

# Kristoffer Sahlin, PhD

## CONTACT INFORMATION

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Web: <http://sahlingroup.github.io/>

## CURRENT APPOINTMENTS

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**Assistant Professor**

*Jan 2020 -*

Institution: Department of Mathematics, Stockholm University

**SciLifeLab Fellow**

*Jan 2020 -*

Institution: Science for Life Laboratory, Solna

## EDUCATION & TRAINING

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**Postdoctoral researcher**

*July 2019 - Dec 2019*

University: University of Helsinki

Mentor: Professor Veli Mäkinen

**Postdoctoral researcher**

*Oct 2015 - June 2019*

University: Pennsylvania State University

Mentor: Associate Professor Paul Medvedev

**Ph.D. in Computer Science**

*Sept 2010 - Sept 2015*

University: Royal Institute of Technology (KTH), Sweden

Thesis: Algorithms and statistical models for scaffolding contig assemblies and detecting structural variants using read pair data

Advisor: Associate Professor Lars Arvestad

Co-advisor: Professor Joakim Lundberg

**M.Sc. in Mathematical Statistics**

*Aug 2008 - Sept 2010*

University: Stockholm University, Sweden

Thesis: Estimating convergence of Markov chain Monte Carlo simulations

Advisor: Assistant Professor Sebastial Höhna

**B.S. in Mathematics**

*Aug 2005 - June 2008*

University: Stockholm University, Sweden

Thesis: Splines: A theoretical and computational study

Advisor: Professor Hans Rullgård

## FUNDING AND AWARDS

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- Start-up funding from Strategiska forskningsområden (SFO) (2020-2025)
- RECOMB Travel fellowship award. (2019)
- Top five grant proposal finalist in the PacBio 2018 Iso-Seq SMRT Grant Program. (2018)
- KTH opportunities fund: Investing in research talent grant. Grant supported travel and accommodation for one month's collaboration in Helsinki, Finland. (2014)

## VISITING RESEARCH EXPERIENCE

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**Helsinki University**  
*Visiting researcher*

September 2014  
*Helsinki, Finland*

- Visiting Veli Mäkinen's lab for work on scaffolding and gapfilling of genome assemblies.

**Penn State University**  
*Visiting researcher*

November 2014  
*State college, PA, USA*

- Visiting Paul Medvedev's lab for work on structural variation detection.

## PRESENTATIONS

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### Conferences

- RECOMB-Seq (Online, 2021). Title: "Accurate spliced alignment of long RNA sequencing reads."
- ISMB, HitSeq track (Online, 2021). Title: "Strobemers: an alternative to k-mers for sequence comparison"
- SMRT Leiden 2021 (Online, May 2021). Title: "uLTRA: Accurate spliced alignment of long RNA sequencing reads." (Invited talk)
- ISMB, HitSeq track (Online, 2020). Title: "Error correction enables use of Oxford Nanopore technology for reference-free transcriptome analysis."
- RECOMB (Washington D.C., 2019). Title: "De novo clustering of long-read transcriptome data using a greedy, quality-value based algorithm."
- ISMB, HitSeq track (Chicago, 2018). Title: "IsoCon: Deciphering highly similar multi-copy gene transcripts from PacBio Iso-Seq data."
- Genome Informatics (Cold Spring Harbour, 2017). Title: "IsoCon: Deciphering highly similar multi-copy gene transcripts from PacBio Iso-Seq data."
- RECOMB (Los Angeles, 2016). Title: "Structural variation detection with read pair information: An improved null-hypothesis reduces bias."
- WABI (Atlanta, 2015). Title: "Genome scaffolding with PE-contaminated mate-pair libraries"

### Seminars/Workshops

- Lille Computational seminar series (Online, 2021) Title: "Strobemers: an alternative to k-mers for sequence comparison"
- SciLifeLab Seminar Series (Stockholm, 2020). Title: Computational methods for analysis of genome and transcriptome sequencing data.
- Bioinformatics afternoon (Helsinki, 2019). Title: Computational methods for analysis of genome and transcriptome sequencing data.
- CCBB (PennState, 2019), workshop on emerging methods for sequence analysis (2018). Title: IsoCon: Deciphering highly similar multi-copy gene transcripts from PacBio Iso-Seq data.
- Weekly Wednesday Wartik Genomics Lecture Series (PennState, 2016). Title: Genome scaffolding with PE-contaminated mate-pair libraries
- Weekly Wednesday Wartik Genomics Lecture Series (PennState, 2014). Title: An investigation of bias in methods using insert size for inference

## TEACHING

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- Lecturer and responsible for course contents

- Programming Techniques for Mathematicians, Stockholm University. Undergraduate level, 7.5 ECTS. (Spring 2020, Autumn 2020)
- Applied programming for Life Science 2, Stockholm University. Graduate level, 1.5 ECTS. (Spring 2020, Spring 2021)
- Teaching assistant
  - Statistical Methods in Applied Computer Science, KTH (graduate level). Teaching assistant 2012, 2013, and 2014.
  - Applied Bioinformatics, KTH (graduate level). Assisting lecturer and teaching assistant. I gave five lectures on introduction of python (2013).
  - Programming Techniques and Matlab, KTH (undergraduate level). Teaching assistant and recitation session lecturer (2013).
  - Programming Techniques and C, KTH (undergraduate level). Teaching assistant and recitation session lecturer (2013).
  - Bioinformatics and Biostatistics, KTH (graduate level). Teaching assistant (2013).

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## ADVISING

- Alexander Petri - PhD student. Date: September 2020 -
- Natasha Stopa - Using Female Alignment Features to Identify Reads from the Y Chromosome in Nanopore Whole Genome Sequencing Data, Master's Thesis. Date: January 2019 - January 2020.
- Josefine Röhss - Analysing k-mer distributions in a genome sequencing project. Bachelor's Thesis, Date: March 2014 - June 2014.

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## ACADEMIC SERVICE

- Program committee appointments:
  - RECOMB-Seq (2020, 2021)
- Panel session:
  - Panel member on 'Round table discussion - Young Investigator experiences' (SMRT Leiden 2021)
- Reviewer for journals (number of manuscripts):
  - Nature Computational Science (1)
  - Nature communications (1)
  - Genome Research (3)
  - Bioinformatics (8)
  - BMC Bioinformatics (2)
  - BMC Genomics (1)
  - Frontiers in Plant Science (1)
  - GigaScience (2)
  - NAR Genomics and Bioinformatics (1)
  - Communications in Statistics - Simulation and Computation (1)

- Reviewer for conferences:
  - Intelligent Systems for Molecular Biology (ISMB) (2017 - 2019)
  - Research in Computational Molecular Biology (RECOMB) (2014, 2016 - 2021)
  - RECOMB-seq (2018 - 2020)
  - Workshop on Algorithms in Bioinformatics (WABI) (2015)