

boneshwar.vk20@gmail.com Academic portfolio in LinkedIn

Summary

My primary research interests lie at the intersection of computational biology and deep learning, using graphs & transformer based models to fuse single-cell and multi-omics with biological priors for interpretable, clinically grounded analysis. My focus is personalized diagnosis and preventive risk stratification in brain cancers—especially gliomas—linking cell-state biology to patient-level decisions.

Education

2020–2025 B.S. in Biological Sciences, Indian Institute of Technology Madras

M.S. in Biological Sciences (Specialization in Computational Biology), Indian Institute of Technology Madras, CGPA: 8.12/10.0

2018–2020 Higher Secondary Education, Velammal Vidyalaya, Central Board of Secondary Education, Percentage: 95.0%

Technical Skills

Programming O Programming Languages: Python, C++, C#, R

- O Software and tools: Gromacs, AutoDock, Weka, Pymol, VMD, Blender
- o Libraries and packages: PyTorch, NetworkX, PyG, DGL, Gensim, Rpy2, Scikit-learn, Pandas, TensorFlow

Wet lab O Techniques: Molecular cloning, Colony PCR, CFU quantification

o Technologies & Instruments: Thermocycler, Gel documentation, Spectrophotometer

Coursework

Protein interactions 2020-2025

Fundamentals of Deep Learning

Molecular biology

Bioinformatics Algorithmic approaches in computational biology

DSA¹² in Biology

Publications

Computers in Biology PAIGT: Position aware graph transformer for single-cell RNA*. Authors: Boneshwar V. K, and Medicine Vijayavallabh Jayamanikandan , Deepesh Agarwal, Bala Natarajan, and Babji Srinivasan. (Under review)

Computers & Inductive graph neural network framework for imputation of single-cell RNA sequencing data. Chemical Engineering Authors: Boneshwar, V. K., Deepesh Agarwal, Bala Natarajan, and Babji Srinivasan.

Research Experience

Project member Ontology-Grounded LKM for disease diagnosis

CRIS Lab Principal Investigators: Prof. Venkat Venkatasubramanian (Columbia University) Prof. Bala Natarajan (K-State) (May 2025 - Present)

- O Building interpretable, ontology-grounded Large Knowledge Model (LKM) for hierarchical diagnosis.
- Prototyping LKM with distilled chain-of-thoughts with rare disease-symptoms and neuro-oncology.
- O Leverages Llama for safe, local & interpretable clinical decision support with transparent reasoning.
- Curated phenomics corpus via GA4GH Phenopackets; engineered weighted disease-phenotype edges and KG1 communities using MONDO/ORDO and HPO2 attributes; implementing KG-aware inference with hierarchical reasoning and standards-aligned actions (LOINC/SNOMED CT/ICD-11).

Research Associate Foundational models using multi-omics based cell trajectory inference for Gliomas

HILCPS Lab Principal Investigators: Prof. Babii Srinivasan, Prof. Bala Natarajan (K-State) (Dec 2023 - May 2025)

- O Built inductive frameworks for scRNA-seq to recover global topology and glioma-relevant patterns.
- Enhanced cell-type annotation & neoplastic-non-neoplastic separation across sparse scRNA atlases.
- Supports early, pre-clinical risk stratification & prevention by unifying glioma microenvironment profiles.
- O Designed position-structure encodings (PSE) and multiscale embeddings integrating neighborhood states with cohort manifolds while preserving lineage and cell-cell communication patterns leveraging graph transformer aimed for recovering trajectories and pseudotime.

Research Intern Inductive scRNA Imputation GNN

CPSWin Lab Principal Investigator: Prof. Bala Natarajan (K-State)

(May 2024 - July 2024)

- Impute scRNA-seq dropouts by reconstructing subgraph neighborhoods gene—gene co-expression.
- \circ SOTA Metrics: +60% Silhouette, +14.9% ARI³, 48% runtime, 4.5% L₁ median error vs baselines.
- Improves downstream cell-state clustering and strengthens scRNA pipelines for tumor biopsies.
- Developed inductive graph neural network for unsupervised clustering across cohorts to handle the inherent data sparsity associated with single-cell technologies.

Al Researcher CSV Agent for QA tasks and Indic OCR

Sarvam Al Principal Investigators: Prof. Pratyush Kumar, Vivek Raghavan

(Nov 2023 – Jan 2024)

- O Built an end-to-end agent for large CSV QA and multilingual OCR to unlock large scale tabular data.
- SOTA on internal and Yahoo Finance datasets; OCR at 0.62-CER/0.71-WER⁴ with 80% layout fidelity.
- Enables scalable digitization and QA of clinical registries and EHRs in low-resource scripts.
- Designed CSV QA agent; built Indic OCR (Hindi/Tamil/Telugu/Malayalam/Kannada); implemented RAG with hybrid retrieval and cross-encoder re-ranking.

Professional Experience

Associate Data Convozen AI (NoBroker.com)

Scientist Data Science Team

(May 2025 - Present)

- Delivered a low-latency speech-to-speech voice-bot with robust VAD in noisy, real-world call flows.
- O VAD false positives reduced by 13%; 900 ms TTFB⁵; 1600 ms average conversation turn.
- Enables AI based realtime conversational voice bots in healthcare and other customer care operations.
- Integrated DeepFilterNet v3 for neural spectral denoising and non-cascaded⁶ Speech-to-speech collaborating with Google and Azure; exploring TTS for human vocal mimics—SOTA commercial voice bots mirroring human vocals and prosody with biomimetic vocal-tract cues (timbre, breath, coarticulation).

Autonomous Systems 10X, IITM Research Park

Engineer Artificial Intelligence Team

(May 2023 - Jul 2023)

- Prototyped an autonomous wheelchair with SLAM navigation & multilingual conversational interface.
- Improved navigation with a success rate of 76%.
- O Improves assistive mobility for neuro-rehabilitation and elder-care settings.
- Implemented mapping and navigation using IMU, encoders, LiDAR with ROS⁷ (Hector SLAM, Cartographer) and a RASA-based chatbot for 5 major indic languages. Live demo to Google India's CEO and the Chief Justice of India: featured in Times of India

Research Projects

Transformer

MaskedAlign: Multiple Sequence Alignment for large scale genomes

Seq-to-Seq MSA Principal Investigator: Prof. Manikanda Narayanan

(Jan 2024 - May 2024)

- O Enhance generalization of MSA8 on unseen gene families using a masked transformer using PyTorch.
- Higher alignment accuracy vs BetaAlign/ClustalW/MAFFT with runtime reduction up to 11%.
- O Speeds comparative genomics and variant discovery relevant to cancer cohort analyses.
- O Built seq-to-seq model with masked language modeling and robust training for long genomic data.

Recombinant Plasmid Recombinant Plasmid Engineering

Assembly Principal Investigator: Prof. Venkitasamy Kesavan

(Jan 2024 - May 2024)

- Construct and verify recombinant plasmids for reliable downstream expression studies.
- Insert verification via restriction mapping; maintained clonal lineage with stable plasmid retention.
- Establishes a robust cloning backbone for functional assays like oncogene perturbation models.
- \circ Executed miniprep, PCR and gel, BamHI/Ndel digests, ligation, and DH5 α transformation with antibiotic selection and colony PCR monitoring clonal lineage across passages.

GANs⁹ for image generation

Principal Investigators: Prof. Babji Srinivasan, Prof. Rajagopalan Srinivasan (Dec 2023 – Mar 2024)

- Disentangle latent factors for controllable image synthesis and RGB→thermal translation.
- Metrics: +6% accuracy and consistent foreground-background separation over baselines.
- Enables privacy-preserving synthetic data for controlled image generation.
- Fine-tuned GANs to separate shape, size, texture & color with edge-guided translation for RGB to thermal.

Dynamics

Microbial Population Microbial Dynamics Profiling

Trajectory and Cell Coursework: BT2112 Microbiology Laboratory

(Jan 2022 - May 2022)

- O Quantify growth trajectories and stress responses across controlled environmental gradients.
 - O Derived stable phenotype panels and MIC¹⁰ profiles across pH/salinity/temperature conditions.
 - O Reinforces reproducible phenotyping workflows transferable to microbiome QC in oncology pipelines.
 - Performed OD tracking, hemocytometer counts, serial-dilution CFU¹¹, Gram staining, biochemical panels, enzyme assays, and antibiotic susceptibility (Kirby-Bauer, MIC).

Leadership and Outreach

Madras

CVI Club, CFI, IIT O Project member 2022-2023, developed an end-to-end traffic violation detection system (helmet, triple riding, over-speeding) using YOLOv5, achieving F1: 0.79 and 89% accuracy on Jetson Nano, deployed via CCTV infrastructure and incubated at Nirmaan, IIT Madras.

SEC, IIT Madras O Actively served on IIT Madras election body, overseeing the Student General Elections 2021-2022 with 10,000+ electorate and upholding integrity in the Academic Affairs Secretary race.

NGOs

- Social Impact & Omdena Al Singapore: Implemented ML based pipeline for analysing workouts (pushups, situps, pullups)
 - Suvidha Foundation (NGO): Designed framework for teaching ML to support weaker economic sections

Achievements and Extra-Curricular Activities

- Martial Arts O Secured 1st place and 2nd place at World Unified Martial Arts Federation 2011 and 2012
 - Active participant of Ryu Kyu Kobudo & Shonrinryu Karate International seminar and Okinawa karate and Kobudo Invitation championship 2013

- Cricket O Lead Narmada hostel, IIT Madras in Schroeter 2024
 - O Lead the high school cricket team in Inter-District school tournament 2018, Tamil Nadu, secured 3^{rd} place in the tournament and being the highest run scorer among the team
 - Represented Tamil Nadu at Nationals 2015 held by united cricket federation of India.

- Academics O Secured All India Rank 9 (AIR 9) in the JEE MAINS B.Arch conducted in 2020.
 - Secured top 1 percentile among 1,30,000 candidates in JEE Mains & Advanced 2020

^{*-} ongoing/submitted, 1- Knowledge Graphs, 2- Human Phenotype Ontology, 3- Adjusted Rand Index, 4- Character Error Rate & Word Error Rate, 5-Time to first byte, 6-Model with no separate STT-LLM-TTS modules, 7- Robot Operating System, 8- Multiple Sequence Alignment, 9- Generative Adversarial Networks, 10- Minimum Inhibitory Concentration, 11- Colony-Forming Units, 12- Data Structures and Algorithms