## SHORT CURRICULUM VITAE

# Dr. Sophie Sacquin-Mora

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### **Research Interests**

After an initial training and PhD in Chemical Physics, I joined the *Laboratoire de Biochimie Théorique* (LBT) in october 2006. The LBT is a CNRS unit where the structure, mechanics, dynamics and interactions of biological macromolecules are studied by developing and applying algorithms for molecular simulations.

Here, I started developing modeling tools to investigate proteins biological function. Proteins are a central feature of the cellular machinery and constitue an increasingly important target for drug design. My goal is to develop new simulation approaches that can help build a bridge between the available structural data for proteins and their activity in the cell. A long term perspective of my fundamental research work is the development of new drugs that can specifically target proteins involved in a given disease.

My projects are mainly focused on coarse-grain models for investigating protein mechanics (which is tightly related to their biological activity) and protein interactions. These coarse-grain simulations are usually coupled with more classic allatom Molecular Dynamics simulations, bioinformatics or experimental approaches in order to obtain complementary information regarding protein function on the atomic level.

## **Permanent Research positions**

### Oct. 2010-

Senior research scientist CNRS (section13)

Laboratoire de Biochimie Théorique, UPR9080 (IBPC, Paris).

#### Oct. 2006-Oct. 2010

Junior research scientist CNRS (section13)

Laboratoire de Biochimie Théorique, UPR9080 (IBPC, Paris).

## **Post-Doctoral positions**

#### Sept. 2005-Sept. 2006

INSERM postdoc, DECRYPTHON program, advisor: Pr. Alessandra Carbone

Analytical Genomics team, INSERM U511, Paris, France

### Nov. 2004-August 2005

Research and Teaching assistant (ATER), advisor: Pr. Daniel Borgis

Physics Department, Université d'Évry Val d'Essonne, France

#### Nov. 2003-Oct. 2004

CNRS Postdoc, advisor: Richard Lavery

Laboratoire de Biochimie Théorique, CNRS UPR9080, Paris, France

#### **Education**

### Dec. 2011 Habilitation à Diriger les Recherches (HDR)

Université Paris 7-Denis Diderot, France

#### Sept. 2000-Oct.2003 Joint french-german PhD in Physical Chemistry

Fluides Nanoconfinés dans des Systèmes de Basse Symétrie : Simulations et Théorie supervised by Pr. Alain Fuchs (Laboratoire de Chimie Physique, CNRS UMR8000, Orsay) and Pr. Martin Schoen (Stranski Lab. für Physikalische und Theoretische Chemie, TU Berlin)

### 2000 Agrégation de Sciences Physiques option Chimie.

#### 1999 M. Sc. research in Molecular Physical-Chemistry

Université Paris XI, Orsay, France

### Prizes, Awards and Fellowships

1997-2001 Scholarship, École Normale Supérieure, Paris, France

**2006** Postdoctoral grant from the Keystone Symposia

Multi-Protein Complexes Involved in Cell Regulation

## Major research grants (2013-2019)

**2017-2021 ANR ENZYMOR (496k€)** *Orienting enzymes on electrochemical surfaces* 

**2012-2019 Labex DYNAMO** (**760k**€) *Energy transduction in membranes* 

**2012-2017** ANR Investissements d'avenir Bio-informatique MAPPING (180k€) Protein interactions

**2010-2014 ANR BIOPAC** (127k€) *Biocatalytic enzymes for biofuel cells* 

### **Institutional responsibilities**

### **January 2018** Head of the French Network for Theoretical Chemistry (RFCT, GDR3333)

**2014-2017** Secretary for the French Network for Theoretical Chemistry (RFCT, GDR3333)

**2018** Recruitment committee for an Associate Professor position in UTC, Compiegne

**2016** Recruitment committee for an Associate Professor position in Université Paris-sud

2012-2016 Elected member of the Comité National pour la Recherche Scientifique (CoNRS),

section 13, Physical-Chemistry, (http://www.cnrs.fr/comitenational/english/UK\_acc.htm)

**Member of Ph.D. Committees** 2013-2019: Member of 9 Ph. D committees and 2 HDR committees.

#### **Reviewing grant proposals**

2019 Isite-NExT, Investissement d'Avenir

2018 Programma per Giovanni Ricercatori-Rota Levi Montalcini

2016 AIC call INRA, 2014 Émergence, Sorbonne Université

**Editorial board** Frontiers in Molecular Biosciences

## Selection of recent peer reviewed publications (10 out of 38 in total)

- 1. *Conformational stability adaptation of a double-stranded RNA binding domain to transfer RNA ligand* C. Bou-Nader, P. Barraud, L. Pecqueur, C. Velours, M. Fontecave, C. Tisné, S. Sacquin-Mora, and D. Hamdane\*, Biochemistry, **58**, 2463-2473 (2019)
- 2. Mechanical changes in proteins undergoing large-scale motions highlight the formation of structural locks S. Sacquin-Mora\*, J. Structural Biology, **203**, 195-204 (2018), available on BioRxiv, doi: https://doi.org/10.1101/221077.
- 3. *Hidden partners : Predicting binding sites for proteins with multiple interaction partners*N. Lagarde, A. Carbone and S. Sacquin-Mora\*, Proteins, **86**, 723-737 (2018), available on BioRxiv,

- doi: https://doi.org/10.1101/244913
- 4. Mobility and core-protein binding patterns of disordered C-terminal tails in  $\beta$ -tubulin isotypes Y. Laurin, J. Eyer, C. Robert, C. Prévost and S. Sacquin-Mora\*, Biochemistry, **56**, 1746-1756 (2017)
- 5. Bridging enzymatic structure and function via mechanics: A coarse-grain approach
  S. Sacquin-Mora\*, G.A. Voth, editor, Methods in Enzymology: Computational Approaches for Studying Enzyme Mechanism Part B,578, 227-248 (2016)
- 6. *Great interactions: How binding incorrect partners can teach us about protein recognition and function* L. Vamparys, B. Laurent, A. Carbone and S. Sacquin-Mora\*, Proteins, **84**, 1408-1421 (2016)
- 7. Fold and Flexibility: What can protein mechanical properties tell us about their folding nucleus? S. Sacquin-Mora\*, J. R. Soc. Interface, **12**, 20150876 (2015)
- 8. *Investigating the structural variability and binding modes of the glioma targeting NFL-TBS.40-63 peptide on tubulin* Y. Laurin, P. Savarin, C. H. Robert, M. Takahashi, J. Eyer, C. Prévost and S. Sacquin-Mora\*, Biochemistry, **54**, 3660-3669 (2015)
- 9. Fluctuations in the dipole moment of membrane-bound hydrogenase from Aquifex aeolicus account for its adaptability to charged electrodes
   F. Oteri, A. Ciaccafava, A. de Poulpiquet, M. Baaden, E. Lojou and S. Sacquin-Mora\*, PhysChem-ChemPhys 16, 11318-11322 (2014)
- 10. Protein-protein interactions in a crowded environment: An analysis via cross-docking simulations and evolutionary information, A. Lopes, S. Sacquin-Mora, V. Dimitrova, E. Laine, Y. Ponty and A. Carbone\*, PLoS Computational Biology 9, e1003369 (2013).

### Selection of conferences and talks

- **April 2019 Invited talk, CECAM 50th anniversary symposium**, IDRIS Orsay, France 50 years of modeling life
- **Sept. 2018 Contributed talk, CECAM workshop on Normal Modes Analysis**, Paris, France *Mechanical variations in proteins with large-scale motions highlight the formation of structural locks*
- **June 2018 Invited talk, École thématique DynaMoPPI**, Nantes, France
  Great interactions and hidden partners: Lessons from protein blind docking on protein binding sites and function.
- June 2017 Invited talk, Gordon Research Conference: Computational Aspects Biomolecular NMR Sunday River USA, *Protein Mechanics as a Bridge Between Structure and Function, a Coarse-Grain Approach.*
- **June 2015 Contributed talk, TheoBio**, Cagliary, Italy *Multiscale modeling of the mechanical nucleus in globular proteins*
- May 2013 Contributed talk, Molecular Perspectives on Protein-Protein Interactions, Pultusk, Poland *High-throughput investigation of protein-protein interactions via cross-docking simulations*
- Nov. 2012 Invited talk, Congrès SFBBM-SFB: Mécanismes moléculaires et processus vitaux intégrés, Grenoble, France Modeling protein mechanics with coarse-grain representations: From structure to function
- Sept. 2011 Invited talk, Workshop, Hands on course on Coarse Grain Methods for Biomolecular Simulations, Institut Pasteur de Montevideo, Coarse-Grain Models for Protein Mechanics

## Organization of scientific meetings

**May 2019 CECAM Workshop , Biomolecular mechanismes at functionalized soid interfaces**, Paris, France, 40 participants

Oct. 2017 CECAM Workshop, Disordered protein segments: revisiting the structure-function paradigm, Paris, France, 50 participants

**April 2017 Understanding Protein Interactions : from Molecules to Organisms**, Lyon, France, 100 participants

Oct. 2014 Protein-Protein Interactions on the Genomic Scale, Paris, France, 100 participants

**Oct. 2011 XVIIème congrès du GGMM**,(Groupe de Graphisme et Modélisation Moléculaire), La Rochelle, France, 120 participants

### Supervision of junior researchers (2013-2019)

### January 2017 - Nicolas Bourassin (1 paper published)

Master Thesis and PhD Thesis (Université Paris 7)

Modeling enzymes orientation on electrode surfaces for green energy production.

### Sept. 2016-Sept. 2017 – Nathalie Lagarde (1 paper published)

Postdoc (ANR project MAPPING)

Prediction of binding sites for proteins with multiple interaction partners

### July 2014-Dec.2015 – Lydie Vamparys-Laurent (1 paper published)

Postdoc (ANR project MAPPING)

Binding incorrect partners to understand protein recognition and function

### Oct. 2013-Sept.2016 – Yoann Laurin (2 papers published)

PhD Thesis (Université Paris 7-LABEX DYNAMO)

Investigating the structural variability and binding modes of the glioma targeting NFL-TBS.40-63 peptide on tubulin

### Sept. 2011-Sept. 2013 - Francesco Oteri (2 papers published)

Postdoc (ANR project BIOPAC)

Modeling biocatalytic enzymes for biofuel cells

## **Teaching activities (2013-2019)**

**Since 2012-** *Kinetics and Thermodynamics* 

Lecturer in chemistry, Université Paris 5-Descartes, France

**2016** Protein mechanics

DynaMol Summer school.

**Since 2015-** *Multiscale modelling for biological systems* 

Theoretical Chemistry label.

## General public communications

**2019** *Top of the Prots*:https://topoftheprots.com

A popularization blog about proteins (in french)

**July 2018** *Making gender equality more than theoretical* 

Comment published in Chemistry World regarding gender balance in scientific events

https://tinyurl.com/ChemWorldSacquin

**Oct. 2017** Biochemistry expert for the general scientific journal *Pour la Science*: https://tinyurl.com/PLScalamar

**March 2017** Highlight of the project regarding the anti-cancerous NFL-TBS.40-63 peptide on the GENCI website for the *Semaine du cerveau*: http://www.genci.fr/fr/node/834.

**Sept. 2016** The ProPHet program for investigating protein mechanics is available online on the RPBS webserver: http://mobyle.rpbs.univ-paris-diderot.fr/cgi-bin/portal.py#forms::ProPHet