MetaGeneTack: ab initio detection of frameshifts in metagenomic sequences

Shiyuyun Tang, Georgia Institute of Technology
Ivan Antonov, Georgia Institute of Technology
Mark Borodovsky, Emory University

Journal Title: Bioinformatics
Volume: Volume 29, Number 1
Publisher: Oxford University Press (OUP) | 2013-01-01, Pages 114-116
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1093/bioinformatics/bts636
Permanent URL: https://pid.emory.edu/ark:/25593/s56ht

Final published version: http://dx.doi.org/10.1093/bioinformatics/bts636

Copyright information:
© The Author(s) 2012. Published by Oxford University Press. This is an Open Access work distributed under the terms of the Creative Commons Attribution 3.0 Unported License (http://creativecommons.org/licenses/by/3.0/).

Accessed March 12, 2019 12:35 AM EDT
MetaGeneTack: ab initio detection of frameshifts in metagenomic sequences

Shiyuyun Tang\(^1\), Ivan Antonov\(^2\) and Mark Borodovsky\(^{2,3,4,5,*}\)

\(^1\)School of Biology, \(^2\)School of Computational Science and Engineering, \(^3\)Center for Bioinformatics and Computational Genomics, Georgia Institute of Technology, Atlanta, GA 30332, USA, \(^4\)Department of Biological and Medical Physics, Moscow Institute of Physics and Technology, Dolgoprudny, Moscow Region, Russia and \(^5\)Joint Georgia Tech and Emory University Wallace H Coulter Department of Biomedical Engineering, Atlanta, GA 30332, USA

ABSTRACT

Summary: Frameshift (FS) prediction is important for analysis and biological interpretation of metagenomic sequences. Since a genomic context of a short metagenomic sequence is rarely known, there is not enough data available to estimate parameters of species-specific statistical models of protein-coding and non-coding regions. The challenge of ab initio FS detection is, therefore, two fold: (i) to find a way to infer necessary model parameters and (ii) to identify positions of frameshifts (if any). Here we describe a new tool, MetaGeneTack, which uses a heuristic method to estimate parameters of sequence models used in the FS detection algorithm. It is shown on multiple test sets that the MetaGeneTack FS detection performance is comparable or better than the one of earlier developed program FragGeneScan.

Availability and implementation: MetaGeneTack is available as a web server at http://exon.gatech.edu/GeneTack/cgi/metagenetack.cgi. Academic users can download a standalone version of the program from http://exon.gatech.edu/license_download.cgi.

Contact: borodovsky@gatech.edu

Supplementary information: Supplementary data are available at Bioinformatics online.

Received on June 8, 2012; revised on October 3, 2012; accepted on October 20, 2012

1 INTRODUCTION

Metagenomic sequences are obtained from environmental microbial communities. The short sequences are usually sequenced using next generation sequencing (NGS) platforms such as Roche 454 and Illumina or traditional Sanger sequencing. Each platform produces reads of different lengths and has different error models. 454 sequencing platforms produce reads of \(\sim 450\) nt; errors are usually insertions and deletions (indels) in homopolymer regions. Illumina platform currently generates sequences of length \(\sim 150\) nt with substitutions constituting almost all errors. Sanger sequencing produces reads that may contain both types of errors, and the read length is \(\sim 900\) nt. Before gene calling, quality control methods are used to reduce errors on raw reads (e.g. trimming the error-prone 20–25 nt in Illumina 3’ read ends). Short reads are assembled into longer (shotgun) sequences and contigs. In assembled sequences, the per-nucleotide error rate can be reduced from 0.5% in raw reads to as low as 0.005%. Still, errors may affect \(\sim 3\)% to \(\sim 4.5\)% of genes in assembled sequences (Luo et al., 2012). Even several thousand nucleotides long metagenomic sequences do not carry enough sequence data to accurately estimate parameters of statistical models for protein-coding and non-coding regions. Moreover, the performance of conventional tools of gene prediction and annotation are impaired by indels in protein-coding regions (Hoff, 2009). On the other hand, tools of comparative genomics that have certain power in detection of frameshifts interrupting evolutionary conserved regions rely entirely on comparison with sequences from existing databases; these tools would not help to analyse novel genes and genes having low similarity with known genes (Kunin et al., 2008).

Previously we have developed an algorithm and software program GeneTack (Antonov et al., 2010), an ab initio tool for finding frameshifts in prokaryotic genomes. Since GeneTack requires a species-specific statistical model, it cannot work with sequences shorter than several hundred kilobases. The long enough sequence is necessary for self-training of GeneMarkS (Besemer et al., 2001), the gene finder providing model parameters and gene predictions to GeneTack. Here we introduce an ab initio frameshift finder, MetaGeneTack, designed for metagenomic sequences. MetaGeneTack uses a heuristic method (Besemer et al., 1999) to infer model parameters suitable for analysis of a short sequence at hand (e.g. 400 nt). A recently developed ab initio gene finder, FragGeneScan (Rho et al., 2010), is also able to detect frameshifts in short sequences. We have assessed the performance of MetaGeneTack in tests on short sequences from 18 prokaryotic species. We have shown that MetaGeneTack performs comparably or better in frameshift detection than FragGeneScan.

2 MATERIALS AND METHODS

The idea of the heuristic method for building models of protein-coding regions is that frequencies of oligonucleotides, if cannot be derived directly owing to insufficient sequence length, can be inferred as functions of the sequence GC content. Thus, the oligonucleotide frequencies could be reconstructed as soon as we compute GC content of the short sequence that may serve as an estimate of GC content of the genome the sequence originated from. MetaGeneTack uses the fifth-order polynomial approximations of dependencies of hexamer frequencies on genome GC content derived from data on 582 annotated prokaryotic genomes (for details see Zhu et al., 2010).
Heuristic method is used in two major steps of the MetaGeneTack pipeline. At the first step the pipeline is running MetaGeneMark (Zhu et al., 2010), an ab initio gene finder using the heuristic method in the algorithm of analysis of short metagenomic sequences. This step identifies an initial set of protein-coding genes and selects a corresponding heuristic model (of bacterial or archaeal type of a particular GC content) for each sequence. The input sequence is then split into fragments where predicted genes are located in the same strand. The fragments assigned to the same model are grouped together. At the second step, the selected heuristic models are used in GeneTack for FS prediction in the grouped fragments.

To reduce the number of false positives, MetaGeneTack uses three post-processing filters for the initial FS predictions. Filter I rational is as follows. In a frameshifted gene with two overlapping protein-coding open reading frames (ORFs), the downstream ORF should not possess a functional ribosomal binding site (RBS). Therefore, if a gene predicted in the downstream ORF has a high RBS score (>2.0), the FS prediction is filtered out. Filter II is based on the following observation: in high GC genomes, true FS is separated by a rather long distance from a stop codon terminating the upstream ORF. Therefore, a predicted FS situated on a short distance from the stop codon (|Dθ| < 0.38 – 40, with θ designating GC content in percentage scale) is filtered out. Filter III works as follows: if an FS is predicted too close (<50 nt) to a border of the putative frameshifted gene or to 3’ or 5’ end of the sequence fragment, such FS prediction is filtered out. Filters II and III are applied to fragments with high GC content (θ > 50) and low GC content (θ ≤ 50), respectively. As a training set for assignment of the filters’ parameters, we used genomic sequences of *Escherichia coli*. To produce the program output, the final set of FS predictions is mapped back to the initial metagenomic sequences and combined with gene predictions of MetaGeneMark, thus producing the full list of genes with or without frameshifts.

To evaluate the accuracy of FS detection, we generated multiple test sets accounting for different sequence lengths and error models. To simulate metagenomic data, 18 prokaryotic genomes with GC content ranging from 28% to 75% were cut into 400-nt, 600-nt and 800-nt fragments (see Supplementary Table S1 for genome names). In a given genome, we selected 2000 fragments of each length. Frameshifts (indels) were simulated in coding regions of a certain fraction of the fragment set: 5%, 10% and 20% of all fragments. Selection of the 400-nt as the minimum fragment length is in agreement with the conventional practice where fragments shorter than 400 nt are used for detecting nucleotide polymorphisms and short functional motifs (Wooley et al., 2010). Selecting 5%, 10% and 20% fractions correspond to current estimates of per-nucleotide error rates in metagenomic fragments assembled from raw reads ranging from 0.0065% to 0.03%.

In the simulations, the indels were made in long stretches of coding regions (>200 nt) at a random location, separated by a distance of at least 50 nt from the fragment boundary. If an FS was predicted in the 20 nt vicinity of the true FS position, it was reported as a true positive, otherwise as a false positive.

### 3 RESULTS

Using $A$ to denote the number of all FS predictions, $T$ to denote the number of predicted true positives and $S$ to denote the number of simulated frameshifts, we calculated sensitivity, $Sn = \frac{S}{A}$ and specificity $Sp = \frac{T}{A}$. Note that this definition of specificity, accepted in bioinformatics literature, corresponds to the definition of precision in machine learning publications. Accuracy of MetaGeneTack was compared with accuracy of FragGeneScan (version 1.15, downloaded from http://omics.informatics.indiana.edu/FragGeneScan). FragGeneScan requires users to select a sequencing method likely used for generating the input sequence along with indication of approximate sequencing error rate. We chose the option Sanger sequencing with 0.5%, as it yielded the best results of FragGeneScan among all available options. The $Sn$ and $Sp$ values averaged on the whole set of genomes are shown in Table 1. To give an example of genome specific values of $Sn$ and $Sp$, we provide Supplementary Table S1 for the set of 400 nt fragments with 20% containing FSs. Results are averaged between sets of fragments with FS made by insertions and FS made by deletions (see also Fig. 1).

In terms of $(Sn + Sp)/2$, MetaGeneTack performed better than FragGeneScan by 7–12%. The values of FragGeneScan $Sn$ and $Sp$ differed by ~55 percentage points, whereas for MetaGeneTack, this gap was much smaller. The differences were likely due to different methods of derivation of sequence models and differences in the hidden Markov model architectures.

To assess how effective the filters were, we evaluated MetaGeneTack’s performance produced with various combinations of filters and compared with performance of FragGeneScan on fragment sets with insertion FS (Fig. 1A) and sets with deletion FS (Fig. 1B). Here we show results for 600-nt-long sequences with 20% fragments containing frameshifts. Without filters, the $Sn$ of MetaGeneTack was close to

<table>
<thead>
<tr>
<th>Fragment length</th>
<th>Fragments having FS (%)</th>
<th>FragGeneScan</th>
<th>MetaGeneTack</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$Sn$</td>
<td>$Sp$</td>
<td>Avg</td>
</tr>
<tr>
<td>400 nt</td>
<td>5</td>
<td>79.6</td>
<td>15.8</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>80.5</td>
<td>27.3</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>81.0</td>
<td>43.2</td>
</tr>
<tr>
<td>600 nt</td>
<td>5</td>
<td>81.2</td>
<td>11.7</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>81.8</td>
<td>21.2</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>81.9</td>
<td>35.1</td>
</tr>
<tr>
<td>800 nt</td>
<td>5</td>
<td>81.9</td>
<td>9.1</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>82.6</td>
<td>16.9</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>82.8</td>
<td>29.4</td>
</tr>
</tbody>
</table>

Values are averaged among genomes and then averaged between insertion and deletion FS sets (see Supplementary Table S1 for details)
FragGeneScan while the Sp was >10% higher in both cases. With the filters, the average Sn and Sp of MetaGeneTack increased by ~5 percentage points. Similar results were observed when an FS prediction was reported as a true positive if located within 10 nt from the simulated FS (data not shown). The distribution of the distance between predicted FS positions and true FS positions is shown in Supplementary Figure S1. The standard deviation is 10.3 and 12.6 for MetaGeneTack and FragGeneScan, respectively.

4 CONCLUSION
The new software program, MetaGeneTack, addresses the challenging question of predicting frameshifts in protein-coding regions of metagenomic sequences without extrinsic knowledge. An advantage of ab initio approach is the ability to detect frameshifts in genes of orphan proteins that do not have known homologs. We have shown that the accuracy of MetaGeneTack is comparable or better than the accuracy of the ab inito gene prediction tool FragGeneScan. Most of the frameshifts predicted by MetaGeneTack are supposed to result from sequencing errors; still like GeneTack, the program is also able to detect frameshifts caused due to indel mutations and ones related to recoding (programmed frameshifts involved in gene regulation). MetaGeneTack could be integrated into pipelines of metagenomic sequence annotation.

ACKNOWLEDGEMENT
Authors would like to thank Alex Lomsadze and Chengwei Luo for valuable discussions.

Funding: This work was supported in part by NIH grant HG00783 to M.B.

Conflict of Interest: none declared.

REFERENCES