BRIAN M. SCHILDER

Passionately pursuing transdisciplinary research to advance human health and knowledge.

Imperial College London PhD Student MPhil, ScB



EDUCATION

2024

Imperial College London / UK Dementia Research Institute /
The Alan Turing Institute

PhD: Clinical Medical Research; Computational Neurogenomics ♥ London, UK

Supervisors: Nathan G. Skene, PhD; Paul M. Matthews, PhD

Thesis: Multi-omic medicine: dissecting the cell type-specific and pleiotropic mechanisms underlying disease genomics at scale

- Aim 1) Dissect the multi-scale mechanisms (e.g. genes, pathways, cell types, phenotypes) underlying all rare disease genomics.
- Aim 2) Decompose the phenome (all diseases and traits) into a unified latent genomic space to identify pleiotropy and disease trajectories at scale.
- *Aim* 3) Demonstrate and facilitate FAIR (Findable, Accessible, Interoperable, Reproducible) practices.

2017

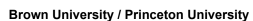
The George Washington University / Georgetown University
MPhil: Human Paleobiology; Evolutionary Neuroscience & Genomics

Supervisors: Chet C. Sherwood, PhD; Brenda J. Bradley, PhD

Thesis: The evolution of the hippocampus and adult neurogenesis: Novel insights into the origins of human memory

- *Aim 1*): Identify human-specific features of hippocampal subfield organization, adult neurogenesis, and their ecological correlates.
- Aim 2) Identify human-specific patterns of hippocampal subfield gene expression.
- *Aim 3*) Identify the genetic mechanisms mediating the evolution of human hippocampal neuroanatomy and gene expression. †

2011



ScB: Cognitive Neuroscience; Neurological Diseases & Disorders

Providence, RI, USA

Supervisors: Sheila E. Blumstein, PhD; David L. Sheinberg, PhD

CONTACT

□ brian_schilder [at] alumni.brown.edu

US

+1 908-268-9859

UK

+44 073-0653-7736

in LinkedIn

ORCID

G Google Scholar

() GitHub

™ Twitter

YouTube

Personal Website

\$\rightarrow\$ Lab Website

SUMMARY

(ii) 15+ years of research

23 publications

> 4 preprints

№ 41 software packages

11 databases & apps

i□ 22 talks

2 13+ years of teaching & team management

TABLE OF CONTENTS

Education

✓ Skills

Publications

Preprints

Acknowledgements

Reviewerships

i□ Internal talks

Invited talks

†□ Conference talks

#□ Posters

(E) Experience

Packages

□ Websites

Databases

\$ Grants

Q Awards

Affiliations

▼ Data visualisation

Extracurricular

Updated May-13-2024

CORE SKILLS

Research

- 15+ years of deep expertise in a genomics, AI, evolutionary biology and biomedicine. Seamlessly fuses ideas and methods across multiple domains.
- Publication record: 23 publications, 4 preprints and 13 awarded grants.
- Reproducibility: Global leader in promoting and enabling High-performance computing: Highly parallelised reproducible scientific practices. E Writes 100% reproducible manuscripts programmatically.
- Bioinformatics: Created 45 Python and R packages to address key challenges in biological research.
- anayses and AI model training (CPUs and GPUs).
 - · Web development: 6+ websites, web apps, and interactive reports.

AI & Machine Learning

Proficient in developing and deploying AI/MI models (PyTorch, tensorflow, Keras and sklearn) to solve complex biological problems. Applied examples include:

- Causal variant prediction: Used functional impact predictions from DNA sequence models (DeepSEA, Basenji, IMPACT) to validate SNPs prioritised with Bayesian fine-mapping.
- · Single-cell omics: Used autoencoders that take RNA expression and ESM protein embeddings (SATURN, CellBLAST) to integrate and embed scRNA-seq atlases from multiple species.
- LLM knowledge extraction: Developed a framework to extract quantitative metrics of phenotype severity from GPT-4.
- Disease genomics embeddings: Developed VAE- and graph-based dimensionality reduction models to create a joint latent representation of genome-wide signatures from all known diseases and phenotypes.
- Topic modeling: Created a suite of proprietary Python packages for NLP-based topic modelling of the PubMed literature to provide business intelligence to the world's largest digital health, biotech, and pharma companies (as a consultant with 120/80 Group).
- · Collaborative AI: Uses generative AI assistants (GitHub Copilot) to accelerate and augment coding.

Project Management

Efficient management strategies to define objectives, track progress and coordinate diverse teams.

- Documentation: Defines obectives and tracks progress with GitHub Projects. Includes useful documentation in Issues, inline code and shareable reports.
- Version control: Extensive and daily use of GitHub, containers (Docker, Singularity, virtual machines),
- environments (conda) and pipelines (Nextflow).
- Team management: Led numerous collaborative research projects and supervised researchers at various career stages.

Soft skills

Advances science through effective problem formulation, collaboration and communcation.

- Problem formulation: Rapid hypothesis generation, project design, and creative problem solving.
- · Collaboration: Diverse and global collaborative networking.
- · Communication: Clear and concise distillation of complex results to a variety of audiences. Presented 25 conference posters.

2023

PUBLICATIONS

2023

rworkflows: automating reproducible practices for the R community

Nature Communications (2023) 15(149); https://doi.org/10.1038/s41467-023-44484-5

BM Schilder, AE Murphy, NG Skene

News

- Featured in Nature Communications Editors' Highlights

Artificial intelligence for neurodegenerative experimental models

Alzheimer's & Dementia (2023) http://doi.org/10.1002/alz.13479

SJ Marzi, BM Schilder, A Nott, C Sala Frigerio, S Willaime-Morawek, M Bucholc, DP Hanger, C James, PA Lewis, I Lourida, W Noble, F Rodriguez-Algarra, JA Sharif, M Tsalenchuk, LM Winchester, U Yaman, Z Yao, DEMON Network, JM Ranson, DJ Llewellyn

2023		Artificial intelligence for dementia genetics and omics Alzheimer's & Dementia (2023) http://doi.org/10.1002/alz.13427
		C Bettencourt, NG Skene, S Bandres-Ciga, E Anderson, LM Winchester, IF Foote, J Schwartzentruber, JA Botia, M Nalls, A Singleton, BM Schilder , J Humphrey, SJ Marzi, CE Toomey, A Al Kleifat, EL Harshfield, V Garfield, C Sandor, S Keat, S Tamburin, C Sala Frigerio, I Lourida, DEMON Network, JM Ranson, DJ Llewellyn
2023		Artificial intelligence for dementia research methods optimization
2023		Alzheimer's & Dementia (2023) http://doi.org/10.1002/alz.13441
		M Bucholc, C James, A Al Khleifat, A Badhwar, N Clarke, A Dehsarvi, CR Madan, SJ Marzi, C Shand, BM Schilder , S Tamburin, HM Tantiangco, I Lourida, DJ Llewellyn, JM Ranson
2023		EpiCompare: R package for the comparison and quality control of epigenomic peak files
		Bioinformatics Advances (2023) 13(1):vbad049; https://doi.org/10.1093/bioadv/vbad049 S Choi, BM Schilder, L Abbasova, AE Murphy, NG Skene
2022		Dissecting the Shared Genetic Architecture of Suicide Attempt, Psychiatric Disorders, and
		Known Risk Factors
		Biological Psychiatry (2022) 91(3):313-327; https://doi.org/10.1016/j.biopsych.2021.05.029
		N Mullins, J Kang, Al Campos, BM Schilder , et al.
2022		Genetic analysis of the human microglial transcriptome across brain regions, aging and disease pathologies
		Nature Genetics (2022) https://doi.org/10.1038/s41588-021-00976-y
		K de Paiva Lopes, G JL Snijders, J Humphrey, A Allan, M Sneeboer, E Navarro, BM Schilder T Raj
		■ News- Microglial transcriptomics meets genetics: new disease leads (Nature Reviews Neurology, 2022)
		- Mighty MiGA: Microglial Genomic Atlas Zeros in on Causal AD Risk Variants (ALZFORUM, 2022)
		- Can a Human Microglial Atlas Guide Brain Disorder Research? (Mount Sinai Health System, 2022)
		- Polygenic Scores Paint Microglia as Culprits in Alzheimer's (ALZFORUM, 2021)
2021		Multi-omic insights into Parkinson's Disease: From genetic associations to functional mechanisms
		Neurobiology of Disease (2021) 105580; https://doi.org/10.1016/j.nbd.2021.105580
		BM Schilder, E Navarro, T Raj
2021		Fine-Mapping of Parkinson's Disease Susceptibility Loci Identifies Putative Causal Variants
202.		Human Molecular Genetics (2021) ddab294; https://doi.org/10.1093/hmg/ddab294
		BM Schilder, T Raj
2021		echolocatoR: An Automated End-to-End Statistical and Functional Genomic Fine-Mapping
		Pipeline Pipeline
		Bioinformatics (2021) btab658; https://doi.org/10.1093/bioinformatics/btab658
		BM Schilder, J Humphrey, T Raj
2021		MungeSumstats: A Bioconductor Package for the Standardisation and Quality Control of Many GWAS Summary Statistics
		Bioinformatics (2021) 37(23):4593-4596; https://doi.org/10.1093/bioinformatics/btab665
		A Murphy, BM Schilder , NG Skene
2021	Ī	Dysregulation of mitochondrial and proteo-lysosomal genes in Parkinson's disease myeloid cells Nature Genetics (2021) https://doi.org/10.1101/2020.07.20.212407
		E Navarro, E Udine, K de Paiva Lopes, M Parks, G Riboldi, BM Schilder T Raj
		■ News
		- Mount Sinai: Fighting Neurodegenerative Disorders (Mount Sinai Health System, 2019)
2021		Phenome-wide and eQTL Associations of COVID-19 Genetic Risk Loci
		iScience (2021) https://doi.org/10.1016/j.isci.2021.102550
		C Moon, BM Schilder , T Raj, K-I Huang

Genome-Wide Association Study of over 40,000 Bipolar Disorder Cases Provides Novel Biological Insights

Nature Genetics (2021) 53:817-829; https://doi.org/10.1038/s41588-021-00857-4

N Mullins, AJ Forstner, KS O'Connell, B Coombes, JRI Coleman...**BM Schilder**... et al.

- Researchers identify 64 regions of the genome that increase risk for bipolar disorder (EurekAlert, 2021)
- Largest Bipolar Disorder Genetics Study Doubles Genetic Risk Factors (Nordic Society of Human Genetics and Precision Medicine, 2021)

Tensor decomposition of stimulated monocyte and macrophage gene expression profiles identifies neurodegenerative disease-specific trans-eQTLs

PLOS Genetics (2020) 16(9):e1008549; https://doi.org/10.1101/499509

S Ramdhani, E Navarro, E Udine, AG Efthymiou, BM Schilder, M Parks, A Goate, T Raj

Evolutionary shifts dramatically reorganized the human hippocampal complex

Journal of Comparative Neurology (2019) 528(17):3143-3170; https://doi.org/10.1002/cne.24822 BM Schilder, HM Petry, PR Hof

• FAIRshake: Toolkit to Evaluate the Findability, Accessibility, Interoperability, and Reusability of Research Digital Resources

Cell Systems (2019) 9; https://doi.org/10.1016/j.cels.2019.09.011

D Clarke, L Wang, A Jones, M Wojciechowicz, D Torre, K Jagodnik, S Jenkins, P McQuilton, Z Flamholz, M Silverstein, BM Schilder...A Ma'ayan

News

2020

2019

2019

2019

2018

2015

2015

2014

2014

- Chosen as 'Featured Frontmatter' article in Cell Systems

Geneshot: search engine for ranking genes from arbitrary text queries

Nucleic Acids Research (2019) 47(W1):W571-W577; https://doi.org/10.1093/nar/gkz393

A Lachmann, **BM Schilder**, ML Wojciechowicz, D Torre, MV Kuleshov, AB Keenan, A Ma'ayan

- Geneshot: Piercing the Literature to Identify and Predict Relevant Genes (University of Pittsburgh Health Sciences Library System Update, 2019)
- The Future of AI at the Hasso Plattner Institute for Digital Health at Mount Sinai (Mount Sinai Health System, 2020)

eXpression2Kinases (X2K) Web: linking expression signatures to upstream cell signaling networks

Nucleic Acids Research (2018) 46(W1):W171-W179; https://doi.org/10.1093/nar/gky458

DJB Clarke, MV Kuleshov, **BM Schilder**, D Torre, ME Duffy, AB Keenan, A Lachmann, AS Feldmann, GW Gundersen, MC Silverstein, Z Wang

I News

- Mount Sinai Faculty Spotlight: Ma'ayan Lab (Mount Sinai Health System, 2018)

Defining elemental imitation mechanisms: A comparison of cognitive and motor-spatial imitation learning across object- and computer-based tasks

Journal of Cognition and Development (2015) 17(2):221-243; https://doi.org/10.1080/15248372.2015.1053483 F Subiaul, L Zimmerman, E Renner, BM Schilder, R Barr

Take the monkey and run

Journal of Neuroscience Methods (2015) 248:28-31; http://doi.org/10.1016/j.jneumeth.2015.03.023

KA Phillips, MK Hambright, K Hewes, **BM Schilder**, CN Ross, SD Tardif **B News**

- Monkeys on a Treadmill? A Conversation with Dr. Kimberley Phillips (Why Social Science?)

Becoming a high-fidelity - super - imitator: what are the contributions of social and individual learning?

Developmental Science (2014) 18(6):1025-1035; http://doi.org/10.1111/desc.12276 F Subiaul, EM Patterson, **BM Schilder**, E Renner, R Barr

Working memory constraints on imitation and emulation

Journal of Experimental Child Psychology (2014) 128:190-200; http://doi.org/10.1016/j.jecp.2014.07.005 F Subiaul, BM Schilder

PREPRINTS

2024

Harnessing AI to annotate the severity of all phenotypic abnormalities within the Human Phenotype Ontology

medRxiv (2024)

KB Murphy, BM Schilder, NG Skene

2023 Fine-mapping genomic loci refines bipolar disorder risk genes

medRxiv (2023) https://www.medrxiv.org/content/10.1101/2024.02.12.24302716v1

M Koromina, A Ravi, G Panagiotaropoulou, **BM Schilder**, ... S Ripke, T Raj, JRI Coleman, N Mullins **B News**

- Currently under journal review

2023

Identification of cell type-specific gene targets underlying thousands of rare diseases and subtraits

medRxiv (2023) https://doi.org/10.1101/2023.02.13.23285820

KB Murphy, R Gordon-Smith, J Chapman, M Otani, BM Schilder, NG Skene

2022

CUT&Tag recovers up to half of ENCODE ChIP-seq peaks

bioRxiv (2022) https://doi.org/10.1101/2022.03.30.486382

D Hu, L Abbasova, BM Schilder, A Nott, NG Skene, SJ Marzi

ACKNOWLEDGEMENTS

2021

eQTL Catalogue: a compendium of uniformly processed human gene expression and splicing QTLs.

Nature Genetics (2021) 53:1290-1299; https://doi.org/10.1038/s41588-021-00924-w N Kerimov, JD Hayhurst, K Peikova et al.

2020

Functionally-informed fine-mapping and polygenic localization of complex trait heritability

Nature Genetics (2020) https://doi.org/10.1038/s41588-020-00735-5

O Weissbrod...AL Price

2019

Wayfinding: The science and mystery of how humans navigate the world.

St. Martin's Press (2019) ISBN-13: 978-1250096968; https://www.amazon.co.uk/Wayfinding-Science-Mystery -Humans-Navigate/dp/1250096960

MR O'Connor

2012

EEG oscillations reveal neural correlates of evidence accumulation

Frontiers in Decision Neuroscience (2012) 6(106):Jan-13; https://doi.org/10.3389/fnins.2012.00106

M van Vugt, P Simen, L Nystrom, P Holmes, J Cohen

2011

Trial-by-trial adaptation of decision making performance: a model-based EEG analysis

Interdisciplinary Perspectives on Cognition, Education, and the Brain (2011) 7; https://www.semanticscholar.org/paper/Trial-by-trial-adaptation-of-decision-making-a-EEG-Vugt-Simen

/330371d08842ecd1bda332dd22351a7135b5cb1f

M van Vugt, P Simen, J Cohen



REVIEWERSHIPS

2024

[Unpublished article]

BMJ Open (2024)

2023		Multi-region brain transcriptomes uncover two subtypes of aging individuals with differences in Alzheimer risk and the impact of APOΕε4 Neuron (2023) https://doi.org/10.1101/2023.01.25.524961
		AJ Lee, Y Ma, L Yu, RJ Dawe, C McCabe, K Arfanakis, R Mayeux, DA Bennett, HU Klein, PL De Jager
2023		Summary statistics-based association test for identifying the pleiotropic effects with set of genetic variants Bioinformatics (2023) https://doi-org.iclibezp1.cc.ic.ac.uk/10.1093/bioinformatics/btad182 D Bu, X Wang, Q Li
2021		Most pathways can be related to the pathogenesis of Alzheimer's Disease Alzheimer's Research & Therapy (2021) https://doi.org/10.3389/fnagi.2022.846902 SL Morgan, P Naderi, K Koler, Y Pita-Juarez, D Prokopenko, IS Vlachos, RE Tanzi, L Bertram, WA Hide
2021		CLIP: accurate prediction of disordered linear interacting peptides from protein sequences using co-evolutionary information Bioinformatics (2021) https://doi.org/10.1093/bib/bbac502 Z Peng, Z Li, Q Meng, B Zhao, L Kurgan
2020		Single-cell transcriptomics and in situ morphological analyses reveal microglia heterogeneity across the nigrostriatal pathway Neurobiology of Disease (2020) https://doi.org/10.3389/fimmu.2021.639613 O Uriarte Huarte, D Kyriakis, T Heurtaux, Y Pires-Afonso, K Grzyb, R Halder, M Buttini, A Skupin, M Mittelbronn, A Michelucci
2019		Deconstructing cerebellar development cell by cell PLOS Genetics (2019) https://doi.org/10.1371/journal.pgen.1008630 MJ van Essen, S Nayler, EBE Becker, J Jacob
2019		Partitioning the genetic architecture of amyotrophic lateral sclerosis Nature Neuroscience (2019) https://doi.org/10.1101/505693 IJ Broce, RS Desikan
	ήū	INTERNAL TALKS
2022		Drug (re)discovery in the age of genomics: multi-omic strategies for identifying disease treatments Seminar Drug Discovery and Trials Optimisation Working Group Deep Dementia Phenotyping Network (DEMON)
2021		We've tagged a lot of cells, and sorted them in wells, some of the reads were double, So we looked into the trouble Seminar UK Dementia Research Institute Imperial College London
2021		Beyond GWAS: getting more out of genomic data in the age of machine learning
		Methods Optimisation Working Group Deep Dementia Phenotyping Network (DEMON)
2021		Interspecies translation of single-cell transcriptomic signatures Seminar Experimental Models Working Group Deep Dementia Phenotyping Network (DEMON)
2020		Automated consensus fine-mapping of neurological disorder genomics Seminar
	1	UK Dementia Research Institute (UK DRI)

忙 INVITED TALKS Harnessing AI to annotate the severity of all phenotypic abnormalities within the Human 2024 Phenotype Ontology **Turing Omics Meeting** Omics Data Generation & Analysis Group The Alan Turing Institute Multi-omics medicine: investigating shared genetic risk factors to better understand 2023 neurodegenerative disease **Turing Omics Meeting** Omics Data Generation & Analysis Group The Alan Turing Institute Decomposing the phenome: learning the latent genomic structure underlying thousands of 2022 diseases and traits Neuroepidemiology of Aging Webinar RUSH Alzheimer's Disease Center (RADC) **RUSH University** Drug (re)discovery in the age of genomics: multi-omic strategies for identifying disease 2022 treatments Department Seminar 3D (Drug, Disease, Delivery) Center / Department of Pharmaceutical Sciences University of South Dakota Statistical and functional genetic fine-mapping across multiple disease 2020 Seminar Alzheimer's Disease Sequencing Project Columbia University / Icahn School of Medicine at Mount Sinai Statistical and functional genetic fine-mapping across multiple disease 2020 Laboratory of Neurogenetics Friday Workshop National Institute on Aging National Institutes of Health **†**ONFERENCE TALKS rworkflows: taming the Wild West of R packages 2023 EuroBioc2023 Bioconductor 45-minute workshop. rworkflows: taming the Wild West of R packages 2023 BioC2023 Bioconductor

10-minute talk within the Infrastructure Track.

2023

2022

Navigating the rare diseases landscape: A comprehensive approach to identify gene therapy targets based on cell type-phenotype associations

Intelligent Systems For Molecular Biology / European Conference on Computational Biology (ISMB/ECCB)

International Society for Computational Biology (ISMB)

20-minute talk within the Bio-Ontologies COSI Track.

Systematic quantification of animal model viability across human diseases

Informatics-Synapse Joint Early Career Researcher Meeting

UK Dementia Research Institute (UK DRI)

2020	Automated genetic fine-mapping of neurological disorders London Genetics Network	
	The Genetics Society 6-minute talk	
2019	Parkinson's disease derived monocytes show alteration in the phago-lysosomal pathway American Society of Human Genetics (ASHG) Annual Meeting	
	American Society of Human Genetics (ASHG) Co-contributor	
2017	Comparative neuroanatomy of navigational maps in primates JB Johnston Club for Evolutionary Neuroscience	
	Society for Neuroscience (SfN) Co-contributor	
2016	The evolution of human hippocampal gene expression JB Johnston Club for Evolutionary Neuroscience	
	Society for Neuroscience (SfN)	
2015	The neurobiological effects of exercise on marmoset models of Multiple Sclerosis Marmoset Social	
	Society for Neuroscience (SfN)	
2015	The neurobiological effects of exercise on marmoset models of Multiple Sclerosis JB Johnston Club for Evolutionary Neuroscience	
	Society for Neuroscience (SfN)	
	CONFERENCE POSTERS	
2024	Navigating the Phenomic Landscape: systematic characterisation of the latent genomic spunderlying all traits and diseases Target to Patient (2024) https://www.ebi.ac.uk/industry/targettopatient/ BM Schilder, NG Skene	oace
2023	CUT&Tag recovers up to half of ENCODE ChIP-seq peaks Connectome (UK Dementia Research Institute) (2023) https://ukdri.ac.uk/	
	J Ismail, D Hu, L Abbasova, BM Schilder, A Nott, NG Skene, SJ Marzi	
2023	Navigating the rare diseases landscape: A comprehensive approach to identify gene thera targets based on cell type-phenotype associations Intelligent Systems For Molecular Biology / European Conference on Computational Biology (ISMB/ECC (2023) https://www.iscb.org/ismbeccb2023 BM Schilder, KB Murphy, R Gordon-Smith, J Chapman, M Otani, NG Skene	
2023	Identification of cell type-specific gene targets underlying thousands of rare diseases and clinical phenotypes Genomics of Rare Diseases (2023) https://coursesandconferences.wellcomeconnectingscience.org/eve/genomics-of-rare-disease-20230424/ BM Schilder, KB Murphy, R Gordon-Smith, J Chapman, M Otani, NG Skene	
2023	Statistical and Functional Fine-Mapping as a Powerful Tool to Unravel the Biological Etiological Disorder Biological Psychiatry (2023) 93(9):S18; https://doi.org/10.1016/j.biopsych.2023.02.063 M Koromina, A Ravi, BM Schilder, B Muller, J Coleman, T Raj	ogy of

2023	UK Dementia Research Institute Scientific Advisory Board (2023) BM Schilder, NG Skene
2022	Systematic quantification of animal model viability across human disease Rising Scientist Day at Imperial College London (2022) BM Schilder, NG Skene
2022	A compehensive statistical and functional fine-mapping pipeline applied to Bipolar Disorder GWAS risk loci European Neuropsychopharmacology (2022) 63:e14; http://dx.doi.org/10.1016/j.euroneuro.2022.07.037 M Koromina, A Ravi, BM Schilder, B Muller, J Coleman, T Raj, N Mullins
2021	Genetic Effects on Human Microglia Transcriptome in Neuropsychiatric Diseases Biological Psychiatry (2021) 89(9):S84-S85; https://doi.org/10.1016/j.biopsych.2021.02.225 G Snijders, K de Paiva Lopes, J Humphrey, S Allan, M Sneeboer, R Navarro, BM Schilder, R Vialle, M Parks, R Missall, W van Zuiden, F Gigase, R Kubler, AB van Berlekom, C Bottcher, J Priller, R Kahn, L de Witte, T Raj
2020	Cell-type-specific reconstruction of primate evolution from genomic positive selection Rising Scientist Day at Imperial College London (2020) K Murphy, BM Schilder, NG Skene
2019	Automated genetic and functional fine-mapping of Parkinson's Disease Loci American Society of Human Genetics (2019) BM Schilder, T Raj
2019	Parkinson's disease derived monocytes show alteration in the phago-lysosomal pathway American Society of Human Genetics (2019) E Udine, E Navarro,BM Schilder,T Raj
2018	Learning X2K: Parameter Optimization via Genetic Algorithms to Calibrate the Expression2Kinases Pipeline Illuminating the Druggable Genome (2018) BM Schilder, A Lachmann, M Kuleshov, A Ma'ayan
2018	Learning X2K: Parameter Optimization via Genetic Algorithms to Calibrate the Expression2Kinases Pipeline Big Data 2 Knowledge - Library of Integrated Network-Based Cellular Signatures (LINCS) (2018) BM Schilder, A Lachmann, M Kuleshov, A Ma'ayan
2017	The evolution of the human hippocampus and neuroplasticity Association for American Physical Anthropologists (2017) https://www.abstractsonline.com/pp8/index.html#! /4071/presentation/4471 BM Schilder, BJ Bradley, CC Sherwood
2016	The molecular evolution of plasticity and the human hippocampus Society for Neuroscience (2016) https://www.abstractsonline.com/pp8/index.html#!/4071/presentation/4471 BM Schilder, BJ Bradley, CC Sherwood
2015	Effects of exercise on disease progression and cognition in the marmoset EAE model JB Johnston Club for Evolutionary Neuroscience (2015) KA Phillips, MK Hambright, K Hewes, BM Schilder, B Jagessar, B t'Hart, SD Tardif
2015	The effects of climatic trends, variability, and rates of change on mammalian brain evolution Association for American Physical Anthropologists (2015) BM Schilder, WA Barr, R Bobe, CC Sherwood

2015 Individual, Observational, and Imitation Learning in Orangutans and Children

Association for American Physical Anthropologists (2015)

E Renner, BM Schilder, F Subiaul

2014 The helper hinderer task revisited: an infant eye tracking study

The George Washington University Research Day (2014)

A Gokhale, BM Schilder, F Subiaul

Dendritic morphology of pyramidal neurons across the visual stream: A direct comparison of chimpanzees and humans

Society for Neuroscience (2013)

BM Schilder, O Adeyo

The striatum in the evolution of learned vocalizations: Understanding the neurobiological precursors to human speech using a chimpanzee model

Society for Neuroscience (2013)

S Bianchi, T Duka, G Muntane, BM Schilder, CD Stimpson, WD Hopkins

Imitation & emulation in a novel box task

Association for Psychological Science (2013)

L Zimmerman, N Brito, C Mendelson, R Barr, E Renner, BM Schilder, F Subiaul

A study of imitation and working memory in 2- to 4- year-olds

Association for Psychological Science (2013)

R Barr, F Subiaul, L Zimmerman, L Renner, BM Schilder, C Mendelson, L Golojuch

The impact of wealth on sharing preferences in children

Child Development Society (2013)

J Miller, BM Schilder, L Peizer, F Subiaul

RESEARCH EXPERIENCE

Lead Data Scientist

120/80 Group

New York, NY, USA

- Offers data-driven consultation services to a wide portfolio of high-profile digital healthcare, pharmaceutical and biotech companies.
- Developed a suite of proprietary softwares to extract customised business intelligence from the published literature to generate customised and interpretable reports to clients.
- Provides clients guidance on strategic Al implementation, data analysis, publication and transparency.

₂₀₂₀ • Bioinformatician II

Icahn School of Medicine at Mount Sinai (Department of Neuroscience / Department of Neurology / Department of Genetics & Genomics / Ronald M. Loeb Center for Alzheimer's Disease)

New York, NY, USA

- Developed machine learning systems to integrate large-scale multi-omics datasets (e.g. whole-genome sequencing, bulk and single-cell RNA-seq, epigenomics, clinical data) to uncover the molecular mechanisms underlying neurodegenerative diseases (e.g. Alzheimer's, Parkinson's, ALS).
- Computationally identified specific disease-causal variants, pathways and cell-types for subsequent functional wet lab validation (e.g. CRISPR-cas9 editing in patient-derived cell cultures, iPSCs and cerebral organoids).

Bioinformatician II

Icahn School of Medicine at Mount Sinai (Department of Pharmacological Sciences)

New York, NY, USA

- Conducted computational systems biology research. Integrated and analyzed large-scale genomic and biomedical data (e.g. Python, R, JavaScript).
- · Developed evolutionary algorithm to optimize gene network kinase regulator prediction (eXpression2Kinases).
- Developed and deployed computational tools, software, databases and web applications for basic and clinical research, resulting in 3 peer-reviewed publications.

2019

2013

2013

2013

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2013

2020 I

| 2018

> 2018 | 2017

> > 10

Participant 2017 Technische Universität Dresden / eMed (Summer School in Systems Medicine) • Frauenchiemsee, Germany Attended lectures and extended skills in extraction and analysis of big data from biomedical and neurogenomic resources. · Developed, performed and wrote manuscript for collaborative bioinformatics research project in less than one week. **Participant** 2016 Icahn School of Medicine at Mount Sinai (Scientific Computing & Data Science) New York, NY, USA · Intensive summer school in high-performance computing, coding, genome database utilization and bioinformatics methods including transcriptomics and genetic association testing. 2017 Collaborator Trinity University / Southwestern National Primate Research Center (Department of Neuroscience) 2014 San Antonio, TX, USA Investigated the neurobiological mechanisms underlying the ameliorating effects of exercise on relapse-remitting Multiple Sclerosis. 2014 Teaching Assistant / Project Leader The George Washington University / Rutgers University (Department of Anthropology) Ileret, Kenya · Served as Teaching Assistant while excavating Lower Paleolithic hominin sites (Homo, Paranthropus). As Project Leader, investigated the running biomechanics of local Daasanach tribespeople while mentoring undergraduate students. Research Assistant 2013 The George Washington University (Department of Anthropology) Washington, DC, USA 2011 Performed dissection, histology, microscopy and quantitative stereology in post-mortem primate brain tissues. · Trained junior and senior personnel on lab protocols. Senior Lab Manager 2013 The George Washington University (Department of Speech, Language & Hearing Sciences) 2011 Washington, DC, USA Organized and trained dozens of undergraduates to conduct weekly cognitive development research; designed and/or directly contributed to over 15 research projects in two years. Volunteer Researcher 2012 University of Winnipeg / University of Belgrade (Department of Anthropology / Department of Archaeology) Sićevo, Serbia Excavated Paleolithic fossils and tools (H. heidelbergensis, H. neanderthalensis) at Mala Balanica, Velika Balanica, and Pešturina sites. Volunteer Researcher 2011 Universidad de Murcia (Department of Zoology & Physical Anthropology) Murcia, Spain • Excavated Paleolithic fossils and tools from Cueva Negra (H. heidelbergensis) and Sima de las Palomas (H. neanderthalensis) with an international research team. Volunteer Research Intern 2011 New York, NY, USA American Museum of Natural History (Division of Anthropology) · Contributed to paleoanthropological research on primate fossils using 3D morphometry imaging equipment including Minolta, Microscribe and CT. Paid Research Intern 2010 Princeton University (Princeton Neuroscience Institute) Princeton, NJ, USA • Investigated the neural basis of decision-making in humans. • Recruited participants, recorded EEG and analyzed data in MATLAB. **Student Researcher** 2010 Brown University (Department of Cognitive, Linguistic & Psychological Sciences) Providence, RI, USA 2009 • Experimental Analysis of Animal Behavior & Cognition: Conducted various operant conditioning experiments on rats. Gained experience in animal behavioral training, data collection, and data analysis in MATLAB. Laboratory in Genes and Behavior: Tested transgenic mice with modified N-type voltage-gated calcium channel subunits in a battery of cognitive and sensorimotor tasks. Results were published.

TEACHING / MENTORING EXPERIENCE

2020

Research Mentor

Imperial College London (Department of Brain Sciences / Department of Life Sciences)

Q London, UK

- · Mentored students and affiliated projects:
- Kitty Murphy (PhD): 'Evolutionary pressures on cell types: leveraging species differences to gain insight into neurodegenerative disease risk'
- · Sheen Lei (BSc): 'Benchmarking cell-type-specific enrichment of genome-wide disease signatures'
- Ted Reese (MSc): 'Computational cell-type annotation of single-cell epigenomics data'
- Xindong Sun (MSc): 'Benchmark of Targeted insertion of promoters sequencing (TIP-seq) on histone modification H3K27ac and H3K27me3 in K562 cell line'
- Shuhan Shen (MSc): 'Evaluation and optimisation of methods for identifying the cell types underlying genetic disease signatures'
- Lusheng Li (MSc): 'Genetic identification of cell types underlying mammalian phenotypes'
- Sera Choi (BSc): 'EpiCompare: R package for QC and benchmarking epigenetic datasets'
- Emilie Cottard (MSc) & Will Lunt (BSc): 'A meta-analysis of selective cell-type vulnerability in Parkinson's Disease neuropathology'
- Jai Chapman (BSc): 'Expression Weighted Cell Type Enrichment as a Tool for Identifying Cell Types Underlying Rare Disease Phenotypes'
- Bobby Gordon-Smith (MSc): 'Identification of cell types involved in rare disease-associated human phenotypes'
- Leyla Abbasova (MSc): 'Analysis and optimisation of CUT&Tag for epigenomic profiling of the brain'
- Barney Hill (BSc): 'Identification of cell-types associated with latent factors inferred from phenome-wide GWAS summary statistics'

2020 | 2019

Research Mentor

Icahn School of Medicine at Mount Sinai (Department of Neuroscience / Department of Neurology / Department of Genetics & Genomics / Ronald M. Loeb Center for Alzheimer's Disease)

New York, NY, USA

• Mentored MS, MD, and PhD students in projects focused on computational exploration of phenotype clustering and genomic regulation of neurodegenerative diseases.

2018

Research Co-mentor

Icahn School of Medicine at Mount Sinai (Department of Pharmacological Sciences)

New York, NY, USA

- · Mentored students and affiliated projects:
- Vivian Utti (BSc): 'ChEA3: Transcription Factor Enrichment Analysis' as part of the Summer Research Training Program in Biomedical Big Data Science.
- Mary Duffy (PhD): 'Predicting upstream kinase regulators from interaction network databases'
- Zach Flamholz (BSc): 'modEnrichr: a suite of gene set enrichment analysis tools for model organisms'

2018

Guest Lecturer

Icahn School of Medicine at Mount Sinai (Department of Pharmacological Sciences)

New York, NY, USA

• Lectured on data visualization in Python and Jupyter notebooks in the PhD/MD course 'Programming for Big Data Biomedicine'.

2017 | 2016

Research Mentor

The George Washington University (Department of Anthropology)

Washington, DC, USA

- · Mentored students and affiliated projects:
- Jamie Kleiner (BSc): 'Animal model simulating MS and exercise's impact on adult hippocampal neurogenesis'

2015

Teaching Assistant

The George Washington University (Department of Anthropology)

Washington, DC, USA

- Course: 'Human Brain Evolution'
- Guest lectured, graded all assignments and exams, and provided additional educational support during office hours.

2014

Teaching Assistant

The George Washington University (Department of Psychology)

Washington, DC, USA

- Course: 'Biological Psychology'
- Led undergraduates in article discussions, graded all assignments and exams, and provided additional educational support during office hours.

2014 • Teaching Assistant

The George Washington University (Department of Anthropology)

Washington, DC, USA

- · Course: 'Biological Anthropology'
- Led undergraduate students in two, 2-hour lab sessions per week, graded lab assignments and exams, and provided additional educational support during office hours.

2013 Research Mentor

2013

2012

2013

2011

The George Washington University (Department of Psychology)

Washington, DC, USA

- · Mentored students and affiliated projects:
- · Anushka Gokhale (BSc): 'Infants' Social Assessment of Characters Through Eye Gaze'

Lab Protocol Trainer

The George Washington University (Department of Anthropology)

Washington, DC, USA

• Trained undergraduate, graduate, and post-doctoral researchers in Social Cognition Lab and Lab for Evolutionary Neuroscience in a variety of methodological research protocols.

مع

SOFTWARE PACKAGES

1. MSTExplorer R

Multi-Scale Targets Explorer: Systematically identify, prioritise and visualise cell-type-specific gene therapy targets across the phenome.

- https://github.com/neurogenomics/MSTExplorer
- https://doi.org/10.1101/2023.02.13.23285820

2. **HPOExplorer R**

Import, annotate and visualise the 18k+ hierarchically structured clinical phenotypes across the Human Phenotype Ontology.

- https://github.com/neurogenomics/HPOExplorer
- https://doi.org/10.1101/2023.02.13.23285820

3. **KGExplorer R** 🖑

Query, construct, and analyse large-scale biomedical knowledge graphs and ontologies.

- https://github.com/neurogenomics/KGExplorer
- 4. autoCV 😱 😇 😇 🧓

Automatically generate and style your CV from tables.

- https://github.com/bschilder/autoCV
- 5. anndataR R 🕏

Bring the power and flexibility of AnnData to the R ecosystem, allowing you to effortlessly manipulate and analyze your single-cell data.

https://github.com/scverse/anndataR

6. gptPhD R

7.

Query Large Language Models for the purposes of systematically extracting biomedical knowledge.

https://github.com/neurogenomics/gptPhD

ThreeWayTest 😱

Summary statistics-based association test for identifying the pleiotropic effects with set of genetic variants.

https://github.com/bschilder/ThreeWayTest

8. SCAVENGE R

Variant to function mapping at single-cell resolution through network propagation.

- https://github.com/sankaranlab/SCAVENGE
- https://doi.org/10.1038/s41587-022-01341-y

9. rworkflows 🕝 🥞

Continuous integration for R packages. Automates testing, documentation website building, and containerised deployment.

- https://github.com/neurogenomics/rworkflows
- https://doi.org/10.21203/rs.3.rs-2399015/v1

TIPseeker R 10. R package for post-processing [single-cell] TIP-seq data. https://github.com/neurogenomics/TIPseeker PeakyFinders R 11. R package for mining, calling, and importing epigenomic peaks. https://github.com/neurogenomics/PeakyFinders graphiti 🖳 12. Extract colour palettes from graffiti artworks. https://github.com/bschilder/graphiti SkillNet R 13. Creates user-specific contribution networks from GitHub Organization repositories. https://github.com/neurogenomics/SkillNet phenoRx 😱 14. Make cell type-specific predictions for patients based on clinical phenotypes and/or risk genes. https://github.com/neurogenomics/phenoRx phenomix 😱 🕏 15. R package for the exploration and analysis of many genotype-phenotype datasets at once. https://github.com/neurogenomics/phenomix MAGMA.Celltyping R 16. Identify cell types underlying the associations found in GWAS summary statistics. https://github.com/neurogenomics/MAGMA_Celltyping **EWCE** 17. Expression Weighted Celltype Enrichment. https://github.com/NathanSkene/EWCE EpiCompare R 18 R package for QC and benchmarking epigenetic datasets. https://github.com/neurogenomics/EpiCompare https://doi.org/10.1101/2022.07.22.501149 19. orthogene R Interspecies gene mapping. https://github.com/neurogenomics/orthogene 20. MungeSumstats R Standardise the format of summary statistics from GWAS. https://github.com/neurogenomics/MungeSumstats https://doi.org/10.1093/bioinformatics/btab665 scNLP 😱 21. Tools for applying natural language processing (NLP) techniques to single-cell (sc) omics data. https://github.com/neurogenomics/scNLP scKirby 😱 柫 22. Automated ingestion and conversion of various single-cell data formats. https://github.com/neurogenomics/scKirby geneshotR R 23. R package for querying and processing results from Geneshot. https://github.com/bschilder/geneshotR templateR 🖳 24. Self-updating template for developing R packages. https://github.com/neurogenomics/templateR https://doi.org/10.21203/rs.3.rs-2399015/v1 25. echoverseTemplate R Self-updating template for creating echoverse R packages. https://github.com/RajLabMSSM/echoverseTemplate/ https://doi.org/10.1093/bioinformatics/btab658

echolocatoR 😱 🗂 26. R package for end-to-end statistical and functional fine-mapping with extensive dataset access. https://github.com/RajLabMSSM/echolocatoR https://doi.org/10.1093/bioinformatics/btab658 27. echodata 😱 Examples of fine-mapped GWAS summary statistics, data formatting functions, and API access to the echolocatoR Fine-mapping Portal. https://github.com/RajLabMSSM/echodata https://doi.org/10.1093/bioinformatics/btab658 echoannot R 28. Functions for annotating genomic data with annotations and epigenomic data. https://github.com/RajLabMSSM/echoannot https://doi.org/10.1093/bioinformatics/btab658 echoplot R 29. R package for LocusZoom-inspired GWAS/QTL visualization, with API access to LD panels. https://github.com/RajLabMSSM/echoplot https://doi.org/10.1093/bioinformatics/btab658 echoconda 😱 🕏 30. Various utility functions to find, build, and use conda environments from within R. https://github.com/RajLabMSSM/echoconda https://doi.org/10.1093/bioinformatics/btab658 echotabix 🗬 🥞 31. Tabix indexing and querying. https://github.com/RajLabMSSM/echotabix https://doi.org/10.1093/bioinformatics/btab658 echoLD 😱 락 32. LD downloading and processing. https://github.com/RajLabMSSM/echoLD https://doi.org/10.1093/bioinformatics/btab658 echofinemap 🤻 🕏 33. Statistical and functional fine-mapping functions. https://github.com/RajLabMSSM/echofinemap https://doi.org/10.1093/bioinformatics/btab658 echodeps R 34. Creates interactive dependency networks for R packages. https://github.com/RajLabMSSM/echodeps https://doi.org/10.1093/bioinformatics/btab658 35. echogithub R Access and process metadata from GitHub. https://github.com/RajLabMSSM/echogithub https://doi.org/10.1093/bioinformatics/btab658 36. devoptera 🤻 Practical tools for R developers. https://github.com/RajLabMSSM/devoptera https://doi.org/10.1093/bioinformatics/btab658 downloadR 🖳 37. Single- and multi-threaded downloading functions.

R package for rapid API-access and colocalization of summary statistics from eQTL Catalogue.

nttps://github.com/RajLabMSSM/downloadR https://doi.org/10.1093/bioinformatics/btab658

nttps://github.com/RajLabMSSM/catalogueR https://doi.org/10.1093/bioinformatics/btab658

catalogueR R

38

15

39. TopicModeler

Proprietary Python package to run advanced topic modeling on text corpuses.

40. LinkReporter 🕏

Proprietary Python package to extract job postings and company employee listings from LinkedIn and generate interactive business intelligence reports.

41. PubReporter 🕏

Proprietary Python package for extract relevant scientific literature, gather citations, and generate interactive business intelligence reports.



1. EpiArchives 😱

Public archive for EpiCompare reports.

- https://github.com/neurogenomics/EpiArchives
- https://doi.org/10.1101/2022.07.22.501149

2. Rare Disease Celltyping Portal R

Web portal connecting to multiple R Shiny apps to explore, visualize, and download cell type-specific enrichment results and systematically prioritised gene targets for over 6,000 rare disease phenotypes.

- https://github.com/neurogenomics/rare_disease_celltyping_apps
- https://neurogenomics.github.io/rare_disease_celltyping_apps/home
- https://doi.org/10.1101/2023.02.13.23285820

3. Parkinson's Disease Omics Review R

Data and code associated with the Parkinson's Disease review paper by Schilder, Navarro & Raj (Neurobiology of Disease, 2021).

- https://github.com/RajLabMSSM/PD_omics_review
- https://rajlabmssm.github.io/PD_omics_review/
- https://doi.org/10.1016/j.nbd.2021.105580

4. Selective Vulnerability Meta-analysis 😱

Selective Vulnerability Meta-analysis: Shiny app dedicated to the exploration and dissemination of metaanalysed cell counts manually curated and harmonised from the Parkinson's Disease literature.

https://github.com/neurogenomics/SelectiveVulnerabilityMetaAnalysis

5. MAGMA Files Public R

7.

Gene enrichment files for hundreds of GWAS generated with Multi-marker Analysis of GenoMic Annotation (MAGMA) for use in downstream analyses.

https://github.com/neurogenomics/MAGMA_Files_Public

6. echolocatoR Fine-mapping Portal 😱

Access to interactive plots and fine-mapping results across many GWAS/QTL datasets using echolocatoR.

- https://github.com/RajLabMSSM/Fine_Mapping_Shiny
- https://rajlab.shinyapps.io/Fine Mapping Shiny
- https://doi.org/10.1093/bioinformatics/btab658

COVID-19 Patient Tracker 🕏

Web app for summarizing and visualizing real-time EHR data of COVID-19 patients within the Mount Sinai Health System.

8. Tensor Decomposition Shiny App R

Interactive application to explore and download all results and plots from Ramdhani et al. (PLOS Genetics, 2020).

- https://github.com/RajLabMSSM/Tensor_myeloid
- https://rajlab.shinyapps.io/Tensor_myeloid
- https://doi.org/10.1101/499509

9. Hippocampal Evolution 😱

Interactive code, results and visualization for the manuscript "Evolutionary selective pressures dramatically expanded and reorganized the human hippocampal complex".

https://github.com/bschilder/Hippo_Eco

https://bschilder.github.io/Hippo Eco/HPsubfield eco

https://doi.org/10.1002/cne.24822

10. **Geneshot** 👙 ট 😈 😈

Flexible tool to identify genes associated with any biomedical term and to predict novel target genes.

http://amp.pharm.mssm.edu/geneshot

https://doi.org/10.1093/nar/gkz393

11. **X2K** 🗐 🖑 🗓 😇

eXpression 2 Kinases (X2K) Web: Automated computational pipeline to infer kinase regulators from weighted or unweighted gene lists.

http://amp.pharm.mssm.edu/X2K

https://doi.org/10.1093/nar/gky458

WEBSITES

1. Personal Website 😇 😇 🗓

nttps://github.com/bschilder/BMSchilder
https://bschilder.github.io/BMSchilder

2. Official Raj Lab Website 😇 💆 🗓

• https://github.com/RajLabMSSM/RajLab_website • http://www.rajlab.org

\$ GRANTS

Total (all grants): \$2,949,872

Total (as primary applicant): \$311,382

₂₀₂₃ • EuroBioc2023 Scholarship,

Bioconductor

Project: 'rworkflows: taming the Wild West of R packages'

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■ News

- Awarded to support attending the EuroBioc2023 meeting.

• Role: Primary applicant

• Amount: \$250

• PI: BM Schilder

2023 PioC2023 Scholarship,

Bioconductor

Project: 'rworkflows: taming the Wild West of R packages'

News

2023

- Awarded to support attending the BioC2023 meeting. Additionally included free lodging.

• Role: Primary applicant • Amount: \$1500

• PI: BM Schilder

Junior Scientist Conference Grant,

The Genetics Society

Project: 'Identification of cell type-specific gene targets underlying thousands of rare diseases and subtraits'

· Role: Primary applicant

• PI: BM Schilder

• Amount: £750

factors' Role: Co-applicant PE NG Skene 1 Turing Community Award, Alan Turing Institute Project: Multi-omic medicine: dissecting the cell-type-specific molecular mechanisms underlying neurodegenerative disease genomics' Role: Primary applicant Prib M Schilder 2022 National Institutes of Health Project: Statistical and functional fine-mapping of bipolar disorder genetic risk loci' Role: Co-applicant Prib M Schilder Collaborative Single Cell and Spatial Transcriptomics Studies Award Programme, UK Dementia Research Institute Project: 'Amplifying genome coverage of single cell epigenetic profiling of the human brain' Role: Co-applicant Role: Role: Co-applicant Role: Role: Primary applicant Role: Role: Primary applicant Role: Role: Primary applicant Role: Role	2023		Imperial UK Research Institute Impact Acceleration Account, Imperial College London Project: 'Creating commercial kit solutions for single cell epigenetic profiling of histone marks and transcription
Alan Turing Institute Project: Wild moir medicine: dissecting the cell-type-specific molecular mechanisms underlying neurodegenerative disease genomics' Role: Primary applicant Project: 'Statistical and functional fine-mapping of bipolar disorder genetic risk loci' Role: Co-applicant Role: Functional Fine-Mapping of LRRK2 Locus Role: Fundee Role: Funder Role: Rol			factors' • Role: Co-applicant • Amount: £80,000
neurodegenerative disease genomics'			Alan Turing Institute
Project: 'Statistical and functional fine-mapping of bipolar disorder genetic risk loci' - Role: Co-applicant - Role: Primary applicant - Role: Fundee - PI: T Raj The Michael J. Fox Foundation - Role: Fundee - PI: T Raj National Science Foundation - Project: 'The evolution of the hippocampus and adult neurogenesis: insights into the origins of human memory' - Role: Primary applicant - Role: Role			• Role: Primary applicant • Amount: £1,500
Collaborative Single Cell and Spatial Transcriptomics Studies Award Programme, UK Dementia Research Institute Project: 'Amplifying genome coverage of single cell epigenetic profiling of the human brain' Role: Co-applicant	2022		Project: 'Statistical and functional fine-mapping of bipolar disorder genetic risk loci'
National Institutes of Health Project: "Cognitive Systems Analysis of Alzheimer's Disease Genetic and Phenotypic Data" • Role: Co-applicant • PI: T Raj, D Knowles UK Dementia Research Institute Project: "UK DRI at Imperial Distinguished Studentship' • Role: Primary applicant • PI: BM Schilder The Michael J. Fox Foundation Project: "The Role of Peripheral Myeloid Cells in Parkinson's Disease" • Role: Fundee • PI: T Raj National Science Foundation Project: "Functional Fine-Mapping of LRRK2 Locus' • Role: Fundee • PI: T Raj National Science Foundation Project: "The evolution of the hippocampus and adult neurogenesis: novel insights into the origins of human memory' • Role: Primary applicant • PI: BM Schilder Wenner-Gren Project: "The evolution of the hippocampus and adult neurogenesis: insights into the origins of human memory' • Role: Primary applicant • PI: BM Schilder Leakey Foundation Project: The evolution of the hippocampus and adult neurogenesis: Novel insights into the origins of human memory' • Role: Primary applicant • PI: BM Schilder Leakey Foundation Project: The evolution of the hippocampus and adult neurogenesis: Novel insights into the origins of human memory' • Role: Primary applicant • PI: BM Schilder Leakey Foundation Project: The evolution of the hippocampus and adult neurogenesis: Novel insights into the origins of human memory' • Role: Primary applicant • Amount: \$19,512	2021		Collaborative Single Cell and Spatial Transcriptomics Studies Award Programme, UK Dementia Research Institute Project: 'Amplifying genome coverage of single cell epigenetic profiling of the human brain'
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• Role: Primary applicant • Amount: \$15,000	2016		Project: 'The evolution of the hippocampus and adult neurogenesis: Novel insights into the origins of human
· I I. DIVI Octilidei		ı	·

2016

COSMOS Club

Project: 'The evolution of adult neurogenesis across primates'

- · Role: Primary applicant • Amount: \$3.250
- PI: BM Schilder



AWARDS

2023

Prize for Computational Reproducibility in Dementia Research,

UK Dementia Research Institute

Project: 'rworkflows: taming the Wild West of R packages'

■ News

- Awarded honourable mention.

2022

Prize for Computational Reproducibility in Dementia Research,

UK Dementia Research Institute

Project: 'MungeSumstats: A Bioconductor package for the standardisation and quality control of many GWAS summary statistics'

■ News

- Awarded honourable mention.

2022

Poster Competition,

Rising Scientist Day

Project: 'Systematic quantification of animal model viability across human diseases'

- Awarded prize for research poster competition.

2022

Award for Outstanding Contribution,

NEUROHACK, Deep Dementia Phenotyping Network (DEMON)

Project: 'Predicting ALS drug targets using integrative multi-modal deep learning'

■ News

- Individually awarded for outstanding contributions during the NEUROHACK 2022, a competitive 4-day hackathon to apply AI in finding ALS therapeutic solutions.

2021

Prize for Computational Reproducibility in Dementia Research,

UK Dementia Research Institute

Project: 'echolocatoR: an automated end-to-end statistical and functional genomic fine-mapping pipeline' News

- Winners announced for UK DRI's first 'Prize for Computational Reproducibility in Dementia Research (UK Dementia Research Institute, 2021)
- Jointly awarded inaugural prize with Kitty Murphy.

2021

Centre Photography Competition,

UK Dementia Research Institute

Project: 'Wildfire Circle, Golden Brain, Wildfire, Geneshot, Geology of Biology, Neon Brain'

- One of the winners of the scientific image competition.

2019

Art of the Brain,

Friedman Brain Institute, Icahn School of Medicine

Project: 'Wildfire'

News

- Awarded 2nd place and Featured on the cover of Biological Psychiatry: Volume 87, Issue 12 (2020). Exhibited and auctioned at the Grady Alexis Gallery (New York City), where all proceeds were voluntarily donated to the Diversity in Neuroscience Initiative.

AFFILIATIONS



- · Synapse Working Group
- · Informatics Working Group



DEMON Network

Data science and AI for dementia

- · Genetics & Omics Working Group
- · Experimental Models Working Group
- Drug Discovery & Trials Optimisation Working Group

The Alan Turing Institute

- Turing Enrichment Scheme
- Turing-Roche Strategic Partnership
- Turing Omics Data Generation & Analysis Interest Group
- Turing Clinical Al Interest Group



· Bipolar Disorder Working Group



- Chair of the Bioconductor Cloud Methods Working Group
- Lead of the Bioconductor GitHub Actions Subgroup



Member

▼ DATA VISUALISATION / ARTWORK PORTFOLIO

2022

echoverse Dependency Graph

♥ London, UK

Hex stickers London, UK 2020 All hex stickers for R packages I've helped develop. 3D Human Phenotype Ontology 2023 **Q** London, UK · 3D force-directed graph of the Human Phenotype Ontology (clouds above) with kernel density estimation projected from the x/y planes (mountains below). Connections represent the hierarchical relationships between rare diseases and their associated symptoms/phenotypes. Associated preprint **Multi-scale Rare Disease Mechanisms** 2023 **Q** London, UK Network of systematically prioritised gene therapy targets for rare diseases Associated preprint **Curriculum Vitae Connexa** 2023 **Q** London, UK • Term co-occurrence network generated by analysing all data that went into this CV. **Experiments with Generative Al** 2021 **Q** London, UK · wombo.art: 'Multi-omic medicine: dissecting the cell-type-specific mechanisms in neurodegeneration genomics' · wombo.art: 'Multi-omic medicine: neurodegenerative disease genomics' · wombo.art: 'Multi-omic medicine: neurodegeneration' · wombo.art: 'Neurodegeneration' Lights in the dark genome: the current state of Parkinson's research 2021 **Q** London, UK • The majority of PD genetics research has focused on a relatively small number of genes. Above, are the top 75 most commonly mentioned genes in the PD literature, extracted using Geneshot. · Associated study **Pacrophage** 2020 **Q** London, UK · Colocalised genetic loci ...but shaped as Pac-Man! Associated study **Circos** 2020 **Q** London, UK Colocalised genetic loci across a variety of neurological disease GWAS and cell-type-specific QTLs. Associated study Wildfire Circle 2019 New York, NY, USA Awarded 2nd place in the 2019 Art of the Brain competition, put on by the Mount Sinai's Friedman Brain Institute. · Exhibited and auctioned at the Grady Alexis Gallery (New York City), where all proceeds were donated to the Diversity in Neuroscience Initiative . · Featured on cover of Biological Psychiatry. Wildfire 2019 New York, NY, USA • Transcriptomic data from 16k+ individual brain cells (shown as points) after reducing the dimensionality with an autoencoder and UMAP. 5 million tracts are shown interconnecting these cells, where shorter tract length represents greater similarity in their molecular profiles. **3D Brain Model** 2019 New York, NY, USA · 3D model of my brain generated from MRI scans.

S EXTRACURRICULAR EXPERIENCE

2003

Competitive Running Career

- 8+ years of varsity and Division I cross country, winter track, and spring track throughout high school and college.
- Year-round, daily training and travel to weekly competitions necessitated a dedicated and regimented lifestyle in order to succeed as a student-athlete
- Running remains a passion of mine and I enjoy training for ultra-marathons in my free time.
- · This passion, work ethic and self-insight have carried over to all aspects of my life, including my career as a researcher.

Earth

1995

Music Production

• Writes, records, produces and performs original music.

Earth

- Instruments: keys, percussion, vocals, etc.
- Proficient in Logic Pro X Digital Audio Workstation (DAW).
- Experiments with generative AI.