#### **CURRICULUM VITAE**

## Xinqiu YAO (Xin-Qiu YAO), Ph.D.

DEPARTMENT OF CHEMISTRY · GEORGIA STATE UNIVERSITY

29 PEACHTREE CENTER AVE NE · ATLANTA, GA 30303-2515, USA
· xinqiu.yao@gmail.com · https://xinqyao.github.io

BIOINFORMATICS • COMPUTATIONAL CHEMISTRY • BIOPHYSICS • MOLECULAR DYNAMICS • COARSE-GRAINED MODEL • DOCKING • FREE ENERGY CALCULATION • MACHINE LEARNING • GENOME INFORMATICS

#### RESEARCH INTERESTS

My research applies bioinformatics and computational chemistry to fundamental questions in molecular biochemistry and biophysics. My general approach is to develop and apply state-of-the-art computational techniques that can be coupled to experiments and be predictive of a wide range of physiological and pathophysiological processes.

EDUCATION	
2008	Ph.D. in Mechanics (Biomechanics and Medical Engineering), Peking University, China
	Title of dissertation: "A Complex Systems Approach to Studies of Protein Structures"
2002	B.Sc. in Applied Physics (Minored in Japanese), Hunan University, China

#### APPOINTMENTS AND RESEARCH EXPERIENCE

## 2017– **Postdoctoral Associate** in Chemistry, Georgia State University

- Developing new computational methods for understanding allosteric regulations
- Elucidating mechanisms underlying human enzymes with molecular dynamics (MD)

# 2012–2017 **Postdoctoral Fellow** in Computational Medicine and Bioinformatics, <u>University of Michigan</u>

- Lead developer of Bio3D, an R package for biomolecular sequence, structure, dynamics, and evolution analysis; Developer of Bio3D-web, a web application for interactive online structural analysis
- Developed the ensemble correlation network analysis method and applied it to elucidate the activation mechanisms in heterotrimeric G proteins
- Developed the ensemble normal mode analysis for exploring dynamical aspects from structural biological data
- Identified the allosteric pathway during activation of GPCR kinases
- Developed a combined approach of molecular docking, MD, and binding free energy calculation to predicting HLA-specific drug hypersensitivities
- Activation and allosteric mechanisms in estrogen receptors

## 2008–2012 **Postdoctoral Fellow** in Biophysics, <u>Kyoto University</u>, Japan

- Elucidated mechanisms underlying drug export in the multidrug transporter AcrB with coarse-grained molecular simulations
- Developed CafeMol, a software package for coarse-grained biomolecular modeling and simulations

## 2002–2008 Research Assistant, Center for Theoretical Biology, Peking University, China

- Developed a novel approach to protein secondary structure prediction using dynamic Bayesian networks and neural networks
- Investigated the folding mechanisms of the small protein Trp-cage with MD
- Developed algorithms for prokaryotic gene identification

## 2004 Internship, Intel China Research Center, Beijing, China

Improved the CE structural alignment algorithm for remote homology detection

#### **PUBLICATIONS** (\*Joint-First; §Co-Corresponding)

## **Book Chapter**

1. Grant BJ, Skjærven L, and <u>Yao XQ</u>. Comparative protein structure analysis with Bio3D-web. *Methods in Molecular Biology* (2019), submitted (invited).

#### Peer-reviewed Journal Articles

- *2019 -*
- 2. Souffrant M, <u>Yao XQ</u>, Momin M, and Hamelberg D. Gaucher disease mutations allosterically alter active site dynamics in acid-β-glucosidase. *Submitted*.
- 2018 -
- 3. Li H, <u>Yao XQ</u>, and Grant BJ. Comparative structural dynamic analysis of GTPases. *PLoS Comput Biol* (2018) **14**:e1006364.
- 4. **Yao XQ**§, Momin M, and Hamelberg D§. Elucidating allosteric communications in proteins with difference contact network analysis. *J Chem Inf Molel* (2018) 58:1325-1330.
- 5. Rodriguez-Bussey I, <u>Yao XQ</u>, Shouaib AD, Lopez J, and Hamelberg D. Decoding allosteric communication pathways in cyclophilin A with a comparative analysis of perturbed conformational ensembles. *J Phys Chem B* (2018) 122:6528-6535.
- Momin MF, <u>Yao XQ</u>, Thor W, and Hamelberg D. Substrate sequence determines catalytic activities, domain-binding preferences, and allosteric mechanisms in Pin1. *J Phys Chem B* (2018) 122:6521-6527.
- 7. Vu PJ\*, <u>Yao XQ\*</u>, Momin M, and Hamelberg D. Unraveling allosteric mechanisms of enzymatic catalysis with an evolutionary analysis of residue-residue contact dynamical changes. *ACS Catalysis* (2018) **8**:2375-2384.
- 2017 -
- 8. <u>Yao XQ\*</u>, Cato MC\*, Labudde E, Beyett TS, Tesmer JJG, and Grant BJ. Navigating the conformational landscape of G protein-coupled receptor kinases during allosteric activation. *J Biol Chem*. (2017) 292:16032-16043.
- 9. Bouley R, Waldschmidt HV, Cato MC, Cannavo A, Song J, Cheung JY, Yao XQ, Koch WJ, Larsen SD, and Tesmer JJG. Structural determinants influencing the potency and selectivity of indazole-paroxetine hybrid G protein-coupled receptor kinase 2 inhibitors. *Mol Pharmacol* (2017) 92:707-717.
- 10. Stender JD, Nwachukwu JC, Kastrata I, Kim Y, Strid T, Yakir M, Srinivasan S, Nowak J, Izard T, Erumbi R, Carlson KE, Katzenellenbogen JA, <u>Yao XQ</u>, Grant BJ, Leong HS, Lin CY, Frasor J, Nettles KW, Glass C. Structural and molecular mechanisms of cytokine-mediated endocrine resistance in human breast cancer cells. *Mol Cell* (2017) 65:1122-1135.e5.
- 11. Nwachukwu JC, Srinivasan S, Bruno NE, Nowak J, Wright NJ, Minutolo F, Rangarajan ES, Izard T, <u>Yao</u> <u>XQ</u>, Grant BJ, Kojetin DJ, Elemento O, Katzenellenbogen JA, Nettles KW. Systems structural biology analysis of ligand effects on ERα predicts cellular response to environmental estrogens and antihormone therapies. *Cell Chem Biol* (2017) 24:35-45.
- 12. Jariwala S, Skjærven L, <u>Yao XQ</u>, and Grant BJ. Investigating protein sequence-structure-dynamics relationships with Bio3D-web. *J Vis Exp* (2017) 125:e55640.
- 2016 -
- 13. Yao XQ, Skjærven L, and Grant BJ. Rapid characterization of allosteric networks with ensemble normal mode analysis. *J Phys Chem B* (2016) 120:8276-8288.
- 14. Yao XQ\*, Malik RU\*, Griggs NW, Skjærven L, Traynor JR, Sivaramakrishnan S, and Grant BJ. Dynamic coupling and allosteric networks in the alpha subunit of heterotrimeric G proteins. *J Biol Chem* (2016) 291:4742-4753.
- 15. Skjærven L, Jariwala S, <u>Yao XQ</u>, and Grant BJ. Online interactive analysis of protein structure ensembles with Bio3D-web. *Bioinformatics* (2016) 32:3510-3512.
- *2008-2015 -*

- 16. Scarabelli G, Soppina V, <u>Yao XQ</u>, Atherton J, Moores C, Verhey K, and Grant BJ. Mapping the processivity determinants of the kinesin-3 motor domain. *Biophys J* (2015) 109:1537-1540.
- 17. Skjærven L, <u>Yao XQ</u>, Scarabelli G, and Grant BJ. Integrating protein structural dynamics and evolutionary analysis with Bio3D. *BMC Bioinformatics* (2014) 15:399.
- 18. Yao XQ and Grant BJ. Domain-opening and dynamic coupling in the  $\alpha$ -subunit of heterotrimeric G proteins. *Biophys J* (2013) 105:L08-L10.
- 19. Yao XQ, Kimura N, Murakami S, and Takada S. Drug uptake pathways of multidrug transporter AcrB studied by molecular simulations and site-directed mutagenesis experiments. *J Am Chem Soc* (2013) 135:7474-7485.
- Kenzaki H, Koga N, Hori N, Kanada R, Li WF, Okazaki K, <u>Yao XQ</u>, and Takada S. CafeMol: a coarse-grained biomolecular simulator for simulating proteins at work. *J Chem Theory Comput* (2011) 7:1979-1989.
- 21. <u>Yao XQ</u>, Kenzaki H, Murakami S, and Takada S. Drug export and allosteric coupling in a multidrug transporter revealed by molecular simulations. *Nature Commun* (2010) 1:117. (The work was highlighted in *Nikkei Business Daily*, *Nikkan Kogyo Shimbun*, and *Kyoto Shimbun*).
- 22. Gao M, Zhu HQ, <u>Yao XQ</u>, and She ZS. Water dynamics clue to key residues in protein folding. *Biochem Biophys Res Comm* (2010) 392:95-99.
- 23. Gao M, <u>Yao XQ</u>, She ZS, Liu ZR, and Zhu HQ. Intermediate structure and slow hydration water dynamics in protein folding process. *Acta Physico-chimica Sinica* (2010) 26:1998-2006.
- 24. Kang H, <u>Yao XQ</u>, She ZS, and Zhu HQ. Water-protein interplay reveals the specificity of alpha-lytic protease. *Biochem Biophys Res Comm* (2009) 385:165-169.
- 25. Yao XQ and She ZS. Key residue-dominated protein folding dynamics. *Biochem Biophys Res Comm* (2008) 373:64-68.
- 26. **Yao XQ**, Zhu HQ, and She ZS. A dynamic Bayesian network approach to protein secondary structure prediction. *BMC Bioinformatics* (2008) 9:49.

GRANTS	
2018	Co-Writer, "Predicting the Impact of Genetic Variations on the Function of Proteins", R01
	[Under review]
2015	Co-Writer, "G Protein Activation Mechanisms", Proposal for allocation of simulation time
	on Anton at Pittsburgh Supercomputing Center [Awarded]

## **TEACHING AND MENTORING**

2013 **Co-Mentor,** Undergraduate Research Opportunity Program (UROP), <u>University of Michigan</u>

Coached an undergraduate student during his summer research project about computer-aided discovery of novel druggable pockets in heterotrimeric G proteins. Helped the student develop proper research strategies, master necessary techniques to conduct the research, and give a presentation about the results on a conference.

Teaching Assistant (Statistical Physics), School of Physics, Peking University, China

- Instructed students on solving problems
- Assisted the lecturer to grade course assignments and the final exam

#### **PRESENTATIONS**

Talks

2003

- "Elucidating allosteric communications in proteins with difference contact network analysis", The
   256th ACS National Meeting & Exposition, Boston (2018)
- "Allosteric modulation of G protein-coupled receptor kinase (GRK) activation," Sigtrans Seminar, University of Michigan (2017)
- "Dynamic coupling and activation mechanisms of heterotrimeric G proteins," Cytoskeleton Seminar,
   University of Michigan (2016)

- "Dynamic coupling and allosteric networks in the alpha subunit of heterotrimeric G proteins," *Sigtrans Seminar*, University of Michigan (2015)
- "G protein allosteric networks from molecular dynamics and normal mode analysis," Sigtrans Seminar, University of Michigan (2014)
- "Probing the activation mechanism of heterotrimeric G proteins with multiscale molecular simulations," Tools and Technology Seminar, University of Michigan (2014)
- "Nucleotide-dependent Transitions of the alpha-subunit of heterotrimeric G proteins probed with computational methods," Sigtrans Seminar, University of Michigan (2013)
- "Drug translocation and allosteric coupling in a multidrug transporter studied by molecular simulations," The 17th International Biophysics Congress (IUPAC), Beijing, China (2011)
- "Drug translocation and allosteric coupling in a multidrug transporter studied by molecular simulations," Center for Theoretical Biology, Peking University, Beijing, China (2011)
- "Functionally rotating mechanism of a multidrug transporter studied by coarse-grained simulations," The 13th Annual Meeting of Chinese Life Science Association, Kyoto, Japan (2011)
- "The uptake and export of drugs in a multidrug transporter studied by coarse-grained simulations," The 49th Annual Meeting of the Biophysical Society of Japan, Himeji, Japan (2011)
- "Drug export and allosteric coupling in a multidrug transporter revealed by molecular simulations,"
   New Era of Biosimulations with Supercomputers, Osaka, Japan (2011)
- "Functionally rotating mechanism of a multidrug transporter studied by coarse-grained simulation,"
   The 47th Annual Meeting of the Biophysical Society of Japan, Tokushima, Japan (2009)
- "Rotatory mechanism of AcrB transporter studied by coarse-grained simulation," Workshop for Next-Generation Integrated Simulation of Living Matters on Molecular Scale, Tokyo, Japan (2009)
- "Protein sequence, structure, and dynamics," Institute for theoretical physics, Chinese Academy of Science, Beijing, China (2007)

#### **Posters**

- Vu PJ, <u>Yao XQ</u>, Momin M, and Hamelberg D. Evolutionarily conserved and divergent residue-residue contact dynamics provide insights into the allosteric regulation of cyclophilins. *The9th Southeast Enzyme Conference*, Atlanta (2018)
- Vu PJ, <u>Yao XQ</u>, Momin M, and Hamelberg D. Evolutionarily conserved and divergent residue-residue contact dynamics provide insights into the allosteric regulation of cyclophilins. *The 255th ACS National Meeting & Exposition*, New Orleans (2018)
- Vu PJ, <u>Yao XQ</u>, Momin M, and Hamelberg D. Evolutionarily conserved and divergent residue-residue contact dynamics provide insights into the allosteric regulation of cyclophilins. *The 62nd Annual Meeting of Biophysical Society*, San Francisco (2018)
- Yao XQ, Jariwala S, Grant BJ. In silico prediction of HLA-associated drug hypersensitivity. *The 61st Annual Meeting of Biophysical Society*, New Orleans (2017)
- Yao XQ, Malik RU, Griggs NW, Skjærven L, Traynor JR, Sivaramakrishnan S, Grant BJ. Dynamic coupling and allosteric networks in the alpha subunit of heterotrimeric G proteins. *The 60th Annual Meeting of Biophysical Society*, LA (2016)
- Skjærven L, Jariwala S, Yao XQ, Ide J, Grant BJ. The Bio3D project: interactive tools for Structural Bioinformatics. The 60th Annual Meeting of Biophysical Society, LA (2016)
- Skjærven L, Yao XQ, Jariwala S, Grant BJ. Bio3D: interactive tools for structural bioinformatics. The 60th Annual Meeting of Biophysical Society, LA (2016)
- Scarabelli G, Soppina V, Yao XQ, Atherton J, Moores CA, Verhey KJ, Grant BJ. Mapping the processivity determinants of the kinesin-3 motor domain. The 60th Annual Meeting of Biophysical Society, LA (2016)
- Li H, Yao XQ, Grant BJ. Comparative structure dynamic analysis of G proteins. The 60th Annual Meeting of Biophysical Society, LA (2016)

- Yao XQ, Skjærven L, Grant BJ. Characterizing nucleotide dependent allostery in G-proteins with molecular dynamics and normal mode analysis. Symposium on Computational Discovery in Complex Systems Biology, University of Michigan (2015)
- Yao XQ, Skjærven L, and Grant BJ. G-protein allosteric networks from molecular dynamics and normal mode analysis. *The 59th Annual Meeting of Biophysical Society*, Baltimore (2015)
- Yao XQ, Scarabelli G, Skjærven L, Grant BJ. The Bio3D package: new interactive tools for Structural Bioinformatics. *The 58th Annual Meeting of Biophysical Society*, San Francisco (2014)
- Yao XQ, Jariwala S, and Grant BJ. Predicting HLA-specific drug hypersensitivity with molecular docking and molecular dynamics simulation. *The 58th Annual Meeting of Biophysical Society*, San Francisco (2014)
- Yao XQ and Grant BJ. Nucleotide-dependent transitions of the alpha subunit of heterotrimeric G-proteins probed with accelerated molecular dynamics simulations. *The 57th Annual Meeting of Biophysical Society*, Philadelphia (2013)
- Yao XQ, Kenzaki H, and Takada S. Functionally rotating mechanism of a multidrug transporter studied by coarse-grained simulation: exploring the binding pathways of drugs. *The 55th Annual Meeting of Biophysical Society*, Baltimore (2011)
- Yao XQ, Kenzaki H, and Takada S. Functionally rotating mechanism of a multidrug transporter studied by coarse-grained simulations. *The 54th Annual Meeting of the Biophysical Society*, San Francisco (2010)
- Kenzaki H, Koga N, Fujiwara S, Hori N, Kanada R, Okazaki K, Yao XQ, Li WF, Takada S. Biomolecular coarse-grained simulation program CafeMol. The 54th Annual Meeting of the Biophysical Society, San Francisco (2010)
- Yao XQ, Kenzaki H, and Takada S. Functionally rotating mechanism of multidrug transporter studied by coarse-grained simulations. *The 48th Annual Meeting of the Biophysical Society of Japan*, Sendai, Japan (2010)
- Yao XQ, Kenzaki H, and Takada S. Functionally rotating mechanism of a multidrug transporter studied by coarse-grained simulation. *The 2nd Biosupercomputing Symposium*, Tokyo, Japan (2010)
- Yao XQ, Kenzaki H, and Takada S. Functionally rotating mechanism of a multidrug transporter studied by coarse-grained simulation. The 4th International Symposium on Molecular Science of Fluctuations toward Biological Functions, Shiga, Japan (2010)

TRAININGS		
2018	"Scalable and Reproducible Structural Bioinformatics Workshop & Hackathon 2018,"	
	San Diego, CA	
2015	"Anton Workshop 2015," Pittsburgh, PA	
2003	"School of Parallel Computing," Chinese Academy of Science, Beijing, China	

# **HONORS AND AWARDS**

2011	The Best Talk, "The 13th Annual Meeting of Chinese Life Science Association,"
	Kyoto, Japan
2010	The Best Poster, "The 2nd Biosupercomputing Symposium," Tokyo, Japan
2008	Presentation Award, "Bio-rad Forum of Graduate Students," Peking University, China
1999–2000	General Scholarship, Hunan University, China

#### **PROFESSIONAL AFFILIATIONS**

2018-	American Chemical Society
2009-	Biophysical Society
2010-2012	Biophysical Society of Japan

## **SKILLS**

OS Linux, Windows

Software Amber, Gromacs, AutoDock, Modeller, VMD, PyMol, etc.

Programming R, Bash, C/C++, FORTRAN, Python, Perl, Tcl

# **REVIEWER SERVICES**

#### Journals:

Nature Communications • Communications Chemistry (Nature Publishing Group)
BMC Bioinformatics • Proteins: Structure, Function, and Bioinformatics • PLoS ONE
The Journal of Physical Chemistry • Journal of Molecular Graphics and Modelling
Chemical Physics Letters • Computational Biology and Chemistry
Journal of Bioinformatics and Computational Biology

## Conferences:

2017's Scientific Computing Day, Atlanta 2018's Scientific Computing Day, Atlanta

#### **REFERENCES**

Available upon requests