# Jaime Rodríguez-Guerra



### 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

### **Positions**

### Postdoctoral researcher in joint project for Prof. Volkamer (Berlin) & Prof. Chodera (NY)

CHARITÉ UNIVERSITY OF MEDICINE & MEMORIAL SLOAN KETTERING CANCER CENTER (MSKCC)

Berlin, Germany Mar 2019 - present

### Postdoctoral researcher in Prof. Maréchal's group

AUTONOMOUS UNIVERSITY OF BARCELONA (UAB)

Barcelona, Spain Dec 2018 - Mar 2019

### Research assistant in Prof. Maseras' group

INSTITUTE OF CHEMICAL RESEARCH OF CATALONIA (ICIQ)

Tarragona, Spain

Apr - Oct 2018

### Education

### PhD in Biotechnology

AUTONOMOUS UNIVERSITY OF BARCELONA (UAB)

Barcelona, Spain

2015 - 2018

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

### **MSc in Bioinformatics**

AUTONOMOUS UNIVERSITY OF BARCELONA

Barcelona, Spain

2013 - 2014

2008 - 2013

### **BSc in Biotechnology**

UNIVERSITY OF SALAMANCA

Salamanca, Spain

## **Highlighted software projects**

### **GaudiMM**

Python 2.7, UCSF Chimera, deap and more

nsilichem/gaudi

- 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

nsilichem/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

### **ESIgen**

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

nsilichem/esigen

- 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

#### **GARLEEK**

Python 2.7/3.4+; Tinker; Gaussian

nsilichem/garleek

- 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

nsilichem/tangram

- 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**

### **PUBLISHED & IN PRESS**

† co-corresponding authorship

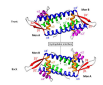


# Discovery of substrate-assisted processivity by an exo-hydrolase with a pocket-shaped catalytic site

2019



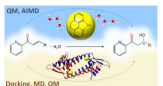
Victor Streltsov, Sukanya Luang, Alys Peisley, Joseph Varghese, James Ketudat Cairns, Sébastien Fort, Marcel Hijnen, Igor Tvaroska, Ana Ardá, Jesús Jiménez-Barbero, Mercedes Alfonso-Prieto, Carme Rovira, Fernanda Mendoza, Laura Tiessler-Sala, José-Emilio Sánchez-Aparicio, Jaime Rodríguez-Guerra, José M. Lluch, Jean-Didier Maréchal, Laura Masgrau, Maria Hrmova



# The effect of cofactor binding on the conformational plasticity of the biological receptors in artificial metalloenzymes: the case study of LmrR

2019

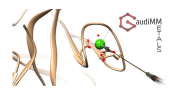
Front. Chem. 7, 211 🚳 10.3389/fchem.2019.00211 Lur Alonso-Cotchico, Jaime Ro<u>dríguez-Guerra</u>, Agustí Lledós, Jean-Didier Maréchal



# Integrated computational study of the Cu-catalyzed hydration of alkenes in water solvent and into the context of an artificial metallohydratase

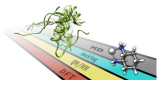
2019

ACS Catal. 9 (5), 4616-4626 © 10.1021/acscatal.8b04919 Lur Alonso-Cotchico, Giuseppe Sciortino, <u>Jaime Rodríguez-Guerra</u>, Ivana Drienovska, Gerard Roelfes, Agustí Lledós, Jean-Didier Maréchal



# Simple coordination geometry descriptors allow to accurately predict metal binding sites in proteins

2019



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

2019

**Metallomics 4** • 10.1039/C8MT00341F

Giuseppe Sciortino, José-Emilio Sánchez-Aparicio, <u>Jaime Rodríguez-Guerra</u>, Eugenio Garribba, Jean-Didier Maréchal



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

2019



### 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

Sci. Total. Environ. 648, 561-571 @ 10.1016/j.scitotenv.2018.07.426

Lara Schmielau, Martin Dvorak, Michael Niederwanger, Nicole Dobieszewski, Veronika Pedrini-Martha, Peter Ladurner, Jaime Rodríguez-Guerra, Jean-Didier Maréchal, Reinhard Dallinger



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

nsilichem/esigen Jaime Rodríguez-Guerra, † Pablo Gómez-Orellana, Jean-Didier Maréchal†



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

nsilichem/pychimera Jaime Rodríguez-Guerra,† Jean-Didier Maréchal†



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

Giuseppe Sciortino, Jaime Rodríguez-Guerra, Agustí Lledós, Eugenio Garribba, Jean-Didier Maréchal



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

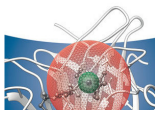
nsilichem/gaudi Jaime Rodríguez-Guerra, † Giuseppe Sciortino, Jordi Guasp, Martí Municoy, Jean-Didier Maréchal†



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

Jon I. Mujika, Jaime Rodríguez-Guerra, 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**

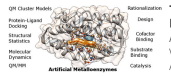
Ch. 4 in "Artificial Metalloenzymes and MetalloDNAzymes in Catalysis: From Design to Applications", 

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



### **Enzyme Design**

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



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

**ACS Catal. 5 (4), 2469-2480** ☑ **REVIEW** ⑤ 10.1021/acscatal.5b00010 Victor Muñoz Robles, Elisabeth Ortega-Carrasco, Lur Alonso-Cotchico, Jaime Rodriguez-Guerra, Agustí Lledós, Jean-Didier Maréchal

### PRE-PRINTS



### **OMMProtocol: A Command Line Application to Launch Molecular Dynamics Simulations with OpenMM**

**ChemRxiv** • 10.26434/chemrxiv.7059263 nsilichem/ommprotocol Jaime Rodríguez-Guerra, † Lur Alonso-Cotchico, Lorea Velasco-Carneros, Jean-Didier Maréchal†

2018

2018

2018

2018

2017

2017

2018

2016

2015

2018

### **Presentations**

**XXXIV XRQTC Annual Meeting** 

THE INSULICHEM 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

Sitges, Spain

June 28, 2017

**XXXVI RSEQ Biennial Meeting** 

A MULTI-OBJECTIVE OPTIMIZATION SOFTWARE FOR STRUCTURAL SKETCHING OF CHEMOBIOLOGICAL COMPOUNDS

Awarded "Best Flash Presentation"

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** 

**Computer-aided Drug Design - Methods and Application** 

SUMMER SEMESTER COURSE CATALOG, FREE UNIVERSITY OF BERLIN

https://www.fu-berlin.de/vv/en/fb

Berlin, Germany

Barcelona, Spain

Modern Python for scientific software development

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

Practicum: Molecular Dynamics simulations in UCSF Chimera

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

2017-2018

MSC INDUSTRIAL CHEMISTRY AND INTRODUCTION TO CHEMICAL RESEARCH, AUTONOMOUS UNIVERSITY OF BARCELONA

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

Barcelona, Spain 2016-2018

Introduction to Python
MSc Bioinformatics, Autonomous University of Barcelona

http://mscbioinformatics.uab.cat

Barcelona, Spain

## **Supervised & co-supervised projects**

2019 B. Adas, KNIME workflows for machine learning pipelines
 2019 M. Wichmann, KNIME workflows for Computer-aided Drug Design
 2018 Lorea Velasco, SAXS-driven homology modelling refinement
 2018 PhD Biomedicine & Molecular Biology, EHU

Daniel Viladrich, Energy-corrected interpolation of molecular trajectories
 Mercè Alemany, TALAIA: visual dictionary for protein residues

2017 **Pablo Orenes**, Ligand-binding pathways in GaudiMM
2017 **Mireia Bertrán**, Inertia tensors for protein-ligand docking

José Emilio Sánchez, Graph-based molecular topology generation
 David Teixé, Web-based hierarchical molecular visualization

2016 Martí Municoy, Evaluation of metal coordination geometries

2016 Daniel Soler, Cationic Dummy Atoms Software
 2016 Jordi Guasp, Normal Modes interfacing with ProDy & UCSF Chimera

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\_

### JProf. Dr. Andrea Volkamer

ASSISTANT PROFESSOR, INSTITUTE OF PHYSIOLOGY, CHARITÉ UNIVERSITY OF MEDICINE

■ andrea.volkamer@charite.de • ⑤ http://www.volkamerlab.org

Berlin, Germany

John D. Chodera, PhD
ASSISTANT MEMBER, COMPUTATIONAL BIOLOGY PROGRAM, MEMORIAL SLOAN-KETTERING CANCER CENTER (MSKCC)

**■** john.chodera@choderalab.org • **②** http://http://www.choderalab.org

Prof. Dr. Jean-Didier Maréchal

Barcelona, Spain

New York, USA

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

**■** jeandidier.marechal@uab.cat • **②** http://www.insilichem.com

Prof. Dr. Feliu Maseras

Tarragona, Spain

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

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