

Giacomo Fiorin - Curriculum Vitae

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Web: <http://giacomofiorin.github.io> <http://scholar.google.com/citations?user=-Z0e2xoAAAAJ>

ACADEMIC APPOINTMENTS

07/2011 – present Assistant Professor of Research, Temple University

09/2009 – 06/2011 Postdoctoral Research Associate, Temple University (Advisor: Michael L. Klein)

05/2007 – 08/2009 Postdoctoral Research Associate, University of Pennsylvania (Advisor: Michael L. Klein)

EDUCATION

10/2002 – 12/2006 PhD in Statistical and Biological Physics, International School for Advanced Studies, Trieste, Italy (Advisor: Paolo Carloni)

10/1997 – 02/2002 Combined BS+MS in Physics, University of Padua, Italy (Advisor: Enrico Maglione)

TEACHING EXPERIENCE

2011 – present Instructor for the *Biostatistics* course for the programs: biology (BA, BS, MS, and PhD), mechanical and biomedical engineering (MS and PhD), kinesiology and health professions (BS, MS, PhD), pharmacy (PhD), Temple University

2010 – 2013 Adjunct Instructor for the *Statistical Thermodynamics* course, chemistry PhD program, Temple University

CURRENT RESEARCH ACTIVITIES

- Structure and dynamics of biological membranes.
- Molecular structure of the stratum corneum of human skin and permeation of small molecules.
- Membrane fusion in eukaryotes and in viruses.
- Development of coarse-grained force fields for membrane lipids and embedded proteins.
- Development of the collective variables software module (<http://colvars.github.io/>) for biased sampling simulations.
- Mechanism of ion conduction through the M2 viral channel of the influenza A virus.
- Analysis of the water structure in protein cavities to guide the design of new small-molecule pharmaceuticals inhibitors.
- Mechanisms of protein-protein interaction and recognition.

PEER-REVIEWED PUBLICATIONS

In reverse chronological order; asterisks indicate equal contributions.

1. MacDermaid CM, DeVane RH, Klein ML, **Fiorin G**.
“Dehydration of multilamellar fatty acid membranes: a molecular dynamics-based model for skin acidification”
J. Chem. Phys. **141**:22D526 (2014) <http://dx.doi.org/10.1063/1.4902363>
2. Gianti E, Carnevale V, DeGrado WF, Klein ML, **Fiorin G**.
“Hydrogen-bonded water molecules in the M2 channel of the influenza A virus guide the binding preferences of ammonium-based inhibitors”
J. Phys. Chem. B (in press) <http://dx.doi.org/10.1021/jp506807y>
3. Wu Y, Canturk B, Jo H, Ma C, Gianti E, Klein ML, Pinto LH, Lamb RA, **Fiorin G**, Wang J, DeGrado WF.
“Flipping in the Pore: Discovery of Inhibitors that bind in different orientations to the Wild-Type versus the Amantadine-Resistant S31N Mutant of Influenza A Virus M2 Proton Channel”
J. Am. Chem. Soc. (in press) <http://dx.doi.org/10.1021/ja508461m>
4. Thomaston JL, Alfonso-Prieto M, Woldeyes RA, Fraser JS, Klein ML, **Fiorin G**, DeGrado WF.
“High resolution structures of the M2 proton channel from influenza A virus reveal dynamic pathways for proton stabilization and transduction.”
(under review)
5. Dong H, **Fiorin G**, DeGrado WF, Klein ML.
“Proton release from the histidine-cluster in the M2 channel of the influenza A virus via molecular dynamics simulations”
J. Phys. Chem. B **118**(44):12644–12651 (2014) <http://dx.doi.org/10.1021/jp5102225>
6. Dong H, Klein ML, **Fiorin G**.
“Counterion-assisted Cation Transport in a Biological Calcium Channel”
J. Phys. Chem. B **118**(32):9668–9676 (2014) <http://dx.doi.org/10.1021/jp5059897>
7. Dewan S, Carnevale V, Bankura A, Eftekhari-Bafrooei A, **Fiorin G**, Klein ML, Borguet E.
“Structure of Water at Charged Interfaces: A Molecular Dynamics Study”
Langmuir **30**(27):8056–8065 (2014) <http://dx.doi.org/10.1021/la5011055>
8. Kumar BK, **Fiorin G**, Klein ML, Balasubramanian S.
“Supramolecular Polymerization of Benzene-1,3,5-tricarboxamide: A Molecular Dynamics Simulation Study”
J. Phys. Chem. B **118**(19):5218–5228 (2014) <http://dx.doi.org/10.1021/jp502779z>
9. Vidossich P, Loewen P, Carpena X, **Fiorin G**, Fita I, Rovira C
“Binding of the Anti-Tubercular Pro-Drug Isoniazid in the Heme Access Channel of Catalase–Peroxidase (KatG). A Combined Structural and Metadynamics Investigation”
J. Phys. Chem. B **118**(11):2924–2931 (2014) <http://dx.doi.org/10.1021/jp4123425>
10. **Fiorin G**, Klein ML, Hénin J.
“Using collective variables to drive molecular dynamics simulations.”
Mol. Phys. **22-23**:3345–3362 (2013) <http://dx.doi.org/10.1080/00268976.2013.813594>
(most downloaded article of the *Mol. Phys.* website)
11. Dong H*, **Fiorin G***, DeGrado WF, Klein ML.
“Exploring histidine conformations in the M2 channel lumen of the influenza A virus via molecular simulations”
J. Phys. Chem. Letters **4**:3067–3071 (2013) <http://dx.doi.org/10.1021/jz401672h>

12. **Fiorin G**, Klein ML, DeVane RH, Shinoda W.
 “Computer Simulation of Self-assembling Macromolecules”
Advances in Polymer Science **262**:93-107 (2013) http://dx.doi.org/10.1007/12_2013_262
13. Dong H, **Fiorin G**, Carnevale V, Treptow W, Klein ML.
 “Pore waters regulate ion permeation in a calcium release-activated calcium channel”
Proc. Natl. Acad. Sci. USA, **110**:17332-17337 (2013) <http://dx.doi.org/10.1073/pnas.1316969110>
14. Ma C*, **Fiorin G***, Carnevale V*, Wu Y, Wang J, Lamb RA, Klein ML, Pinto LH, DeGrado WF.
 “Asp44 stabilizes the Trp41 gate of the M2 proton channel of influenza A virus”
Structure **21**:2033-2041 (2013) <http://dx.doi.org/10.1016/j.str.2013.08.029>
15. Wang J, Ma C, Wang J, Jo H, Canturk B, **Fiorin G**, Pinto LH, Lamb RA, Klein ML, DeGrado WF.
 “Discovery of Dual Inhibitors of WT and the Amantadine-Resistant Mutant, S31N of M2 from Influenza A Virus”
J. Med. Chem. **56**(7):2804–2812 (2013). <http://dx.doi.org/10.1021/jm301538e>
16. Wang J, Wu Y, Ma C, **Fiorin G**, Wang J, Pinto LH, Lamb RA, Klein ML, DeGrado WF
 “Structure and inhibition of the drug-resistant S31N mutant of the M2 ion channel of influenza A virus”
Proc. Natl. Acad. Sci. USA **110**(4):1315-20 (2013) <http://dx.doi.org/10.1073/pnas.1216526110>
17. Wang J, Ma C, **Fiorin G**, Carnevale V, Wang T, Hu F, Lamb RA, Pinto LH, Hong M, Klein ML, DeGrado WF.
 “Molecular dynamics simulation directed rational design of inhibitors targeting drug-resistant mutants of influenza A virus M2.”
J. Am. Chem. Soc. **133**(32):12834–41 (2011). <http://dx.doi.org/10.1021/ja204969m>
18. Donald JE*, Zhang Y*, **Fiorin G***, Carnevale V, Slochower DR, Gai F, Klein ML, Degrado WF.
 “Transmembrane orientation and possible role of the fusogenic peptide from parainfluenza virus 5 (PIV5) in promoting fusion.”
Proc Natl Acad Sci USA **108**(10):3958–63 (2011). <http://dx.doi.org/10.1073/pnas.1019668108>
19. **Fiorin G**, Carnevale V, DeGrado WF.
 “The flu's proton escort.” (perspective)
Science **330**:456-8 (2010). <http://dx.doi.org/10.1126/science.1197748>
20. Carnevale V*, **Fiorin G***, Levine BG*, DeGrado WF and Klein ML.
 “Multiple Proton Confinement in the M2 Channel from the Influenza A Virus.”
J. Phys. Chem. C **114**(48):20856–20863 (2010). <http://dx.doi.org/10.1021/jp107431g>
21. Acharya R*, Carnevale V*, **Fiorin G***, Levine BG*, Polishchuck AL*, Balannik V, Samish I, Lamb RA, Pinto LH, Klein ML, DeGrado WF.
 “Structure and mechanism of proton transport through the transmembrane tetrameric M2 protein bundle of the influenza A virus.”
Proc Natl Acad Sci USA **107**(34):15075-80 (2010). <http://dx.doi.org/10.1073/pnas.1007071107>
22. Vidossich P, **Fiorin G**, Alfonso Prieto M, Derat E, Shaik S, Rovira C.
 “On the role of water in peroxidase catalysis: a theoretical investigation of HRP compound I formation.”
J. Phys. Chem. B **114**(15):5161-9 (2010). <http://dx.doi.org/10.1021/jp911170b>
23. Balannik V, Carnevale V, **Fiorin G**, Levine BG, Lamb RA, Klein ML, DeGrado WF, Pinto LH.
 “Functional studies and modeling of pore-lining residue mutants of the influenza A virus M2 ion channel.”
Biochemistry **49**(4):696-708 (2010). <http://dx.doi.org/10.1021/bi901799k>

24. Hénin J, **Fiorin G**, Chipot C, Klein ML.
 “Exploring Multidimensional Free Energy Landscapes Using Time-Dependent Biases on Collective Variables.”
J. Chem. Theory Comput. **6**(1):35-47 (2010). <http://dx.doi.org/10.1021/ct9004432>
25. **Fiorin G**, Pastore A, Carloni P, Parrinello M.
 “Using metadynamics to understand the mechanism of calmodulin/target recognition at atomic detail.”
Biophys. J. **91**(8):2768-2777 (2006). <http://dx.doi.org/10.1529/biophysj.106.086611>
26. **Fiorin G**, Biekofsky RR, Pastore A, Carloni P.
 “Unwinding the helical linker of calcium-loaded calmodulin: a molecular dynamics study.”
Proteins **61**(4):829-39 (2005). (Cover article) <http://dx.doi.org/10.1002/prot.20597>
27. **Fiorin G**, Maglione E, Ferreira LS.
 “Theoretical description of deformed proton emitters: nonadiabatic quasi-particle method.”
Phys. Rev. C **67**(5):054302 (2003). <http://dx.doi.org/10.1103/PhysRevC.67.054302>

INVITED TALKS, PRESENTATIONS AND POSTERS

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| 11/14/2014 | “ <i>Confined water molecules used as a template for drug design and as a tool to study the (dis)assembly of human skin</i> ”, Dept. of Biochemistry, Rowan University (invited talk) |
| 10/15/2014 | “ <i>Confined water molecules used as a template for drug design and as a tool to study the (dis)assembly of human skin</i> ”, Physics Colloquium, Drexel University (invited talk) |
| 09/16/2014 | “ <i>Confined water molecules used as a template for drug design and as a tool to study the (dis)assembly of human skin</i> ”, Department of Chemistry faculty research seminar, Temple University (invited talk) |
| 09/03/2014 | “ <i>Simulation studies of co-aggregation in the mechanism of vesicle fusion and the assembly of human skin</i> ”, Biomembrane Days Symposium, Berlin, Germany (invited talk) |
| 08/10/2014 | “ <i>Deconstructing the pathways of ion conduction to describe the geometry of inhibition sites of the flu's proton channel</i> ”, American Chemical Society National Meeting, San Francisco, CA (poster) |
| 08/10/2014 | “ <i>Water molecules work in concert to direct the traffic of protons in a viral channel</i> ”, American Chemical Society National Meeting, San Francisco, CA (invited talk) |
| 04/01/2014 | “ <i>Advanced modeling of the human skin barrier</i> ”, Interdisciplinary Research Group seminar, University of Pennsylvania, Philadelphia, PA (invited talk) |
| 02/18/2014 | “ <i>Role of model proteins in membrane fusion</i> ”, Biophysical Society National Meeting, San Francisco, CA (poster) |
| 04/17/2013 | “ <i>Proton conduction in biological membranes</i> ”, Workshop on “Frontiers in Neutron Structural Biology”, Oak Ridge National Laboratory, TN (invited talk) |
| 08/21/2012 | “ <i>Molecular dynamics simulation-directed rational design of inhibitors targeting drug-resistant mutants of influenza A virus M2</i> ”, American Chemical Society National Meeting, Philadelphia, PA (talk) |
| 02/29/2012 | “ <i>Mapping water density to design new blockers against a viral proton channel</i> ”, Biophysical Society Annual Meeting, San Diego, CA (poster) |

10/03/2011	<i>“Mapping water density to design new blockers against a viral proton channel”</i> , Workshop on “Innovative Approaches to Computational Drug Discovery”, CECAM, Lausanne, Switzerland (invited talk)
03/03/2011	<i>“Role of the fusogenic peptide from parainfluenza virus 5 in promoting fusion”</i> , Biophysical Society National Meeting, Baltimore, MD (poster)
02/24/2010	<i>“electrostatics of water clusters in the M2 channel of the influenza A virus”</i> , Biophysical Society National Meeting, San Francisco, CA (poster)
03/01/2009	<i>“Collective variable-based calculations in NAMD”</i> , Biophysical Society National Meeting, Boston, MA (poster)
08/13/2008	<i>“Amplitude and time scale of large scale motions in calmodulin”</i> , American Chemical Society Fall National Meeting, Philadelphia, PA (poster + SciMix)
06/19/2008	<i>“Amplitude and time scale of large scale motions in calmodulin”</i> , International School for Advanced Studies, Trieste, Italy (invited talk)
06/08/2008	<i>“Amplitude and time scale of large scale motions in calmodulin”</i> , Meeting on “Pushing the Boundaries of Biomolecular Simulations”, Ascona, Switzerland (poster)
11/29/2007	<i>“Using metadynamics to understand the mechanism of calmodulin/target recognition at atomic detail”</i> , University of Illinois-Urbana Champaign, IL (invited talk)
10/05/2006	<i>“Calmodulin dynamics and NMR properties from MD simulations”</i> , Slovenian-Italian NMR Consortium Inaugural Symposium, Trieste, Italy (invited talk)
10/04/2005	<i>“Calmodulin: dynamics and target recognition by molecular dynamics simulations”</i> Bunsen International discussion meeting, Tuzting - Munich, Germany (invited talk)
08/15/2005	<i>“Calmodulin: dynamics and target recognition by molecular dynamics simulations”</i> School on Computer Simulations in Condensed Matter, Erice, Italy (poster)
04/25/2005	<i>“Calmodulin: dynamics and target recognition by molecular dynamics simulations”</i> Swiss-Italian University, Lugano, Switzerland (invited talk)

SYNERGISTIC ACTIVITIES AND OUTREACH

2014 – present	Member of the American Association for the Advancement of Sciences (AAAS)
2014 – present	Member of the Temple University Institute for Regenerative Medicine and Engineering (TIME)
2012 – present	Member of the American Chemical Society (ACS)
2008 – present	Reviewer for the journals: <i>PNAS</i> , <i>PLOS Comp Biol</i> , <i>PLOS One</i> , <i>J Phys Chem B</i> , <i>Comp Phys Comm</i> , <i>Biophys J</i> , <i>Scientific Reports</i>
2008 – present	User support on the NAMD, LAMMPS and VMD software mailing lists
2007 – present	Member of the Biophysical Society
10/16/2014	Guest lecturer, Physics Colloquium (undergraduate program), Drexel University
8/18 – 8/22/2014	Instructor and co-organizer, School on “ <i>Molecular Dynamics for Biomolecules and Nanomaterials</i> ”, Temple University

03/18/2014	Guest lecturer, course on “ <i>Free Energy Calculations</i> ”, City University of New York, chemistry and biology PhD programs
12/2013	Reviewer of proposals for the Human Brain Project competitive call, European Union
4/16 – 4/18/2013	Invited speaker and panelist, Workshop on “Frontiers in Neutron Structural Biology”, Oak Ridge National Laboratory, TN
2/24 – 3/10/2013	Visiting instructor, course on “ <i>Free Energy Calculations and Advanced Molecular Dynamics Simulations</i> ”, Jawaharlal Nehru Centre for Advanced Scientific Research (JNCASR), Bangalore, India
04/2012	Guest lecturer on “ <i>Experimental determination of protein structure</i> ”, course on “ <i>Introduction to Structural Bioinformatics</i> ”, Chemistry and Biology PhD programs, Temple University
10/2009	Guest lecturer, course in “ <i>Physical chemistry II</i> ”, Chemistry BA and BS programs, Temple University
6/11 – 6/15/2007	Lecturer, workshop on “ <i>High Performance Computing</i> ”, University of Pennsylvania
11/21/2005	Guest lecturer, United World College of the Adriatic (high school), Duino, Italy

GRANTS, ADMINISTRATION AND RESEARCH SUPPORT

01/2015 – 12/2015	Co-PI with Michael Klein (PI) and Russell DeVane of the project “ <i>Shutdown and recovery of the barrier function of human skin</i> ”, DOE-INCITE supercomputing program – 92,000,000 core-hours
01/2014 – 12/2014	Co-PI with Michael Klein (PI) and Russell DeVane of the project “ <i>Assembling and sustaining the acid mantle of the human skin barrier</i> ”, DOE-INCITE supercomputing program – 75,000,000 core-hours
07/2012 – present	Co-PI with Michael Klein (PI) and Axel Kohlmeyer of the NSF grant 1212416 “ <i>Building Computational Models to Probe Membrane Fusion</i> ” – \$405,999 (2012 – 2015)
01/2013 – 12/2013	Co-PI with Michael Klein (PI) and Russell DeVane on the project “ <i>Advanced modeling of the human skin barrier</i> ”, DOE-INCITE supercomputing program – 65,000,000 core-hours
01/2011 – 12/2012	Co-PI with Michael Klein (PI), Russell DeVane, Vincenzo Carnevale and Axel Kohlmeyer of the project CHM045 “ <i>Coarse grained molecular dynamics studies of vesicle formation and fusion</i> ” under the DOE INCITE supercomputing program – 48,000,000 core-hours over 2 years on OLCF Jaguar
07/2011 – present	Supercomputing allocations manager for ICMS (NSF-XSEDE, DOE-INCITE and DOE-ERCAP programs)