Carlos Vigil Vásquez

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EXPERIENCE

Research Assistant - Molecular Design Lab

Hybrid

Pontificia Universidad Catolica de Chile, Santiago, Chile

July 2017 - December 2022

- Proposed and implemented a novel predictive model called SimSpread that combines network-based inference derived from social-studies with run-of-the-mill chemical similarity measures which resulted in a drug discovery predictive model that exceeded the predictive performance of state-of-the-art network-based inference predictive models.
- o Implemented and validated a drug-discovery pipeline based in the SimSpread predictive model that resulted in the identification of N novel compounds with antifungal activity for M clinically relevant fungi organism.

Research Assistant - Psychophysiology Lab

Remote

Pontificia Universidad Católica de Chile, Chile

January 2022 - December 2022

o Implemented a machine-learning predictive and statistical modelling pipeline of human psychological states that resulted in the identification of the effect of different contemplative practices into the subject well-being as measured by 7 distinct well-being metrics.

Research Assistant - REFRACT

Hybrid

Universidad Pablo de Olavide, Sevilla, Spain

September 2022 - December 2023

• Implemented a bioinformatic protocol for the search of protein homology using a three-step "search, detect, and enrich" model using a combination of structural and sequence aligners working in tandem to filter and enrich conservation signals that helped shed light on the homologous nature of repeat proteins related to the membrane and nuclear pore organization.

EDUCATION

B.S. Biochemistry

Santiago, Chile

2015 - 2022

- Pontificia Universidad Católica de Chile
 - o Licenciature grade 5.6/7.0 (3.1/4.0 in GPA scale)
- o Graduated with 2 of 3 votes of distinction
- Research thesis grade 7.0/7.0 (4.0/4.0 in GPA scale)

Publications

- C. Vigil-Vásquez, A. Schüller; De novo prediction of drug targets and candidates by chemical similarity-guided network-based inference. IJMS (2022). DOI:10.3390/ijms23179666
- C. Vigil-Vásquez, M. Jimenez-Socha, P. Ortiz-Bermudez, A. Schüller; Antifungal drug discovery by chemical similarity-guided network-based inference In preparation
- Psychophysiology Lab paper In preparation
- REFRACT paper In preparation

Presentations

- Antifungal drug discovery by chemical similarity-guided network-based inference: Sociedad Chilena de Bioinformática (January, 2022)
- DDTNBI: de novo target prediction using a social network-derived method: ISMB/ECCB (August, 2021)
- A computational chemogenomics method for the prediction of off-target interactions with coagulation factor Xa: EHA (August, 2020)
- Limits and potential of in silico target prediction by chemical similarity: ISCB-LA (October, 2018)

SKILLS

- Human languages: Spanish (native), English (TOEFL score Tools: Git, GitHub, MySQL, SQLite 101 of 120)
- Machine languages: Python, Julia, LaTeX, Bash, SQL
- Frameworks: Graphs.jl, CUDA.jl, Plots.jl, Scikit-Learn, Pandas, NumPy, Matplotlib, Seaborn, NetworkX, Pingouin
- Platforms: Linux, MacOS
- Practical experience: Recommender systems, graph theory, statistics, data visualization, REST API

Awards

- Undergraduate Research Contest Winter 2017: Project title: "In silico prediction and prioritization of novel drug targets."
- Undergraduate Research Contest Summer 2020: Project title: "Use of biochemical networks for the prediction of novel drugs for coagulation factor Xa"