Lluis Artus | Dr.

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Born: 16th of December, 1992

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https://lluisartussuarez.github.io/



Personal Statement

PhD in computational chemistry, specializing in the simulation of molecular structures and properties. I have been studying large chemical systems for the last two. Using folding models, classical molecular dynamics, density functional theory, and microkinetic modeling, I can study the intra- and inter-molecular interactions. That information is essential to identify favorable and detrimental factors that determine the molecule's stability and reaction rates and allow the development of faster and more stable drugs.

Machine learning applied to theoretical chemistry is rising in popularity. Therefore, on 2023, I started training myself in it. I enrolled in IBM's online course on Machine Learning. There, I learned about exploratory data analysis, regression, classification, and unsupervised models, genetic models and deep learning. I have been attending ML conferences applied to computational chemistry and I am reading about state-of-the-art ML techniques. I look forward to implementing what I learned into a practical application.

Professional experience

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Science	
NEC Oncolmmunity AS	Oslo, Norway
Computational scientist Project: "Vaccine development"	2025-present
NCMM Norwegian Centre for Molecular Medicine	Oslo, Norway
Laboratory technician	2024-2024
Project: "Caretaker of the fish facility"	
KTH Royal Institute of Technology	Stockholm, Sweden
Postdoctoral researcher	2022-2024
Project: "Forcefields for electrochemistry in metal-organic frameworks" Project: "Reaction mechanisms of enantioselective hydrogenation of olefines"	
Nordic Consortium for CO ₂ Conversion	Scandinavia
Postdoctoral researcher	2022-2024
Project: "Modelling of CO2 electroreduction in metal-organic frameworks"	
PhD student	2018-2021
Project: "Rational catalyst design for the hydrogenation of CO ₂ derivates"	
Lecturer in	
Workshop in Microkinetic Modeling with COPASI Stockholm, Sweden. April 28-29	2022

Workshop in Microkinetic Modeling with COPASI Tromsø, Norway. March 26-27	2019
Internships	
Iceland University, Iceland Group of Egill Skúlason. Ab-initio molecular dynamics. November	2022
Valladolid University, Spain Group of Martin Jaraiz. Microkinetic modelling. June – July	2018
ICIQ, Spain Group of Feliu Maseras. Modelling of single electron transfers.	2015-2016
GMMF, Spain (Not paid) Group of E. Carolina Sañudo. Synthesis and characterization of magnetic	2014-2015 and fluorescent molecules.
Others	
Frenchie Bistro Waiter	Oslo, Norway Nov. 2024 – Feb. 2025
Frenchie Bistro Waiter	Oslo, Norway Sept. 2021 – Feb. 2022
Skogstad Hotel *** Waiter and housekeeper	Hemsedal, Norway Summer. 2021
Hotel Port-Bó***	Palafrugell, Spain
Receptionist	Summers 2013 – 2015
Hotel & Spa Mas de Torrent ***** Receptionist	Torrent, Spain Summers 2009 – 2010

Education

Academic education	
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Oslo University

Ph.D. Chemistry

2016-2021

Thesis: "Computational study on the deaminative hydrogenation of amides catalyzed by base metal complexes"

Rovira i Virgili University Tarragona, Spain

M.Sc. Chemistry 2015-2016

Thesis: "Computational characterization of the mechanism of the reaction between Fe(I) complexes and aryl halides"

Barcelona UniversityB.Sc. Chemistry

Barcelona, Spain
2010-2015

Thesis: "Synthesis and characterization of coordination compounds with a fluorescent ligand"

Courses and Certifications

IBM's Machine Learning Professional Certificate
Online. IBM course of >134h. February

Machine Learning Capstone 2025

Online. IBM course of >20h. February

Oslo Bioinformatics Workshop Week 2024 2024

Oslo, Norway. December 9-13

Unsupervised Machine Learning Online. IBM course of >23h. June	2024
Supervised Machine Learning: Classification Online. IBM course of >24h. April	2024
Singularity workshop Stockholm, Sweden. February 12	2024
Supervised Machine Learning: Regression Online. IBM course of >20h. February	2024
Exploratory Data Analysis for Machine Learning Online. IBM course of >14h. December	2023
Introduction to GPUs course Stockholm, Sweden. October 12-13	2023
Alpha fold v2.0 and RoseTTAFold Workshop Online. Aug. 31-Sep. 1	2021
8 th MOLCAS Users' Workshop Uppsala, Sweden. Nov. 20-24	2017



Computational chemistry

Drug discovery and material science

Software: AlphaFold, RoseTTA, Amber, Gromacs, CP2K Forcefield parametrization of organometallic systems

Free energy calculations: FEP, EVB, umbrella sampling, metadynamics Properties calculations: binding energies, diffusion coefficients, viscosities...

Reaction mechanism exploration

Software: Gaussian, Jaguar (Schrodinger), COPASI, Acuchem

Rational catalyst design of organometallic catalysts.

Microkinetic modelling for the design and interpretation of experimental kinetic experiments and for the evaluation of theoretical models.

Cheminformatics

Software: RDKit, OpenBabel, molSimplify

Automatic interpretation of chemical databases

Automatic generation of molecular structures for future processing (e.g. machine learning)

Others

Software: NBO6, NCIPLOT, OpenMolcas, Chemcraft, PHI, ioChem-BD

Experimental chemistry.....

Basic training in

Good laboratory practices, NMR, MS, UV-Vis, FT-IR, Spectrofluorometry, HPLC, GC and SQUID.

Informatics.....

I have worked with supercomputers and computer clusters for eight years, at a user level.

I am comfortable with python, and have basic knowledge of C++, Fortran, bash, awk and seq.

The python libraries I use most are os, sys, shutil, pandas, numpy, scipy, matplotlib, seaborn, sklearn, rdkit, molSimplify and OpenBabel.

Most of my programming has been scripting for workflow automation or data analysis. I have recently started studying machine learning. So far, I have completed IBM online courses on "Exploratory Data Analysis" and "Machine Learning regression models".

Publications

Seven articles published in scientific journals. H index of five. Cited more than a hundred times. Five first-authorships.