REGINA H REYNOLDS

Molecular biologist turned bioinformatician, with experience translating largescale omics data into actionable insights for drug discovery. I've led crossfunctional analyses across transcriptomics, genomics, and functional screens, collaborating closely across teams to build reproducible pipelines that support impactful R&D decisions. I'm driven by a commitment to scientific rigour, integrity, and transparency — and by a broader desire to make a meaningful contribution to the world through thoughtful, data-driven science.

View this resume online with links at https://rhreynolds.github .io/cv

WORK EXPERIENCE

Present 2022

Lead Bioinformatician

CoSyne Therapeutics

♀ London, UK

- Provide scientific and strategic leadership as co-lead of a 5-person computational biology team, setting research priorities in collaboration with team members, experimental and AI teams, and senior leadership.
- · Drive company-wide 'omics initiatives spanning whole-genome sequencing, transcriptomics and CRISPRi screens to support target discovery in glioblastoma multiforme.
- · Establish scalable, robust, and reproducible computational workflows through adoption of tools such as Nextflow and Segera Cloud for workflow orchestration, Docker for containerisation, R targets for reproducible exploratory data analysis, and Cruft templates for project scaffoldina.
- · Contribute to shaping company culture through active involvement in recruitment and development of CoSyne's personal development framework

2022 2021

Research Fellow

University College London

♀ London, UK

- · Lead analyst involved in processing and analysing transcriptomic data generated with the aim of identifying molecular signatures of Parkinson's disease progression. Work performed using R, Nextflow and
- · Co-lead of Code and Pipeline Alignment Working Group in the Aligning Sciences Across Parkinson's initiative. This group aimed to maximise the value of data generated from finite post-mortem brain tissues through code alignment, which would enable eventual meta-analysis.
- · Published 1 co-first author research article.

2016 2016

Research Assistant

University of Copenhagen

• Copenhagen, Denmark

· Led project exploring the interactions between miR-34a, Sirt1 and p53 in a Huntington's disease mouse model, which culminated in a first author publication².

CONTACT

G GitHub

in LinkedIn

ResearchGate

PROGRAMMING LANGUAGES

The long-form version of my CV, with a list of conferences attended, teaching experience and voluntary work is available here.

Made with the R packages datadrivency and pagedown.

The source code is available GitHub.

Last updated on 2025-06-15.



2021 2016

PhD, Bioinformatics

University College London

♀ London, UK

- Thesis: Exploring the importance of cell-type-specific gene expression regulation and splicing in Parkinson's disease³.
- · Integrated GWAS summary statistics with bulk/single-cell transcriptomic, eQTL, and chromatin accessibility data to identify celltype-specific regulatory mechanisms in Parkinson's disease. Methods used included partitioned heritability, COLOC, and MAGMA.
- · Applied transcriptomic methods, including cell-type deconvolution, differential splicing analysis, and RNA-binding protein motif analysis, to investigate splicing alterations and their relevance to Lewy body diseases.
- Published 3 first/co-first author research articles and 1 first author review. Successfully secured £10,000 from Signe og Peter Gregersens Mindefond to undertake transcriptional profiling of Parkinson's disease brain tissue.

2016 2014

MSc, Molecular Biomedicine

University of Copenhagen

Ocenhagen, Denmark

- · Thesis: Changes in the miR-34a-SIRT1 axis in Huntington's disease
- · Grade: A (92.5%)

2013 2010

BSc. Molecular Biomedicine

University of Copenhagen

Openhagen, Denmark

- · Thesis: Pro-apoptotic factors in Huntington's disease: a study in the R6/2 transgenic mouse model
- · Grade: A (96.7%)



KEY PUBLICATIONS

2023

Local genetic correlations exist among neurodegenerative and neuropsychiatric diseases4

NPJ Parkinson's disease

- · Reynolds, RH, Wagen, AZ, Lona-Durazo, F, Scholz, SW, Shoai, M, Hardy, J, Gagliano Taliun, SA, Ryten, M
- · Role: Co-first author, lead analyst and corresponding author.
- · Analysis⁵ of local genetic correlations between neurodegenerative and neuropsychiatric disorders, with the aim of identifying genomic regions and genes that may drive pleiotropy.

A full list of publications is available online at https:// rhreynolds.github.io/cv

2021

Cross-platform transcriptional profiling identifies common and distinct molecular pathologies in Lewy body disorders⁶

Acta Neuropathologica

- · Feleke, R, Reynolds, RH, Smith, A, Tilley, B, Gagliano Taliun, SA, Hardy, J, Matthews, PM, Gentleman, S, Owen, D, Johnson, MR, Srivastava, P, Ryten,
- · Role: Co-first author and analyst.
- Transcriptomic analysis of cell-type-specific changes in the Lewy body diseases.

2019

Informing disease modelling with brain-relevant functional genomic annotations⁸

Brain

- · Reynolds, RH, Hardy, J, Ryten, M, Gagliano Taliun, SA
- · Role: First author.
- Review of conceptual advances in the generation of brain-relevant functional genomic annotations and among tools that allow integration of these annotations with genome-wide association summary statistics.

2019

Moving beyond neurons: the Role of cell type-specific gene regulation in Parkinson's disease heritability⁹

NPJ Parkinson's disease

- Reynolds, RH, Botía, JA, Nalls, MA, International Parkinson's Disease Genomic Consortium (IPDGC), System Genomics of Parkinson's Disease (SGPD), Hardy, J, Gagliano Taliun, SA, Ryten, M
- · Role: First author and lead analyst.
- Analysis of Parkinson's disease common variation, with the aim of identifying cell types and pathways of importance to disease risk.



- 1: https://parkinsonsroadmap.org/research-network/pd-functional-genomics/
- 2: https://pubmed.ncbi.nlm.nih.gov/29289683/
- 3: https://discovery.ucl.ac.uk/id/eprint/10119171/
- 4: https://pubmed.ncbi.nlm.nih.gov/37117178/
- 5: https://rhreynolds.github.io/neurodegen-psych-local-corr/
- 6: https://pubmed.ncbi.nlm.nih.gov/34309761/
- 7: https://rhreynolds.github.io/LBD-seq-bulk-analyses/
- 8: https://pubmed.ncbi.nlm.nih.gov/31603214/
- 9: https://pubmed.ncbi.nlm.nih.gov/31016231/