# HALFDAN RYDBECK

#### **Profile**

Recently developed a pipeline<sup>7</sup> of rscripts to perform pathway and correlation network analysis of cord blood plasma proteomics data. Developed pipelines for automatic preprocessing and delivery of Illumina and Pacbio DNA sequencing output. Set up server infrastructure for NGS data handling. Manged NGS file servers, user access and data delivery. Examples of downstream analysis: Blasted an exon sequence against multiple assembled fish genomes to look for systematic loss in phylogeny. Analyzed bacterial methylomes, sequenced with Pacbio in, respect to virulens. PhD project<sup>2</sup>. Developed an R package for clustering of genomic tracks. Scripts developed in R and The Genomic Hyperbrowser<sup>3</sup> was used to study the dependencies and select genes, based on genomic, epigenomic and transcriptomic alterations in samples from osteosarcoma and immune cells. The gene expression, promoter methylation and DNA copy number data were acquired by oligonucleotide microarray technology, and the histone modification data was acquired by technology based on chromatin immune precipitation and next generation sequencing, ChIP-sea.



# RESEARCH EXPERIENCE

2022 2020

#### Researacher bioinformatics

Core Facilities Bioinformatics vid Sahlgrenska Akademin, Univeristy of Gothenburg

**♀** Göteborg, Sweden

· Worked with Rstudio and the R package collections Bioconductor and Tidyverse. Reports where generated with R notebooks and the R package Bookdown. Github and GitKraken is used for code version control. Recently applied R packages are DEqMS for differential expression analysis of mass spectrometry data, WGCNA for proteins correlation analysis and ReactomeGSA for pathway analysis

2017 2014

### Senior engineer bioinformatics

Norwegian Sequencing Center

**Q** University of Oslo, Norway

· System setup, user management, data preprocessing, delivery and down stream analysis of Illumina and Pacbio sequence data at the Norwegian DNA sequencing center. Work

2008 2007

#### Principal engineer

Bioinformatics core facility

Rikshospitalet HF, Norway

- · Microarray gene expression analysis
- · Developing network-based methods to investigate and visualize clinically relevant patterns in data.



View this CV online with links at https://rpubs.com/halryd/cv

### CONTACT

- github.com/hrydbeck
- halfdanrydbeck.me
- in https://www.linkedin.com /in/halfdan rydbeck

## LANGUAGE SKILLS

R	
Bash	
Python	
SQL	

Made with the R package pagedown.

The source code is available on github.com/nstrayer/cv.

Last updated on 2022-03-19.

Research assistant 2007 Children's Hospital of Philadelphia, USA Human Genetics division 2006 · Bash and scripting for data wrangling and testing Solar, Plink, TDT, Merlin, STATA, SAS, R · Work primarily done in R Shiny and Javascript (node and d3js). Senior research assistant 2005 • A. I duPont Hospital for Children, USA Dept. of genetic epidemiology 2004 · Used software for genetic linkage and association studies of diseases/syndroms Velocardiofacial Syndrome, ADHD, Neuroblastoma, Osteoporosis, Cornelia de Lange's syndrome and Myopia · Solar, Plink, TDT, Merlin, STATA, SAS, R **Database coordinator** 2004 Children's Hospital of Philadelphia, USA **Human Genetics division** 2002 · Manged a database for SNP information Progeny2000 ♣■ TEACHING EXPERIENCE 2018 R for reprodicible scientific programming Oslo Norway Software carpentry 2017 · Covered introduction to R language for statistics applications · Graduate level class I like when teaching and learning is relaxing and joyfull Version controlling with git and github 2018 Oslo Norway Software carpentry 2017  $\cdot$  Topics covered from penalized regression to boosted trees and neural · Highest level course offered in department 2018 Unix Oslo Norway Software carpentry 2017 · Covered modern statistical computing algorithms · 4th year PhD level class **EDUCATION** PhD bioinformatics 2012 Oslo, Norway Medical faculty, University of Oslo 2007  $\cdot \cdot$  Thesis title: Integrative epigenome analysis  $\cdot$  courses mostly statistics

(30p) · Analysed gene expression, promoter methylation, histone

· University Graduate Fellow

occupancy and copynumber variation data with R and The Genomic

Hyperbrwser. Developed and R package for clustering of genomic tracks.

Sometimes I am almost tireless. This means that I fight hard so that the tasks or projects I undertake are completed on time and I do not give up until I've achieved the intended result





# ■ SELECTED PUBLICATIONS, POSTERS, AND TALKS

2022 2022 The proteome signature of cord blood plasma with high hematopoietic stem and progenitor cell count

Stem Cell Research

· Authored with Anders K.Nilsson, Annika Thorsell, Sofia Frändberg, Helena Barreto Henriksson, Camilla Hesse, Gunnel Hellgren, Pia Lundgren, AnnHellström

References are availabel upon request



- 1: https://github.com/halryd/Neored
- 2: https://www.duo.uio.no/handle/10852/39398?locale attribute=en
- 3: https://hyperbrowser.uio.no/hb/#!mode=basic
- 4: https://www.sciencedirect.com/science/article/pii/S1873506122001015