# SIDDHARTH S. TOMAR

## PERSONAL INFORMATION

Born in India, 08 May 1994

email siddharth.tomar@scilifelab.se

website https://www.scilen.se

github siddharthst

phone (Swe) +46-702759580 · (Ind) +91-8894636427

EDUCATION

2016-2018 KTH Royal Institute of Technology

Master of Science Master's programme in Molecular Techniques in Life Science, a joint program with Karolinska Institutet and

Stockholm University Duration: 2 years (full time) Date of graduation: 31-May-2018

Country: Sweden

2012-2016 Jaypee University of Information Technology

Bachelor of GPA: 8.6 out of 10 · Specialization in Bioinformatics

Technology Duration: 4 years (full time)

Date of graduation: 09-July-2016

Country: India

Additional electives: Cryptography · High Performance Computing · Systems Biology

BIOINFORMATICS SKILLS

Workflows Genome/Transcriptome assembly, Quantification, Differential expression, Novel gene discovery

Programming C++/C (INCLUDING CUDA AND MPI), JAVA, PYTHON, PERL, R

Libraries Pandas, Dask, Matplotlib, Altair

Databases MySQL

Machine learning Scikit-learn, Tensorflow

Documentation LATEX, MARKDOWN

Version control GIT (Including automated testing and CI), Snakemake

Workflows Time series analysis, Weighted correlation network analysis

Model organisms Human, Mice, P. abies

LAB SKILLS

Gene cloning using bacteria, Gene disruption and gene replacement in bacteria

Engineering General

DNA isolation, RNA isolation, Plasmid and genomic DNA isolation, PCR, Sanger and Illumina sequencing

COURSEWORK - KEYWORDS

Computer science Algorithm design and analysis, High performance computing, Cryptography, Object Oriented Programming,

Relational Database Management Systems, Machine Learning - Support Vector Machines and Artificial Neural

Networks

Molecular biology

Systems biology, Genetics, translational research in cardiology, neurodegenerative disorders, homology search, multiple sequence alignment, phylogeny reconstruction, protein topology prediction, biophysical chemistry, molecular dynamics, comparative genomics, proteomics, mass spectrometry, antibody assays, short and long read sequencing, mircoarrays, bead based assays, droplet barcoding, single cell sequencing, ChIP sequencing

#### RESEARCH EXPERIENCE

## Jun'18-present Karolinska Institutet - Sweden

SciLifeLab -Stockholm I am currently working as a research assistant in Science For Life laboratory, Stockholm. My current work builds upon my experience in my master's thesis and is more focused on the developmental dynamics of PIWI-interacting RNA (piRNA) in mice and their interplay with transposons. I mainly use Pandas for storing and operating on sequence alignment data, coupled with matplotlib and Altair for static and interactive visualisations. I have also repurposed algorithms and methods used in signal processing for the genomic annotation and discovery of novel elements in mice genome.

# Jun–Aug '17 Karolinska Institutet - Sweden

SciLifeLab -Stockholm After being awarded the SciLifeLab Summer Fellowship, I joined Dr Claudia Kutters group at the Karolinska Institute/SciLifeLab. My work focused on annotating various RNA classes from mice liver at different developmental stages by using total RNA sequencing data. This project involved methods for discovering novel developmental stage-specific noncoding RNAs and correlation with other meta-resources to mark functionality. I mainly used Pandas along with Python REST interface to automatically acquire and create relational data links for my genes, thereby annotating them using multiple meta-resources in using a robust and reproducible framework. During this time I also explored the use of Python interactive visualisation library Bokeh for exploratory analysis. Reference: Hassan Foroughi Asl +46-702759580 · hassan.foroughi@scilifelab.se

## Jun-Aug '15 Institute of Oncology Research - Switzerland

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I worked under Dr Francesco Bertoni's group at Institute of Oncology Research, Bellinzona, Switzerland on performance profiling of short read aligners (using Nvidia CUDA framework) for testing scalability in GPGPU environment. I studied the impact of GPU based tools on the next generation sequencing analysis pipeline and included cost versus performance analysis based on scaling.

Master's thesis

Micro-evolution of regulatory RNAs

In this master thesis project, I investigated the functional contribution of regulatory RNA in speciation in closely related mice species. This involved studying the dynamics of their species-specific gene expression by utilising ChIP sequencing to define active promoters along with short and total RNA sequencing.

Supervisor: Claudia Kutter

Bachelor's thesis

Improving the computation of positive selection using codon models

In my bachelor's project, I was working on improving the computation of positive selection using codon models. In this project, I implemented the Branch-Site Model on GPU architecture, and test the efficiency gains which are potentially possible. I changed the underlying algorithm to reduce the redundancy and increase the sensitivity by optimising the subtree pruning. I mainly used (CUDA)C++ for this project. The primary challenge of this project was to implement the current model as it is, without any heuristics involved, and still archive exceptionally low runtimes.

#### **PROJECTS**

2017 KTH ROYAL INSTITUTE OF TECHNOLOGY

Course Project

Assembling reference transcriptome for *P. abies* using massively parallel and PacBio real time sequencing data. This project involved utilization of large amount of data (>700 million reads) and optimization techniques for transcriptome assembly.

Reference: Lars Arvestad · +4687906436 · lars.arvestad@scilifelab.se

2017 STOCKHOLM UNIVERSITY

Course Project

Constructing a SVM based predictor for classifying topological features of  $\beta$  barrels based on their sequence.

2014 Jaypee University of Information Technology

Course Project

Identification of novel leads against dimeric interface(DBP) in *P. vivax* using computer-aided drug design. For this project, I investigated the structural dynamics of Duffy binding proteins with respect to the phylum Apicomplexa.

2013 Jaypee University of Information Technology

Course Project

Created an application written in Java for searching repeat content in a given genome. Simple machine learning technique was used, applying the Hidden Markov Model. Lempel–Ziv–Welch algorithm was also used and modified according to requirements. Perl was used as a wrapper and as text acquisition/annotation search engine.

POSTER PRESENTATION AND CONFERENCES

2018 2<sup>nd</sup> Uppsala Transposon Symposium

Poster Micro-evolution of regulatory RNA in M. domesticus and M. castaneus

GRANTS

OTHER INFORMATION

Scholarships 2017 · Science for Life Laboratory summer fellowship

 $2017~\cdot~$  Karolinska Institutet Foundation scholarship

Languages English · Fluent (C2/ TOEFL iBT score 108)

HINDI · Native

Swedish · Beginner

Last updated on May 3, 2019