CURRICULUM VITAE

Xin-Qiu YAO, Ph.D.

2011

2010

2008

2000

Georgia State University Department of Chemistry 29 Peachtree Center Ave NE Atlanta, GA 30303-2515, USA Cell: (734) 355-1268 E-Mail: xinqiu.yao@gmail.com Website: https://xinqyao.github.io

APPOINTMENTS		
05. 2017 – present	Postdoctoral Research Associate (The Hamelberg Lab), Dept Chemistry, Georgia State University	
09. 2012 – 04. 2017	Postdoctoral Research Fellow (The Grant Lab), Dept Computational Medicine & Bioinformatics, the University of Michigan	
09. 2008 – 07. 2012	Postdoctoral Research Fellow (The Takada Lab), Dept Biophysics, Grad School of Science, Kyoto University	
EDUCATION		
09. 2005 – 06. 2008	Ph.D., Biomechanics and Medical Engineering, College of Engineering, Peking University (Advisors: Zhen-Su She & Huaiqiu Zhu. Title of dissertation: "A complex systems approach to study of protein structures")	
09. 2002 – 06. 2005	Graduate student, Condensed Matter Physics, School of Physics, Peking University	
09. 1998 – 06. 2002	B.S. , Applied Physics (Minored in Japanese), Dept Applied Physics, Hunan University	
TRAINING		
07. 2002 – 07. 2008	RA, Center for Theoretical Biology (The She Lab), Peking University, Beijing, China	
07. 2003	Trainee, School of Parallel Computing, Chinese Academy of Science, Beijing, China	
10. 2004 – 12. 2004	Internship, Intel China Research Center, Beijing, China	
TEACHING AND MENTORING		
2003 (Fall) 2013 (Summer)	TA, Statistical Physics, School of Physics, Peking University Co-Mentor, Undergraduate Research Opportunity Program (UROP), the University of Michigan	
GRANTS		
2015	Co-I , "G Protein Activation Mechanisms" (Proposal for allocation of simulation time on Anton at Pittsburgh Supercomputing Center, Grant No. MCB150033P) [Awarded]	
HONORS AND AWARDS		

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Association," Kyoto, Japan

Tokyo, Japan

The Best Talk, "The 13th Annual Meeting of Chinese Life Science

The Best Poster, "The 2nd Biosupercomputing Symposium,"

General Presentation Award, "Bio-rad Forum of Graduate

General Scholarship, Hunan University, Changsha, China

Students," Peking University, Beijing, China

AFFILIATIONS AND SERVICES

Academic 2009 – present 2010 – 2012	Membership, Biophysical Society Membership, Biophysical Society of Japan
Non-academic 2009 – 2012 1999 – 2002 1998 – 1999	General Secretary, Chinese Scholars Association in Kyoto Student Committee, Dept. Applied Physics, Hunan University President Assistant, Philately Society, Hunan University

SKILLS

OS	Linux, Windows
Software	Amber, Gromacs, AutoDock, Modeller, VMD, PyMol, etc.
Programming	R, C/C++, FORTRAN, Bash, Perl, Python, Tcl, etc.
Language	Mandarin Chinese (native), English (fluent), Japanese (daily
	communication)

Expertise molecular dynamics, coarse-grained molecular modeling and simulation, normal mode analysis, free energy calculation (FEP, umbrella sampling, MM-GB/SA, LIE), homology modeling, molecular

docking, PCA, HMM, Bayesian network, dynamic Bayesian network,

neural network

REVIEWER

Journals:

Proteins, BMC Bioinformatics, PLoS ONE, J Mol Graph Model, J Phys Chem, J Bioinform Comput Biol, Comp Biol & Chem

Conferences:

2017's Scientific Computing Day, Atlanta

PUBLICATIONS

Peer-reviewed Journal Articles (*Joint-First Authors)

- Vu PJ*, Yao XQ*, Momin M, and Hamelberg D. Unraveling allosteric mechanisms of enzymatic catalysis with an evolutionary analysis of residue-residue contact dynamical changes. PNAS (under review)
- Yao XQ*, Jariwala S*, and Grant BJ. In silico prediction of HLA-associated drug hypersensitivity. In preparation.
- Yao XQ*, 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. [link]
- 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. [link]
- Yao XQ*, Malik RU*, Griggs NW, Skjaerven 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. [link]
- Yao XQ and Grant BJ. Domain-opening and dynamic coupling in the α-subunit of heterotrimeric G proteins. Biophys J (2013) 105:L08-L10. [link]
- 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. [link]

- 6. Yao XQ, 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 reported by public media: *Nikkei Business Daily, Nikkan Kogyo Shimbun*, and *Kyoto Shimbun*). [link]
- 7. Yao XQ and She ZS. Key residue-dominated protein folding dynamics. *Biochem Biophys Res Comm* (2008) 373:64-68. [link]
- 8. Yao XQ, Zhu HQ, and She ZS. A dynamic Bayesian network approach to protein secondary structure prediction. *BMC Bioinformatics* (2008) 9:49. [link]
- Li H, <u>Yao XQ</u>, and Grant BJ. Comparative structural dynamic analysis of GTPases. To be submitted.
- 9. Bouley R, Waldschmidt HV, Cato MC, Cannavo A, Song J, Cheung JY, <u>Yao XQ</u>, 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), in press. https://doi.org/10.1124/mol.117.110130.
- 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. [link]
- 11. Nwachukwu JC, Srinivasan S, Bruno NE, Nowak J, Wright NJ, Minutolo F, Rangarajan ES, Izard T, <u>Yao 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 anti-hormone therapies. *Cell Chem Biol* (2017) 24:35-45. [link]
- 12. Jariwala S, Skjærven L, **Yao XQ**, and Grant BJ. Investigating protein sequence-structure-dynamics relationships with Bio3D-web. *J Vis Exp* (2017) 125:e55640. [link]
- 13. 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. [link]
- 14. 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. [link]
- 15. Skjaerven L, <u>Yao XQ</u>, Scarabelli G, and Grant BJ. Integrating protein structural dynamics and evolutionary analysis with Bio3D. *BMC Bioinformatics* (2014) 15:399. [link]
- 16. 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. [link]
- 17. 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. [link]
- 18. 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. [link]
- 19. 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. [link]

Talks

- "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)
- "Functionally rotating mechanism of a multidrug transporter studied by coarse-grained simulations," The 13th Annual Meeting of Chinese Life Science Association, Kyoto, Japan (2011)
- (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

- 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, Skjaerven 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)
- Skjaerven 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)
- Skjaerven 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, Skjaerven 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, Skjaerven 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, Skjaerven 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)

References are available upon request.