### Ilan E. Chemmama

ichemmama@gmail.com (786)262-6090 Google Scholar Profile

### **QUALIFICATION SUMMARY**

Structural biologist, programmer, and software engineer experienced in using Bayesian inference for biology

- Inferring structure macromolecular assemblies of various sizes and scales by integrative modeling
  - Contributed to the determination of eight macromolecular assemblies structures
  - Maximized accuracy and precision by simultaneously utilizing all available information, including experimental data, physical principles, statistical analyses, and other prior models
  - Used sample-based Bayesian inference to sample models
  - Used high-performance scientific computing to sample those models
  - Estimated uncertainty originating from information that is sparse, noisy, ambiguous, or derived from heterogeneous samples
  - Developed and applied a pipeline to assess sampling convergence for stochastic sampling
  - Developed, implemented, and contributed code for data analyses in C++ and Python to the open-source software *Integrative modeling platform* (IMP) package
  - Attempted to improve the classification of low signal-to-noise images using convolution neural network.
- Strong communication and collaboration skills
  - Collaborated with large and small multidisciplinary teams of scientists to produce peer-reviewed publications, conference presentations, and successful competitive grant proposals
  - Mentored high school, undergraduate, and graduate students

### RELEVANT RESEARCH EXPERIENCES

#### **Computational Structural Biology**

University of California - San Francisco, San Francisco, CA USA

2013 to Present

Inferring structures, free energy differences, and kinetic rates of biological macromolecular assemblies by integrative modeling

Principal Investigator: Prof. Andrej Sali

- Computing integrative atomic structures based on medium-resolution EM density maps: Biomolecular structures are routinely computed based on their electron microscopy (EM) density maps. The accuracy and precision of such modeling depends critically on a scoring function that quantifies a match between a model and the map. We introduce a Bayesian posterior model density for a model given its map, explicitly taking into account the resolution and other uncertainties in the map.
- Structural dynamics of the human Cop9 signalosome: The COP9 signalosome (CSN) is an evolutionarily conserved eight-subunit protein complex that controls protein ubiquitination by deneddylating Cullin-RING E3 ligases. The activation and function of CSN hinges on its structural dynamics, which has been challenging to decipher by conventional tools. We determined the structural dynamics of the human Cop9 signalosome complex by integrative modeling, demonstrating how the synergy between multi-chemistry cross-linking mass spectrometry and integrative modeling can map the structural dynamics of macromolecular assemblies.
- Validation of stochastic sampling in integrative modeling: Modeling of macromolecular structures involves structural sampling guided by a scoring function, resulting in an ensemble of good-scoring models. By necessity, the sampling is often stochastic, and must be exhaustive at a precision sufficient for accurate modeling and assessment of model uncertainty. We present an objective and automated method for validating and assessing stochastic sampling in integrative modeling.

- Inferring free-energy differences and kinetic rates of biomolecules: Mapping the functional cycle of a biomolecular machine requires determining the structures of key states, their stabilities, and the kinetics of their interconversions under native conditions. We describe a method that outputs these free-energy differences and kinetic rates based on time-dependent negative-stain electron microscopy particle images for a given set of structural states and transitions between them. We illustrate our method by applying it to the yeast Hsp90 chaperone engaged in its ATPase cycle in the presence of two ATP analogs, the non-hydrolysable AMP·PNP or the slowly hydrolysable ATP $\gamma$ S. I have also attempted to improve negative-stain EM images into pre-determined classes using convolution neural network in PyTorch. The signal-to-noise ratio in the images limited the usefulness of the approach.
- 3D structures determination of biological assemblies of various sizes and scales by integrative modeling: Leveraging multiple sources of information (e.g. cross-linking mass spectrometry, 2D EM images, 3D EM maps, among others), I contributed to the structure determination of · the yeast exocyst complex, · the human ECM29-Proteasome assembly under oxidative stress, · the yeast nuclear pore complex (NPC) cytoplasmic mRNA export platform, · the major membrane ring component of the yeast NPC, · the entire yeast NPC, and · the reconstruction of 3D structures of MET antibodies.

#### THEORETICAL BIOPHYSICS

2008-2013

Florida International University, Miami, FL USA

Physics of small peptide folding and the formation of secondary structures

**Principal Investigators:** Profs. Bernard S. Gerstman and Prem P. Chapagain

- Structural propensities and entropy effects in peptide helix-coil transitions: The helix-coil transition in peptides is a critical structural transition leading to functioning proteins. Peptide chains have a large number of possible configurations that must be accounted for in statistical mechanical investigations. Using hydrogen bond and local helix propensity interaction terms, we develop a method for obtaining and incorporating the degeneracy factor that allows the exact calculation of the partition function for a peptide as a function of chain length.
- Pairwise amino acid secondary structural propensities: We investigate the propensities for amino acids to form a specific secondary structure when they are paired with other amino acids. Our investigations use molecular dynamics (MD) computer simulations, and we compare the results to those from the Protein Data Bank (PDB). Proper comparison requires weighting of the MD results in a manner consistent with the relative frequency of appearance in the PDB of each possible pair of amino acids.

### **EDUCATION**

PH.D. IN BIOPHYSICS March 2020

University of California - San Francisco, San Francisco, CA USA

**Thesis**: "Inferring structures, free-energy differences, and kinetic rates of biological macromolecular systems by integrative modeling." *Advisor*: Andrej Sali

National Science Foundation Graduate Research Fellow

#### **BACHELOR OF SCIENCE IN PHYSICS**

May 2011

Florida International University, Miami, FL USA

**Honors Thesis**: "Theoretical Quantification of Self-Organization of Secondary Interactions in Protein Folding." *Advisors*: Prem P. Chapagain and Bernard S. Gerstman

Graduated with minors in Chemistry and Mathematics

Graduated Magna Cum Laude, Honors Course of Study in Physics, and through The Honors College.

### **SELECTED PUBLICATIONS (OUT OF 13)**

## Using Integrative Modeling Platform to compute, validate, and archive a model of a protein complex structure

Protein Science Jan 2021, 30 (1) 250-261

Daniel J Saltzberg, Shruthi Viswanath, Ignacia Echeverria, **Ilan E Chemmama**, Ben Webb, Andrej Sali **Integrative structure and function of the yeast exocyst complex** 

Protein Science June 2020, 29 (6) 1486-1501

Sai J Ganesan, Michael J Feyder, **Ilan E Chemmama**, Fei Fang, Michael P Rout, Brian T Chait, Yi Shi, Mary Munson, Andrej Sali

## Structural dynamics of the human COP9 signalosome revealed by cross-linking mass spectrometry and integrative modeling

Proceedings of the National Academy of Sciences Feb 2020, 117 (8) 4088-4098

Craig Gutierrez\*, **Ilan E Chemmama**\*, Haibin Mao, Clinton Yu, Ignacia Echeverria, Sarah A Block, Scott D Rychnovsky, Ning Zheng, Andrej Sali, Lan Huang

#### Modeling biological complexes using integrative modeling platform

Biomolecular Simulations 353-377 (2018) Humana, New York, NY

Daniel Saltzberg, Charles H. Greenberg, Shruthi Viswanath, **Ilan E. Chemmama**, Ben Webb, Riccardo Pellarin, Ignacia Echeverria, Andrej Sali

#### **Integrative Structure and Functional Anatomy of a Nuclear Pore Complex**

Microscopy and Microanalysis 24 S1, 1212-1213 (2018)

Ilona Nudelman, Seung Joong Kim, Javier Fernandez-Martinez, Yi Shi, Wenzhu Zhang, Barak Raveh, Thurston Herricks, Brian D. Slaughter, Joanna Hogan, Paula Upla, **Ilan E. Chemmama**, Riccardo Pellarin, Ignacia Echeverria, Manjunatha Shivaraju, Azraa S. Chaudhury, Junjie Wang, Rosemary Williams, Jay R. Unruh, Charles H. Greenberg, Erica Y. Jacobs, Zhiheng Yu, M. Jason de la Cruz, Roxana Mironska, David L. Stokes, John D. Aitchison, Martin F. Jarrold, Jennifer L. Gerton, Steven J. Ludtke, Christopher W. Akey, Brian T. Chait, Andrej Sali, and Michael P. Rout

#### **Structure and Functional Anatomy of a Nuclear Pore Complex**

*Nature* 555, 475-482 (2018)

Seung Joong Kim, Javier Fernandez-Martinez, Ilona Nudelman, Yi Shi, Wenzhu Zhang, Barak Raveh, Thurston Herricks, Brian D. Slaughter, Joanna Hogan, Paula Upla, **Ilan E. Chemmama**, Riccardo Pellarin, Ignacia Echeverria, Manjunatha Shivaraju, Azraa S. Chaudhury, Junjie Wang, Rosemary Williams, Jay R. Unruh, Charles H. Greenberg, Erica Y. Jacobs, Zhiheng Yu, M. Jason de la Cruz, Roxana Mironska, David L. Stokes, John D. Aitchison, Martin F. Jarrold, Jennifer L. Gerton, Steven J. Ludtke, Christopher W. Akey, Brian T. Chait, Andrej Sali, and Michael P. Rout

## Assessing exhaustiveness of Stochastic sampling for Integrative Modeling of Macro-molecular Structures

Biophys. J. 113, 2344-2353 (2018)

Shruthi Viswanath\*, **Ilan E. Chemmama**\*, Peter Cimermancic, and Andrej Sali

## The Proteasome-interacting Ecm29 Protein Disassembles the 26S Proteasome in Response to Oxidative Stress

J. Biol. Chem. 292, 16310-16320 (2017)

Xiaorong Wang, **Ilan E. Chemmama**, Clinton Yu, Alexander Huszagh, Yue Xu, Rosa Viner, Sarah A. Block, Peter Cimermancic, Scott D. Rychnovsky, Yihong Ye, Andrej Sali, and Lan Huang

## Reconstruction of 3D Structures of MET Antibodies from Electron Microscopy 2D Class Averages

PLoS One 12, e0175758 (2017)

Qi Chen, Michal Vieth, David Timm, Christine Humblet, Dina Schneidman-Duhovny, **Ilan E. Chemmama**, Andrej Sali, Jirong Lu, Ling Liu, and Wei Zeng

## Molecular Architecture of the Major Membrane Ring Component of the Nuclear Pore Complex

Structure 25, 434-445 (2017)

Paula Upla, Seung Joong Kim, Parthasarathy Sampathkumar, Kaushik Dutta, Sean M. Cahill, **Ilan E. Chemmama**, Rosemary Williams, Jeffrey B. Bonanno, William J. Rice, David L. Stokes, David Cowburn, Steven C. Almo, Andrej Sali, Michael P Rout, and Javier Fernandez-Martinez

### Structure and Function of the Nuclear Pore Complex Cytoplasmic mRNA Export Platform *Cell* 167, 1215-1228 (2016)

Javier Fernandez-Martinez, Seung Joong Kim, Yi Shi, Paula Upla, Riccardo Pellarin, Michael Gagnon, **Ilan E. Chemmama**, Junjie Wang, Ilona Nudelman, Wenzhu Zhang, Rosemary Williams, William J. Rice, David L. Stokes, Daniel Zenklusen, Brian T. Chait, Andrej Sali, and Michael P. Rout

#### **Pairwise Amino Acid Secondary Structural Propensities**

Phys. Rev. E 91, 042709 (2015)

Ilan E. Chemmama, Prem P. Chapagain, and Bernard S. Gerstman

#### Structural Propensities and Entropy Effects in Peptide Helix-Coil Transitions

Phys. Rev. E 86, 031915 (2012)

Ilan E. Chemmama, Adam Colt Pelea, Yuba R. Bhandari, Prem P. Chapagain, and Bernard S. Gerstman

### **SELECTED LIST OF AWARDS**

#### **National Science Foundation Graduate Research Fellow**

University of California - San Francisco, 2015

#### The Phi Beta Kappa Society

Epsilon Chapter of Florida, 2011

### SELECTED PRESENTATION AND CONFERENCES ATTENDED

# Towards modeling multiple states of biomolecules satisfying spatial restraints from single-particle EM images

Ilan E. Chemmama 6-10 May 2017

World Molecular Engineering Conference, Cabo San Lucas, Mexico

### Towards modeling multiple states of biomolecules by satisfaction of spatial restraints from single-particle electron microscopy images

llan E. Chemmama, Charles H. Greenberg, and Andrej Sali

11-15 February 2017

61st Annual Meeting of the Biophysical Society, New Orleans, LA USA

## Theoretical and computational quantification of self-organization of secondary interactions in protein folding

*Ilan E. Chemmama, Prem P. Chapagain, Bernard S. Gerstman, Jose L. Parra* 31 March to 2 April 2011 National Conference for Undergraduate Research, Ithaca, NY USA

#### **OUTREACH AND TEACHING EXPERIENCES**

#### **Volunteer for JUMA Ventures**

San Francisco, CA USA

January 2016 to Present

Juma strives to break the cycle of poverty by paving the way to work, education, and financial capability for youth across America. I am a STEM tutor where I mentor students in science and mathematics (Algebra, Statistics, and Calculus, Chemistry) as well as SAT/ACT prep.

#### **Volunteer for the Overtown Youth Center**

Miami, FL USA

June 2010 to August 2013

I was a volunteering tutor for a partnership established by Dr. Jayne Klein between the Honors College at Florida International University - Biscayne Bay Campus and the Overtown Youth Center. Overtown is Miami's most economically disadvantaged neighborhood. The partnership provides youth attending the center opportunities to build competency in English, mathematics, and critical thinking skills, with the aim of improving their access to higher education. I often used science as an anchor for students to stimulate their interest in science and mathematics.

#### **Research Mentorship**

UCSF, CA USA

2015 to present

I mentored two first-year graduate students rotating in the Sali lab and two high-school students doing summer internships

#### **Teacher Assistant**

UCSF, CA USA

2014-2017

I was a teacher assistant for Macromoleculess, Structures, and Interactions A and the NSF Writing Class

#### **Instructor for the Department of Physics**

FIU, FL USA

January 2010 to July 2013

I taught various sections of classes for first and second-year level astronomy and physics courses and laboratories.