

# Dylan Lawless



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## Experience

2023– Senior scientist. Bioinformatics, intensive care/neonatology unit, [University of Zurich](#).

2018–2023 Senior postdoctoral scientist. Precision medicine, translational genomics, host-pathogen immunology, EPFL, [Fellay lab](#).

2017–2023 Consultancy and genomic analysis via [LawlessGenomics.com](#).

2015–2018 Bioinformatics, genetic discovery in rare PID, University of Leeds, [Savic lab](#).

2014–2015 Analytical Scientist, clinical trials, [ACM global](#).

2014 Intracellular innate immunity, EPFL, [Ablasser lab](#).

2013 Viral immunology, Trinity College Dublin, [Bowie lab](#).

2012 Innate immunity in microbiology, University College Cork [Morgan lab](#).

## Education

2015–2020 PhD University of Leeds, [School of Medicine](#) and [St. James's University Hospital](#). Novel genetic discoveries in rare primary immunodeficiencies.

2013–2014 MSc Immunology, [Trinity College Dublin \(1st class hon\)](#).

2009–2013 BSc Microbiology, [University College Cork \(Honours\)](#).

2022 Drug/device product development and regulation in Europe and US, [epfl.ch/educate4life](http://epfl.ch/educate4life).

## Expertise

- **Leadership in genomics and bioinformatics:** Orchestrated precision medicine studies, clinical and translational genomics, and design of statistical and visualization tools. Developed collaborations with [SPHN](#), [PHRT](#), [Genomics England](#), [UKBB](#), [SPHN](#), [NIHR-RD](#). Supervised numerous international research projects, MSc and PhD students, teaching in EPFL.
- **Contribution in rare diseases:** Directly involved in diagnosing severe rare diseases, leading to discovery of novel mechanisms and genetic disorders, enabling precise clinical interventions and enhancing patient care.
- **Proficiency in multi-omics:** Hands-on experience with [DNAseq](#), [RNAseq](#), [methylation seq](#), proteomics, [cytokine](#) and [antibody](#) profiling, transcriptomics, [GWAS](#), and [FACS and flow cytometry](#) single-cell analysis. Developed various statistical techniques for multi-modal data analysis and predictive modeling.

- **Pathogen genomics and immunology:** Parallel analyses of host, bacterial, viral, and fungal mechanisms in various contexts including neonatal sepsis and [viral](#) infection. Comprehensive understanding of immunodeficiencies, host-pathogen interactions, and immunogenomics.
- **Programming and statistics:** Strong command over R, bash, and other Unix/Linux tools, with ongoing interest in other languages and big data software. Proficient in diverse statistical techniques including mixed models, multivariable modelling, and sensitivity analysis.
- **Data visualization and regulation:** Ability to present [complex genomic data](#) to varied audiences. Knowledge of drug/device development, regulation, quality compliance, and project management from FDA, EUdralEx, and ICH.
- **Democratized genomics and wet-lab expertise:** Genomic analysis lead through [LawlessGenomics](#) including collaboration and cloud-based pipeline development. Significant wet-lab experience, including *in vitro* and *in vivo* models, molecular assays, and clinical cohort design.

## Selected Publications

- *Journal of Allergy and Clinical Immunology*. Oct 2023; doi: [10.1016/j.jaci.2023.09.023](#). You AIn't using it right—artificial intelligence progress in allergy. - [pdf](#)
- *Journal of Allergy and Clinical Immunology*. Feb 2023; doi: [10.1016/j.jaci.2023.01.035](#). Prevalence of *CFTR* variants in PID patients with bronchiectasis - an important modifying co-factor. - [pdf](#)
- *The Journal of Infectious Diseases*. Nov 2022; doi: [10.1093/infdis/jiac442](#). Viral genetic determinants of prolonged respiratory syncytial virus infection among infants in a healthy term birth cohort - [pdf](#)
- *Blood*. Jun 2020; doi: [10.1182/blood.2020005844](#). Germline *TET2* loss-of-function causes childhood immunodeficiency and lymphoma. - [pdf](#)
- *Journal of Clinical Immunology*. 2019 Aug; doi: [10.1007/s10875-019-00670-z](#). Predicting the occurrence of variants in *RAG1* and *RAG2*. - [pdf](#)
- *Frontiers in Immunology*. Jul 2018; doi: [10.3389/fimmu.2018.01527](#). A case of AOSD caused by a novel splicing mutation in *TNFAIP3* successfully treated with tocilizumab. - [pdf](#)
- *Journal of Allergy and Clinical Immunology*. Feb 2018; doi: [10.1016/j.jaci.2018.02.007](#). Prevalence and clinical challenges among adult PID patients with recombination-activating gene deficiency. - [pdf](#)
- *eLife*. Dec 2021; doi: [10.7554/elife.72559](#). Biallelic mutations in calcium release activated channel regulator 2A (*CRACR2A*) cause a primary immunodeficiency disorder - [pdf](#)
- *Arthritis & Rheumatology*. Sep 2020; doi: [10.1002/art.41531](#). A novel *RELA* truncating mutation in familial Behçet's Disease-like mucocutaneous ulcerative condition. - [pdf](#)
- *Journal of clinical immunology*. Dec 2019; doi: [10.1007/s10875-019-00735-z](#). Expanding Clinical Phenotype and Novel Insights into the Pathogenesis of ICOS Deficiency. - [pdf](#)
- *Blood*. Dec 2018; doi: [10.1182/blood-2018-07-866939](#). A novel *RAG1* mutation reveals a critical in vivo role for HMGB1/2 during V(D)J recombination. - [pdf](#)
- *Science Translational Medicine*. 2016 Mar; doi: [10.1126/scitranslmed.aaf1471](#). Familial autoinflammation with neutrophilic dermatosis reveals a regulatory mechanism of pyrin activation. - [pdf](#)

Dylan Lawless,  
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