Sina Rüeger

(Genomic) Data Scientist

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Areas of experience

studies (GWASs)

Genome-wide association Creating GWAS pipelines, quality control procedures, GWAS for unconventional endpoints (time-to-event data, proxy cases), rare variant association analysis, summary statistics follow-up methods (colocalization, fine-mapping, causal inference, heritability estimation).

Statistical modelling Regression modelling techniques, model selection, data dimension reduction, classification, unsupervised clustering.

Data visualisation Visual representation of concepts and results.

Reproducibility

Version control, workflow automation, documentation with literate programming and best coding practices.

Domain knowledge

Genomics, Parkinson's disease, bariatric surgery, Hepatitis C, Tuberculosis & other infections, solid organ transplantation, multiple sclerosis, human height, critical care

Data sets Genomic data, brain imaging data, clinical data, questionnaire data, drugbank data.

Employment

Feb-2021 – present **Data Scientist**, data42 at Novartis, Basel, Switzerland.

Sep-2019 – Jan-2021 Data Analyst / Bioinformatician, FinnGen, Institute for Molecular Medicine Finland (FIMM), University of Helsinki, Helsinki, Finland.

Extending PheWAS with novel statistical genetics methods.

Sep-2018 - Aug-2019

Postdoctoral Researcher, Fellay Lab, Global Health Institute, EPFL, Lausanne, Switzerland.

Analysis of human and pathogen genomes for different infections.

Feb-2013 - Sep-2013 Internship, Statistical Genetics Group, Department of Medical Genetics, University of Lausanne, Lausanne, Switzerland.

Rare genetic variant analysis.

Feb-2012 - Mar-2013 Biostatistician, Division of Biostatistics, ISPMZ (now EBPI), University of Zurich, Zurich, Switzerland.

Statistical consulting.

Mar-2009 - Jan-2012 Research Assistant, Institute of Data Analysis and Process Design, Zurich University of Applied Sciences, Winterthur, Switzerland. Statistical modelling for a mix of industry projects.

Education

Oct-2013 - Sep 2018 PhD in Life Sciences, University of Lausanne, Lausanne, Switzerland, supervised by Prof. Zoltán Kutalik, Statistical Genetics Group, Institute of Social and Preventive Medicine, Lausanne University Hospital (CHUV).

Topic: Integrative statistical analysis of -omics and genome-wide association studies data.

Feb-2009 - Jun-2011

Master of Science in Engineering with Specialisation in Information and Communication Technologies, Zurich University of Applied Sciences, Winterthur,

Oct-2005 - Oct-2008 Bachelor in Engineering, Degree Program in Data Analysis and Process Design, Zurich University of Applied Sciences, Winterthur, Switzerland.

Computing skills

Statistical programming R [highly proficient], Python [familiar], SQL [familiar]

Genomics tools Plink, QCTOOL, GTCA, ...

Developer tools Unix/Linux, git, GNU make

R Daily used for statistical computing and data visualisation; familiar with the architecture of R, package development and best practices.

Computing experience Experience using cloud computing / high performance clusters for analysis and storage of large datasets.

Other activities

RStudio Tidyverse Instructor Certified since July 2020

SatRday Neuchatel 2020 Co-organiser of SatRday Neuchatel in 2020, a one day conference for R users.

R-Ladies community Founding member and former co-organiser of R-Ladies Lausanne, a bi-monthly R user group. Former community manager of R-Ladies Slack.

Languages

German [native speaker], **English** [fluent, C1], **French** [good command, B1], **Finnish** [beginner]

Awards

Apr-2017 The 'Human Heredity' Best Student Platform Presentation Award at the European Mathematical Genetics Meeting in Tartu, Estonia.

Mar-2017 *Poster prize* at the ePerMed Workshop about 'Functional annotation of genome-wide variants' in Lausanne, Switzerland.

May-2016 Lodewijk Sandkuijl Award for an outstanding talk in the field of complex genetics and statistical genetics at the European Society of Human Genetics Conference, Barcelona, Spain.

Main publications

Sina Rüeger, Christian Hammer, Alexis Loetscher, Paul J. McLaren, Dylan Lawless, Olivier Naret, Daniel P. Depledge, Sofia Morfopoulou, Judith Breuer, Evgeny Zdobnov, and Jacques Fellay. The influence of human genetic variation on Epstein–Barr virus sequence diversity. *Scientific Reports*, 11(1):4586, February 2021. ISSN 2045-2322. doi: 10.1038/s41598-021-84070-7. URL https://www.nature.com/articles/s41598-021-84070-7. Number: 1 Publisher: Nature Publishing Group

Sina Rüeger, Aaron McDaid, and Zoltán Kutalik. Evaluation and application of summary statistic imputation to discover new height-associated loci. *PLOS Genetics*, 14(5):1–32, 05 2018a

Sina Rüeger, Aaron McDaid, and Zoltán Kutalik. Improved imputation of summary statistics for realistic settings. *bioRxiv*, page 203927, 2018b

Sina Rüeger et al. Impact of common risk factors of fibrosis progression in chronic hepatitis C. *Gut*, 64(10):1605–1615, 2015