# Justice (Hsiu-Yi) Lu

hyjusticelu@gmail.com | +1.312.623.5370 | Justice-lu.github.io

#### **Summary**

Computational biologist with a strong foundation in human and model organism biology, applying large-scale multi-omics analysis, structural modeling, and machine learning to tackle fundamental questions in cell biology and disease. Experienced in single-cell and bulk transcriptomics, proteomics, protein structure—function studies, and scalable ML workflows.

## **Work Experience**

## Graduate Student Researcher, Duke University, Hiroaki Matsunami Lab

2020 - Present

- Applied single-cell, bulk RNAseq and proteomic analysis pipeline and identified gene candidates to enhance olfactory receptor cell surface expression for *in vitro* genetic screen.
- Developed convolutional neural network (CNN) models to predict receptor–ligand binding from protein structures, integrating structural biology with machine learning.
- Created geLine, a web tool for gene co-expression analysis and maturation trajectory tracing, enabling researchers to interrogate maturation trajectories across cell type.
- Mentored trainees in both computational and experimental projects, fostering independent design and test skills.

### ML Data Science Intern, Kriya Therapeutics

2023

- Engineered scalable ML pipelines for high-throughput drug target prediction, increasing 10× data throughput.
- Optimized data preprocessing and model training workflows using Python, scikit-learn, and cloud computing resources.
- Partnered with biology teams to incorporate omics-derived features into predictive models, improving biological relevance.

#### Research Technician II, Duke University, John Rawls Lab

2017 - 2019

- Conducted studies on host-microbe interactions in zebrafish, uncovering protective gut responses to pathogenic bacteria.
- Performed a genetic screen identifying novel regulators of lipid metabolism and adipose tissue morphogenesis, with implications for metabolic disease biology.

#### **Education**

## **Duke University** (Expected graduation, Fall 2025)

2019 - Present

Ph.D. in Molecular Genetics and Microbiology (Advisor: Hiroaki Matsunami)

IBIEM Computational Biology Fellow | Taiwan-Duke Fellowship Recipient

Purdue University, B.S in Biological Science

2013 - 2016

### **Selected Publications (\* First author)**

• Identifying Key Regulators in Odorant Receptor Trafficking.

Lu, HY\*. & Matsunami, H. Journal of Neuroscience (in revision)

Physiology of olfactory sense.

Lu, HY\*., Meesa, P. & Matsunami, H., Digital Technologies in Olfaction, 7-14 (2025)

- Role of iRhom2 in Olfactory Sensory Neurons: Implications for Odorant Receptor Regulation.
  Azzopardi, S\*., Lu, HY\*., Monette, S., Rabinowitsch, A., Salmon, J., Matsunami, H., & Blobel, C. IJMS (2024)
- Sex steroid hormones synthesis, metabolism and the effects on the mammalian olfactory system.

Abaffy, T., Lu, HY., & Matsunami, H. Cell and tissue research (2023).

- An odorant receptor that senses four classes of musk compounds
  Yoshikawa, K., Deguchi, J., Jieying, H., Lu, HY., & Matsunami, H. Current Biology (2022).
- Enteroendocrine cells sense bacterial tryptophan catabolites to activate enteric and vagal neuronal pathways. Ye L, Bae M, Cassilly CD, Jabba SV, Thorpe DW, Martin AM, Lu HY., ... & Rawls JF. Cell Host Microbe (2021).

#### Skills

Computer Programming Language: Python, R, Bash, Git

Bioinformatics: omics analysis (single-cell/bulk RNA-seq, proteomics), protein modeling

ML & Data Science: PyTorch, scikit-learn, data visualization, CNNs, regression/classification

**Collaboration**: Bench-to-computation project design, cross-disciplinary communication

**Interest**: Building pipelines and tools for optimization