### JACK GISBY

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#### **EDUCATION**

### Imperial College London

July 2020 - August 2023

PhD Candidate - Department of Immunology and Inflammation

- · Thesis: Using Multi-Omics to Understand Inflammatory Disease
- · Supervised by Dr James Peters and Dr Jacques Behmoaras
- · Assessed by Professor Marc Chadeau and Dr Jessica Strid

### University of Birmingham

September 2016 - June 2020

Biochemistry with Professional Placement (MSci) - First Class

· Dissertation: A Prior Knowledge-Based Computational Workflow for de novo Structural Elucidation of Small Molecules in Mass Spectrometry Metabolomics

### **PUBLICATIONS**

# Longitudinal proteomic profiling of dialysis patients with COVID-19 reveals markers of severity and predictors of death

First author

*eLife* 2021 - doi:10.7554/eLife.64827

- Lead the analysis of a high-dimensional proteomics (Olink) dataset with a complex cohort consisting of repeated measurements at inconsistent time points.
- · Applied linear mixed models and joint models to identify key proteins that changed over time following COVID-19 symptom onset.
- · Utilised supervised learning algorithms to identify biomarkers of severe disease.

# The widespread nature of Pack-TYPE transposons reveals their importance for plant genome evolution

First author

bioRxiv 2021 - doi:10.1101/2021.06.18.448592

- · Designed an algorithm for the specific annotation of repetitive elements that capture chromosomal DNA; implemented this as an R package available as part of the Bioconductor project.
- · Mined open source genetic data to demonstrate the abundance of these elements for multiple superfamilies and genomes. Found that recent insertions of these elements have impacted the evolution of genes.

### Mendelian randomisation identifies alternative splicing of the FAS death receptor as a mediator of severe COVID-19

Equal first contributor

 $medRxiv \ 2021 - doi:10.1101/2021.04.01.21254789$ 

- $\cdot$  Developed a pipeline to apply two sample Mendelian Randomisation comparing COVID-19 GWAS to Olink pQTLs.
- · Used fine mapping and colocalisation tools in tandem with public data repositories to investigate causal variants of disease.

### Plasma Lectin Pathway Complement Proteins in Patients With COVID-19 and Renal Disease

Equal first contributor

Frontiers in Immunology 2021 - doi:10.3389/fimmu.2021.671052

· Provided statistical support for the analysis of a repeated measures study design.

· Generated data visualisations to investigate biological hypotheses.

#### CONFERENCES AND PRESENTATIONS

- · Presentation Biomarkers of the Future 2021 Longitudinal proteomic profiling of dialysis patients with COVID-19 reveals markers of severity and predictors of death
- · Poster HUPO Reconnect 2021 Longitudinal proteomic profiling of dialysis patients with COVID-19 reveals markers of severity and predictors of death
- · Poster UK-CIC Immunology 2021 Longitudinal proteomic profiling of dialysis patients with COVID-19 reveals markers of severity and predictors of death
- · Presentation Longitudinal Studies 2021, Wellcome Genome Campus Longitudinal proteomic profiling of dialysis patients with COVID-19 reveals markers of severity and predictors of death

#### ADDITIONAL RESEARCH EXPERIENCE

## A Prior Knowledge-Based Computational Workflow for de novo Structural Elucidation of Small Molecules in Mass Spectrometry Metabolomics

Dissertation project, Dr Ralf Weber

October 2019 - June 2020

- · Developed and optimised a Python package, *MetaboBlend*, to propose candidate molecules using tandem mass spectra .
- · Processed large, high-dimensional mass-spectrometry datasets for the validation of *Metaboblend*.
- · Used Python, SQLite and Bash to implement the *MetaboBlend* workflow on a Linux high-performance computing cluster for the generation of large structure databases.

## Development of an Immunoturbidimetric Assay for Serum Amyloid A for an Automated Clinical Analyser

Industrial Placement, The Binding Site

August 2018 - August 2019

- · Developed assays for inflammatory biomarkers, such as Serum Amyloid A, for use in clinical laboratories.
- · Interacted with, and regularly delivered presentations to, researchers from diverse fields and non-scientific staff to ensure the timely completion of my development project.
- · Took responsibility for planning and carrying out experiments and statistical analyses in a research environment.

### TEACHING EXPERIENCE

- · Lecture: RNA-seq in Molecular Epidemiology Developed a two-part lecture explaining the basics of RNA sequencing and its application to investigate genetic variants. Set homework and final exam questions for the module.
- · Supervision: Co-supervisor of a medical undergraduate on-going bioinformatics project.
- · Volunteer: Code Club Developed my teaching abilities and provided help with coding skills to 9-13 year-olds as part of the Code Club voluntary initiative.

### SPECIFIC COMPUTATIONAL SKILLS

Operating Systems Windows, Ubuntu

**Programming Languages** Python, R, SQLite, Bash, working knowledge of C++ & Julia

Other Tools Git, Linux HPC, Conda, Docker, Nextflow

General Software MS Office, LATEX

Data types Olink & SOMA proteomics, RNA-seq, MS metabolomics,

whole genome sequences, GWAS

Bioinformatics/Statistics Package development in R and Python, Linear and mixed models,

Mendelian randomisation & colocalisation, Joint models,

Supervised learning (scikit-learn, tensorflow, pytorch), WGCNA

### ADDITIONAL COURSES

· From large datasets to biological insight by Wellcome Connecting Science & EMBL-EBI

- · Introduction to Assessment and Feedback by Imperial College London
- · Introduction to Teaching and Learning by Imperial College London
- · Profiling and optimisation in Python by Imperial College London
- · Computational Systems Biology: Deep Learning in the Life Sciences by MITx
- · Software Development with C++ by University of Birmingham Research Computing
- · The Biostars Handbook: Bioinformatics Data Analysis by Istvan Albert
- · Introduction to Computer Science and Programming Using Python by MITx
- · Introduction to Computational Thinking and Data Science by MITx
- · Data Analysis for Life Sciences and Genomics Data Analysis by HarvardX