# HALFDAN RYDBECK

#### **Profile**

Expert in applied bioinformatics with 8+ years of experience in providing transparent and reproducible solutions for diverse areas of biological and medical research. Deep knowledge of underlying issues in need of bioinformatics support. Experience includes genetic epidemiology of rare diseases, integrative omics analysis of data from cancer tissue and immune cells. microarray analysis, preprocessing to final analysis of NGS output, sequence alignment and assembly working with non-human genomes from bacteria to fish species and wolf in evolutionary research and finally integrative analysis of proteomics, clinical and brain imaging data. Focused on keeping at the forefront of the latest within computational omics analysis. I seek to understand the healthcare environment and external trends. I have emotional intelligence, strong communication skills and am a great teammate. I am dynamic and work very hard to achieve the intended result which has been decided from the start on time. I am not pretentious. I speak my mind and make it safe for others to do so. My collegues say about me that I am hardworking, joyful and problemsolving. I love being creative and come up solutions that wasn't thought of in advance but am also careful that no negative consequences have an impact on anyone else.

### Objective

To further develop my skills in developing reproducible bioinformatics pipelines in a commercial setting and to contibute to the annotation of genomes and to the increased understanding of functional genomics.

### Relevant competences

- · NGS preprocessing and analysis
- · R, git, Github, Nexflow, Docker
- · GWAS
- · scientific communication and writing
- · project management



### RESEARCH EXPERIENCE



Researacher bioinformatics



- Developed a pipeline of rscripts to perform pathway and correlation network analysis of cord blood plasma proteomics data.
- · 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



View as html: https://rpubs.com /halryd/cv\_hr\_2022\_04\_24\_Bioinf \_OlsAro\_Crop\_Biotech

### **CONTACT**

**♀** Göteborg, Sweden

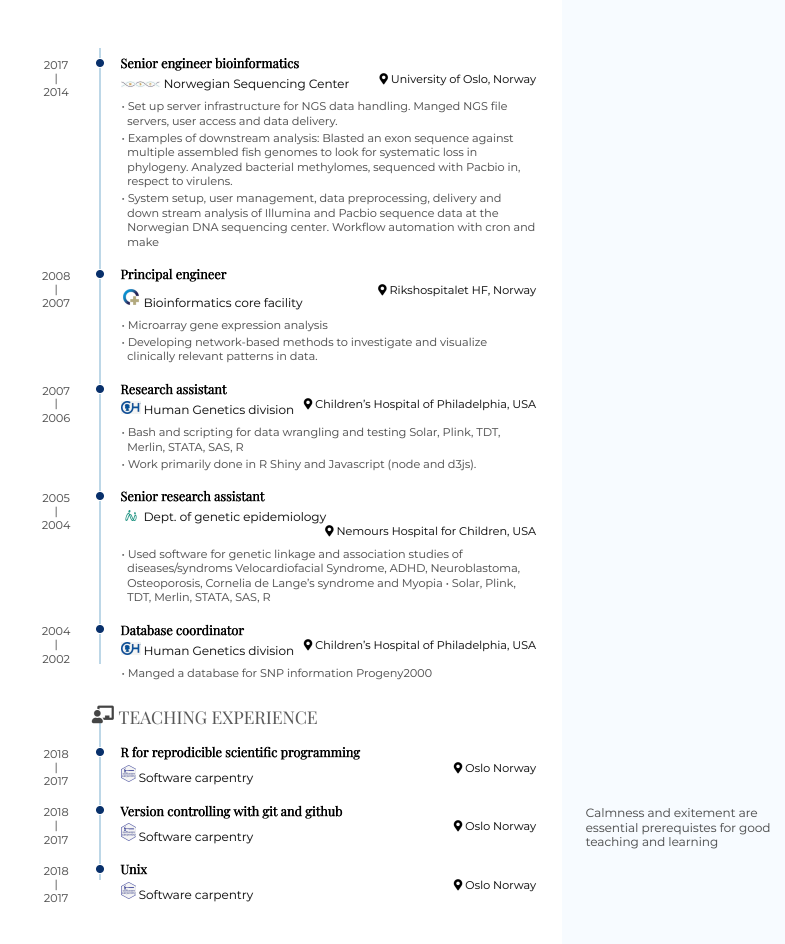
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in https://www.linkedin.com

/in/hallfalala vvytdhotetekR package pagedown.

The source code is available here.

Last updated on 2022-04-24.



## **EDUCATION**

2012 2007 PhD bioinformatics



Medical faculty, University of Oslo

Oslo, Norway

- · PhD project with thesis title: Integrative epigenome analysis<sup>7</sup>. Scripts developed in R and The Genomic Hyperbrowser 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-
- · Developed and R package for clustering of genomic tracks.

Halfdan did a "very good job as a senior engineer at NSC and is a responsible and pleasant person to work with" Professor Kjetil Sigurd Jakobssen, founder of Norwegin DNA Sequencing Center

2001 2000 **Master of Science in Bioinformatics** 



Chalmers University of Technology

Oothenburg, Sweden

1999 1993 Master of Science in Molecular biology



University of Gothenburg /Uppsala

**♀** Göteborg/Uppsala, Sweden

## ■ SELECTED PUBLICATIONS, POSTERS, AND TALKS

2022

The proteome signature of cord blood plasma with high hematopoietic stem and progenitor cell count<sup>2</sup>

Stem Cell Research

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

2015

ClusTrack: Feature Extraction and Similarity Measures for Clustering of Genome-Wide Data Sets3

Plos One

· Authored with Geir Kjetil Sandve, Egil Ferkingstad and Eivind Hovig

2010

The Genomic HyperBrowser: inferential genomics at the sequence level<sup>4</sup> Genome biology

· Authored with Geir K Sandve, Sveinung Gundersen, Ingrid K Glad, Lars Holden, Marit Holden, Knut Liestøl, Trevor Clancy, Egil Ferkingstad, Morten Johansen, Vegard Nygaard, Eivind Tøstesen, Arnoldo Frigessi & Eivind Hovig

https://www.researchgate.net /profile/Halfdan Rydbeck; ORCID identification number: 0000-0003-1606-38420000

## SPECIAL COMPETENCES

Preprocessing of NGS output

FASTQC, etc

• Downstream analysis of NGS output

Gene centric(DEG), GO and pathway centric, correlation module centric

Programming

R, shell, Python

 Version control git,Github

A Z LANGUAGES

Swedish (native)

English (native)

Norwegian (good)

☐ CONFERENCE PARTICIPATION

Pangenomes Evolution and Computation<sup>5</sup>

♣ PEDAGOGIC EXPERIENCE

Tutoring PhD students



- 1: https://www.duo.uio.no/handle/10852/39398?locale attribute=en
- 2: https://www.sciencedirect.com/science/article/pii/S1873506122001015
- 3: https://www.researchgate.net/publication/277618279\_ClusTrack\_Feature \_Extraction\_and\_Similarity\_Measures\_for\_Clustering\_of\_Genome Wide\_Data\_Sets
- 4: https://link.springer.com/article/10.1186/gb 2010 11 12 r121
- 5: https://pgec2021.schlieplab.org/PGEC2021 Program.pdf

References are available on request