Linqing Hu

+44(0)7849294300 linqinghu120120@outlook.com linkedin.com/in/linqing-hu-6836b7239 https://github.com/Luna120120

#### **EDUCATION**

The University of Manchester

09/2023 - 09/2024, Manchester, UK

MSc Bioinformatics and Systems Biology (Distinction)

Core modules: Bioinformatics, Computational Approaches to Biology, Experimental Design and Statistics
The University of Edinburgh

09/2019 – 05/2023, Edinburgh, UK

BSc Hons Biological Sciences (Biotechnology) (UK 2:1)

**Core modules:** Cell Biology; Molecular Biology, Genome and Genetics; Gene Expression and Microbial Regulation, Enzymology and Biological Production, Biotechnology; Novel Approaches, Quantitative Skills

### **RESEARCH EXPERIENCE**

Master's Project 1: Prof Gisela Orozco's Lab

2024

# Using GWAS, RNA-seq and ATAC-seq combined functional genomic annotation to discover the TF-binding regulatory mechanism responsible for Psoriatic Arthritis (PsA)

Autoimmune diseases (ADs) pose significant global healthcare challenges due to their complex nature and chronic progression. Psoriatic Arthritis (PsA), a subtype of AD, impacts patient quality of life and poses economic burdens. PsA's pathogenesis involves a combination of genetic predispositions and environmental factors. Understanding the role of transcription factors (TFs) in regulating gene expression is crucial for decoding PsA's molecular mechanisms. TFs bind to genetic regulatory elements such as enhancers and promoters, influencing immune responses and inflammation. This study integrates multi-omics data from ATAC-seq, GWAS, and RNA-seq to help identify regulatory elements and provide insight into PsA-related mechanisms, thereby enabling the development of new diagnostic tools and treatment approaches. From the comparison, a new motif was found at the BCL2L11 promoter-associated enhancer SNP rs13401811, which provides a hint at the potential regulatory mechanism of this gene and the pathogenesis of PsA.

#### Master's Project 2: Prof Simon Hubbard's Lab (paper is under publication)

2024

# Co-translational Protein Complex Assembly: integrating structural and quantitative mass spectrometry data to identify candidates

Multimeric protein complexes assembly through protein-protein interactions are important for cellular functions across all domains of life. Other than the conventional thought that complexes are formed from random collisions of fully synthesised and folded subunits, the process of co-translational assembly (Co-TA) was suggested and found to happen prevalently for both the homomeric and heteromeric complexes. Previous studies have raised three factors (1) the Co-TA processing orders: simultaneous (Co-co) vs sequential (Co-post), (2) the ribosome translation locations: on the same mRNA (cis) vs on different mRNAs (trans), and (3) the subunits assembly orders: directional vs symmetrical that is important to Co-TA, overall, give 6 possible modes of the Co-TA mechanism, and was found favoured by homomeric and heteromeric complexes differently.

Since studies showed that the relative location (either towards the N-terminus or the C-terminus of the protein sequence) of the subunit interactive (buried) surface is evolution-selective and important for the two types of complexes in adopting different modes/mechanisms of Co-TA, this study focused on heteromeric complexes, raised a concept of subunit Moment (M), hypothesised that the Co-TA mechanisms may be distinguishable by the relative location of the buried surfaces in the component subunits of a protein complex through relative Moment (M-rel). Through the use of a novel in vivo experiment, 62 new

heteromeric complexes were identified that may undergo Co-TA. By developing a computational method to calculate the relative Moment (M) of subunits by their residual solvent accessibility (ASA) generated from PDB structural data. This study confirmed the potential of using relative M-rel to suggest the possible Co-TA mechanism for complexes and raised its significance in helping tackle diseases.

## **Undergraduate Honours project: Dr Sander Granneman's lab**

2023

Performing Differential Gene Expression (DGE) analysis and GO term enrichment analysis of the Nanopore RNA sequencing data to decipher the role of the endonuclease RNase III in regulating Methicillin-resistant *Staphylococcus aureus* (MRSA) gene expression.

Methicillin-resistant Staphylococcus aureus (MRSA) is a globally prevalent pathogen that can cause severe infections and result in death, if have no effective treatment. The mobile genetic elements ruched highly plastic genome acquires MRSA the ability to generate antibiotic resistance rapidly, leaving us to lack sufficient treatments in response to infections caused by multi-antibiotic resistance strains in a post-antibiotic era. Given that MRSA endoribonuclease RNase III plays a key role in the regulation of gene expression at transcription and post-transcription levels. This study aimed to decipher the function and new RNA and protein targets of RNase III, to discover potential new drug targets for developing new treatments. After using the bioinformatics approach to perform differential gene expression analysis an GO term environment analysis between the transcriptome sequence of wild type and the RNase III knockout generated through Oxford Nanopore sequencing. We identified a series of up and down-regulated genes enriched into important cellular pathways, including nitrogen regulation, de novo pyrimidine biosynthesis, and host immune system escaping. We suggest RNase III plays a role in these pathways; however, the functioning mechanism was not studied. Therefore, this project provides future studies with a direction to reveal RNase III functioning mechanism in these pathways.

## **COMPETITION AND WORKING EXPERIENCES**

iGEM Competition, team Edinburgh-UHAS Ghana

06/2022 - 10/2022, Edinburgh, UK

Building a biodegradation device containing a PETase and MHETase-involved enzyme cocktail to solve water pollution problem in Ghana

- https://2022.igem.wiki/edinburgh-uhas-ghana/index.html
- Gold Award, Best Track Nomination
- Independently designed team logo, wiki page and video to promote the team internationally
- Deeply involved in social engagement activities to increase the publicity of the team and synthetic biology, led fundraising of £4,000 to send the whole team to attend the Paris Giant Jamboree.

#### iDEC Competition, team Edinburgh-UHAS Ghana

06/2022 - 10/2022, Edinburgh, UK

Random mutagenesis and silver nitrate (AgNO3) based negative selection of different metallothioneins (MTs) exploring the potential for functional MT directed evolution

- https://idec-teams.github.io/2022 Edinburgh-UHAS Ghana/
- Gold Award, Best Community, Best Presentation, Best Target Molecule Nomination, Funniest Evolution Nomination, Best Molecular Evolutionary Machines Nomination and Best Molecular Evolutionary Outcomes Nomination

#### Bluepha Co., Ltd. iGEM advisor and Summer programs instructor 05/2021 – 08/2021, Shenzhen, China

- Supervised 13 high school students on web lab experiments and social engagement activities on the project 'Design an improved ligand-nanoparticle system which utilises pH-sensitive aptamer to improve specificity on targeting HER2 proteins'. The team achieved a Gold Award. <a href="https://2021.igem.org/Team:GreatBay">https://2021.igem.org/Team:GreatBay</a> SCIE
- Independently developed teaching materials for teenager-oriented summer programs 'Building a microscope with Lego Bricks', demonstrating excellent independent task execution ability.
- Independently taught 4 middle school students on a synthetic biology-based summer school costing £10000 'Constructing plasmid for fluorescent expression in *E. coli*' and 'Constructing plasmid to express Polyhydroxyalkanoates (PHAs) biodegradable plastics in *E. coli*'. My teaching received significant recognition from the students, parents and the company.

#### **SKILLS**

Wet lab: Animal and bacterial cells culturing and screening, High-throughput sequencing library preparation, CRISPR-Cas9 library preparation, PCR, Plasmid construction, transmission and DNA assembly, Centrifugation techniques, Spectrophotometer analysis, Agarose gel electrophoresis, Western blotting, Chromatography for protein extraction and purification, ELISA, Light and fluorescence microscopy Dry lab: Genome, Transcriptome and ATAC-seq data processing; differential expression and GO term

**Dry lab**: Genome, Transcriptome and ATAC-seq data processing; differential expression and GO term enrichment analysis; protein structure analysis; PCA and clustering; Network and logic modelling; Mathematical modelling, Statistics, Machine Learning

**Programming language and tools:** Python3, R, Bash, Git, PyTorch, Snakemake, DESeq2, TOBIAS **Languages:** Mandarin (native), English (professional), Japanese and Korean (basic proficiency)

Others: Affinity Designer2, Adobe Illustrator, Procreate, Adobe Premiere Pro

### SELECTED ACADEMIC AWARDS AND HONOURS

2023 SynBioBeta Synthetic Biology Global Conference (Student representative), Oakland, US	2023
The 12th Summer Research Program on Biomedicine, Medical Facility, Tsukuba University	2022
Quantitative Biology Summer Training Program, Peking-Tsinghua University Life	<i>2021</i>
Science Joint Centre, Peking University	
Silver Award from Intermediate Biology Olympiad, Royal Society of Biology	<i>2018</i>
Silver Award from Cambridge Chemistry Challenge, University of Cambridge	2018

### SELECTED LEADERSHIP EXPERIENCE AND OTHER AWARDS

British Kendo Association (BKA) Kendo Level 1 Coach

2024

- Demonstrates a good understanding and dedication to providing professional training in kendo.

Edinburgh University Kendo Club (EUKC), Health and Safety Officer, Active member

*2021 - 2023* 

- Represented the university in local and national competitions and achieved awards.
- Provided training advice and first aid help to club members and maintained a safe training environment.
- Determined, persistent and resilient, has a strong spirit that does not stop until the goal is achieved.

## **Edinburgh Model United Nations**

*2023* 

- Participated in the 2023 Scot MUN to debate on 'the legitimacy of commercial and altruistic surrogacy and egg donation', great at understanding problems from multiple positions.

The University of Edinburgh Chinese Debating Society, Active member

2020

- Excellent in critical thinking, language construction and argument presentation in public speaking
- Has a high level of language fluency, articulation and clarity in effective communication

High School Jiulian Business Society, Committee member, President of Central Bank

2016 - 2017

- Represented the high school in the 5th National Business Challenge. Assisted in the organisation of the 4th Shanxi province business competition with 120+ attendees.
- Has knowledge in auditing and finance, great at brainstorming and seeking business opportunities.

#### Professional training in fine art for 5 years

2012 - 2017

- Has a good aesthetic. Skilled in drawing and making designs and figures at the publication level.