René



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778.386.4192 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

Group Leader

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

Bioinformatics Coordinator

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

Technical Officer

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

EDUCATION

Certificate Computer Science | Concordia U.

MSc Biochemistry & Molecular Biology | UBC -

BSC Biochemistry (Honours) | U. de Montréal -

ACCOLADES

2015, 16 Awarded John Jambor Knowledge Fund
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	Pac. Symp. Biocomputing Hawai	i posters
2010	SFAF Santa Fe	talk
2007	Synthetic Biology Zürich	talk

PUBLICATIONS

Selected from 63 peer-reviewed [23 lead author]

Warren RL, et al. 2019. ntEdit: scalable genome sequence polishing. Bioinformatics. doi: 10.1093/bioinformatics/btz400

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

IT Skills: Python, PERL, R, MySQL, HTML/js, Git, unix/mac/win, MSoffice Projects: SAM, SSAKE, TASR, HLAminer, LINKS, XMV, RAILS, ARCS, ntEdit