

Authors' Instructions: Preparation of Camera-Ready Contributions to SCITEPRESS Proceedings

First Author Name¹, Second Author Name¹ and Third Author Name²

¹*Institute of Problem Solving, XYZ University, My Street, MyTown, MyCountry*

²*Department of Computing, Main University, MySecondTown, MyCountry*
{f_author; s_author}@ips.xyz.edu, t_author@dc.mu.edu

Keywords: The paper must have at least one keyword. The text must be set to 9-point font size and without the use of bold or italic font style. For more than one keyword, please use a comma as a separator. Keywords must be titlecased.

Abstract: The abstract should summarize the contents of the paper and should contain at least 70 and at most 200 words. The text must be set to 9-point font size.

1 INTRODUCTION

Algorithms that can efficiently and accurately identify and classify bacterial taxonomic hierarchy have become a focus in computational genetics. The idea that secondary structure of genomic sequences is sufficient for solving the detection and classification problems lies at the heart of many tools (Rivas and Eddy, 2000; Knudsen and Hein, 1999; Yuan et al., 2015; Dowell and Eddy, 2004). The secondary structure can be specified in terms of formal grammars. The sequences obtained from the real bacteria usually contain a huge number of mutations and “noise” which renders precise methods impractical. Probabilistic grammars and covariance models (CMs) are a way to take the noise into account (Durbin et al., 1998). For example, CMs are successfully used in the Infernal tool. Neural networks is another way to deal with “noisy” data. The works (Sherman, 2017; Higashi et al., 2009) utilize neural networks for 16s rRNA processing and demonstrate promising results.

2 PROPOSED SOLUTION

We combine neural networks and ordinary context-free grammars to detect genomic sequences. We extract features by using the ordinary (not probabilistic) context-free grammar and use the dense neural network for features processing. Features can be extracted by any parsing algorithm and then presented as a boolean matrix M such that $M[i, j] = 1$ iff $S \Rightarrow_G^*$

$w[i, j]$ where w is the input sequence and G is context-free grammar with the start nonterminal S .

```
s1: stem<s> any

a_0_7 : any*[2..10]

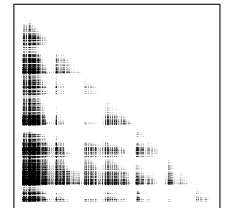
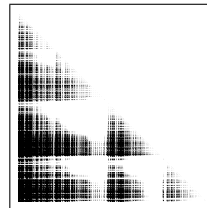
s0: a_0_7 | a_0_7 stem<s> s0

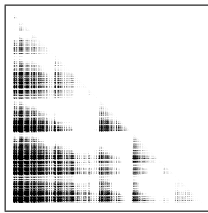
any: A | U | C | G

stem1<s>: A s U | G s C | U s A | C s G

stem2<s>: stem1< stem1<s> >

stem<s>:
    A stem<s> U
    | U stem<s> A
    | C stem<s> G
    | G stem<s> C
    | stem1< stem2<s> >
}
```





3 EVALUATION

We evaluate the proposed approach for 16s rRNA detection. We specify context-free grammars which detect stems with the height of more than two pairs and their arbitrary compositions. For network training we use dataset consisting of two parts: random subsequences of 16s rRNA sequences from the Green Genes database form positive examples, while the negative examples are random subsequences of full genes from the NCBI database. All sequences have the length of 512 symbols, totally up to 310000 sequences. After training, current accuracy is 90% for validation set (up to 81000 sequences), thus we conclude that our approach is applicable.

4 FUTURE WORK

The presented is a work in progress. The ongoing experiment is finding all instances of 16s rRNA in full genomes. Also we plan to use the proposed approach for the filtration of chimeric sequences and the classification. Composition of our approach with other methods and tools as well as grammar tuning and detailed performance evaluation may improve the applicability for the real data processing.

5 DISCUSSION

Protenomics
Different lengths
Other types of NNs

5.1 Manuscript Setup

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ACKNOWLEDGEMENTS

The research was supported by the Russian Science Foundation grant 18-11-00100 and a grant from Jet-Brains Research.

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