

# AMIRTESH RAGHURAM

Email: [amirtesh21.5@gmail.com](mailto:amirtesh21.5@gmail.com) | Phone: +91 9353724370

GitHub: [github.com/amirtesh](https://github.com/amirtesh) | LinkedIn: [in.linkedin.com/in/amirtesh-raghuram-90161828a](https://in.linkedin.com/in/amirtesh-raghuram-90161828a)

Vellore Institute of Technology, Vellore

---

## Education

---

### B.Tech Biotechnology

*Vellore Institute of Technology, Vellore*

2023 - 2027 (Expected) | CGPA: 8.99/10

## Professional Experience

---

### Summer Research Intern

*Strand Life Sciences, Bangalore* | May 2025 - July 2025

- Worked on "Identification of Homologous Regions for the Purpose of Variant Verification in NGS Tests"
- Developed automated Bash scripts for NGS data processing pipelines and homology detection workflows
- Utilized BLAST command-line suite for sequence alignment, variant verification, and homologous region identification
- Implemented bioinformatics solutions for high-throughput genomic data analysis and quality control

## Ongoing Research Projects

---

### Natural Plant-Based Inhibitors for Diabetes Treatment (Manuscript in Submission)

- Collaborative research with PhD scholar identifying natural plant compounds as alpha-amylase inhibitors
- Implementation of computational screening methods, molecular docking, and binding affinity calculations
- Molecular dynamics simulations to evaluate stability of protein-ligand complexes
- Identification of lead compounds with potential therapeutic applications through ADMET profiling

### Multi-Target Inhibitors for Triple Negative Breast Cancer Treatment (Manuscript in Submission)

- Independent research screening natural drug-like compounds as multi-target inhibitors for TNBC therapeutic targets
- Performed triplicate molecular dynamics simulations with MMGBSA binding free energy calculations for robust validation
- Evaluated lead compounds for multi-target efficacy using computational screening and pharmacokinetic profiling
- Assessment of compound stability, binding interactions, and drug-likeness properties

## Natural Inhibitors for Glaucoma Treatment (Manuscript in Submission)

- Independent research identifying plant-based inhibitors of human carbonic anhydrase II
- Development of computational pipeline for high-throughput virtual screening
- In-depth molecular dynamics simulations to evaluate stability and effectiveness
- Characterization of binding mechanisms and inhibitory potential of lead compounds

## Technical Skills

---

### Programming & Data Science

- Python (NumPy, Pandas, Scikit-learn, BioPython)
- R (dplyr, ggplot2, Bio3D)
- PyTorch & TensorFlow
- Matplotlib/Seaborn
- Statistical Analysis & Machine Learning
- Bash Scripting & Workflow Automation

### Structural Bioinformatics

- Protein Structure Analysis (ProDy, Bio3D)
- Protein-Ligand Docking (AutoDock Vina, Smina, QVina, PyRx, AMDock)
- Molecular Dynamics Simulations (GROMACS)
- Binding Free Energy Calculations (gmx\_MMPBSA, MMGBSA)
- Protein Modeling (AlphaFold, SWISS-MODEL, I-TASSER, Phyre2)
- Structural Visualization (PyMol, Chimera, Discovery Studio)
- Cheminformatics (RDKit)
- Virtual Screening Workflows

### Genomics & Next-Generation Sequencing

- RNA-Seq Analysis
- Single-Cell RNA-Seq (Seurat)
- ChIP-Seq Analysis
- Variant Calling & Genome Assembly
- DNA Methylation Analysis
- GWAS (PLINK)
- Sequence Alignment (BLAST)
- Galaxy, HOMER, BioConductor

### Cancer Genomics

- Somatic Mutation Analysis (maf-tools, oncoplot, lollipopPlot)
- Tumor Heterogeneity and Clonality Assessment
- Mutation Signature Analysis (COSMIC signatures)
- Co-occurrence and Mutual Exclusivity Analysis
- VAF-based Clonal Population Identification
- Oncogenic Driver Identification (Oncodrive)
- rainfallPlot for Mutation Visualization

### Computational Drug Discovery & ADMET

- Virtual Screening Pipelines
- In Silico Vaccine Design
- SwissDock, SwissADME, Swiss Target Prediction
- Swiss Similarity, pkCSM, ProTox 3.0
- MolSoft Drug-likeness Assessment
- KEGG Pathway Analysis

## Software Contributions

---

- **Torchify:** Python library simplifying PyTorch workflows with enhanced model API functions; published on PyPI with 2,500+ downloads ([github.com/Amirtehs/Pytorch-Torchify](https://github.com/Amirtehs/Pytorch-Torchify))
- **Automated-Virtual-Screening:** Binary executable automating ligand screening using Vina, Smina, and QVina with features for file conversion, parallel docking, and results management ([github.com/Amirtehs/Automated-Virtual-Screening](https://github.com/Amirtehs/Automated-Virtual-Screening))

## Core Competencies & Achievements

---

- Complete computational drug discovery pipelines from virtual screening to molecular dynamics simulations and binding free energy calculations
- Next-generation sequencing data analysis and interpretation including variant calling and clinical genomics
- Cancer genomics data analysis including somatic mutation profiling, tumor evolution assessment, and clonality studies
- Protein structural analysis, comparative modeling, and structure-function relationship investigations
- Development of custom computational workflows for high-throughput data processing and automation
- Implementation of machine learning approaches for biological data analysis and predictive modeling

## Bioinformatics Tools & Resources

---

SwissDock, SwissADME, Swiss Target Prediction, Swiss Similarity, pkCSM, ProTox 3.0, MolSoft, AlphaFold, SWISS-MODEL, I-TASSER, Phyre2, BLAST, Galaxy, HOMER, KEGG, BioConductor, rainfallPlot, oncoplot, Oncodrive, maftools, ProDy, Bio3D