Sergey Nurk

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INFORMATION Homepage: http://cab.spbu.ru/employees/sergey-nurk/

Phone: +1-416-824-24-12

 ${\bf Biographical}$

Date of birth: 8th March 1989

Data Citizenship: Russia Marital status: Married

EDUCATION

Saint-Petersburg State University

Ph.D candidate

• Expected degree: Ph.D. in Bioinformatics

• Advisor: Pavel A. Pevzner

• Area of study: de novo microbial (single-cell) and metagenomic assembly

• Expected defense date: Summer 2018

St. Petersburg State University, Mathematics and Mechanics Faculty, Software Engineering Department

M.Sc. in Computer Science, 2006–2011

• Thesis: "Development of de Bruijn graph processing algorithms for genome assembly problem" (Advisor: Nikolay I. Vyahhi)

• Diploma cum laude

Lyceum of Mathematics and Physics #239, GPA 5.0 (2002–2006)

ADDITIONAL

Academy of Modern Software Engineering

EDUCATION

 $\begin{array}{c} {\rm Student} \\ 2008-2010 \end{array}$

EMPLOYMENT

• Center for Algorithmic Biotechnology

St. Petersburg State University

Junior research fellow January 2015-present

• Algorithmic Biology Laboratory

St. Petersburg Academic University

Junior research fellow February 2011–January 2015

• Yandex company

Search engine company in Russia, developing various Internet-based services

Structured Web Mining Department

Software Engineer (Java)

2009 – 2011

• Digital SPomics

Co-founder

January 2014-present

Areas of interest

- genome assembly
- metagenomics

- comparative genomics
- software design
- data analysis

Research Papers

1. Sergey Nurk, Yury Gorshkov and Pavel A. Pevzner

"Reconstructing individual genomes from multiple metagenomic samples with MTS pipeline."

Manuscript in preparation

 Sergey Nurk*, Dmitry Meleshko* (*equal contribution), Anton Korobeynikov, and Pavel A. Pevzner

"metaSPAdes: a new versatile metagenomic assembler." Genome Research, 27(5), 824-834, 2017

3. Yu Lin, Sergey Nurk and Pavel A. Pevzner

"What is the difference between the breakpoint graph and the de Bruijn graph?".

BMC genomics, 15(6), S6, 2014

4. Gabriela I Guzman, Jose Utrilla, **Sergey Nurk**, Elizabeth Brunk, Jonathan M Monk, Ali Ebrahim, Bernhard Palsson, Adam M. Feist

"Model-driven discovery of underground metabolic functions in Escherichia coli".

Proceedings of the National Academy of Sciences, 112(3), 2015

 Andrey D. Prjibelski, Irina Vasilinetc, Anton Bankevich, Alexey Gurevich, Tatiana Krivosheeva, Sergey Nurk, Son Pham, Anton Korobeynikov, Alla Lapidus and Pavel A. Pevzner

"ExSPAnder: a universal repeat resolver for DNA fragment assembly". Bioinformatics, 30(12), i293-i301, 2014

6. Sergey Nurk*, Anton Bankevich* (*equal contribution), Dmitry Antipov, Alexey A. Gurevich, Anton Korobeynikov, Alla Lapidus, Andrey D. Prjibelski, Alexey Pyshkin, Alexander Sirotkin, Yakov Sirotkin, Ramunas Stepanauskas, Scott R. Clingenpeel, Tanja Woyke, Jeffrey S. Mclean, Roger Lasken, Glenn Tesler, Max A. Alekseyev, and Pavel A. Pevzner

"Assembling Single-Cell Genomes and Mini-Metagenomes From Chimeric MDA Products".

Journal of Computational Biology 20(10), 2013

7. Jeffrey S. McLean, Mary-Jane Lombardo, Michael G. Ziegler, Mark Novotny, Joyclyn Yee-Greenbaum, Jonathan H. Badger, Glenn Tesler, Sergey Nurk, Valery Lesin, Daniel Brami, Adam P. Hall, Anna Edlund, Lisa Z. Allen, Scott Durkin, Sharon Reed, Francesca Torriani, Kenneth H. Nealson, Pavel A. Pevzner, Robert Friedman, J. Craig Venter and Roger S. Lasken

"Genome of the pathogen Porphyromonas gingivalis recovered from a biofilm in a hospital sink using a high-throughput single cell genomic platform".

Genome research, 23(5), 867-877

8. Anton Bankevich*, **Sergey Nurk*** (*equal contribution), Dmitry Antipov, Alexey Gurevich, Mikhail Dvorkin, Alexander Kulikov, Valery Lesin, Sergey Nikolenko, Son Pham, Andrey Prjibelski, Alexey Pyshkin, Alexander Sirotkin, Nikolay Vyahhi, Glenn Tesler, Max Alekseyev and Pavel Pevzner

"SPAdes: a New Genome Assembler and its Applications to Single Cell

Sequencing".

Journal of Computational Biology 19(5), 2012

9. Sergey Nurk

"An $O(2^{0.4058m})$ Upper Bound for Circuit SAT". PDMI preprint 10/2009, 2009

ATTENDED AND WORKSHOPS

- STUDENT SCHOOLS Systems Biology Workshop. Saint-Petersburg, Russia, 2015
 - Systems Biology Workshop. Saint-Petersburg, Russia, 2014
 - Microsoft School on Algorithms for Massive Data (ALMADA). Moscow, Russia, 2013
 - Russian Summer School in Information Retrieval (RuSSIR). Voronezh, Russia, 2010
 - Microsoft Data Structures and Algorithms School (MIDAS). Saint-Petersburg, Russia, 2010
 - NoNA Summer School on Complexity Theory. Saint-Petersburg, Russia, 2009
 - Joint Advanced Student School (JASS). Saint-Petersburg, Russia, 2009.

Topic: Propositional Proof Complexity

Talks

Talk "metaSPAdes: a New Versatile Metagenomic Assembler"

 20^{th} Conference on Research in Computational Molecular Biology (RECOMB-2016). Los Angeles, USA, 2016.

- Talk "Probabilistic models in molecular sequence analysis"
 - Bioinformatics summer school
 - St. Petersbrug, Russia, 2014
- Talk "De Bruijn graphs and genome assembly algorithms"

Bioinformatics summer school

Moscow, Russia, 2013

• Talk "Assembling Genomes and Mini-metagenomes from Highly Chimeric Reads"

17th Conference on Research in Computational Molecular Biology (RECOMB-2013). Beijing, China, 2013.

- Talk about a web-harvesting tool developed by my department at Yandex Yet Another Conference (YaC'2010)
 - Moscow, Russia, 2010
- Talk "Upper bound for Circuit SAT"

Estonian Theory Days 2009

Palmse, Estonia, 2009

- Talk "Lower bounds for k-DNF resolution on random 3-CNFs" Joint Advanced Student School (JASS'2009)
 - St. Petersburg, Russia, 2009

TEACHING

"Molecular sequence analysis" course

EXPERIENCE

St. Petersburg Academic University, Spring 2015

"Comparative genomics" student seminar Supervisor

St. Petersburg Academic University, Spring 2014

"Molecular sequence analysis" course Lecturer

St. Petersburg Academic University, Spring 2014

"Java programming language" course Teaching assistant

St. Petersburg Academic University, 2012 - 2013

RELEVANT SKILLS

- C++
- Java
- Python
- R
- \bullet SQL
- NGS genomics stack
- LATEX, bash, git, vim

LANGUAGES

English: fluentRussian: native