

# 16s rRNA Detection by Using Neural Networks

Neural networks for secondary structure information processing

Semyon Grigorev<sup>1</sup>, Polina Lunina<sup>1</sup>

<sup>1</sup>Saint Petersburg State University, JetBrains, St. Petersburg, Russia **E-mail:** semen.grigorev@jetbrains.com



#### Motivation

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 [?, ?, ?, ?]. 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 [?]. For example, CMs are successfully used in the Infernal tool. Neural networks is another way to deal with "noisy" data. The works [?, ?] utilize neural networks for 16s rRNA processing and demonstrate promising results.

#### Results

- We propose the graph parsing algorithms based on different parsing techniques [?, ?, ?].
- We solve some problems of existing approaches (such as cycles processing problem, [?]).
- Our solution provides an ability to use GPGPU and multi-core systems for graph parsing which can be useful for large biological data analysis.

Performance comparison of context-free querying algorithms

Graph	#edges	#results	GLL(ms)	GPGPU(ms)
$g_1$	8688	141072	1926	82
$g_2$	14712	532576	6246	185
(In	15840	449560	7014	127

## Context-free path querying

#### Grammar

s1: stem<s0> any

 $a_0_7 : any*[2..10]$ 

s0: a\_0\_7 | a\_0\_7 stem<s0> s0

any: A | U | C | G

stem1<s>: AsU | GsC | UsA | CsG

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>>

Fixed context-free grammar describes features of secondary structure.

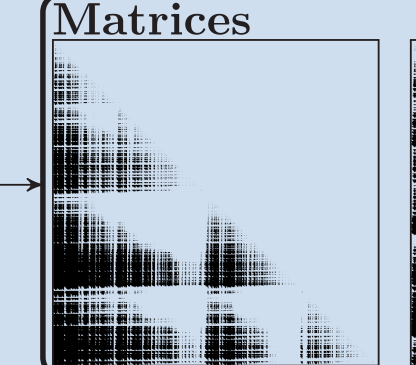
### Sequences

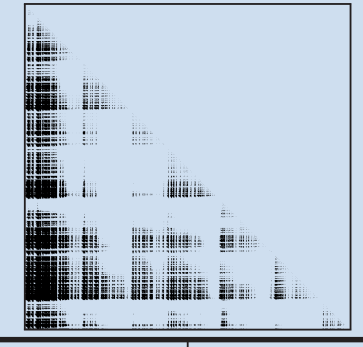
Parts of genom of fixed lengs. Current length is 512.

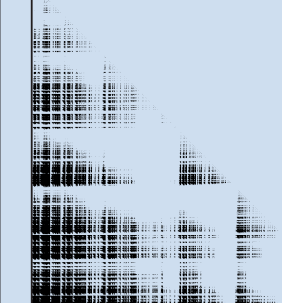
Result of classification

Parser

→Extracts features of secondary strcture. Utilize GPGPU.







Currently just binary classifier that separates -16s and non-16s sequences.

-DNN←

Vectors Line-by-line compressed matrix representation. Shoulb be treated as vector of features.

### Database querying

One of the examples of database querying is an analysis of graphs where vertices correspond to entities and concepts such as gene or phenotype while edges represent the known relationships such as "codes for", "interacts with", etc.

Example of graph structured data [?] is presented below.

Querying paths with special constraints may shed light upon unknown before links between vertices, forming the basis for new hypotheses.

### References

#### Metagenomic assemblies analysis

Metagenomic assemblies can be presented as graph structured data. Some sequences have specific secondary structure, which can be described in terms of a context-free grammar, and this grammar can be used for searching and classification.

## Acknowledgments

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#### Information

All materials available on GitHub: https://github.com/YaccConstructor