# Phu T. Van, PhD

Computational Biologist

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#### **SUMMARY**

I'm interested in scientific or technical positions on collaborative teams working to solve complex problems in computational biology.

#### TECHNICAL STRENGTHS

ProgrammingR, Python, Matlab, C/C++, bash, JavaScript, PerlData Analysisdimension reduction, clustering, regression/classificationStatisticslinear models, variable selection, hypothesis testing

DatabasesMySQL/MariaDB, PostgreSQL, SQLServerWorkflowsNextflow, WDL, Slurm, Docker, GitHub

Markup/Interfaces LaTeX, Markdown, Shiny, HTML

**Bioinformatics** BWA, STAR, RSEM, BLAST, SAMtools, BCFtools

VardictJava, varscan, delly, limma, edgeR

#### **EXPERIENCE**

#### TwinStrand Biosciences

November 2021 - Present

Bioinformatics Solutions Manager

Seattle, WA

- · Served as bioinformatics expert on company panel for approving new projects.
- · Spearheaded "à la carte" bioinformatics offering to customers to increase revenue.
- · Provided bioinformatics guidance to customers and co-authored customer communication SOPs with Field&Technical Support department.
- · Mentored Bioinformatics Solutions scientists.
- · Oversaw development of new R data analysis packages.
- $\cdot$  Initiated and later supervised data analyses that expanded TwinStrand DuplexSeq<sup>TM</sup> support to two new sequencing platforms.
- · Produced whitepapers and reviewed marketing materials for technical accuracy.

#### TwinStrand Biosciences

January 2021 - October 2021

Seattle, WA

Bioinformatics Scientist II

- Performed analyses on duplex sequencing data for diverse projects including mutagenesis and cancer MRD.
- · Reviewed and contributed R code to internal bioinformatic pipelines and customer-facing reports.
- $\cdot$  Prepared data products and presented findings to industry and academic clients.

#### Fred Hutchinson Cancer Research Center

October 2015 - December 2020

Bioinformatics Analyst, Analyst Group Leader

Seattle, WA

- · Performed differential gene expression and geneset enrichment analysis on transcriptomic datasets.
- · Performed variable selection and sensitivity analyses on flow cytometry datasets.
- · Led the lab's PhD-level analysts in brainstorming sessions and journal club.
- · Coordinated analyses on projects with external collaborators.
- · Authored Statistical Analysis Plans and peer-reviewed manuscripts.
- · Created bioinformatic analysis pipelines using R, Nextflow and WDL.
- · Created interactive data visualizations using RMarkdown, igraph and leaflet.

· Mentored PhD students and bench scientists on statistics, advised hiring and performed reviews of junior analysts' code.

#### Fred Hutchinson Cancer Research Center

October 2014 - October 2015

Postdoctoral Fellow

Seattle, WA

- · Developed R code to normalize mass cytometry data across experiments using multi-mixture models.
- · Worked with engineers from FlowJo Inc. (Ashland, OR) to develop parsing in of XML-encoded flow cytometry data in FlowJo v10.
- · Performed dimension reduction on mass cytometry data and identified correlates of disease.

## Carnegie Mellon University

August 2009 - May 2014

PhD Student

Pittsburgh, PA

- · 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 processes that improved reduction and alkylation of complex protein samples in preparation for mass spectrometric sequencing.
- · Developed workflows to quantify protein abundances in 2DE gel images using bash.

## Institute for Systems Biology

May 2006 - August 2009

Research Associate

Seattle, WA

- · Identified factors that affect peptide detectability in mass spectrometry experiments using R.
- · Constructed a SQLServer database with web frontend for exploring mass spectrometry data and planning proteomic experiments.
- · Developed a regression algorithm in R to model regulators of stress response in microarray data.

### **EDUCATION**

## Carnegie Mellon University

2014

PhD, Biological Sciences

#### University of Washington

2007

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

### CONFERENCES, PATENTS AND PUBLICATIONS

- Eunnara Cho, Carol D Swartz, Andrew Williams, Miriam Rivas, Leslie Recio, Kristine L Witt, Elizabeth K Schmidt, Jeffry Yaplee, Thomas H Smith, Phu Van, et al. Error-corrected duplex sequencing enables direct detection and quantification of mutations in human tk6 cells with strong inter-laboratory consistency. Mutation Research/Genetic Toxicology and Environmental Mutagenesis, page 503649, 2023
- 2. Annette E Dodge, Danielle LeBlanc, Andrew Williams, Phu Van, Jake Higgins, Fang Yin Lo, Jeffry Yaplee, Clint C Valentine, Jesse J Salk, Carole L Yauk, et al. Characterisation of procarbazine-induced mutation spectrum in the bone marrow of mutamouse males using duplex sequencing. In ENVIRONMENTAL AND MOLECULAR MUTAGENESIS, volume 63, pages 92–93. WILEY 111 RIVER ST, HOBOKEN 07030-5774, NJ USA, 2022

- 3. E Cho, F Marchetti, P Van, J Higgins, FY Lo, J Yaplee, CC Valentine III, J Salk, and CL Yauk. Optimal methodological design for duplex sequencing (tm) in tk6 cells determined through a time and concentration response analysis following enu treatment. In *Environmental and Molecular Mutagenesis*, volume 62, page 36, 2021
- 4. D LeBlanc, M Meier, A Williams, J Buick, J Higgins, J Yaplee, P Van, FY Lo, CC Valentine, JJ Salk, et al. Duplex sequencing reveals an attenuated mutation frequency increase in the germ cells of mutamouse males exposed to n-ethyl-n-nitrosourea and benzo [a] pyrene relative to the tgr lacz assay. In *Environmental and Molecular Mutagenesis*, volume 62, page 87, 2021
- 5. AE Dodge, D LeBlanc, A Williams, P Van, J Higgins, FY Lo, J Yaplee, CC Valentine, JJ Salk, and CL Yauk. Duplex sequencing (tm) reveals increases in mutation frequencies and c; t transitions in the bone marrow of mutamouse males exposed to procarbazine. In *Environmental and Molecular Mutagenesis*, volume 62, page 87, 2021
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- 9. M. Blundon, V. Ganesan, B. Redler, P. T. Van, and J. S. Minden. Two-Dimensional Difference Gel Electrophoresis. *Methods Mol. Biol.*, 1855:229–247, 2019
- 10. D. F. E. Ker, S. Eom, S. Sanami, R. Bise, C. Pascale, Z. Yin, S. I. Huh, E. Osuna-Highley, S. N. Junkers, C. J. Helfrich, P. Y. Liang, J. Pan, S. Jeong, S. S. Kang, J. Liu, R. Nicholson, M. F. Sandbothe, P. T. Van, A. Liu, M. Chen, T. Kanade, L. E. Weiss, and P. G. Campbell. Phase contrast time-lapse microscopy datasets with automated and manual cell tracking annotations. Scientific Data, 5:180237, 11 2018
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- 13. P. T. Van, V. Ganesan, V. Bass, A. Parthasarathy, D. Schlesinger, and J. S. Minden. In-gel equilibration for improved protein retention in 2DE-based proteomic workflows. *Electrophoresis*, 35(20):3012–3017, Oct 2014
- 14. P. T. Van, V. Bass, D. Shiwarski, F. Lanni, and J. Minden. High dynamic range proteome imaging with the structured illumination gel imager. *Electrophoresis*, 35(18):2642–2655, Sep 2014
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