## TIMOTHY M. FARRELL

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OBJECTIVE SUMMARY Full-time bioinformatics engineer position in biotech industry.

Biotech engineer with computational/ software engineering skill. Most interested in the

intersection of bioengineering and automation/ computation.

**EDUCATION** 

Master of Science, Bioinformatics, Boston University, 2014-2016

Project: Clinical sequencing classifier for highly-polymorphic genes

Activities: West End Boys and Girls Club STEM Tutor

GPA: 3.02/4.0

Bachelor of Science, Biomedical Engineering, Rutgers University, 2008-2012

Project: Optimization of localizable stem-cell immunotherapeutic
Activities: Army ROTC, Rutgers Future Scholars Mentorship Program

GPA: 3.31/4.0

**S**KILLS

Laboratory: genomic library prep, PCR, ELISA, confocal microscopy, cell culture (some)

Math/ Stats/ CS: graph theory, inference/ statistical modeling, linear algebra, machine learning,

stochastic modeling, time series

Bioinformatics: standard tools (samtools, bedtools, etc.), NGS/3GS experiment design/ data

analysis, pipeline development/ management

Programming:

Data management: SQL, PostgresSQL, SQLite Operating systems: Unix-like (preferred), Windows

Data analytics: pandas/ matplotlib, R/ ggplot, high-performance computing

Languages: Python, shell, R, Java/C++, Haskell/ ML, Javascript/ HTML/ CSS (some)

RELEVANT EXPERIENCE

Biomedical Informatics Intern, Aug 2015 - May 2016

Dept. of Biomedical Informatics (DBMI), Harvard Medical School (HMS)

Worked to develop a clinical sequencing assay/ classifier for structurally-variant antigens, using both NGS/3GS technologies. Supervised by Peter Tonellato, PhD (DBMI, HMS) and William Lane, MD PhD (Dept. of Pathology, Brigham Women's Hospital).

Research Assistant, Sept 2015 - May 2016

Dept. of Health Sciences, Boston University

Built preprocessing and preliminary analysis pipeline for dynamic functional connectivity study of Human Connectome Project data (Jason Bohland, PhD). Produced a fast, memory-efficient system that processed 2000 brain images in under 6 hours on a HPC cluster. See repository.

Bioinformatics Intern, May - Aug 2015

Genomic Research Divison, New England Biolabs (NEB)

Investigated development of error mitigation applications for emerging third-generation sequencing technologies (3GS). Setup and streamlined sequencing and computational workflows, facilitating future related work at NEB.

Student-Researcher, Aug 2011 - May 2012

Dept. of Biomedical Engineering, Rutgers University

Worked to optimize the immunomodulatory function of a mesenchymal stem cell-based therapeutic, designed to treat localized inflammatory disease.