Nur M Shahir, PhD

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EDUCATION

University of North Carolina at Chapel Hill, Chapel Hill, NC, USA

• Ph.D. in Bioinformatics and Computational Biology

Aug 2014 - May 2020

- Dissertation: Inflammatory Bowel Disease Differentially Affects Region Specific Composition and Aerotolerance Profiles of Mucosally-Adherent Bacteria
- · Adviser: Dr. Terrence S. Furey, Dr. Shezad Z. Sheikh
- Committe: Dr. Michael I. Love, Dr. Ian Carroll, Dr. Yufeng Liu
- Focus: Inflammatory Bowel Disease, gut microbiota, bioinformatics, 16S amplicon sequencing.

University of Maryland, Baltimore County, Baltimore, Maryland, USA

M.S. in Statistics

Aug 2011 – Dec 2013

- Track: Biostatistics
- Thesis: Longitudinal Analysis of Urea Cycle Disorder Patients
- Adviser: Dr. DoHwan Park

Massachusetts Institute of Technology, Springfield, Pennsylvania, USA

■ B.S. in Mathematics 2006 – 2010

SKILLS

PROGRAMMING

R, Python, SQL, MATLAB, Machine Learning, Statistical Modeling, bash/shell scripting, LATEX

TOOLS & PLATFORMS

AWS, GCP, Nextflow, snakemake, Git, CI/CD, Containerization (Docker, Singularity), HPC clusters

BIOINFORMATICS

16S rRNA-seq, bulk RNA-seq, scRNA-seq, WGS, GWAS, pathogen genomics

RELEVANT INDUSTRY EXPERIENCE

Booz Allen Hamilton Remote

■ Lead Scientist Aug 2022 – Oct 2024

- Served as a federal contractor bioinformatician, contributing to public health genomic surveillance, pathogen genomics, and large-scale human genomics projects across CDC and NIH.
- Redesigned and optimized public health genomics workflows to align with Nextflow nf-core standards, implementing
 rigorous pipeline validation and automated unit testing. Streamlined collaboration through Git, ensured reproducibility
 via Docker containerization, and enhanced project delivery efficiency by integrating Agile tracking in JIRA.
- Engineered reproducible bioinformatics pipelines with Snakemake on high-performance computing (HPC) clusters, enabling robust rare variant detection and uncovering genetic associations with ulcer development in sickle cell disease.
- Led the design and implementation bioinformatics pipelines in R and Python on Google Cloud Platform (GCP)
 via the NIH All of Us Researcher Workbench, spanning genomic data ingestion, QC, and advanced downstream
 analysis. Delivered cloud-native, production-ready workflows that accelerated large-scale genomic insights for
 precision medicine applications.

RESEARCH EXPERIENCE

Davenport Lab, Pennsylvania State University

Postdoctoral Fellow

Jun 2020 – Jul 2022

- Piloted a benchmarking study on computational approaches to identify viral transcripts from bulk and single-cell RNA sequencing data, assessing the accuracy and precision of transcript identification.
- Designed bioinformatics workflows in snakemake for efficient data processing on HPC computing environments, employing tools including samtools, bwa, bowtie2, Kraken2, and STAR, and utilizing R for downstream analysis and visualization.
- Mentored undergraduate students, enhancing their research skills and academic performance.

Furey Lab, University of North Carolina at Chapel Hill

Graduate Research Assistant

Nov 2014 – May 2020

- Identified key microbial associations using R for data analysis and visualization of 16S rRNA amplicon data from IBD patients and controls.
- Developed a consensus analysis method in R with DEseq2 and Lefse, significantly enhancing the understanding of microbial dysbiosis in IBD.
- Presented research findings at various conferences, including the American Society of Human Genetics, to engage the scientific community.
- Authored a peer-reviewed journal article, contributing to the advancement of knowledge in the field of microbial dysbiosis.

PUBLICATIONS

Shahir, NM, et.al, "Crohn's Disease Differentially Affects Region-Specific Composition and Aerotolerance Profiles of Mucosally Adherent Bacteria." *Inflammatory bowel diseases*, vol. 26, no. 12, pp. 1843–1855, 2020.

PRESENTATIONS EXTERNAL TALKS

 Crohn's Disease Differentially Affects Intestinal Region Composition and Aerotolerance Profiles of Mucosally-Adherent Bacteria, Remote
 Virtual Microbiome Summit

 IBD differentially affects region specific composition and aerotolerance profiles of mucosal-adherent bacteria, Remote
 May 2020

MIT and UNC Joint Virtual Microbiome Seminar Series

UNIVERSITY OF NORTH CAROLINA AT CHAPEL HILL

• Crohn's Disease and the Intestinal Microbiota,

Chapel Hill, NC Dec 2016

Genetics Research Colloquium

Alterations in the Mucosal-Adherent Enteric Microbiota Between CD and nonIBD,

Chapel Hill, NC

Translations Medicine Closed Door Talks

A distinct microbiota signature characterizes patients with penetrating Crohn's disease,
 Chapel Hill, NC

Oct 2015

Center for Gastrointestinal Biology and Disease

Analysis of the Composition and Diversity of the Colonic Mucosa Microbiota in Crohn's Disease,
 Chapel Hill, NC
 May 2015

Bioinformatics and Computational Biology Curriculum New Student Talks

Identification of SERPINA1 Splice Variants from Next-Gen Sequencing Data,
 Chapel Hill, NC

Bioinformatics and Computational Biology Research in Progress Talks

POSTERS EXTERNAL

 Analysis of mucosal adherent 16S rRNA reveals altered microbial composition and decreased diversity in patients with Crohns disease,

Baltimore, MD Oct 2015

American Society for Human Genomics

UNIVERSITY OF NORTH CAROLINA AT CHAPEL HILL

• Characterizing the Intestinal Mucosal Landscape in Inflammatory Bowel Disease,

Chapel Hill, NC Genetics Department Retreat Aug 2017

Oct 2016

Oct 2014

Analysis of the Composition and Diversity of the Colonic Mucosa Microbiota in Crohn's Disease,
 Chapel Hill, NC
 Genetics Department Retreat

Analysis of the Composition and Diversity of the Colonic Mucosa Microbiota in Crohn's Disease,
 Chapel Hill, NC

Center for Gastrointestinal Biology and Disease Poster Session

Analysis of the Composition and Diversity of the Colonic Mucosa Microbiota in Crohn's Disease,
 Chapel Hill, NC

May 2015

Information Technology Services Research Computing Symposium

HONORS & AWARDS

■ NIH T32 Training Fellow

2015

Bioinformatics and Computational Biology Predoctoral Training Grant

OTHER RESEARCH National Human Genome Research Institute, Bethesda, MD

EXPERIENCE

Mentors: Dr. Julie Segre, Dr. Sean Conlan

Jun 2012– Aug 2012

Summer Fellow

- Piloted a study on the viral diversity of the human skin through the use of metagenomic datasets acquired from the human microbiome project.
- Assisted with fungal speciation of Malassezia species through bioinformatic tools
- $\bullet\,$ Applied various bioinformatic tools including: Clustal, BioPython, BioPerl, and Bowtie
- · Extracted full viral genomes from metagenomic datasets

• Presented research at NHGRI and NIH poster sessions.

National Human Genome Research Institute, Bethesda, MD

Mentors: Dr. Ellen Sidransky, Dr. Nahid Tayebi

Jun 2009- Aug 2009

- Summer Fellow
- Worked on defining the association between glucocerebrosidase mutations and Parkinsons disease.
- · Learned and applied biological methods: including sequencing, PCR, westerns, RNA and protein extractions.
- Performed statistical analysis on gene expression data via Excel
- Trained incoming fellows in lab protocols
- Presented research at NHGRI and NIH poster sessions

National Cancer Institute, Bethesda, MD

 Mentors: Dr. David D. Roberts, Dr. Michael Pendrak Summer Fellow Jun 2007- Aug 2007

- Worked on the development of a modification of a tetracycline-regulating system for developmental regulation in Candida albicans.
- Learned and applied methods in molecular biology and bioinformatics including: PCR, gel electrophoresis, BLAST, ClustalW, DNA purification.
- Trained incoming high school intern in lab protocols.

LANGUAGES

- English: Native language.
- French: Intermediate (reading); basic (speaking, writing).