# Jaime Rodríguez-Guerra

■ jaime.rogue@gmail.com ・ ② rjai.me ・ У @jaime\_rgp 



#### Skills

**Programming** Python, JavaScript, LaTeX, PHP **Web development** Flask, Hugo, React, Joomla **Code maintenance** GitHub, Readthedocs, Travis, AppVeyor **Deployment** conda, constructor, pip, npm Vultr, Heroku, Netlify, GitHub Pages

#### Scientific software

Molecular Dynamics OpenMM, MDTraj, AmberTools, ParmEd **Quantum Mechanics** Gaussian, NWChem Homology Modelling Modeller

**Cheminformatics** RDKit, ase, openbabel

**Docking** GOLD, AutoDock Vina

**Visualization** UCSF Chimera, VMD, PyMol, NGLView

#### **Education & Positions**

#### Research assistant in Prof. Maseras' group

INSTITUTE OF CHEMICAL RESEARCH OF CATALONIA (ICIQ)

Tarragona, Spain Apr - Oct 2018

Barcelona, Spain

2015 - 2018

#### PhD in Biotechnology

AUTONOMOUS UNIVERSITY OF BARCELONA (UAB)

- Government of Catalonia FI 3-year predoctoral fellowship
- Supervisor: Jean-Didier Maréchal
- Committee: Gerald Monard, Sílvia Osuna, Xavier Solans

**MSc in Bioinformatics** Barcelona, Spain

AUTONOMOUS UNIVERSITY OF BARCELONA 2013 - 2014

**BSc in Biotechnology** Salamanca, Spain UNIVERSITY OF SALAMANCA 2008 - 2013

## **Highlighted software projects**

#### **GaudiMM**

Python 2.7, UCSF Chimera, deap and more

- Multi-objective genetic algorithm optimization platform for 3D molecular sketching
- · Modular architecture on top of UCSF Chimera to support 3rd party libraries (OpenMM, ProDy, RDKit, NWChem, Vina, IMP)
- Open development supported by GitHub, Readthedocs, and TravisCI

#### **OMMProtocol**

Python 2.7/3.4+, OpenMM, ParmEd, MDTraj, openmoltools

☑ O insilichem/ommprotocol

- Command-line application to create MD protocols with YAML input files
- · Built on top of OpenMM and enhanced with loaders, reporters and integrators found in ParmEd, MDTraj and openmoltools
- · Open development and deployment supported by GitHub, Readthedocs, TravisCI, AppVeyor and conda constructor

Python 2.7/3.4+, cclib, Jinja, Flask; JavaScript

- Web application to generate technical reports from computational chemistry calculations
- Exposes cclib parsers through the Jinja templating engine and serves it with Flask
- · Deployed to Heroku and connected with online platforms (GitHub, FigShare, Zenodo) through JavaScript hooks

Python 2.7/3.4+; Tinker; Gaussian

- Command-line application to extend Gaussian's ONIOM engine with external MM force fields
- Modular design to accept any MM library. Tinker compatibility already available, including polarizable force fields
- · Designed as a replacement for an existing collection of Bash scripts with poor maintainability

#### **Tangram**

Python 2.7, UCSF Chimera and more

- Collection of graphical interfaces for molecular modelling in UCSF Chimera
- Setup external calculations (Gaussian, OpenMM, NCIPlot, ProDy...)
- Visualize results from external tools (PLIP, GOLD, GaudiMM, PoPMuSiC, PropKa...)

#### **Publications**

#### SUBMITTED & UNDER REVIEW

† co-corresponding authorship

## A Multiscale Computational Study of the Cu-catalyzed Hydration of Alkenes in Water Solvent and into the Context of an Artificial Metallohydratase

2018

SUBMITTED

Lur Alonso-Cotchico, Giuseppe Sciortino, <u>Jaime Rodríguez-Guerra Pedregal</u>, Ivana Drienovska, Gerard Roelfes, Agustí Lledós, Jean-Didier Maréchal

#### Computational insight on the interaction of oxaliplatin with insulin

2018

SUBMITTER

Giuseppe Sciortino, Jaime Rodríguez-Guerra Pedregal, José-Emilio Sánchez-Aparicio, Eugenio Garribba, Jean-Didier Maréchal

## Pillar[5]arene glyco(mimetic)rotaxanes for the functional interrogation of multivalency responsive glycosidases

2018

SUBMITTED

Iwona Nierengarten, Manuel González-Cuesta, <u>Jaime Rodríguez-Guerra Pedregal</u>, Uwe Hahn, Sonia Romero-Téllez, Jean-Didier Maréchal, Laura Masgrau, José Manuel Garcia Fernández, Jean-François Nierengarten, Carmen Ortiz Mellet

#### **PUBLISHED & IN PRESS**



#### **GARLEEK: Adding an Extra Flavor to ONIOM**

2018

☑ ♠ insilichem/garleek

<u>Jaime Rodríguez-Guerra Pedregal</u>, Ignacio Funes-Ardoiz, Giuseppe Sciortino, José-Emilio Sánchez-Aparicio, Gregori Ujaque, Agustí Lledós, Jean-Didier Maréchal, Feliu Maseras



## Differential response to Cadmium exposure by expression of a two and a three-domain metallothionein isoform in the land winkle Pomatias elegans: Valuating the marine heritage of a land snail

2018

Sci. Total. Environ. 648, 561-571 ☑ ❷ 10.1016/j.scitotenv.2018.07.426 Lara Schmielau, Martin Dvorak, Michael Niederwanger, Nicole Dobieszewski, Veronika

Pedrini-Martha, Peter Ladurner, <u>Jaime Rodríguez-Guerra Pedregal</u>, Jean-Didier Maréchal, Reinhard Dallinger



## ESIgen: Electronic Supporting Information Generator for Computational Chemistry Publications

2018

2018

2018

J. Chem. Inf. Model. 58 (3), 561-564 ☑ ⑤ 10.1021/acs.jcim.7b00714 ☑ ۞ insilichem/esigen Jaime Rodríguez-Guerra Pedregal, † Pablo Gómez-Orellana, Jean-Didier Maréchal†



#### PyChimera: Use UCSF Chimera modules in any Python 2.7 project

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Bioinf. 34 (10), 1784-1785 ☑ ⑤ 10.1093/bioinformatics/bty021 ☑ ⑤ insi

Jaime Rodríguez-Guerra Pedregal, † Jean-Didier Maréchal†



## Prediction of the interaction of metallic moieties with proteins: An update for protein-ligand docking techniques

J. Comput. Chem. 39 (1), 42-51 🛭 🍪 10.1002/jcc.25080

Giuseppe Sciortino, <u>Jaime Rodríguez-Guerra Pedregal</u>, Agustí Lledós, Eugenio Garribba, Jean-Didier Maréchal



## GaudiMM: A modular multi-objective platform for molecular

**J. Comput. Chem. 38 (24), 2118-2126 1 1** 10.1002/jcc.24847 Jaime Rodríguez-Guerra Pedregal,† Giuseppe Sciortino, Jordi Guasp, Martí Municoy, Jean-Didier



#### Elucidating the 3D structures of Al(III)-Aβ complexes: a template free strategy based on the pre-organization hypothesis

2017

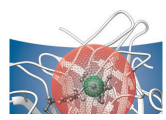
2017

Chem. Sci. 8. 5041-5049 

© 10.1039/C7SC01296A

Jon I. Mujika, Jaime Rodríguez-Guerra Pedregal, Xabier Lopez, Jesus M. Ugalde, Luis Rodríguez-Santiago, Mariona Sodupe, Jean-Didier Maréchal

#### **REVIEWS & BOOK CHAPTERS**

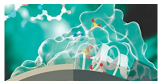


#### **Computational Studies of Artificial Metalloenzymes: From Methods and Models to Design and Optimization**

2018

Ch. 4 in "Artificial Metalloenzymes and MetalloDNAzymes in Catalysis: From Design to Applications", Wiley & Sons 🛛 🚳 10.1002/9783527804085.ch4

Jaime Rodríguez-Guerra Pedregal, Lur Alonso-Cotchico, Giuseppe Sciortino, Agustí Lledós, Jean-Didier Maréchal



#### **Enzyme Design**

2016

Ch. 15 in "Simulating Enzyme Reactivity", Royal Society of Chemistry 🛛 🚭 10.1039/9781782626831-00481 Lur Alonso-Cotchico, Jaime Rodríguez-Guerra Pedregal, Agustí Lledós and Jean-Didier Maréchal

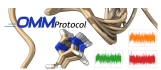


#### **Toward the Computational Design of Artificial Metalloenzymes:** From Protein-Ligand Docking to Multiscale Approaches

2015

Victor Muñoz Robles, Elisabeth Ortega-Carrasco, Lur Alonso-Cotchico, Jaime Rodriguez-Guerra Pedregal, Agustí Lledós, Jean-Didier Maréchal

#### **PRE-PRINTS**



## OMMProtocol: A Command Line Application to Launch Molecular

2018

Dynamics Simulations with OpenMM **ChemRxiv** ☑ **1**0.26434/chemrxiv.7059263

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Jaime Rodríguez-Guerra Pedregal, † Lur Alonso-Cotchico, Lorea Velasco-Carneros, Jean-Didier Maréchal<sup>†</sup>

#### **Presentations**

#### **XXXIV XRQTC Annual Meeting**

THE INSILICHEM MOLECULAR MODELLING SUITE

Barcelona, Spain July 12, 2018

#### X International School on Organometallic Chemistry Marcial Moreno Mañas

MULTICRITERIA OPTIMIZATION OF CHEMOSTRUCTURAL DRAFTS WITH A MODULAR SOFTWARE PLATFORM

Ciudad Real, Spain July 7, 2017

#### **XXXVI RSEQ Biennial Meeting**

A MULTI-OBJECTIVE OPTIMIZATION SOFTWARE FOR STRUCTURAL SKETCHING OF CHEMOBIOLOGICAL COMPOUNDS Awarded "Best Flash Presentation"

Sitges, Spain June 28, 2018

#### 4th COST MC Meeting "Dynamics of Biomacromolecular Machines"

A MODULAR MULTI-OBJECTIVE PLATFORM FOR MOLECULAR MODELING

Bugibba, Malta March 28, 2017

#### IX International School on Organometallic Chemistry Marcial Moreno Mañas

SOLVING COORDINATION GEOMETRIES FOR COMPLEX BIOMETALLIC SYSTEMS

San Sebastián, Spain July 7, 2016

### **Teaching**

#### Modern Python for scientific software development

SGB - Servei de Genòmica i Bioinformàtica, Autonomous University of Barcelona

♦ http://sct.uab.cat/genomica-bioinformatica/en

#### **Practicum: Molecular Dynamics simulations in UCSF Chimera**

MSc Industrial Chemistry and Introduction to Chemical Research, Autonomous University of Barcelona

http://pagines.uab.cat/chemistry-master/

**Introduction to Python** 

MSc Bioinformatics, Autonomous University of Barcelona

http://mscbioinformatics.uab.cat

Barcelona, Spain

2017-present

Barcelona, Spain

2016-present

Barcelona, Spain

2016-present

#### Co-supervised projects

**Lorea Velasco**, SAXS-driven homology modelling refinement

Daniel Viladrich, Energy-corrected interpolation of molecular trajectories

2018 Mercè Alemany, TALAIA: visual dictionary for protein residues

Pablo Orenes, Ligand-binding pathways in GaudiMM 2017

Mireia Bertrán, Inertia tensors for protein-ligand docking 2017

2017 José Emilio Sánchez, Graph-based molecular topology generation

**David Teixé**, Web-based hierarchical molecular visualization 2017

Martí Municoy, Evaluation of metal coordination geometries 2016 **Daniel Soler**, Cationic Dummy Atoms Software 2016

Jordi Guasp, Normal Modes interfacing with ProDy & UCSF Chimera

PhD Biomedicine & Molecular Biology, EHU

BSc Physics, UAB

MSc Bioinformatics, UAB MSc Bioinformatics, UAB

BSc Mathematics, UAB

MSc Bioinformatics, UAB

Double BSc Chemistry + Physics, UAB

Double BSc Chemistry + Physics, UAB

Double BSc Chemistry + Physics, UAB

BSc Mathematics, UAB

#### References

#### Dr. Jean-Didier Maréchal

ASSOCIATE PROFESSOR, DEPARTMENT OF CHEMISTRY, AUTONOMOUS UNIVERSITY OF BARCELONA (UAB)

■ jeandidier.marechal@uab.cat • ♦ http://www.insilichem.com

#### Dr. Feliu Maseras

GROUP LEADER, INSTITUTE OF CHEMICAL RESEARCH OF CATALONIA (ICIQ)

Barcelona, Spain

Tarragona, Spain