# Phu T. Van, PhD

Bioinformatics Solutions Manager

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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 STAR, RSEM, BLAST, SAMtools/BCFtools, DEG, GSEA, KEGG, STRING

#### **EXPERIENCE**

#### TwinStrand Biosciences

November 2021 - Present Seattle. WA

Bioinformatics Solutions Manager

- · Provided bioinformatics guidance to external clients through Field&Technical Support departments.
- · Supported Custom Applications department in creating new genomic panels.

· Served as bioinformatics expert for the company in approving new projects.

- · Supervised Advanced Sequencing Technology projects in collaboration with Laboratory Sciences department.
- · Oversaw production of new R software packages in collaboration with Bioinformatics Development department.
- · Spearheaded data mining collaborations with the Data Engineering department and co-led joint Machine Learning Interest Group.
- · Reviewed company marketing and training materials for technical accuracy.
- · Initiated and later supervised data analyses that expanded DuplexSeq(TM) support to two new sequencing platforms.

#### TwinStrand Biosciences

January 2021 - October 2021

Bioinformatics Scientist II

Seattle, WA

- Performed analyses on duplex sequencing data for diverse projects including mutagenesis and cancer MRD.
- · Contributed R code to internal bioinformatic pipelines and customer-facing reports.
- · Presented findings to commercial and academic clients.
- · Performed code reviews of R packages.

#### 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 REPRESENTATIVE PUBLICATIONS

- 1. J. Simmons, K. Dill-McFarland, C. Stein, P. Van, Chihota V., Ntshiqa T., Maenetje P., Peterson G., Benchek P., Velen K., Fielding K., Grant A., Gottard R., Mayanja-Kizza H., Wallis R., Churchyard G., Boom W.H., and Hawn T.R. Monocyte transcriptional responses to Mycobacterium tuberculosis associate with clinical resistance to TST/IGRA conversion in Uganda and South Africa. mSphere, 2022. in press
- G. Moncunill, J. Carnes, C. Young, L. Carpp, S. de Rosa, J. Campo, A. Nhabomba, M. Mpina, C. Jairoce, G. Finak, P. Haas, C. Murie, P. Van, H. Sanz, S. Dutta, Mordmuller B., S. Agnandji, N. Diez-Padrisa, N. Williams, J. Aponte, C. Valim, D. Neafsey, C. Daudenberger, J. McElrath, C. Dobano, K. Stuart, and R. Gottardo. Transcriptional correlates of malaria in rts,s/as01-vaccinated african children: A matched case-control study. eLife, 2021. in press

- 3. J. Simmons, P. Van, C. Stein, Chihota V., Velen K., Fielding K., Grant A., Ntshiqa T., Peterson G., Graustein A., Nguyen F., Gottard R., Mayanja-Kizza H., Wallis R., Churchyard G., Boom W.H., and Hawn T.R. Monocyte metabolic transcriptional programs associate with resistance to tuberculin skin test/interferon- release assay conversion. *Journal of Clinical Investigations*, June 2021
- 4. J. S. Minden, F. Lanni, and P. T. Van. USPTO US10362237: Structured illumination system for increased dynamic range in quantitative imaging, July 2019
- 5. P. Van, W. Jiang, R. Gottardo, and G. Finak. ggCyto: next generation open-source visualization software for cytometry. *Bioinformatics*, 34(22):3951–3953, 11 2018
- 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
- 7. 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
- 8. U. K. Iheagwara, P. L. Beatty, P. T. Van, T. M. Ross, J. S. Minden, and O. J. Finn. Influenza virus infection elicits protective antibodies and T cells specific for host cell antigens also expressed as tumor-associated antigens: a new view of cancer immunosurveillance. *Cancer Immunology Research*, 2(3):263–273, Mar 2014
- 9. A. Kaur, P. T. Van, C. R. Busch, C. K. Robinson, M. Pan, W. L. Pang, D. J. Reiss, J. DiRuggiero, and N. S. Baliga. Coordination of frontline defense mechanisms under severe oxidative stress. *Molecular Systems Biology*, 6:393, Jul 2010
- T. Koide, D. J. Reiss, J. C. Bare, W. L. Pang, M. T. Facciotti, A. K. Schmid, M. Pan, B. Marzolf,
  P. T. Van, F. Y. Lo, A. Pratap, E. W. Deutsch, A. Peterson, D. Martin, and N. S. Baliga.
  Prevalence of transcription promoters within archaeal operons and coding sequences. *Molecular Systems Biology*, 5:285, 2009
- P. T. Van, A. K. Schmid, N. L. King, A. Kaur, M. Pan, K. Whitehead, T. Koide, M. T. Facciotti, Y. A. Goo, E. W. Deutsch, D. J. Reiss, P. Mallick, and N. S. Baliga. Halobacterium salinarum NRC-1 PeptideAtlas: toward strategies for targeted proteomics and improved proteome coverage. *Journal of Proteome Research*, 7(9):3755-3764, Sep 2008
- 12. A. K. Schmid, D. J. Reiss, A. Kaur, M. Pan, N. King, P. T. Van, L. Hohmann, D. B. Martin, and N. S. Baliga. The anatomy of microbial cell state transitions in response to oxygen. *Genome Research*, 17(10):1399–1413, Oct 2007
- 13. 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