Metagenomic Clustering in Search of Common Origin

Jolanta Kawulok and Michal Kawulok
Institute of Informatics, Silesian University of Technology, Gliwice, Poland

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Abstract: Analysis of metagenomic samples is aimed at extracting relevant information on these samples, including their composition and origin. To determine where a sample comes from, it is commonly compared with a set of reference samples extracted from known locations. However, if such reference samples are unavailable or when the origins of the investigated samples are not covered by the reference set, it may be helpful to identify groups of similar samples that may have a common origin. In this paper, we tackle this problem with hierarchical clustering applied to analyse a matrix of mutual similarities obtained using the Mash and our CoMeta programs. We report initial, yet encouraging results of our experimental study performed for the metagenomic data extracted from two large metropolises, downloaded from the Sequence Read Archive repository. The obtained results indicate that the proposed approach is effective, which justifies further exploration of the topic using more extensive data.

1 BACKGROUND

In recent years, analysis of metagenomic reads (collections of genome fragments derived from microbes living in a given location) has become a hot research topic. Such analysis has a large potential, as it is no longer necessary to isolate and culture organisms in laboratory conditions to study them (Simon and Daniel, 2011; Handelsman, 2004). The majority of the research works are aimed at discovering the composition of the metagenomic samples. They consist in identifying the species of the organisms (taxonomic classification) or in determining the functions that can be performed by the microorganisms from the sample (functional classification) (Bengtsson-Palme, 2018). There are many metagenomic software tools for 16S analysis and shotgun metagenomic analysis (Oulas et al., 2015). The latter data can be analyzed following two kinds of methodological approaches: read-based and assembly-based (Breitwieser et al., 2017). Metagenomic reads may also be subject to binning (Li et al., 2012; Wang et al., 2015), which commonly consists in clustering the reads. This process is aimed at identifying artificial duplicates or grouping similar sequences into species or operational taxonomic units.

Furthermore, metagenomic analysis can be used to predict the place where the samples come from and to create a profile of that place. Walker et al. (2018) used the 16S gene profile for taxonomic classification prior to building the city profiles. Taxonomic analysis for classifying samples to the most probable environment was proposed by Qiao et al. (2018), whose MetaBinG2 program allows for decomposing the complete genome sequence into short substrings composed of k symbols (k-mers). The use of functional classification was explored by Casimiro-Soriguer et al. (2019) and Zhu et al. (2019). Zolfo et al. (2018) used both taxonomic and functional classification for this purpose. For the metagenomic classification, various machine learning techniques are also tested, including random forests, linear discriminant analysis, and support vector machines (Harris et al., 2019; Walker and Datta, 2019).

The aforementioned research works were focused on comparing the query samples with those extracted from known locations. In addition to that, dimensionality reduction techniques, including principal component analysis and t-distributed stochastic neighbor embedding, were employed to visualize the relation between the samples based on their identified species or functions (treated as highly-dimensional features of these samples).

In this paper, we are focused on clustering the metagenomic samples to identify those that may have a common origin. Recently, we investigated such an unsupervised scenario (Kawulok et al., 2019) with no reference samples available, and we exploited our
CoMeta program (Kawulok and Deorowicz, 2015) to determine mutual similarities between the samples. In our earlier research (Kawulok and Kawulok, 2018), we demonstrated that CoMeta can be successfully used to classify the samples by comparing them with entire metagenomic collections derived from reference samples, which allowed us to determine their origin. Contrary to other approaches, we proposed to compare the metagenomic samples by measuring their similarity directly in the space of the reads, which means that it is not necessary to identify the species of organisms that are present in the samples to compute their similarity—hence a reference database with species or functions of microorganisms is not required. After obtaining the mutual similarities, we formed the groups of similar samples manually. The metagenomic samples are a mixture of diverse DNA fragments. Thus, for a number of samples derived from several different locations, it may be expected that appropriate clustering would help identify those that come from the same location.

Compared to our earlier research (Kawulok et al., 2019), here we perform automatic (rather than manual) clustering of the samples to determine those that have a common origin. For this purpose, we employ hierarchical clustering (Rokach and Maimon, 2005) (it is important advantage is that it does not require the number of clusters to be provided in advance); and we consider two different approaches toward analyzing the similarity matrix. Furthermore, in addition to using CoMeta, we also exploit the Mash program to determine the similarities between the samples. The reported results indicate that the clusters can be correctly identified in an automatic way without the necessity of performing taxonomic or functional classification.

2 MATERIALS AND METHODS

2.1 Metagenomic Data

In our experiments, our intention was to verify, if we could cluster the samples, even if their origins are geographically similar to each other. Therefore, we aimed at selecting samples extracted from large cities located relatively close to each other, in which there are many travellers carrying microbes from other places. From the Sequence Read Archive repository 1 (SRA), we selected two projects that provide data of urban metagenome. The first dataset was derived from New York City MTA subway (Afshinnekoo et al., 2015). For our experiments, we have chosen 100 samples from them, each of which contains 0.8 – 11.7 million paired-end reads (together 105.2G bases). The dataset from the second project contains sequences from train cars and subway stations across the Boston subway system (Hsu et al., 2016). For our experiments, we used 23 samples, each of which contains 0.9 – 58.6 million paired-end reads with 102 bp length.

2.2 Data Preprocessing

The SRA repository stores raw sequencing data, therefore it can be expected that the samples acquired from various cities contain highly-similar fragments of the human genome. Therefore, we removed human DNA from the investigated samples. The GRCh38_latest_genomic.fna.gz file (containing human reference genome) was downloaded from the NCBI Website. We filter each metagenome sample using the kmctools software (Deorowicz et al., 2015) —if at least one human k-mer ($k$ = 24) appears in a read, then that read is removed from the sample.

2.3 Research Methodology

The clustering of the metagenomic samples is performed on the basis of their mutual distances. For determining the distances between the samples, we have considered two programs.

The first one is the Mash program which estimates the similarity between two genomes or metagenomes. The program uses the MinHash dimensionality reduction technique to compress k-mer sets of whole genomes (Ondov et al., 2019). In the program, the reads in the sample $i$ must be first sketched with $s$ hashes ($s$ is termed the sketch size). Then, the similarity between two samples is determined using these sketched files by counting the number of overlapping k-mers among all the s hashes.

Apart from Mash, we also used the our CoMeta program to determine the similarities between the samples. First, CoMeta creates k-mer databases for all the reference samples that the query sample is to be compared against. Subsequently, each read derived from a query sample is compared against each other sample (represented by a k-mer database). For each $i$th read and $j$th sample, their similarity is computed as the number of the nucleotides in the k-mers which are present both in the read and in the database (associated with that sample), divided by the length of the query read. For clustering, a k-mer database must be built for every sample, and then the similarity of each sample (treated as a set of reads) to other sample (rep-

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Comparison using CoMeta or Mash

Removal of human DNA from the samples using the kmc tools software

S1 S2 S3 SN

Comparison using CoMeta or Mash

Hierarchical clustering

Grouping of the most similar samples

Figure 1: The processing pipeline for metagenomic reads clustering.

A simplified diagram of our clustering scheme is shown in Figure 1. At the beginning (as described in Section 2.2), the human fragments (S_H) are subtracted from the original metagenomic samples (S^0) using the kmc tools software. As a result, we obtain N samples (S_i) which are smaller than the original ones. The next step is to compare the samples between each other using CoMeta or Mash. From these comparisons, we build a square matrix of similarities (SSM) between the samples. It is worth noting that the Mash program compares the samples using two sketched files, therefore the similarity is symmetrical (SIMS_1S_2 = SIMS_2S_1). Contrary to that, the CoMeta algorithm compares each sample in a read-wise manner to a k-mer database built from another sample. Hence, the similarities are not symmetrical (SIMS_1S_2 ≠ SIMS_2S_1).

The Mash program sketches each file using the same size of a sketch, so despite the fact that the files with reads are of different sizes, the size of each sample is the same after sketching. The CoMeta program builds a k-mer database using the whole sample, therefore the sizes of these databases differ significantly from each other. In the reported research, we test CoMeta program using whole k-mer databases and using reduced databases. The latter are built after reducing each sample to the size of 0.8 million paired-end reads (which is the size of the smallest sample), therefore each sample is represented by a k-mer database of the same size.

For each sample, we obtain a set of N similarities between that sample and the remaining samples. However, the distributions of the similarity values differ significantly for individual samples. This is particularly visible for the scores obtained with CoMeta, where even the values of self-similarity (i.e., the similarity between the sample and a k-mer database created from that sample) are varied. To address that problem, we normalize the similarities in the following way. First, we substitute each value on the diagonal (which contains the self-similarities) with the highest value from the given row:

$$SIMS_iS_i \leftarrow \max\{SIMS_iS_j : i, j \in \{1, N\}, i \neq j\}. \quad (1)$$

Subsequently, each value in the row is divided by that highest value to obtain the distance (DST) between the samples:

$$DST_iS_k = 1 - \frac{SIMS_iS_k}{SIMS_iS_i} : i, k \in \{1, N\}. \quad (2)$$

In this way, we convert the SSM matrix into the square distance matrix (SDM). While for CoMeta we always exclude the self-similarities (1), we treat it as an optional step for Mash, considering two versions here: with self-similarities (WSS) and after excluding self-similarities (ESS).

The distance matrix is subsequently used to identify the groups of samples which are supposed to have the same origin. We consider two variants of exploiting the hierarchical clustering, namely: (original dst)—the distances from the SDM are used as an input for clustering; (recomputed dst)—the values in columns are treated as individual attributes for the samples in the rows (hence each sample is represented with an N-dimensional feature vector containing the distances of that sample to all the samples). In the latter variant, the Euclidean distances between the samples’ feature vectors are treated as the distances between the samples, which forms a new SDM that is subject to hierarchical clustering.

Then, the samples are grouped using hierarchical clustering analysis (HCA). The HCA algorithm starts by treating each sample as a singleton cluster. Then, the following two steps are repeatedly executed: (1) determine a pair of the closest clusters, and
(2) merge them together. This searching-and-merging process is continued until all the clusters (samples) are merged together. The relationship between the clusters is represented by the dendrogram plot of the hierarchical binary cluster tree. In our work, for determining the distances between sets of samples, we use single-linkage clustering criteria, which is the shortest distance.

3 EXPERIMENTAL VALIDATION

Our experimental study was performed using two programs: CoMeta and Mash. For the CoMeta program, we use whole $k$-mer databases and reduced $k$-mer databases, as explained earlier in this paper. The SSM matrix is normalized for both programs, and for Mash we report the results obtained without and with excluding the self-similarities. For HCA, we use original SDM, as well as the recomputed distance matrix.

3.1 Evaluation of Clustering

The clustering outcome can be evaluated taking into account internal or external criteria (Rokach and Maimon, 2005). As an internal quality criterium, for $M$ clusters, we use a sum of squared error (SSE):

$$ D = \frac{1}{2} \sum_{m=1}^{M} N_m D_m, $$

where $N_m = |C_m|$ is the number of instances (here, samples) belonging to the cluster $C_m$, and:

$$ D_m = \frac{1}{N^2_m} \sum_{S_i, S_j \in C_m} d^2(S_i, S_j), $$

where $d(S_i, S_j)$ is the distance between the samples $S_i$ and $S_j$.

The external quality criteria can be useful for examining whether the structure of the clusters matches some predefined classification of the samples. One of the simplest metrics here is the Rand index, which consists in determining the ratio between matched and unmatched observations among two clustering structures—$C_1$, which is an induced clustering structure and $C_2$, which is a given (ground-truth) clustering structure. This index is defined as:

$$ RAND = \frac{a + d}{a + b + c + d}, $$

where $a$ is the number of pairs of samples that are assigned to the same cluster in both structures ($C_1$ and $C_2$); $b$ is the number of pairs of samples that are in the
same cluster in $C_1$, but not in the same cluster in $C_2$; $c$ is the number of pairs of samples that are in the same cluster in $C_2$, but not in the same cluster in $C_1$; and $d$ is the number of pairs of samples that are available in different clusters in $C_1$ and $C_2$. The Rand index value lies between 0 and 1, and it equals 1 when the samples are perfectly separated.

In addition, we inspect the appearance of the ob-

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Figure 3: The dendrogram plots for hierarchical clustering. The dark blue color indicates the samples from Boston, the other colors indicate the samples from New York.
Table 1: The clustering quality scores. $D_{BOS}$ and $D_{NY}$ are cluster-wise SSEs (4), computed for individual clusters of Boston and New York samples, respectively; $D$ is the overall SSE (3); and RAND is the Rand index (5). The three best scores in each column are bolded.

<table>
<thead>
<tr>
<th>Name:</th>
<th>$D_{BOS}$</th>
<th>$D_{NY}$</th>
<th>$D$</th>
<th>RAND</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) CoMeta (whole kmer db, original dst)</td>
<td>0.23</td>
<td>0.44</td>
<td>0.33</td>
<td>0.68</td>
</tr>
<tr>
<td>(b) CoMeta (whole kmer db, recomputed dst)</td>
<td>0.09</td>
<td>0.23</td>
<td>0.16</td>
<td>0.68</td>
</tr>
<tr>
<td>(c) CoMeta (reduced kmer db, original dst)</td>
<td>0.19</td>
<td>0.44</td>
<td>0.32</td>
<td>1.00</td>
</tr>
<tr>
<td>(d) CoMeta (reduced kmer db, recomputed dst)</td>
<td>0.08</td>
<td>0.23</td>
<td>0.15</td>
<td>1.00</td>
</tr>
<tr>
<td>(e) Mash (WSS, original dst)</td>
<td>0.47</td>
<td>0.49</td>
<td>0.48</td>
<td>1.00</td>
</tr>
<tr>
<td>(f) Mash (WSS, recomputed dst)</td>
<td>0.45</td>
<td>0.47</td>
<td>0.46</td>
<td>1.00</td>
</tr>
<tr>
<td>(g) Mash (ESS, original dst)</td>
<td>0.26</td>
<td>0.34</td>
<td>0.30</td>
<td>1.00</td>
</tr>
<tr>
<td>(h) Mash (ESS, recomputed dst)</td>
<td>0.12</td>
<td>0.22</td>
<td>0.17</td>
<td>1.00</td>
</tr>
</tbody>
</table>

The clustering quality scores obtained using data shown in Figure 3 are reported in Table 1. The SSE is the sum of squared error defined in (3). It is computed using additional minimum variance criteria (4), whose values for single clusters are $D_{BOS}$ and $D_{NY}$ for Boston and New York, respectively. The smaller the value is, the greater is the homogeneity within the cluster. RAND is the Rand index defined in (5). The best scores for each parameter are bolded.

Analyzing all plots, we can observe that the use of small $k$-mer databases as opposed to the whole ones allows for correct identification of clusters for the CoMeta program. When whole $k$-mer databases are used, then some additional clusters are induced within the New York samples which can be seen in Figures 2(a) and 3(a, b). Hence, for these two sets of data, the value of Rand index is below 1 in Table 1. This means that unbalanced samples lead to identifying false positive clusters in the results.

From the plots obtained for Mash, we can notice that excluding the self-similarities allows us to separate the samples from both cities more clearly. This could also be noticed in Table 1—the values of $D_{BOS}$, $D_{NY}$ and SSE are smaller for the Mash data after excluding the self-similarities. For more sophisticated data with more ground-truth clusters, this can be crucial for correct cluster identification, as it could be difficult to clearly separate the individual clusters.

Comparing the left and right dendrograms in Figure 3, it can be seen that the recomputed distances between the samples reduce the distances between the samples within each cluster, and it has a similar effect to excluding the self-similarities for Mash. These observations are also confirmed by the scores in Table 1—the homogeneity within the clusters is larger when the distances are refined taking into account the distance features of each sample.

Overall, the presented work clearly indicates that it is possible to automate the process of clustering the samples without identifying the microorganisms de-
rived from them. The best results have been obtained using Mash based on the recomputed distances after excluding the self-similarities (Figure 3(h)). Also, the operation of balancing the samples by reducing the size of the databases allows for obtaining similar results with the CoMeta program (Figure 3(d)). It is worth noting here that such an operation is indirectly performed by Mash, as it builds sketches of a constant size, independently on the sample size.  

4 CONCLUSIONS AND FUTURE WORK

In this paper, we proposed a new approach toward clustering metagenomic reads in search of the samples that have common origin. The results of our experimental study indicate that the presented method allows for separating the samples based on their mutual similarity.

An important advantage of the reported approach lies in determining the sample similarity at the reads level without the necessity to understand the contents of these samples. Therefore, our methodology does not require large databases (taxonomical and functional) of annotated reads. Here, we used two programs (CoMeta and Mash) for comparing the samples prior to clustering, and the results obtained for the best variants of both programs were similar. Importantly, we show that clustering of the metagenomic samples can be automated, which may be extremely important when a larger number of samples is to be processed.

In the presented preliminary research, we used the samples from two large cities located relatively close to each other—Boston and New York. While based on that limited dataset it is difficult to indicate which program is more suitable for clustering, we have demonstrated how important it is to deal with the problem of imbalanced data as well as to preprocess the similarity scores. In our future work, we will extend the database used for evaluation to verify this approach for a larger number of clusters (i.e., ground-truth locations) and increase their diversity.

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