objects are similar then there should be a simple description of how to transform each one of them into the other one. And conversely, if all descriptions for transforming each one of them into the other one are complex, then the objects should be dissimilar.

More formally the *normalized information distance* between two strings  $\mathbf{x}$  and  $\mathbf{y}$  is defined as

$$NID(\mathbf{x}, \mathbf{y}) = \frac{\max\{\mathsf{K}(\mathbf{x}|\mathbf{y}), \, \mathsf{K}(\mathbf{y}|\mathbf{x})\}}{\max\{\mathsf{K}(\mathbf{x}), \, \mathsf{K}(\mathbf{y})\}} \,, \tag{1}$$

where K(x|y) is the length of the shortest program that outputs x on input y, and K(x) is the length of the shortest program that outputs x on the empty input. It is beyond the scope of the present paper to discuss the technical details of the definition of the NID. We refer the reader to Vitányi *et al.* [21]. The NID has nice theoretical properties, the most important of which is universality. The NID is called *universal*, since it accounts for the dominant difference between two objects (cf. Li *et al.* [12] and Vitányi *et al.* [21] and the references therein).

In a sense, the NID captures all computational ways in which the features needed in the traditional approach could be defined. Since its definition involves the Kolmogorov complexity  $K(\cdot)$ , the NID cannot be computed. Therefore, to apply this idea to real-world data mining tasks, standard compression algorithms, such as gzip, bzip2, or PPMZ, have been used as approximations of the Kolmogorov complexity. This yields the *normalized compression distance* (NCD) as approximation of the NID (cf. Definition 1).

In a typical data mining scenario we are given some objects as input. The pairwise NCDs for all objects in question form a distance matrix. This matrix can be processed further until finally standard algorithms, e. g., clustering algorithms can be applied. This has been done in a variety of typical data mining scenarios with remarkable success. Works of literature and music have been clustered according to genre or author; evolutionary trees of mammals have been derived from their mitochondrial genome; language trees have been derived from several linguistic corpora (cf., e.g., [9, 11, 6, 7, 3]).

As far as virus data are concerned, Cilibrasi and Vitányi [8] used the SARS TOR2 draft genome assembly 120403 from Canada's Michael Smith Genome Sciences Centre and compared it to other viruses by using the NCD. They used the bzip2 compressor and applied their quartet tree heuristic for hierarchical clustering. The resulting ternary tree showed relations very similar to those shown in the definitive tree based on medical-macrobiological genomics analysis which was obtained later (see [8] for details).

In the present paper we aim at a detailed analysis of the general method outlined above in the domain of influenza viruses. More specifically, we are interested in learning whether or not specific gene data for the hemagglutinin of influenza viruses are *correctly* classifiable by using the concept of the NCD. For this purpose we have chosen a set of 106 gene sequences from the National Center for Biotechnology Information for which the correct classification of the hemagglutinin is known. As explained in Section 3, there are 16 subtypes commonly called H1, ..., H16. For these 106 gene sequences (or subsets thereof) we then compute the NCD by using the CompLearn Toolkit (cf. [5]) as done in [8].

This computation returns a symmetric matrix D such that  $d_{ij}$  is the NCD between the data entries i and j (henceforth called distance matrix). Furthermore, we study the influence of the compressor chosen and restrict ourselves here to the zlib and bzip2 compressors which are the standard two built-in compressors for the CompLearn Toolkit.

The next step is the clustering. Here of course the variety of possible algorithms is large. Note that the CompLearn Toolkit contains also an implementation of quartet tree heuristic for hierarchical clustering. However, this heuristic is computationally quite expensive and does currently not allow to handle a matrix of dimension  $106 \times 106$ . Therefore, we have decided to try the *hierarchical clus* 

tering algorithm from the R package (called hclust) with the average option. In this way we obtain a rooted tree showing the relations among the input data.

The second clustering algorithm used is *spectral clustering* via kLines (cf. Fischer and Poland [10]). We have successfully applied this method before (cf. [19, 18]) in settings where the NID is approximated by the so-called Google distance or Web distance. In such settings we are given non-literal objects, i.e., essentially names and not the the literal objects themselves as in the present paper. The Web distance is then based on computing probabilities by determining the frequency of web pages for the individual names and those containing simultaneously two of the given names. We refer the reader to [21] for a comprehensive explanation.

It should be noted that spectral clustering generally requires the transformation of the distance matrix into an adjacency matrix of pairwise similarities (henceforth called similarity matrix). The clustering is then done by analyzing its spectrum.

The results obtained for our data are generally very promising. Since we know the true subtype of the hemagglutinin from the description of the gene sequences used, we could determine the quality of the clustering obtained. Quite often, we arrived at a *perfect* clustering independently of the compressor and of the clustering method used. On the other hand, when including all data or a rather large subset thereof, the clustering obtained is not perfect but the number of errors made is still sufficiently small to make the results interesting. Without going into details here, it can be said that the zlib compressor seems more suitable in this setting than the bzip2 compressor (see Subsection 3.2 for details).

### 2 Background and Theory

As explained in the Introduction, the theoretical basis for computing the distance matrix is deeply based in Kolmogorov complexity theory. In the following we assume the definition of the NID as shown in Equation (1). The definition of the NID depends on the function K which is *uncomputable*. Thus, the NID is *uncomputable*, too.

Using a real-word compressor, one can approximate the NID by the NCD (cf. Definition 1). Again, we omit details and refer the reader to [21].

**Definition 1.** The normalized compression distance between two strings x and y is defined as

$$NCD(\mathbf{x}, \mathbf{y}) = \frac{C(\mathbf{x}\mathbf{y}) - \min\{C(\mathbf{x}), C(\mathbf{y})\}}{\max\{C(\mathbf{x}), C(\mathbf{y})\}}$$

where C is any given data compressor.

Common data compressors are gzip, bzip2, zlib, etc. Note that the compressor C has to be computable and *normal* in order to make the NCD a useful approximation. This can be stated as follows. **Definition 2** ([21]). A compressor C is said to be normal if it satisfies the following axioms for all strings x, y, z and the empty string  $\lambda$ .

(1) $C(xx) = C(x)$ and $C(\lambda) = 0$ ;	(identity)
(2) $C(xy) \ge C(x);$	(monotonicity)
(3) C(xy) = C(yx);	(symmetry)
$(4) \ C(xy) + C(z) \leqslant C(xz) + C(yz);$	(distributivity)

up to an additive  $O(\log n)$  term, with n the maximal binary length of a string involved in the (in)equality concerned.

These axioms are in various degrees satisfied by good real-world compressors like bzip2, PPMZ and gzip, where the latter did not perform so well, as informal experiments have shown (cf. [9]). Also note that in all cases the compressorspecific window or block size determines the maximum usable length of the arguments. As a matter of fact, for our data these axioms seem to be fulfilled.

For our investigations we used the built-in compressors bzip2 and zlib and the ncd function from the CompLearn Toolkit (cf. [5]). After having done this step, we have a distance matrix  $D = (d^{ncd}(x,y))_{x,y \in X}$ , where  $X = (x_1, \ldots, x_n)$  is the relevant data list.

Next, we turn our attention to clustering. First, we shortly outline the hierarchical clustering as provided by the R package, i.e., by the program hclust (cf. [2]). Input is the  $(n \times n)$  distance matrix D. The program uses a measure of dissimilarity for the objects to be clustered. Initially, each object is assigned to its own cluster and the program proceeds iteratively. In each iteration the two most similar clusters are joint, and the process is repeated until only a single cluster is left. Furthermore, in every iteration the distances between clusters are recomputed by using the Lance–Williams dissimilarity update formula for the particular method used.

The methods differ in the way in which the distances between clusters are recomputed. Provided are the *complete linkage method*, the *single linkage method*, and the *average linkage clustering*. In the first case, the distance between any two clusters is equal to the greatest similarity from any member of one cluster to any member of the other cluster. This method works well for compact clusters but causes sensitivity to outliers. The second method pays attention solely to the area where the two clusters come closest to one another. The more distant parts of the clusters and the overall structure of the clusters is not taken into account. If the total number of clusters is large, a messy clustering may result.

The *average linkage clustering* defines the distance between any two clusters to be the average of distances between all pairs of objects from any member of one cluster to any member of the other cluster. As a result, the average pairwise distance within the newly formed cluster, is minimum.

Heuristically, the average linkage clustering should give the best results in our setting, and thus we have chosen it (see also Manning *et al.* [14] for a thorough exposition). Note that for hierarchical clustering the number k of clusters does *not* to be known in advance.

Next, the *spectral clustering* algorithm used is shortly explained. Spectral clustering is an increasingly popular method for analyzing and clustering data by using only the matrix of pairwise similarities. It was invented more than 30 years ago for partitioning graphs (cf., e.g., Spielman and Teng [20] for a brief history and Luxburg [22] for a tutorial). Formally, spectral clustering can be related to approximating the normalized min-cut of the graph defined by the adjacency matrix of pairwise similarities [24]. Finding the exactly minimizing cut is an NP-hard problem.

The transformation of the distance matrix into a similarity matrix is done by using a suitable kernel function. In our experiments we have used the Gaussian kernel function, i.e.,

$$\mathbf{k}(\mathbf{x},\mathbf{y}) = \left(\exp(-\frac{1}{2}\mathbf{d}(\mathbf{x},\mathbf{y})^2/(2\cdot\sigma^2))\right)\,,\tag{2}$$

where  $\sigma$  is the kernel width. As pointed out by Perona and Freeman [17], there is nothing magical with this function. Moreover, it is most commonly used. An advantage of using the Gaussian kernel function is that the resulting similarity matrix is positive definite.

So, the remaining problem is a suitable choice for  $\sigma$ . Unfortunately, the performance of spectral clustering heavily depends on this  $\sigma$ . In the experiments, we compute the mean value of the entries of the distance matrix D and then set  $\sigma = \text{mean}(D)/\sqrt{2}$ . In this way, the kernel is most sensitive around mean(D). Though we are not aware of a theoretical result supporting this choice, it worked remarkably well and further studies are needed to explore the properties of this choice.

The final spectral clustering algorithm for a known number of clusters k is stated below.

#### Algorithm Spectral Clustering

Input: data list  $X=(x_1,x_2,\ldots,x_n),$  number of clusters k Output: clustering  $c\in\{1\ldots k\}^n$ 

- 1. for  $x, y \in X$ , compute the distance matrix  $D = (d^{ncd}(x, y))_{x,y \in X}$
- 2. compute  $\sigma = \text{mean}(D)/\sqrt{2}$
- **3.** compute the similarity matrix  $A = \left(\exp(-\frac{1}{2}d(x,y)^2/(2\cdot\sigma^2))\right)$

4. compute the Laplacian  $L=S^{-\frac{1}{2}}AS^{-\frac{1}{2}},$  where  $S_{ii}=\sum_jA_{ij}$  and  $S_{ij}=0$  for  $i\neq j$ 

5. compute top k eigenvectors  $V \in \mathbb{R}^{n \times k}$ 

6. cluster V using kLines [10]

### **3** Experiments and Results

In this section we describe the data used, the experiments performed and the results obtained.

#### 3.1 Influenza Viruses - The Data Set

We shortly describe the data set used. For any relevant background concerning the biological aspects of the influenza viruses we refer the reader to Palese and Shaw [16] and Wright *et al.* [23].

Influenza viruses were probably a major cause of morbidity and mortality world wide. Large segments of the human population are affected every year. The family of *Orthomyxoviridae* is defined by viruses that have a negative-sense, single-stranded, and segmented RNA genome. There are five different genera in the family of *Orthomyxoviridae*: the influenza viruses A, B and C; *Thogotovirus*; and *Isavirus*. Influenza A viruses have a complex structure and possess a lipid membrane derived from the host cell (cf. Figure 1).

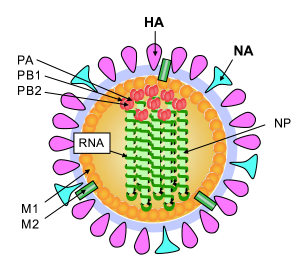


Fig. 1. Influenza A virus

Biologists classify influenza A viruses primarily by their hemagglutinin (HA) subtypes and neuraminidase (NA) subtypes. So far, 16 subtypes of HA are known and commonly denoted by H1, ..., H16. In addition to these HA types, biologists distinguish 9 NA subtypes denoted by N1, ..., N9.

Influenza A viruses of all 16 hemagglutinin (H1-H16) and 9 neuraminidase (N1-N9) subtypes are maintained in their nature host, i.e., the duck. Of these duck viruses, H1N1, H2N2 and H3N2 subtypes jumped into human population, and caused three pandemics in the last century. Therefore, in the experiments performed we have exclusively selected data of influenza viruses that have been obtained from viruses hosted by the duck.

The complete genome of these influenza viruses has 8 segmented-genes. Of these 8 genes, here we are only interested in their HA gene, since HA is the major target of antibodies that neutralize viral infectivity, and responsible for binding the virus to the cell it infects. The corresponding gene is found on segment 4.

Each datum consists of a sequence of roughly 1800 letters from the alphabet  $\{A, T, G, C\}$ , e.g., looking such as

AAAAGCAGGGGAATTTCACAATTAAA....TGTATATAATTAGCAAA.

These gene sequences are publicly available from the National Center for Biotechnology Information (NCBI) which has one of the largest collections of such sequences (cf. [15]).

When analyzed by biologists the definite method to determine the correct HA subtype is based on the antiserum that prevent the docking of the virus. Sometimes biologists also compare the actual sequence to already analyzed sequences and produce a guess based on the Hamming distance of the new sequence to the analyzed ones.

As explained in the Introduction, the primary goal of the investigations undertaken is to cluster the sequences correctly with respect to their HA subtype. In order to achieve this goal with collected from each subtype up to 8 examples. The reason for choosing at most 8 sequences from each type has been caused by their availability. While for some subtypes there are many sequences, there are also subtypes for which only very few sequences are available. The extreme case is the subtype H16 for which only one sequence is in the data base. Figure 2 shows the number of sequences chosen.

I	H1	H2	H3	H4	H5	H6	H7	H8	H9	H10	H11	H12	H13	H14	H15	H16
	8	8	8	8	8	8	8	7	8	8	8	8	2	4	4	1

Fig. 2. Number of sequences for each subtype

It should be noted that most of these sequences are marked as complete cds, but some are also marked as partial cds by the NCBI. For a complete list of the data description we refer the reader to

http://www-alg.ist.hokudai.ac.jp/106Data\_description.html .

For the ease of presentation, below we use the following abbreviation for the data entries. Instead of giving the full description, e.g.,

>gi|113531192|gb|AB271117| /Avian/4 (HA)/H10N1/Hong Kong/1980/// Influenza A virus (A/duck/Hong Kong/938/80(H10N1)) HA gene for hemagglutinin, complete cds.

we refer to this datum as H10N1AB271117 for short.

Among the available files, there were two files containing only a very short partial sequence of the gene, i.e., H7N1AM157391 and H10N4AM922160 (483 and 80 letters, respectively). So, we did not consider these two files, since they do not seem to contain enough information.

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#### 3.2 Results

All experiments have been performed under SuSE Linux. As already mentioned, for the hierarchical clustering we used the open source R package (cf. [2]).

The Algorithm Spectral Clustering from Section 2 has been realized by performing Step 1 via the CompLearn function ncd (cf. [5]). Steps 2 through 6 have been implemented in GNU Octave, version 2.1.72 (cf. [1]). It should be noted that ncd assigns 0.000000 to all elements on the main diagonal of the distance matrix (Version 1.1.5).

By performing our experiments we aimed to answer the following questions. First, does the NCD provide enough information to obtain a correct clustering for the virus data? Second, does the rather large number of clusters (recall that we 16 HA types) cause any problems? Third, do the answers to the first and second question depend on the compressor and clustering, respectively, chosen?

To get started and for the sake of comparison, we used the subset containing all data belonging to H1, H2, and H3, i.e., a total of 24 sequences (cf. Figure 2).

Using the maketree program from the CompLearn Toolkit, we get the following clustering (cf. Figures 3 and 4). As Figures 3 and 4 show, the data are clearly and correctly separated into three clusters. However, the intra-cluster dissimilarities clearly differ from inter-cluster dissimilarities in Figure 3, i.e., for the zlib compressor, while there is no such clear difference for the bzip2 compressor (cf. Figure 4).

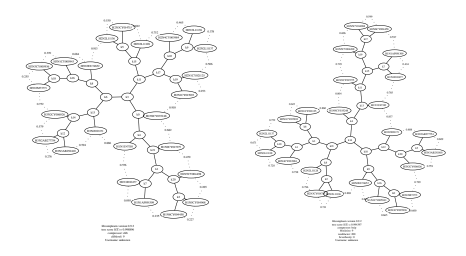


Fig. 3. Classification of HA sequences; Fig. 4. Classification of HA sequences; compr.: zlib compr.: bzip2

Using hclust we obtained the trees shown in Figure 5 and 6 for the matrix D computed for the compressor zlib and bzip2, respectively. As Figures 5 and 6

show, we obtained a correct clustering into three clusters independently of the compressor used.

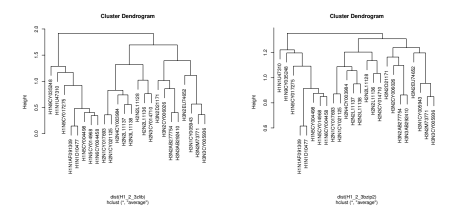


Fig. 5. Clustering all HA sequences for H1 Fig. 6. Clustering all HA sequences for H1 through H3 via hclust; compr.: zlib through H3 via hclust; compr.: bzip2

Next, we tried our algorithm *Spectral Clustering* for the same data set. After having computed the matrix D, we get the following order of the data

```
H2N4CY003984, H3N1CY005943, H3N2AB277754, H1N9CY017275,
H1N9CY035248, H3N3CY005936, H2N2L11128, H2N2L11136,
H2N2L11137, H2N1CY017693, H2N1CY021125, H2N3L11138,
H1N6CY004458, H1N1D10477, H2N3CY014710, H3N2EU74652,
H3N2CY006026, H1N1AF091309, H1N1U47310, H3N3AB292410,
H3N2D21171, H3N2M73771, H1N5CY004498, H1N5CY014968
```

Since spectral clustering is a hard clustering method, it has to return for each data entry just one class label. Assigning canonically the clusters 1, 2, and 3 to the HA subtypes H1N..., H2N..., and H3N..., respectively, we therefore should get the sequence

#### $2 \ 3 \ 3 \ 1 \ 1 \ 3 \ 2 \ 2 \ 2 \ 2 \ 2 \ 1 \ 1 \ 2 \ 3 \ 3 \ 1 \ 1 \ 3 \ 3 \ 3 \ 1 \ 1$

which was indeed returned for both compressors. Note that  $\sigma = 0.56078$  and  $\sigma = 0.57329$  for the zlib and bzip2 compressor, respectively.

Next, we tried all HA sequences for H1 through H8 and from H9 through H16. The reason for this partition has been caused by the different number of sequences available. Recall that there are only two sequences for H13 and only one sequence for H16 (cf. Figure 2).

For H1 through H8 the hierarchical clustering was error free for the zlib compressor but not for bzip2 compressor (1 error) (see Figures 8 and 9 in the

Appendix). Interestingly, for H9 through H16 the tree obtained for the zlib compressor contains 4 errors, while the one obtained for bzip2 compressor has only one error.

Our spectral clustering algorithm returned a perfect clustering for all HA sequences for H1 through H8 for both compressors. On the other hand, for all sequences from H9 through H16 the results differed with respect to the compressor used.

For the **zlib** compressor we obtained 5 errors and for the **bzip2** compressor the number of errors was 7 when using for  $\sigma$  the mean as described above. However, it is well-known that spectral clustering is quite sensitive to the kernel width  $\sigma$ . So, we also tried to vary it a bit around the mean by rounding it to two decimal digits and then changing the second one. For **zlib** the mean was 0.60873 and after two variations we found  $\sigma = 0.59$  which resulted in just one error, i.e., H16 was classified as H13. For the **bzip2** compressor such an improvement could not be obtained.

As a possible explanation we conjecture that one needs a certain minimum of available sequences in order to arrive at a correct spectral clustering. Trying all HA sequences for H1 through H12 kind of confirmed this conjecture, since we again obtained a perfect spectral clustering for both compressors.

For the hierarchical clustering, the tree obtained for the **zlib** compressor is correct, but the the one obtained for the **bzip2** compressor has one error. These trees are shown in the Appendix.

Finally, we tried all data. Again hierarchical clustering was best for the zlib compressor and showed only 2 errors. For the bzip2 compressor, we obtained 3 errors (see the Appendix for details). On the other hand, the best result we could obtain for spectral clustering had 5 errors (for both compressors). In Figure 7 we show the clustering obtained for the zlib compressor for  $\sigma = 0.63$ , where c0 is the desired classification and sp the one returned from the spectral clustering algorithm (partitioned into six groups).

c0 =	7	7	14	2	11	12	12	3	7	10	10	5	9	9	9	3	1	1	9	11
sp =	1.	7	14	2	11	12	12	3	7	10	10	5	9	9	9	3	1	1	9	11
<b>c</b> 0 =	11	5	3	7	5	2	2	2	4	10	5	8	12	2	2	4	4	4	11	9
sp =	11	5	3	7	5	2	2	2	4	10	5	8	12	2	2	4	4	4	11	9
c0 =	10	2	6	6	6	5	1	1	4	10	7	4	8	15	2	9	9	16	10	14
sp =	10	2	13	13	6	5	1	1	4	10	7	4	8	15	<b>2</b>	9	9	3	10	14
c0 =	14	7	7	6	14	7	8	8	12	12	11	15	3	15	5	11	3	1	1	8
sp =	14	7	7	6	14	7	8	8	12	12	11	15	3	15	5	11	3	1	1	8
c0 =	4	3	3	6	12	10	4	5	3	6	13	13	12	1	1	11	12	8	11	10
sp =	4	3	3	6	12	10	4	5	3	6	13	13	12	1	1	11	12	8	11	10
<b>c</b> 0 =	5	9	15	8	6	6														
sp =	5	9	15	8	13	13														

Fig. 7. Clustering all HA sequences via Spectral Clustering; compr.: zlib

So, the errors occur at positions 43, 44, 58, 105, and 106 and affect H6 which is four times assigned to H13 and one time H16 which got in the H3 cluster. We omit further details due to the lack of space.

Note that one can also compute the sum square error (s.s.e.) of all eigenvalues with respect to their means in order to determine quite reliably from the eigenvalues of the Laplacian the number k of clusters (cf. Poland and Zeugmann [19] for details).

### 4 Conclusions

The usefulness of the normalized compression distance for clustering the HA type of virus data for the HA gene for it (segment 4) has been demonstrated. Though we just used the built-in compressors zlib and bzip2 the results are (almost) correct when clustering the resulting distance matrix for the whole data set with hclust or spectral clustering via kLines. What is also remarkable in this context is the robustness with respect to the completeness of the data. As mentioned above, some data contain only a partial cds but this did not influence the quality of the clustering as the results, e.g., H1N1U47310 and H3N2D21171 have only 1000 letters.

We have not reported the running time here, since it is still in the range of several seconds. Though the quartet tree algorithm by Cilibrasi and Vitányi [8] returns a high quality classification, it lacks scalability, since it tries to optimize a quality function, a task which is NP-hard. So, even for the small example including the 24 data for H1, H2, and H3 resulting in  $(24 \times 24)$  distance matrix, it took hours to find the resulting (very good) clustering. In contrast, the clustering algorithms used in this study scale nicely at least up to the amount of data for which the distance matrix is efficiently computable, since they have almost the same running time as the ncd algorithm.

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## Appendix

Here we show the results obtained for the remaining data.

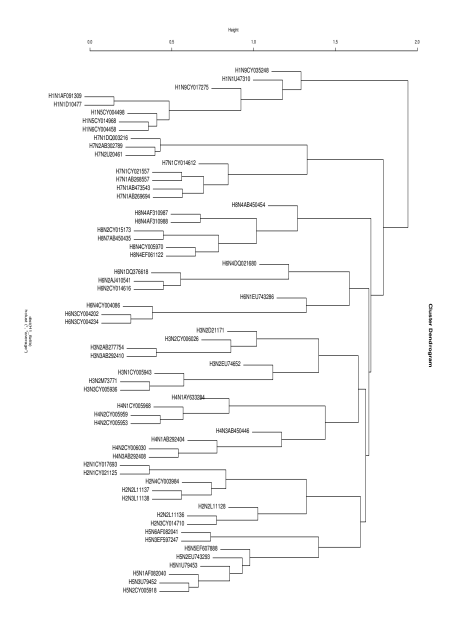
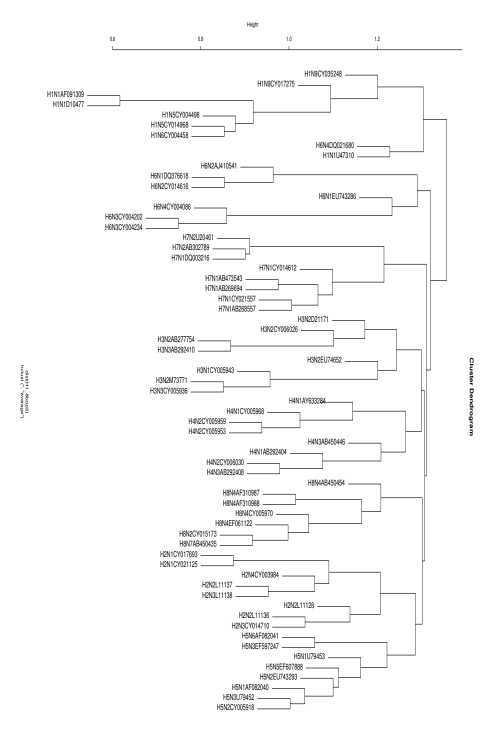


Fig. 8. Clustering of all HA sequences for H1 through H8 via hclust; compr.: zlib



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Fig. 9. Clustering of all HA sequences for H1 through H8 via hclust; compr.: bzip

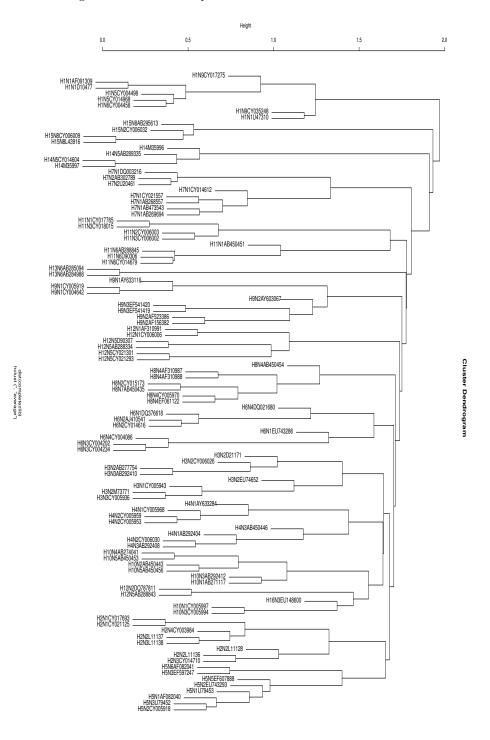
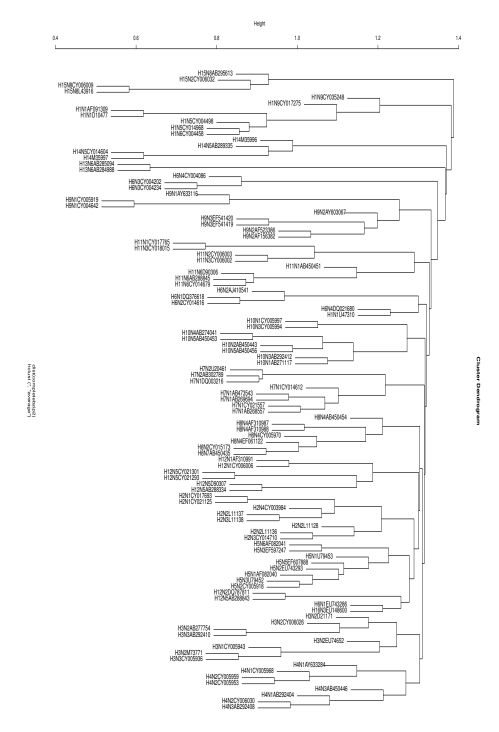


Fig. 10. Clustering of all HA sequences via hclust; compr.: zlib



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Fig. 11. Clustering of all HA sequences via hclust; compr.: bzip