## Phu T. Van, PhD

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## WORK EXPERIENCE

2021Nov -**Bioinformatics Solutions Manager** 

**TwinStrand Biosciences**  Served as bioinformatics SME for the company in approving new projects. Spearheaded "à la carte" bioinformatic offering to customers to increase revenue. Provided bioinformatics guidance to customers with Field&Technical Support department. Assisted Custom Applications department in creating new genomic panels. Supervised R&D collaborations between bioinformatic and wetlab technicians. Oversaw development of new R data analysis packages. Initiated and later supervised data analyses that expanded TwinStrand DuplexSeq(TM) support to two new sequencing platforms. Produced whitepapers and reviewed marketing materials for technical accuracy. Co-authored customer communication SOPs. Reference: Clint Valentine III, MS · clint@twinstrandbio.com

2021 Jan - Oct Bioinformatics Scientist II

**TwinStrand Biosciences**  Performed analyses on duplex sequencing data for numerous mutagenesis and cancer diagnostic projects. Reviewed and contributed R code to internal bioinformatic pipelines and customer-facing reports. Prepared data products and presented findings to industry and academic clients. Reference: Fang Yin Lo, PhD · fangyinlo@gmail.com

2014 - 2020 Postdoc → Bioinformatics Analyst

Fred Hutchinson Cancer Research Center

Coordinated projects among PhD-level analysts, led brainstorming sessions and journal clubs. Performed analyses on flow cytometry and transcriptomic datasets. Created data analysis pipelines & interactive data visualizations. Authored Statistical Analysis Plans & peer-reviewed manuscripts. Reviewed junior analysts' code, mentored PhD students and bench scientists on statistics. Developed R code to normalize mass cytometry data across experiments using multi-mixture models. Performed dimension reduction on mass cytometry data and identified correlates of disease using regression.

Reference: Amy Paguirigan, PhD · apaguiri@fredhutch.org

2009 - 2014 **Doctoral Student** 

Carnegie Mellon University Designed and built a patented high-dynamic-range protein gel imager with robotic gel cutting arm for capturing rare proteins. Developed SIGILab, a C++ GUI application controlling gel imager's acquisition of high-dynamic-range images. Developed bash scripts to quantify protein abundances in 2DE gel images. Lectured in Modern Biology course and mentored junior students. Reference: Jonathan Minden, PhD · minden@cmu.edu

TECHNICAL SKILLS

Programming R, Python, Matlab, C/C++, bash, JavaScript, Perl

Data Analysis / dimension reduction, clustering, regression/classification, linear models, variable Statistics

selection, hypothesis testing

Databases MySQL/MariaDB, PostgreSQL, SQLServer

Workflows Nextflow, Slurm, Docker, GitHub BWA, STAR, RSEM, BLAST, SAMtools, BCFtools, VardictJava, varscan, delly, limma, edgeR

## EDUCATION

2009 - 2014 Carnegie Mellon University

PhD, Biological Sciences

2001 - 2007 University of Washington

BS, Biology (Physiology specialization); BS, Wildlife Sciences

## SELECTED PUBLICATIONS & PATENTS

complete list: https://scholar.google.com/citations?user=IDadFEkAAAAJ

2023 May Duplex Sequencing Provides Detailed Characterization of Mutation Frequencies and Spectra in the Bone Marrow of MutaMouse Males Exposed to Procarbazine Hydrochloride

Archives of Toxicology This manuscript characterizes genetic mutations in procarbazine-exposed mice using Duplex Sequencing as a potential replacement for the gold-standard Ames test. As the contract analyst to HealthCanada, I performed sequence alignment, variant calling and downstream analyses on the data. I also provided some data analysis assistance to the first author. This collaboration with HealthCanada also resulted in the manuscript below (Cho *et al.*).

Authors: Annette Dodge, Danielle LeBlanc, Andrew Williams, Phu T. Van, et al.

2023 May Error-corrected Duplex Sequencing enables direct detection and quantification of mutations in human TK6 cells with strong inter-laboratory consistency

Mutation Research

This manuscript characterizes genetic mutations in ENU-exposed cultured human cells as part of the same HealthCanada collaboration above. I performed sequence alignment, variant calling and downstream analyses on the data. I also provided some data analysis assistance to the first author.

Authors: Eunnara Cho, Carol Swartz, Andrew Williams, Miriam Rivas, Leslie Recio, Kristine Witt, Elizabeth Schmidt, Jeffry Yaplee, Thomas Smith, Phu T. Van, et al.

2021 Jun Monocyte metabolic transcriptional programs associate with resistance to tuberculin skin test/interferon- $\gamma$  release assay conversion

Journal of Clinical Investigations

This manuscript reports a potential link between oleic metabolism and *Tuberculosis* resistance in humans. As the primary analyst on the project, I aligned the raw transcriptomic data to the genome, quantified gene expression, created data visualizations and made recommendations on downstream statistical/machine learning methods. I also wrote the computational parts of the manuscript. Authors: Jason Simmons, Phu T. Van, *et al.* 

2019 Jul US10362237: Structured illumination system for increased dynamic range in quantitative imaging

United States Patent "The systems disclosed herein employ an iterative image collection strategy that utilizes structured illumination to achieve greater than 1,000,000-fold dynamic range measurements, representing a dramatic improvement over the prior art." Inventors: Jonathan Minden, Frederick Lanni, Phu T. VAN