# Nikhil Mark Lakra

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## Experience

#### **Bioinformatics Scientist**

Delhi

Elucidata

Oct 2022 - Present

- Collaborated with major pharmaceutical clients on diverse research projects, leveraging Bulk RNA-Seq and single-cell transcriptomics data to derive high-quality insights for drug discovery and development.
- Standardized and automated workflows for preprocessing, differential expression analysis, and pathway enrichment, ensuring reproducible and efficient delivery of results.
- Developed and deployed a meta-analysis pipeline with KNN-based gene imputation for differential expression analysis, and pathway enrichment using Python and R, enabling the discovery of key therapeutic targets.
- Built a dockerized gene similarity pipeline to identify genes with similar expression signature patterns across single-cell RNA-seq datasets, utilizing pandas, scikit-learn, Scanpy, and scvi-tools for efficient and reproducible analysis.
- Developed a workflow using pre-trained machine learning models (e.g., scVI-tools, CellTypist) to automate cell-type annotation in scRNA-Seq data, reducing the time to identify and accurately annotate cell-types by 20%.
- Conducted data audits on public repositories (e.g., GEO, Single Cell Portal, and ArrayExpress) to identify high-quality datasets of interest to support client-specific analyses and projects.
- Led an internal training session on preprocessing Bulk RNA-seq data, covering the nuances of normalization methods and the underlying algorithms of differential expression tools (LIMMA, edgeR, DESeq2), significantly enhancing the team's proficiency in downstream analysis workflows.

#### Education

## Indraprastha Institute of Information Technology Delhi (IIITD)

Oct 2019 - Jan 2022

M. Tech in Computational Biology

- o GPA: 7.96/10.0
- o Coursework: Machine Learning, Data Science in Genomics, Biostatistics, Computer Aided Drug Design

### Projects and Thesis

# Thesis: Predicting Selection Pressure on SNPs in Human Populations

Link to Thesis 🗹

- Developed a machine learning pipeline to predict single nucleotide polymorphisms (SNPs) under selection pressure across 17 populations from the 1000 Genomes Project.
- Implemented a Random Forest classifier, taking advantage of population genetic characteristics such as allele frequency, F statistics, and linkage disequilibrium for accurate predictions.
- Achieved high classification accuracy, contributing to the identification of genetic variations of evolutionary significance.

## AQI Forecasting Using Deep Learning (LSTM)

GitHub Link 🗹

• Developed an LSTM-based deep learning model in PyTorch to predict the next day's Air Quality Index (AQI) using historical AQI data from CPCB and climate data from NASA Power.

## **Technologies**

Languages: Python, R, SQL, Bash.

**Technologies Frameworks:** Pandas, NumPy, scikit-learn, TensorFlow, PyTorch, scVI-tools, Scanpy, Seurat, DESeq2, edgeR, LIMMA.

Cloud Development Tools: Docker, Git, AWS, Sagemaker.