## René



# Warren

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http://renewarren.ca

20+ years experience in biotechnology, genomics, informatics

**Developed** the first *de novo* genome assembly software (SSAKE) with short DNA sequences Discovered Fusobacterium in colon cancer, one of Time Magazine's 2011 top ten breakthrough **Coordinated** bioinformatics analyses of *Rhodococus, Cryptococcus*, Bullfrog, Spruce genomes

### seeking new challenges & additional leadership

#### **EXPERIENCE ACCOLADES** now

#### **Group Leader**

BC Cancer - Genome Sciences Centre, Vancouver Lead group, project management, guidance Conceptualize, lead, develop genome technologies Interview, supervise, mentor staff / students

#### Coordinator

BC Cancer – Genome Sciences Centre, Vancouver Lead bioinformatics R&D Published research (science journals, conferences) Interviewed, taught, trained, supervised staff

### Officer

NRC Biotechnology Research Institute, Montréal Engineered gene expression regulation technology Designed, fabricated, tested DNA "gene switch" Collaborated with stakeholders, scientists

#### **EDUCATION**

Certificate Comp. Science | Concordia U.

I UBC **MSc** Biochemistry & Molecular Biology

**BSc** Biochemistry (*Honours*) | U. de Montréal

Awarded the John Jambor Knowledge Fund 2015, 16 2011 NTN24 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 for honour's research project 1995 Worked at NASA, CMIX-4 payload protein crystal.

#### **PRESENTATIONS**

Selected from 17 lead author

2017, 18	RECOMB, Hong Kong / Paris	talks
2015, 16, 19	ISMB, Dublin / Orlando / Basel	talks
2008, 12, 15	Pacific Symposium Biocomputing,	Hawaii
2010	SFAF, Santa Fe USA	talk
2007	Synthetic Biology, Zürich	talk

#### **PUBLICATIONS**

61 peer-reviewed [23 lead author], \*co-first

Warren RL, et al. 2019. ntEdit: scalable genome sequence polishing. Bioinformatics. btz400

Warren RL, et al. 2015. LINKS: Scalable scaffolding of 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 in colorectal carcinoma. Genome Res. 22:299

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

Projects: SAM, SSAKE, TASR, HLAminer, LINKS, XMV, RAILS, ARCS, ntEdit

Skills: Python, PERL, R, MySQL, HTML/js, unix/mac/win, MSOffice, Git