Probabilistic Model for Purity Values of Bacterial Genome Sequences

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Abstract: In sequence analysis, quantifying characteristics of a genomic region is helpful to identify functions of the region, and many computational measures have been proposed. Purity measure is a computational measure, and its potential to characterize horizontally transferred genes has been shown in the literatures. However, in the previous studies, only repeating regions were studied, and also the statistical property of purity values, evaluation values of the measure, has not been uncovered. In this paper, we propose a generative model for \( k \)-mers and evaluate the accuracy of distributions of purity values predicted by the probabilistic model. We train the model for each of 14 bacterial genomes, and our analysis shows that our model predicts the distributions successfully for six genome sequences. It is also shown that our model makes better predictions for shorter \( k \)-mers.

Key words: Sequence analysis, purity measure, generative model, \( k \)-mers, prediction.

1. Introduction

Sequence analysis is a widely used approach for understanding genomes. Traditional methods such as sequence alignments and hidden Markov Models have been successfully applied to this end [1]. For identifying functions of regions of a genome sequence, existing studies have also employed the characteristics of regions and proposed many computational measures for quantifying the characteristics.

One of the approaches to quantify characteristics of genomic regions is to use biological knowledge, and there exist many methods based on various domain knowledge [2]-[4]. Another approach focuses on the compositional characteristics of a sequence and does not need any domain knowledge. This approach includes nucleotide-level composition [5], di-nucleotide abundance [6], probability [7] and complexity [8], [9].

Purity measure [10] is another compositional measure, which was proposed in the field of text mining for finding unusual regions of an input string. It has been shown [11] that this measure certainly characterize genes such as mobile elements, phages, RNAs and transposons which can be considered as horizontally transferred genes [12]-[14] as well as existing measures [15]-[17]. Since horizontal gene transfer is considered as one of the primary reasons of bacterial genetic diversity [18], finding such genes could lead to evidences of the
hypothesis. However, in the previous studies of purity measure, only repeating regions, which appear at least twice in an entire sequence, were examined with the measure. Moreover, the statistical property of the evaluation values of the measure, that we call purity values, has not been uncovered. Thus, currently, there is no definitive way to discriminate regions of horizontal transferred genes from others using purity values. Revealing statistical properties of purity values would enable us, for example, to decide a threshold of purity values for the discrimination in an objective fashion.

In this paper, we aim to find out statistical properties of purity values. To this end, we propose a generative model for \( k \)-mers and evaluate the accuracy of distributions of purity values predicted by the model. Making histograms of the purity values, we have found that they form a bell curve-like distribution. Then we propose a probabilistic model for \( k \)-mers based on a binomial model. We trained our model for each of 14 bacterial genome sequences that we chose, and evaluate the accuracies of predicted distributions of purity values comparing them to the observed distributions.

2. Method

First we explain the definition of purity measure before proposing a probabilistic model for its evaluation values.

2.1. Purity Measure

Given a string \( T \) and a substring \( x \) of \( T \), purity measure quantifies how many substrings of \( x \) appear the same number of times as \( x \). In other words, how many substrings of \( x \) appear only as substrings of \( x \) in \( T \). Obviously, most substrings of \( x \) are much shorter than \( x \), and such substrings are considered to appear more frequently than \( x \) in \( T \). Hence, we could consider \( x \) is unusual if most substrings of \( x \) appear only as parts of \( x \). Yamada et al. [10] proposed three different definitions of purity measure called probability, entropy and difference. In this study, we only use the probability definition as with the previous study [11]. We call the measure based on the definition just purity measure in the rest of this paper.

Formally the purity measure is defined as follows. Let \( N \) be the set of non-negative integers. Let \( \Sigma \) be a finite set of characters. We call \( \Sigma \) an alphabet. We denote a set of finite sequences of zero or more characters by \( \Sigma^* \) and call its element a string. The length of a string \( x \in \Sigma^* \) is denoted by \( |x| \). For a string \( x = a_1 a_2 \ldots a_n \in \Sigma^* \) of length \( n \), the \( i \)-th character \( a_i \) of \( x \) is denoted by \( x[i] \) for a positive integer \( i \), and a contiguous part \( a_i \ldots a_j \) of \( x \) is denoted by \( x[i : j] \) for positive integers \( i \) and \( j \) such that \( i \leq j \) and called a substring of \( x \).

For a string \( x \in \Sigma^* \), \( \text{sub}(x) \) is defined as follows:

\[
\text{sub}(x) = \{(i, j) \in \mathbb{N}^2 \mid 1 \leq i \leq j \leq |x|\}.
\]

For a string \( T \) and a string \( x \in \Sigma^* \), we define \( \text{pos}_T(x) \) as follows:

\[
\text{pos}_T(x) = \{(i, j) \in \text{sub}(T) \mid T[i : j] = x\}.
\]

For a string \( T \) and a string \( x \in \Sigma^* \), \( \text{freq}_T(x) \) is defined as \( \text{freq}_T(x) = |\text{pos}_T(x)| \). Intuitively, \( \text{sub}(x) \) represents a set of the all substrings of \( x \), \( \text{pos}_T(x) \) is a set of occurrences of \( x \) in \( T \), and \( \text{freq}_T(x) \) is the frequency of \( x \) in \( T \).

**Definition 1.** Given an input string \( T \) and a substring \( x = T[i : j] \) of \( T \), the purity value of \( x \) on \( T \) is defined as follows:
This definition of the purity measure quantifies a characteristic of $x$ in $T$ as the fraction of the substrings of $x$ that only appear as parts of $x$ in $T$.

The definition requires us to compute the frequencies of a target string $x$ and its substrings. Employing a naïve way, that is to search a genome sequence for these substrings and to count their occurrences, can easily become impractical when we try to evaluate many regions of a sequence. Special data structures such as suffix trees and suffix arrays provide efficient algorithms that can solve this problem [19], and we can construct a practical algorithm to compute purity values.

2.2. Probabilistic Model

In this section, we discuss the distribution of purity values. Since purity values highly depend on the length of a target substring, we focus only on fixed-length substrings of length $k$, which are called $k$-mers, in this paper. We have found that distributions of the purity values of $k$-mers form bell curve-like distributions which are slightly different from Gaussian distributions. Hence, we propose a probabilistic model of purity values on top of binomial models rather than Gaussian models based on the observations.

Suppose we are given an input string $T$ and an integer $k$, where $0 \leq k \leq |T|$. Let $x$ be a substring of $T$ such that $1 \leq |x| \leq k$, and let $y$ be a substring of $T$ such that $|y| = k$. We call $x$ a specific substring of $y$ if and only if $x$ appears only as a part of $y$ in $T$. Now, the purity value of the substring $y$ can be described as the ratio of specific substrings of $y$ to all the substrings of $y$. Since a $k$-mer always has $n = k(k + 1)/2$ substrings, a distribution of purity values of $k$-mers depends only on the number of specific substrings of every $k$-mer.

Our basic idea is to regard a $k$-mer as a set of $n$ independent substrings. Let’s suppose we randomly choose every element of such a set from either specific substrings or non-specific ones. Assuming choice probability $p$ for a specific substring and $(1 - p)$ for a non-specific substring, a final number $X$ of specific substrings in a assembled set follows a binomial distribution $B(n, p)$, and the probability that the number becomes $l$ is written as $\Pr[X = l] = \binom{n}{l} p^l (1 - p)^{n-l}$. Let $Z$ a $k$-mer chosen from a sequence randomly. The probability that $Z$’s purity value $\text{purity}_T(Z)$ is equal to $q$ is written as follows:

$$\Pr[\text{purity}_T(Z) = q] = \Pr[X = nq] = \binom{n}{nq} p^{nq} (1 - p)^{n-nq}.$$ 

Then, given a set of purity values $\{q_i\}$ of $k$-mers, we can determine the parameter $p$ by maximum likelihood estimation as follows:

$$\hat{p} = \arg\max_p \prod_i \Pr[\text{purity}_T(Z) = q_i] = \frac{\sum_i q_i}{\sum_i 1}.$$ 

Furthermore, we replace the parameter $p$ with another variable $m$ to make the model parameter more intuitive. As a longer substring has more possibility to be a specific substring of some other substrings, we assume that every substring of any $k$-mers whose length is longer than $m$ is a specific substring of the $k$-mer. Therefore, we can denote $p = (k - m)(k - m + 1)/k(k + 1)$. Solving the equation for $m$, we finally obtain an optimal parameter value $\hat{m}$ as

$$\hat{m} = k + \frac{1}{2} \left[\frac{1}{2} \sqrt{1 + 8n \sum_i q_i} \right].$$ 

Finally, our probabilistic model only has a single parameter $m$. We can train our model for a genome sequence by giving a set of purity values of $k$-mers included by the sequence.
3. Result and Discussion

We train our probabilistic model for various bacterial genomes with four different lengths of $k$-mers. Table 1 shows 14 bacterial genome sequences for which our model is trained. We chose the genomes from popular ones so that both gram-positive and gram-negative genomes, and sequences with various lengths and G+C contents are included. We only use a single strand of a genome sequence which is included in a GenBank file retrieved from NCBI’s RefSeq database [20]. We tried $k = 30, 50, 70, 90$ for the lengths of $k$-mers.

Table 1. Bacterial Genome Sequences Used in Our Analysis

<table>
<thead>
<tr>
<th>Accession No.</th>
<th>Length</th>
<th>Organism</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC_000117.1</td>
<td>1,042,519</td>
<td>Chlamydia trachomatis D/UW-3/CX</td>
</tr>
<tr>
<td>NC_000911.1</td>
<td>3,573,470</td>
<td>Synechocystis sp. PCC 6803</td>
</tr>
<tr>
<td>NC_000913.2</td>
<td>4,639,675</td>
<td>Escherichia coli str. K-12 substr. MG1655</td>
</tr>
<tr>
<td>NC_000962.2</td>
<td>4,411,532</td>
<td>Mycobacterium tuberculosis H37Rv</td>
</tr>
<tr>
<td>NC_000964.3</td>
<td>4,215,606</td>
<td>Bacillus subtilis subsp. subtilis str. 168</td>
</tr>
<tr>
<td>NC_002695.1</td>
<td>5,498,450</td>
<td>Escherichia coli O157:H7 str. Sakai</td>
</tr>
<tr>
<td>NC_002946.2</td>
<td>2,153,922</td>
<td>Neisseria gonorrhoeae FA 1090</td>
</tr>
<tr>
<td>NC_003228.3</td>
<td>5,205,140</td>
<td>Bacteroides fragilis NCTC 9343</td>
</tr>
<tr>
<td>NC_007517.1</td>
<td>3,997,420</td>
<td>Geobacter metallireducens GS-15</td>
</tr>
<tr>
<td>NC_008261.1</td>
<td>3,256,683</td>
<td>Clostridium perfringens ATCC 13124</td>
</tr>
<tr>
<td>NC_009882.1</td>
<td>1,257,710</td>
<td>Rickettsia rickettsii str. ‘Sheila Smith’</td>
</tr>
<tr>
<td>NC_010572.1</td>
<td>8,545,929</td>
<td>Streptomyces griseus subsp. griseus NBRC 13350</td>
</tr>
<tr>
<td>NC_012973.1</td>
<td>1,576,758</td>
<td>Helicobacter pylori B38</td>
</tr>
<tr>
<td>NC_015431.1</td>
<td>1,153,998</td>
<td>Mycoplasma mycoides subsp. capri LC str. 95 010</td>
</tr>
</tbody>
</table>

For Each Sequence, an accession number in NCBI’s RefSeq database, sequence length and organism name are shown.

Fig. 1 shows the parameter values of $\hat{m}$ obtained by training for every combination of genome and value of $k$. In the figure, we cannot see clear relationship between $\hat{m}$ and $k$. To see the relationship from a different perspective, we made another plot shown in Fig. 2. This figure shows the same for every combination of sequence length and value of $k$. However, there is still no clear relationship in the figure. For
the most genome sequences, it seems that $\bar{m}$ depends on genomes more than on the lengths of $k$-mers, and thus we could use the same model for prediction of the purity distribution regardless of $k$ for some cases.

In Fig. 3, for every genome sequence, a plot of observed distribution and theoretical one of purity values of 30-mers is shown. It is shown in the figures that our model successfully predicts the distributions of six genomes, NC_000117.1, NC_000911.1, NC_000913.2, NC_000964.3, NC_002695.1, and NC_003228.3, out of 14 genomes. For the rest of sequences, although mean values of purity values are predicted well, variance of the distributions are very different from those of theoretical ones.

![Fig. 2. Relationships between $\bar{m}$ and sequence length.](image)

![Fig. 3. Predicted distributions (red lines) and observed distributions (black areas) of purity values for 30-mers of every genome sequence. We can see that relatively good predictions are made on genomes of NC_000117.1, NC_000911.1, NC_000913.2, NC_000964.3, NC_002695.1, and NC_003228.3. These accession numbers are emphasized in the above figure.](image)

We also emphasized the names of genomes which look better than the others in the previous figure.

We quantitatively evaluate how well our model predicted the distributions. We computed three quantities to measure accuracy of the predictions: overlap ratio, mode gap and variance gap. An overlap ratio is the fraction of common area on a plot. A mode gap and a variance gap are the absolute difference of modes and variances of theoretical distribution and observed one respectively. Fig. 4 shows those three quantities for every genome and $k$ values. We can see the accuracies of the predictions get worse as $k$ gets bigger. It is also shown that the accuracy depends on genome very much.
Fig. 4. Plots of three quantities for measuring accuracy of predictions made by our model.

From the above analysis, we conclude that our model captures the statistical properties of purity values very well for about half of genome sequences we tested though there is a room for improvement in cases of long \( k \)-mers and the rest of genome sequences.

4. Conclusion

We proposed a generative model for \( k \)-mers of genome sequences and analyzed the accuracy of distributions of purity values predicted by the model. Our model is based on a binomial distribution and has only a single parameter which can be easily determined from a set of purity values. We train our model for 14 bacterial genome sequences with different lengths of \( k \)-mers, and the accuracies of the predictions made by the model are investigated with three quantitative measures. Our analysis shows that our model can predict the distribution of purity values very well for six genome sequences and it makes better prediction for shorter \( k \)-mers. We conclude that we have successfully uncovered the statistical properties of purity values at least for a part of bacterial genome sequences. We believe our result will become the foundation of future development of the discriminator of horizontal transferred genes.

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References


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