**Hirak Sarkar**

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**OBJECTIVE**

I am interested in applying machine learning techniques such as Bayesian inference and deep learning to analyze and extract information from large corpus in the field of genomics, social and computer networks.

**EDUCATION**

**Stony Brook University (SBU) Stony Brook,NY**

*Ph.D in Computer Science (3.99/4) 2014-2019(exp)*

**Indian Statistical Institute Calcutta, India***M.Tech in Computer Science (1st class Hons.) 2011-2013*

**West Bengal University of Technology Calcutta, India  
(Kalyani Govt. Engg. College)***B.Tech in Computer Science (8.88/10) 2007-2011*

**EXPERIENCE**

COMBINE-Lab (Stony Brook University)  
*Research Assistant (*[*https://github.com/COMBINE-lab*](https://github.com/COMBINE-lab)*) Jan 2015-Present*

* Application of machine learning methods for publicly available massive genomic databases. (Python, sklearn)  
  - Development of SeaDragon (under development) involved application of different dimensionality reduction techniques, and gradient boosted trees for detection of population type from GEUVADIS dataset.
* Development of graph based k-mer mapper, Pufferfish (C++)   
  - Genome sequences (string in the order of gigabytes) are difficult to index and search in bounded memory, used a minimum perfect hash based, rank-select algorithm to implement a fast query scheme for nucleotide sequences.
* Developed an intermediate solution for accurate mapping of read sequences. (C++)   
  - Alignments involves dynamic programming and therefore are costly. Mapping of reads are first yet not accurate, to carry best of the both worlds we developed a selective-alignment based algorithm.
* Developed alignment free methods for sequence reads. (C++)   
  - We developed *RapMap,* an ultra fast mapper which builds a suffix array over transcriptomic sequences.
* Graph based clustering for novel organisms. (C++, Python)
* State of the art compression tool for RNA-seq reads. (C++)

Collaboration Siepel-Lab (Cold Spring Harbor Lab) *June 2016-Aug 2016*

* Developed probabilistic graphical model for inferring transcription rate from multi-assay dataset. (Python)

Collaboration with Wings Lab (Stony Brook University) *Aug 2017-Present*

* Data driven inference models for spectrum sensing. (Python)

**PUBLICATION**

* Journals
* Bioinformatics’16,17 (*impact factor*: 7), Journal of Theoretical Computer Science’15 (*impact factor*: 0.8)
* Conferences
* ISMB’16, RECOMB-seq’16, WALCOM’13
* Posters
* BioData’16, WABI’17

**AWARDS**

* Special CS Chair Fellowship ($10K) from *SBU*, Post-Graduate Scholarship from *Govt. Of India*

**SKILLS**

* Programming: *C++, Python, R*
* Data analysis: *Jupyter,* *Pandas*
* Machine Learning Tools: *sklearn, tensorflow*