# Rodrigo Dorantes-Gilardi

#### ⊠ rodgdor@gmail.com

Telcel Born: April 24 1990

Mexico City Nationality: Mexican/Spanish

Web: https://rodogi.github.io GitHub: https://github.com/rodogi

#### Work

June 2018– | Data Scientist at Telcel Present |

#### Education

April 24 | PhD Defense "Bio-mathematical aspects of the plasticity of protein folding" 2018

Thesis committee:

Frédéric Cazals (INRIA),

Lashuel Hilal (École Polytechnique Fédérale de Lausanne), Kavé Salamatian (Université de Savoie Mont-Blanc),

2014–2018 | PhD Applied Mathematics

Under supervision of Claire Lesieur and Laurent Vuillon, IXXI Complex Systems Institute, ENS-Lyon, France. LAMA Mathematics Laboratory, Le Bourget-du-Lac, France. "Bio-mathematical aspects of the plasticity of protein folding"

2014 Master thesis supervised by Gelasio Salazar,

Universidad Autónoma de San Luis Potosí, Mexico

"An algorithm in python on the minium crossing number of complete graphs".

2012–2014 Master degree in applied mathematics,

Universidad Autónoma de San Luis Potosí, Mexico.

2008–2012 Bachelor degree in Economics,

Université de Toulouse Capitole, France.

2005–2008 High School diploma, option: General Sciences,

Instituto Tecnológico de Estudios Superiores de Monterrey, Mexico.

#### **Publications**

#### Articles

2018 | R. Dorantes-Gilardi, L Bourgeat, L Vuillon, and C Lesieur

(October) In proteins, the structural responses of a position to mutation rely on the Goldilocks

principle: not too many links, not too few.

Phys. Chem. Chem. Phys., 20, 25399 (2018)

2016 M. Achoch, R. Dorantes-Gilardi, C. Wymant, G. Feverati, K. Salamatian, L.

Vuillon, and C. Lesieur

(December) | Protein structural robustness to mutations: an in silico investigation.

Phys. Chem. Chem. Phys., 18, 13770 (2016)

#### Chapters

2018 | C. Lesieur, R. Dorantes-Gilardi, and L. Vuillon

(July) Chapter: Induced Graphs: surfing on protein structures.

Book: Allostery and protein dynamics: from physical chemistry to drug discovery

To appear in Springer Methods in Molecular Biology

#### **Proceedings**

2017 | R. Dorantes-Gilardi, L. Vuillon, and C.Lesieur

(December) Perturbation of amino acid networks: A statistical study of the defects introduced in proteins by mutations.

The 6th International Conference on Complex Networks and Their Applications

#### **Talks**

2016 Workshop on Mechanisms underlying local to global signals in networks

(May) Amino-acid network as a model of the protein's structure, an in silico investigation. IXXI, École Normale Supérieure de Lyon, France.

2016 Workshop on Advanced mathematics for network analysis

(March) Amino-acid networks used to capture protein structural changes caused by mutations.

Luchon, France.

2015 Workshop on protein fibers: from pathology to nanomaterial

(June) | Protein Graphs.

École Normale Supérieure de Lyon, France.

2015 | IXXI Seminar

(April) Tentative to relate functional and structural changes in protein, caused by muta-

tions (perturbations) using amino-acid networks.

École Normale Supérieure de Lyon, France.

2014 Theoretical Approaches for the Genome and the proteome

(December) What impact to expect on a whole protein from geometrical changes produced by local amino acid side chain perturbation (in silico amino acid mutation): resilience

and innovation.

Bourget-du-Lac, France.

#### Posters

2017 The 6th International Conference on Complex Networks and Their Applications (December)

 Perturbation of amino acid networks: A statistical study of the defects introduced in proteins by mutations.
 Lyon, France.

 2015 (May) Statistics On Protein Graphs.

 Grenoble, France.

#### Conferences and Schools

2016 Spring school of theoretical informatics. CIRM, Marseille, France. (April) 2016 School for young researchers in mathematical informatics. (March) IMJ-PRG, Paris, France. 2015 School Algorithms and Heuristics for Large-scale Data Sets. École Normale Supérieure de Lyon, Lyon, France. (January) 2014 Lyon systems biology. École Normale Supérieure de Lyon, Lyon, France. (November)

## Organization

2014 Theoretical Approaches for the Genome and the proteome (December) Bourget-du-Lac, France.

#### Skills

#### **Programing**

Python Advanced (pandas, numpy, sklearn, pyspark, matplotlib, networkx, biopython) R Intermediate (data.table) Tex Advanced Elisp Intermediate (Emacs)  $\mathbf{C}$ basic SQLIntermediate (Oracle and Netezza) Git Intermediate Linux/GNU Advanced (Bash) HTML, CSS Intermediate

#### **Bioinformatics**

Pymol Intermediate
YASARA Intermediate
FoldX Intermediate
Bio Protein Data Bank: query (requests module in python), cleanse, analyze (numpy and pandas)
Bio-Structure
Bio-Space Computational Algorithms using Delaunay triangulations and Convex hulls (Scipy spatial module)

## Open Source Software

I have contributed to the following projects:

- networkx (https://networkx.github.io/)
- biopython (http://biopython.org/)

### Languages

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Spanish Reading, Writing, Speaking: Native language.
English Reading, Writing, Speaking: Fluent.
French Reading, Writing, Speaking: Fluent.
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#### **Recommendation Contacts**

- Claire Lesieur Laboratoire Ampère, Université Claude Bernard Lyon 1/ IXXI, ENS-Lyon 

   ⊠ claire.lesieur@ens-lyon.fr
- Laurent Vuillon Laboratoire de Mathématiques, Université de Savoie Mont-Blanc ⊠ laurent.vuillon@gmail.com
- Gelasio Salazar Instituto de Física, Universidad Autónoma de San Luis Potosí 

  ⊠ gelasio.salazar@gmail.com