# Hector Roux de Bézieux

2615 California Avenue - Berkeley, CA 94703  $\bigcirc$  +1 (510) 345-9460 •  $\bigcirc$  +33 (0)679215150 

# Education

## Ph.D in Biostatistics

University of California, Berkeley, USA

Dissertation adviser: Sandrine Dudoit, GPA 4.0

2017 - 2021 (expected)

Degree Emphasis in Computational Biology

- Conceived and implemented machine-learning methods and software for genomic data analyses. Publications include:
- 1. Roux de Bézieux, H., Van den Berge, K., Street, K. Dudoit, S. (2021). Trajectory inference across multiple conditions with condiments: differential topology, progression, differentiation, and expression. BioRxiv
- 2. Van den Berge, K., Roux de Bézieux, H., ... Dudoit, S., Clement, L. (2020). Trajectory-based differential expression analysis for single-cell sequencing data. Nature Communications, 11(1), 1201.
- Collaborated with applied biology teams to design experiments and analyze genomic data, including on.
- 1. Brann, D. H., ... Roux de Bézieux, H.,... (2020). Non-neuronal expression of SARS-CoV-2 entry genes in the olfaory system suggests mechanisms underlying COVID-19-associated anosmia. Science Advances, eabc5801.
- 2. Yao, Z., Liu, ... Roux de Bézieux, H.,... (2020). An integrated transcriptomic and epigenomic atlas of mouse primary motor cortex cell types. BioRxiv

Master of Science in Biology & Bioengineering

École polytechnique and UC Berkeley

Fung Institute Fellowship, Outstanding Investment Award, GPA 3.86

2015 - 2017

 Designed and conducted experiments for bacteriophage discovery in the context of antibiotic-resistant bacteria in a year-long project with SmartPhage, a UC Berkeley-based start-up.

**Bachelor of Science in Mathematics & Physics** 

**École polytechnique, France** 

Received with rank 1/4,500 at the national entry exam, GPA 3.94

2011-2015

# **Experience**

# **Pendulum Therapeutics** Computational Biologist

San Francisco, USA

June 2018 - Ongoing

• Analyzed clinical trial data and designed future clinical trial: Roux de Bézieux, H.,..., Perraudeau, F (2021) A Pilot Medical Food Study using a Smartphone Application with Continuous Glucose Monitoring Sensors JMIR Form Res

Developed computational methods for the analysis of metagenomic datasets: Roux de Bézieux, H.... Jacob, L. (2020). Finding All Significant Closed Connected Subgraphs. Machine Learning in Computational Biology (MLCB)

## Institut Jacques Monod, Cellular Adhesion and Mechanism Lab

Paris, France

Research Intern, Benoit Ladoux and René-Marc Mège's team

March-July 2016

• Designed and conducted in vivo experiments on protein-protein interactions.

#### PSA Peugeot Citroën

Shanghaï, China

Intern, Strategic Partnership Department

June-August 2015

#### French Army, Elite Mountain Troop

France

2<sup>nd</sup> Lieutenant

September 2013- April 2014

• Led a group of 8 professional soldiers during a NATO exercise in great Northern Norway.

• **Programming**: Python, Bash (Intermediate), Latex (Expert)

Author of several open-source R packages, with over 5,000 downloads since 2020 on Bioconductor

• Language: English (fluent), French (native), German (intermediate)

First Aid Certification by the Red Cross: 2011, 2015 Rowing: 6 years at national level

**Triathlon:** 1000 miles annually, Ironman 70.3 (Santa Cruz, 2019)

# **Publications**

- Roux de Bézieux, H., Van den Berge, K., Street, K. Dudoit, S. (2021). Trajectory inference across multiple conditions with condiments: differential topology, progression, differentiation, and expression. *BioRxiv*
- Roux de Bézieux, H.,..., Perraudeau, F (2021) A Pilot Medical Food Study using a Smartphone Application with Continuous Glucose Monitoring Sensors. *JMIR Form Res*
- Van den Berge, K., Chou, H. J., **Roux de Bézieux, H.**,..., Dudoit, S. (2021). Normalization benchmark of ATAC-seq datasets shows the importance of accounting for GC-content effects. *bioRxiv*.
- Roux de Bézieux, H.,... Jacob, L. (2020). Finding All Significant Closed Connected Subgraphs. *Machine Learning in Computational Biology (MLCB)*
- Brann, D. H., ... Roux de Bézieux, H.,... (2020). Non-neuronal expression of SARS-CoV-2 entry genes in the olfaory system suggests mechanisms underlying COVID-19-associated anosmia. *Science Advances, eabc5801*.
- Adkins, R. S., ... Roux de Bézieux, H.,... (2020). A multimodal cell census and atlas of the mammalian primary motor cortex. *BioRxiv*.
- Roux de Bézieux, H., Street, K. . . . Dudoit, S. (2020). Improving replicability in single-cell RNA-Seq cell type discovery with Dune. *BioRxiv*
- Yao, Z., Liu, ... Roux de Bézieux, H.,... (2020). An integrated transcriptomic and epigenomic atlas of mouse primary motor cortex cell types. *BioRxiv*
- Van den Berge, K., **Roux de Bézieux**, **H**., ... Dudoit, S., Clement, L. (2020). Trajectory-based differential expression analysis for single-cell sequencing data. *Nature Communications*, 11(1), 1201.
- Croizat, G., Kehren, A., **Roux de Bézieux, H**., Barakat, A. (2018). Influence of pulsatile blood flow on allometry of aortic wall shear stress. *arXiv*

# **Talks**

## **European Bioconductor Meeting 2020**

**Europe (Virtual)** 

90mn workshop

December 2020

Trajectory inference across conditions: differential expression and differential progression.

BioC 2020

USA (Virtual)

60mn workshop

July 2020

Trajectory inference across conditions: differential expression and differential progression.

# JBIMS Data Sciences Workshop

Berkeley, California

15mn Talk

February 2020

Bacterial and metagenome GWAS

## **Statistics and Genomics Seminar**

Berkeley, California

1h talk

May 2019

Trajectory inference, Differential Expression and clustering along developmental trajectories in single cell RNA-Sequencing.

#### **Berkeley Stem Cell Center Retreat**

Monterey, California

15mn talk

Mav 2019

Unlocking the power of continuity in single cell RNA-Seq - Differential gene Expression along developmental trajectories.

#### FAST-BIG: Workshop on High-dimensional statistics

INRIA, Paris

30mn talk

March 2019

Beyond DBGWAS: exploring assembly graphs in an efficient manner