# René



# Warren

warrenlr [at] gmail [dot] com

https://warrenlr.github.io

20+ Years Experience - Biotechnology | Genomics | Informatics

**Developed** first *de novo* genome assembly software (SSAKE) with short DNA sequences **Discovered** *Fusobacterium* in colon cancer, Time Magazine's 2011 top 10 breakthrough **Coordinated** bioinformatics analyses of *Rhodococus*, *Cryptococcus*, bullfrog & spruce genomes

Seeking: New Challenges | Additional Leadership

### **EXPERIENCE**

# Group Leader

**BC Cancer – Genome Sciences Centre**, Vancouver Research project concept, management, guidance Interview, supervise, mentor staff / students

## **Bioinformatics Coordinator**

**BC Cancer – Genome Sciences Centre**, Vancouver Lead bioinformatics R&D Interviewed, taught, trained, supervised staff

## **Technical Officer**

NRC – Biotechnology Research Institute, Montréal Engineered gene expression regulation technology Collaborated with stakeholders / scientists

#### **EDUCATION**

**Certificate** Concordia University

**Computer Science** 

MSC University of British Columbia
Biochemistry & Molecular Biology

**BSC** Université de Montréal Biochemistry (Honours)

#### **ACCOLADES**

2015, 16 Awarded John Jambor Knowledge Fund

2011 Interview Fusobacterium cancer discovery

2009 Genome Technology interview, next-gen. seq.

2007 GenomeWeb interview, SSAKE development

1998 Awarded MSc UBC Graduate Fellowship

1997 **Awarded** BSc *FRSQ* bursary

1996 Awarded BSc FRSQ honour's research project

1995 Worked at NASA, CMIX-4 payload

### **PRESENTATIONS**

Selected from 17 lead author

talks

talk

2017, 18 **RECOMB** Hong Kong / Paris

2015, 16, 19 ISMB Dublin / Orlando / Basel talks

2008, 12, 15 Pac. Symp. Biocomputing Hawaii posters

2010 SFAF Santa Fe talk

2007 Synthetic Biology Zürich

# **PUBLICATIONS**

Selected from 63 peer-reviewed 23 lead author

**Warren RL**, *et al.* 2019. ntEdit: scalable genome sequence polishing. *Bioinformatics*. 35:4430

**Warren RL**, et al. 2015. LINKS: Scalable, alignment-free scaffolding [...] genomes with long reads. *GigaScience*. 4:35

**Warren RL**, et al. 2012. Derivation of HLA types from shotgun sequence datasets. *Genome Med.* 4:95

Castellarin M\*, Warren RL\*, et al. 2012. Fusobacterium infection [...] in colorectal carcinoma. Genome Res. 22:299

**Warren RL**, et al. 2007. Assembling millions of short DNA sequences using SSAKE. *Bioinformatics*. 23:500

Projects SAM SSAKE TASR HLAminer LINKS XMV RAILS ARCS ntEdit

IT Skills Python PERL R MySQL HTML/js Git unix/mac/win Office

References Available