

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
 - Committee: 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.” <i>Inflammatory bowel diseases</i> , vol. 26, no. 12, pp. 1843–1855, 2020.	
PRESENTATIONS	EXTERNAL TALKS	
	<ul style="list-style-type: none"> ▪ <i>Crohn’s Disease Differentially Affects Intestinal Region Composition and Aerotolerance Profiles of Mucosally-Adherent Bacteria</i>, Remote May 2020 Virtual Microbiome Summit ▪ <i>IBD differentially affects region specific composition and aerotolerance profiles of mucosal-adherent bacteria</i>, Remote May 2020 MIT and UNC Joint Virtual Microbiome Seminar Series 	
	UNIVERSITY OF NORTH CAROLINA AT CHAPEL HILL	
	<ul style="list-style-type: none"> ▪ <i>Crohn’s Disease and the Intestinal Microbiota</i>, Chapel Hill, NC Dec 2016 Genetics Research Colloquium ▪ <i>Alterations in the Mucosal-Adherent Enteric Microbiota Between CD and nonIBD</i>, Chapel Hill, NC Oct 2016 Translations Medicine Closed Door Talks ▪ <i>A distinct microbiota signature characterizes patients with penetrating Crohn’s disease</i>, Chapel Hill, NC Oct 2015 Center for Gastrointestinal Biology and Disease ▪ <i>Analysis of the Composition and Diversity of the Colonic Mucosa Microbiota in Crohn’s Disease</i>, Chapel Hill, NC May 2015 Bioinformatics and Computational Biology Curriculum New Student Talks ▪ <i>Identification of SERPINA1 Splice Variants from Next-Gen Sequencing Data</i>, Chapel Hill, NC Oct 2014 Bioinformatics and Computational Biology Research in Progress Talks 	
POSTERS	EXTERNAL	
	<ul style="list-style-type: none"> ▪ <i>Analysis of mucosal adherent 16S rRNA reveals altered microbial composition and decreased diversity in patients with Crohns disease</i>, Baltimore, MD Oct 2015 American Society for Human Genomics 	
	UNIVERSITY OF NORTH CAROLINA AT CHAPEL HILL	
	<ul style="list-style-type: none"> ▪ <i>Characterizing the Intestinal Mucosal Landscape in Inflammatory Bowel Disease</i>, Chapel Hill, NC Aug 2017 Genetics Department Retreat ▪ <i>Analysis of the Composition and Diversity of the Colonic Mucosa Microbiota in Crohn’s Disease</i>, Chapel Hill, NC Aug 2016 Genetics Department Retreat ▪ <i>Analysis of the Composition and Diversity of the Colonic Mucosa Microbiota in Crohn’s Disease</i>, Chapel Hill, NC Jul 2015 Center for Gastrointestinal Biology and Disease Poster Session ▪ <i>Analysis of the Composition and Diversity of the Colonic Mucosa Microbiota in Crohn’s Disease</i>, Chapel Hill, NC May 2015 Information Technology Services Research Computing Symposium 	
HONORS & AWARDS	<ul style="list-style-type: none"> ▪ NIH T32 Training Fellow 2015 Bioinformatics and Computational Biology Predoctoral Training Grant 	
OTHER RESEARCH EXPERIENCE	National Human Genome Research Institute, Bethesda, MD <ul style="list-style-type: none"> ▪ Mentors: Dr. Julie Segre, Dr. Sean Conlan Jun 2012– Aug 2012 Summer Fellow <ul style="list-style-type: none"> • 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 <i>Malassezia</i> species through bioinformatic tools • 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

Jun 2007– Aug 2007

Summer Fellow

- 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).