



Ernesto García Alfonso

PhD student



November 14th, 1996



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[https://github.com/
ErnestoGarciaALfonso](https://github.com/ErnestoGarciaALfonso)



[https://www.
researchgate.net/profile/
Ernesto-Garcia-Alfonso-2](https://www.researchgate.net/profile/Ernesto-Garcia-Alfonso-2)

Skills

Spanish (mother tongue)

English (B2)

French (B1)

About me

During my free time, I like reading mystery and fiction books and practising some sports such as football, volleyball and biking. I like to spend time with my family and friends. I enjoy team work.



Education



PhD: October 1st 2021-current

Title: Dynamics of pure and doped superfluid Helium droplets using ⁴He-TDDFT simulations.

Lab: Laboratoire Collisions Agrégats Réactivité (LCAR)

Thesis advisor: Nadine Halberstadt

■ Superfluid ⁴He_N nanodroplets

□ Helium Time Dependent Density Functional Theory (⁴He TD-DFT)

■ Clusterization of foreign atoms within ⁴He nanodroplets.

□ Quantum Vortices

■ Coalescence of ⁴He_N nanodroplets

□ Coulomb explosion of Ak₂ on ⁴He_N nanodroplets

During my PhD, I have developed and/or improved some skills such as teamwork, Fortran 90, C++ and Python programming as language, handling GIT-HUB repositories, and Unix system. I have delivered a certain amount of speeches both international and national. I have written papers which they have been very useful when writing in Latex code (scientific writing program)



2015–2020

Master in physics. Havana University, Physics Faculty. Title: "Study of the Vibrational Predissociation of the NeBr₂ Complex by Computational Simulation Using the Trajectory Surface Hopping Method".

Thesis advisor: Maykel Marquez Mijares and Jesús Rubayo Soneira

- Van der Waals complexes
- Quasiclassic Method
- Trajectory Surface Hopping (TSH)
- C++ language (I built my own package for calculating TSH)
- Wolfram Mathematics

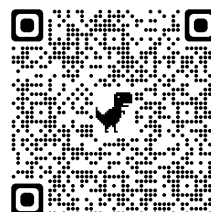


Conferences

Participation in national/international Conferences

[Poster (4/5) and Oral Presentation (5/8)]

[https://github.com/ErnestoGarciaALfonso/
Experience/blob/main/Professional_experience.pdf](https://github.com/ErnestoGarciaALfonso/Experience/blob/main/Professional_experience.pdf)



Publications

In international journals with peer review (8) [https://github.com/ErnestoGarciaALfonso/
List-of-Papers/blob/main/
Papers.pdf](https://github.com/ErnestoGarciaALfonso/List-of-Papers/blob/main/Papers.pdf)



Stay Abroad

★[07/2019–10/2019] Participation. Project "Theoretical study of helium nanodroplets dynamics: alkali dopants and quantum Vortex; rare gas dopants and cluster formation" in Toulouse, France. Participants: Nadine Halberstad, Manuel Barranco and Martí Pi from University of Barcelona.

★[04/03-27/2024] "Photodissociation of Ak₂ induced by the helium environment. A DIM (Diatomics in Molecules) model to describe the interactions between the dialkali and the ⁴He_N nanodroplet and the couplings between the electronics states of the dialkali molecule induced by helium". Consejo Superior de Investigaciones Científicas(CSIC), Madrid Spain.



Membership

2023 Confined Molecular Systems: From a new Generation of Materials to the Starts (COSY) Work Group 4

Publications

- 2024 Ernesto García-Alfonso, Manuel Barranco, Nadine Halberstadt and Martí Pi. "*Time-resolved solvation of alkali ions in superfluid helium nanodroplets*". *J. Chem. Phys.* 160, 164308 (2024)
- 2023 Loidel Puentes-Milián, Ernesto García-Alfonso, Maykel MARQUEZ-MIJARES, and Jesús Rubayo-Soneira. "*NeI2 Photofragmentation Dynamics Through Quasiclassical and Semiclassical Studies*". *ChemPhysChem* e202300406, 2023
- Ernesto García-Alfonso, Francesco Ancilotto, Manuel Barranco, Martí Pi, Nadine Halberstadt. "*Quantized vortex nucleation in collisions of superfluid nanoscopic helium droplets at zero temperature*". *J. Chem. Phys.* 159, 074305 (2023)
- 2022 Ernesto García-Alfonso, Maykel Márquez-Mijares, Jesús Rubayo-Soneira, Nadine Halberstadt, Kenneth C. Janda and Craig C. Martens. "*Photofragmentation dynamics study of ArBr₂ ($\nu = 16 \dots, 25$) using two theoretical methods: trajectory surface hopping and quasiclassical trajectories*". *Eur. Phys. J. D.* 76, 79. 2022.
- Ernesto García-Alfonso, Manuel Barranco, David A. Bonhommeau, Nadine Halberstadt, Martí Pi and Florent Calvo. "*Clustering, collision, and relaxation dynamics in pure and doped helium nanoclusters: Density - vs particle -based approaches*". *The Journal of Chemical Physics.* 157, 014106. 2022.
- 2020 Ernesto García-Alfonso, Francois Coppens, Manuel Barranco, Martí Pi, Frank Stienkemeier and Nadine Halberstadt. "*Alkali atoms attached to vortex-hosting helium nanodroplets*". *The Journal of Chemical Physics.* 152, 194109. 2020.
- Ernesto García-Alfonso, Maykel Márquez-Mijares, Jesús Rubayo-Soneira, Nadine Halberstadt, Kenneth C. Janda and Craig C. Martens. "*Study of the Vibrational Predissociation of the NeBr₂ Complex by Computational Simulation using the Trajectory Surface Hopping Method*". *Sigma Mathematics.* 8, 2029. 2020.
- 2019 C. Calvo-Mola, S. López Pérez, E. García Alfonso, and J. Cerutti-Torres. "*Determination of the Planck Constants through the use of LEDs*". *Revista Cubana de Física*, 36, 125. 2019.



- 01/03-09/2024 **Poster** Molecular and Ionic Clusters. Gordon Research Conference (GRC). Ventura (California). United States. (Due to U.S policy aim to Cuban people I was unable to get a visa,so Nadine Halberstadt presented my poster on site)
- 07/03-07/2023 **Oral Presentation+Poster** Congrès Général des 150 ans de la Société Française de Physique. Paris,France
- 06/20-21/2023 **Poster** AtmChemClimate workshop in Montpellier. France
- 06/01-02/2023 **Oral Presentation** XVèmes Journées de l'ED. Toulouse
- 03/01-03/2023 **Oral Presentation** First COSY General Meeting, Cadiz,Spain.
- 01/24/2023 **Oral Presentation** First meeting of the CONFINED MOLECULAR SYSTEMS: FROM A NEW GENERATION OF MATERIALS TO THE STARS (COSY), COSY Working Group 4 (Helium nanodroplets in science and engineering) (online)
- 11/11/2022 **Poster** (online) PHOTODYNAMICS 2022, XI International Meeting of Photodynamics, La Havana,Cuba
- 11/7-8/2022 **Oral Presentation** Les Toulousaines du Calcul Atomique et Moléculaire (TouCAM), Toulouse, France
- 10/19/2022 **Oral Presentation** at Instituto de Física Fundamental (IFF-CSIC) Departamento de Procesos Atómicos, Moleculares y en Superficies Consejo Superior de Investigaciones Científicas, Madrid Spain. *"Coulomb Explosion of Alkali Dimers on Helium Droplets. Is Really Triplet State Dominant?"*
- 09/05-09/2022 **Participation in training school** "Machine Learning and Quantum Computing for Quantum Molecular Dynamics" (CECAM-FR-MOSER). Gustave Eiffel Université, (Paris), France.
- 08/7-12/2022 **Poster** Molecular and Ionic Clusters. Gordon Research Conference (GRC) , Lucca (Barga), Italy.
- 08/6-7/2022 **Oral Presentation** Molecular and Ionic Clusters. Gordon Research Seminar (GRS) , Lucca (Barga), Italy.
- 04/18-23/2022 **Oral Presentation + Poster** The 14th International Conference on Quantum Fluid Clusters, Erice, Sicily, Italy.
- 03/10-11/2022 **Poster** NanoX-FerMI days. Toulouse, France.
- 11/2021 **Oral Presentation** Réunion annuelle du GDR THEMES. Toulouse, France.
- 06/2021 **Oral Presentation** (online). XV Taller de Física de la Materia Condensada y Molecular. Cuernavaca, Morelos, Mexico.
- 10/2020–08/2021 I taught Atomic and Molecular Physics online (on-site classes were forbidden because of COVID-19) to two groups of 3rd year (20 students each)
- 03/2020 **Poster**. XV Simposio y XIII Congreso de la Sociedad Cubana de Física. La Habana, Cuba.
- 03/11-15/2019 **Oral Presentation** VIII Taller Iberoamericano de Enseñanza de la Física Universitaria. ("Determination of the Planck Constants through the use of LEDs") La Havana, Cuba
- 2018 Presentation Scientific day ICIMAF ("Study of the Vibrational Predissociation of the NeBr₂ Complex by Computational Simulation using the Trajectory Surface Hopping Method") Cuba
- 2018–2021 **Participation** . Research Project (National Basic Sciences Program). *"Study of confinement, relaxation and energy transfer processes of atomic and molecular systems"*. Department of Atomic and Molecular Physics. Higher Institute of Technologies and Applied Sciences (InSTEC), University of Havana. Cuba
- 2016–2017 **Participation**. InSTEC Research Project (Financier: InSTEC). Physico-chemical study of atomic and molecular systems with interest for the environment. Department of Atomic and Molecular Physics. Higher Institute of Technologies and Applied Sciences (InSTEC). Cuba