BIOGRAPHICAL SKETCH

Provide the following information for the Senior/key personnel and other significant contributors. Follow this format for each person. **DO NOT EXCEED FIVE PAGES.**

NAME: Lee, David John

eRA COMMONS USER NAME (credential, e.g., agency login): DJL002

POSITION TITLE: Postdoctoral Researcher

EDUCATION/TRAINING (Begin with baccalaureate or other initial professional education, such as nursing, include postdoctoral training and residency training if applicable. Add/delete rows as necessary.)

INSTITUTION AND LOCATION	DEGREE (if applicable)	Start Date MM/YYYY	Completion Date MM/YYYY	FIELD OF STUDY
University of California, Santa Cruz	B.S.	09/2005	06/2009	Biology
University of California, Santa Cruz	B.S.	09/2005	06/2009	Chemistry
University of California, San Diego	M.S.	09/2010	06/2012	Chemistry
University of California, San Diego	Ph.D.	06/2012	12/2016	Chemistry
University of California, San Diego	Postdoctoral	12/2016	10/2017	Structural Biology
University of California, San Francisco	Postdoctoral	11/2017	Current	Structural Biology

A. Personal Statement

My philosophical approach to science aligns with my greater philosophical approach to life—I like to know how things work. I like to take things apart, and learn how they function. I have found great satisfaction in my graduate work exploring how modular synthases produce a multitude of natural products. Using small molecules to study protein structure and protein function is analogous to using a wrench to work on a small engine, and philosophically draws me to chemical and structural biology.

During my undergraduate years, I was introduced to organic chemistry. I performed a substrate study, and enjoyed pushing a simple reaction to its limits to learn what guiding principles dictated its applicability. Similarly I was tasked with producing a family of glucosamine analogs, ultimately for incorporation into biological systems. I was especially interested in this project, and it was my first true introduction to chemical biology.

During graduate school and a brief postdoctoral position, I expanded on this interest in Professor Michael Burkart's laboratory at UC San Diego. The Burkart laboratory studies carrier protein dependent modular synthases and synthetases using structural and biophysical approaches. Collaborating with Professor Stan Opella, I learned solution state protein NMR techniques, solving multiple NMR structures of several different carrier proteins. I truly enjoy mentoring, and trained three undergraduate researchers and multiple graduate students in protein NMR. Together, we used NMR approaches to reveal specific information about proteins—specific pulse sequences to study dynamics, others to measure distance constraints for structural calculations, etc. We found that the carrier protein's cargo had significant ramifications on the carrier protein's structure, likely a structural handle to direct molecules through the pathway by tuning reaction partner protein affinity; small molecules with significant effects on protein structure. We focused on the mechanisms of substrate delivery from the carrier protein to the appropriate reaction partner, with minor conformations proving to be crucially important but difficult to study.

Ultimately, interest in minor conformational states and allostery guided my search for post-doctoral laboratories. I am engaged in my continuing training in Prof. James Fraser's laboratory, with additional technical training in Prof. Yifan Cheng's laboratory and collaboration with Prof. Ian Seiple's synthetic chemistry laboratory. This synergistic set of labs provides the ideal complement of skills to learn and study conformational heterogeneity, structural motions, and how function arises from structure in human-health

relevant systems. CryoEM studies on inhibitor-bound ribosomes, with the specific intent to understand and counteract antimicrobial resistance, perfectly tailor my continued training in chemical and structural biology. I have continued to learn and teach within the positive and collaborative environment at UCSF, rapidly advancing through CryoEM training and now guiding my labmates through CryoEM data collection and processing techniques. Additionally, UC San Francisco provides many career development opportunities, both academic and industrial interactions, and leadership and mentorship opportunities.

B. Positions and Honors

Positions and Employment

2007–2007	Intern, Medicinal Chemistry, University of Utah
2010-2010	Intern, Synthetic Chemistry, Intel Labs, Santa Clara
2010-2014	Teaching Assistant, UC San Diego
2016-2017	Professional Consultant, Biochemistry, Jones Day
2016-2017	Postdoctoral Researcher, UC San Diego
2017-	Postdoctoral Researcher, UC San Francisco

Other Experience and Professional Memberships

2014–2017	Co-organizer, Natural Products Affinity Group, UC San Diego
2014–	Member, Royal Society of Chemistry

2015- Member, American Chemical Society

<u>Honors</u>

2007	American Heart As	sociation Summer	Research Fellowship
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- 2009 Dean's Award in Physical and Biological Sciences, UC Santa Cruz
- 2010 Harold Urey Award, UC San Diego
- 2014 Bruno Zimm Award, UC San Diego
- 2015 President's Dissertation Year Fellowship, UC San Diego

C. Contributions to Science

- 1. Organic synthesis efforts towards chiral glucosamine analogs and asymmetric reduction of α,β-unsaturated ketones: My early introduction to laboratory science and research, as an undergraduate, was predominantly through synthetic organic chemistry. As an American Heart Association Undergraduate fellow, I interned with Prof. Kuberan Balagurunathan at the University of Utah, synthesizing glucosamine analogs for future incorporation into heparan sulfate glycosaminoglycan chains. Heparan sulfate chains are crucial for intercellular communication and developmental processes, and we were attempting to modulate sulfation patterns of the chains through perturbation of the glucosamine monomers. After the end of the internship, I volunteered with Prof. Bakthan Singaram at the University of California, Santa Cruz, executing a substrate study to quantify and validate the efficacy of a chiral director, "Tar-B-NO2". Tar-B-NO2 facilitates asymmetric reduction of α,β-unsaturated ketones via directed hydride delivery. Prof. Singaram instilled in me the importance of applying science to benefit human health, ultimately culminating in my undergraduate thesis, "Efforts towards an enantioselective synthesis of the HIV inhibitor abacavir".
 - a. Kim J, Bruning J, Park KE, **Lee DJ**, Singaram B. Highly enantioselective and regioselective carbonyl reduction of cyclic α,β -unsaturated ketones using TarB-NO2 and sodium borohydride. *Organic Letters*. 2009; 11(19):4358–61.
- 2. Graduate work on Fatty Acid Synthases: Carrier protein structure During graduate school, I refocused my studies towards chemical biology and structural biology. Joining Prof. Michael Burkart's laboratory at the University of California, I began studying protein structure by NMR. I performed many studies of carrier proteins from carrier protein dependent modular synthases, including fatty acid synthases (FAS). Several studies of the bacterial FAS were carried out, with two primary focuses. Carrier proteins sequester and protect intermediates as they transport them from partner to partner, but the nature of sequestration is poorly understood. We characterized cargo-induced structural changes to the carrier protein. First, we turned to solution-state NMR methods, observing significant chemical shift

perturbations upon binding cargo corresponding to a global tightening of the carrier protein around the cargo. Additionally, we found that unnatural probes could be appropriately sequestered, encouraging us to explore covalent crosslinking by attaching modified cargo to carrier proteins.

- a. Ishikawa F, Haushalter RW, **Lee DJ**, Finzel K, Burkart MD. Sulfonyl 3-alkynyl pantetheinamides as mechanism-based cross-linkers of acyl carrier protein dehydratase. *Journal of the American Chemical Society*. 2013; 135(24):8846–9. PMCID: PMC3713789
- b. **Lee DJ***, Finzel K*, Burkart MD. Using modern tools to probe the structure-function relationship of fatty acid synthases. *ChemBioChem.* 2015; 16(4): 528–547. PMCID: PMC4545599
- 3. Graduate work on Fatty Acid Synthases: Protein-protein interactions The protein-protein interactions required for fatty acid biosynthesis were observed by exploiting mechanism-based crosslinking probes. These probes allowed us to covalently trap the transient carrier protein-partner protein interaction for crystallographic and solution-state NMR evaluation. This approach was applied specifically to the E. coli acyl carrier protein and a partner dehydratase, yielding the first structural observations of an interaction between two partners within a fatty acid biosynthetic pathway. Additionally, NMR was used to observe the non-crosslinked in vitro interaction by titration, with regions of the protein perturbed in these titration experiments matching well with the interacting residues identified crystallographically. Molecular Dynamics simulations, supported by Residual Dipolar Coupling measurements, was used to quantify the flexibility of the carrier protein before and during interaction with the partner dehydratase. Together, these efforts allowed conclusive mapping of the acyl carrier protein and dehydratase interaction, with significant implications in future engineering and inhibition efforts.
 - a. Sztain T, Patel A, **Lee DJ**, Davis T, McCammon JA, Burkart MD. One atom matters: modifying the thioester linkage affects structure of the acyl carrier protein. *Angewandte Chemie International Edition*. 2019, in press.
 - b. **Lee DJ***, Nguyen C*, Haushalter RW*, Markwick PRL, Bruegger J, Caldara-Festin G, Finzel K, Jackson DR, Ishikawa F, O'Dowd B, McCammon JA, Opella SJ, Tsai S-C, Burkart MD. Trapping the dynamic acyl carrier protein in fatty acid biosynthesis. *Nature*. 2014; 505(7483): 427–31. PMCID: PMC4437705
 - c. Beld J, **Lee DJ**, Burkart MD. Fatty acid biosynthesis revisited: structure eluctidation and metabolic engineering. *Molecular Biosystems*. 2015; 11(1): 38-59. PMCID: PMC4276719
- 4. Graduate work on Hybrid polyketide synthase/non-ribosomal peptide synthetases Several pyrrole containing hybrid PKS/NRPS systems were studied and structurally characterized to engineer cross-pathway activity, between carrier proteins of closely related synthetases and non-cognate adenylation enzymes. Specifically, the pyoluteorin and prodigiosin carrier proteins were structurally characterized using traditional NOE solution-state NMR methods. The pyoluteorin carrier protein was also characterized bearing cargo, demonstrating structurally, for the first time, sequestration of cargo in a hybrid PKS/NRPS system. Computational and mutagenesis efforts were employed to produce a mutant prodigiosin carrier protein that could be acted upon by pyoluteorin biosynthetic partner proteins. Together, these efforts advance and highlight engineering opportunities in these modular synthases and synthetases, ideally to eventually allow development of custom synthases.
 - a. Jaremko MJ, **Lee DJ**, Opella SJ, Burkart MD. Structure and substrate sequestration in the pyoluteorin type II peptidyl carrier protein PltL. *Journal of the American Chemical Society*. 2015; 137(36): 11546–9. PMCID: PMC4847951
 - b. Jaremko MJ, **Lee DJ**, Patel A, Winslow V, Opella SJ, McCammon JA, Burkart MD. Manipulating protein–protein interactions in nonribosomal peptide synthetase type II peptidyl carrier proteins. *Biochemistry*. 2017; 56(40): 5269–73. PMCID: PMC5873958
- 5. Graduate work on Polyketide Synthases Polyketide synthases, the secondary-metabolic relatives of fatty acid synthases, use similar proteins and pathways as fatty acid synthases but produce secondary metabolites by various alterations to tailoring and iterative chain extensions. Understanding the substrate recognition and sequestration of cargo is critical to the success of engineering efforts. Unfortunately, many polyketides are formed by repetitive elongation steps yielding an elongated, highly reactive polyketide that can spontaneously cyclize. Studying carrier protein sequestration in the biosynthesis of the antimicrobial polyketide actinorhodin required the preparation of atom-replaced geometrically and electronically similar substrate mimics. Both linear and cyclized atom-replaced

mimics were prepared. Subjecting these mimics to sequestration studies by solution-state protein NMR revealed that only the full-length and cyclized mimics were well sequestered, suggesting that the elongating polyketide remains within the partner ketosynthase until full elongation.

- a. **Lee DJ***, Milligan JC*, Jackson DR*, Schaub AJ, Beld J, Barajas JF, Hale JJ, Luo R, Burkart MD, Tsai S-C. Molecular basis for interactions between an acyl carrier protein and a ketosynthase. *Nature Chemical Biology*. 2019; 15(7): 669-71. PMID: 31209348
- b. Shakya G, Rivera H, **Lee DJ**, Jaremko MJ, La Clair JJ, Fox DT, Haushalter RW, Schaub AJ, Bruegger J, Barajas JF, White AR, Kaur P, Gwozdziowski ER, Wong F, Tsai S-C, Burkart MD. Modeling linear and cyclic PKS intermediates through atom replacement. *Journal of the American Chemical Society*. 2014; 136(48): 16792–9. PMCID: PMC4277753

Complete List of Published Work in My Bibliography:

https://www.ncbi.nlm.nih.gov/sites/myncbi/david.lee.7/bibliography/57262207/public/?sort=date

D. Additional Information: Research Support and/or Scholastic Performance

YEAR	COURSE TITLE	GRADE
	University of California, San Diego	
2010	Enzyme Catalyzed Reactions	Α
2010	Synthetic Methods/Organic Chemistry	A-
2010	Mechanisms/Organic Reactions	B-
2011	Synthesis of Complex Molecules	B+
2011	Structure and Properties of Organic Molecules	Α
2011	Natural Products Chemistry	Α
2011	Applied Spectroscopy	B+
2012	Protein NMR	Α

Biographical Sketch and Relevant Publications

NAME Vanja Stojkovic	POSITION Postdocto		
EDUCATION/TRAINING (Begin with baccala	ureate or othe	er initial profess	sional education,
INICTITUTION AND LOCATION	ב	VEADO	FIELD OF OTLIDY

EDUCATION TRAINING (Begin with baccalaureate of other Initial professional education,			
INSTITUTION AND LOCATION	DEGREE	YEAR(s)	FIELD OF STUDY
Northland College	B.S.	2001-2005	Chemistry
University of Iowa	Ph.D.	2005-2012	Chemistry
University of California, San Francisco	Postdoc.	2013-	Biochemistry

A. Positions and Honors.

Positions and Employment

2004	Research Fellowship at University of Arkansas, Advisor: Robert Gawley
2005-2007	Graduate Student at University of Iowa, Advisor: Sonya Franklin
2007-2012	Graduate Student at University of Iowa, Advisor: Amnon Kohen
2013-present	Postdoctoral researcher at University of California, San Francisco

Academic Honors

2002-2004	Scholarship for sciences and engineering, Northland College, 2002-2004
2005	ACS regional award
2008	University of Iowa Graduate College Summer Fellowship
2008	Outstanding Teaching Award, Department of Chemistry, University of Iowa
2011	ACS Biological Chemistry Travel Award, ACS national meeting, Denver, CO
2012	University of Iowa Lynn Anderson Award for Research Excellence
2012	University of Iowa Strategic Initiative Fund Fellowship
2014	EMBO Short-term Fellowship
2017	PBBR Postdoctoral Research Grant

B. Peer-reviewed publications.

- 1. **Stojković, V.**, Chu, T., Therizols, G., Weinberg, D.E., Fujimori, D.G. miCLIP-MaPseq, a Substrate Identification Approach for Radical SAM RNA Methylating Enzymes. *J Am Chem Soc.* **2018**, 140(23), 7135-7143.
- 2. **Stojković, V.,** Garcia-Noda, L., Tawfik, D.S., Fujimori, D.G. Antibiotic resistance evolved via inactivation of a ribosomal RNA methylating enzyme. *Nucleic Acids Res.*, **2016**, 44(18), 8897-8907.
- 3. **Stojković**, **V.**, Fujimori, D.G. Radical SAM-mediated methylation of ribosomal RNA. *Methods Enzymol.*, **2015**, 560, 355-376.
- 4. Doron, D., **Stojković**, **V**., Gakhar, L., Vardi-Kilshtain, A., Kohen, A., Major, D.T. Free energy simulations of active-site mutants of dihydrofolate reductase. *J. Phys. Chem. B.*, **2015**, 119(3), 906-916.
- 5. Francis, K.,* **Stojković, V**.,* Kohen, A. Preservation of protein dynamics in dihydrofolate reductase evolution. *J. Biol. Chem.* **2013**, 288(50), 35961-35968. *Contributed equally
- 6. Sen, A.,* **Stojković, V**.,* Kohen, A. Synthesis of radiolabeled nicotinamide cofactors from labeled pyridines: versatile probes for enzyme kinetics. *Anal. Biochem.*, **2012**, 430(2), 123-129.
 - *Contributed equally

- 7. **Stojković, V.**, Perissinotti, L.L., Willmer, D., Benkovic, S.J., Kohen, A. Effects of the donor acceptor distance and dynamics on hydride tunneling in the dihydrofolate reductase catalyzed reaction. *J Am Chem Soc*, **2012**, 134(3), 1738-1745.
- 8. **Stojković, V**., Perissinotti, L.L., Lee, J., Benkovic, S.J., Kohen, A. The effect of active-site isoleucine to alanine mutation on the DHFR catalyzed hydride-transfer. *Chem Comm*, **2010**, 46(47), 8974-8976.

C. Research support

Postdoctoral Research Grant 06/30/2018

Stojkovic (PI)

7/01/2017 -

UCSF Program for Breakthrough Biomedical Research

The proposed research aims to develop an RNA-seq method, based on miCLIP-seq, to identify *in vivo* substrates for any member of radical-SAM RNA methylating enzyme family. This family contains several enzymes that have been implicated in antibiotic resistance to several important classes of clinically used antibiotics.

BIOGRAPHICAL SKETCH

NAME: Fujimori, Danica Galonić

era commons user name: **Danica_Galonic**

POSITION TITLE: Professor of Cellular and Molecular Pharmacology and Pharmaceutical Chemistry

EDUCATION/TRAINING

INSTITUTION AND LOCATION	DEGREE	Completion Date	FIELD OF STUDY
University of Belgrade, Belgrade, Serbia	BSc	07/2000	Chemistry
University of Illinois, Urbana, IL	PhD	05/2005	Chemistry
Harvard Medical School, Boston, MA	Postdoc	06/2008	Biochemistry

A. Personal Statement

My research focuses on mechanisms, regulation, and biological function of post-transcriptional and post-translational modifications. In the area of protein methylation, we are investigating histone demethylases, a class of epigenetic eraser proteins that antagonize chromatin methylation. While it is well established that activities of these enzymes are regulated by chromatin environment and metabolic state of the cell, mechanisms by which this is achieved are poorly understood. Our work on defining the functional cross-talk between chromatin recognition and demethylation led to identification of new mechanisms by which chromatin reader domains modulate catalytic activity of histone demethylases. We uncovered a positive feedback regulation in demethylase KDM5A, enabled by allosteric communication between a chromatin reader domain and the catalytic domain in this enzyme. Furthermore, we are developing chemical probes for histone demethylases with a goal of utilizing these molecules for pharmacological target validation.

In area of infectious diseases, we investigate how modifications of the bacterial ribosome confer resistance to ribosome targeting antibiotics, using enzymology, microbiology and structural biology approaches. One such resistance mechanism is enabled by methylation of the ribosome by RNA methylating enzyme Cfr.

Our work combines biochemical reconstitution, mechanistic enzymology, chemical synthesis and cell biology, which provides an ideal multidisciplinary training ground for graduate students and postdoctoral scholars. My background combines chemical synthesis and mechanistic enzymology. I received a Ph.D. in organic chemistry from the University of Illinois at Urbana-Champaign in 2005, where I worked on chemical synthesis under the direction of David Gin and Wilfred van der Donk. As a Damon Runyon Cancer Research Foundation postdoctoral fellow in the lab of Chris Walsh lab at Harvard Medical School I gained expertize in mechanistic enzymology of complex systems.

Significant publications:

- a. Ortiz Torres I, Kuchenbecker KM, Nnadi CI, Fletterick RJ, Kelly MJS, **Fujimori DG.** Histone Demethylase KDM5A is Regulated by its Reader Domain Through a Positive-Feedback Mechanism. *Nat Commun* 6: 6204 2015. PMID 25686748, PMCID: PMC5062987.
- b. Longbotham JE, Chio CM, Dharmarajan V, Trnka MJ, Torres I, Goswami D, Ruiz K, Burlingame AL, Griffin PR, **Fujimori DG**. Histone H3 binding to the PHD1 domain of histone demethylase KDM5A enables active site remodeling, Nature Communications, 10: 94, 2019.
- c. Korczynska M, Le DD, Younger N, Gregori-Puigjané E, Tumber A, Krojer T, Velupillai S, Gileadi C, Nowak RP, Iwasa E, Pollock SB, Ortiz Torres I, Oppermann U, Shoichet BK, **Fujimori DG**. Docking and Linking of Fragments to Discover Jumonji Histone Demethylase Inhibitors. *J Med Chem*, 59: 1580-98, 2016. PMCID: PMC5080985.

B. Positions and Honors

Positions and Employment:

2005-2008	Postdoctoral Fellow, Department of Biological Chemistry and Molecular Pharmacology, Harvard
	Medical School
2008-2014	Assistant Professor, Departments of Cellular and Molecular Pharmacology & Pharmaceutical
	Chemistry, University of California San Francisco
2013-present	Member, Hellen Diller Family Comprehensive Cancer Center, University of California San
	Francisco
2014-2018	Associate Professor, Departments of Cellular and Molecular Pharmacology & Pharmaceutical
	Chemistry, University of California San Francisco
2018-present	Professor, Departments of Cellular and Molecular Pharmacology & Pharmaceutical Chemistry,
•	University of California San Francisco
2018-present	Associate Director, Chemistry and Chemical Biology Graduate Program, University of California
•	San Francisco

Other Experience and Professional Memberships:

2011	Ad hoc reviewer, NIH K99/R00
2011-Pres	Grant review panelist and ad hoc reviewer, NSF
2012	Ad hoc reviewer, NIH MSFE study section
2015-2016	Ad hoc reviewer, NIH SBCB study section
2017-Pres	Member, NIH SBCB study section

Honors and Awards:

Damon Runyon Cancer Research Foundation Postdoctoral Fellowship
NIH Pathway to Independence Award
Sandler Program in Basic Sciences Opportunity Award
Kimmel Scholar Award
V Foundation Scholar Award
Basil O'Connor Starter Scholar Research Award
NSF Career Award
Searle Scholar Award
PBBR New Frontier Research Award
UCSF Haile T. Debas Academy of Medical Educators Excellence in Teaching Award
Chauncey D. Leake Lectureship in Cellular and Molecular Pharmacology
Raymond and Beverly Sackler Sabbatical Exchange Program Award, UC Berkeley
Byers Award, UCSF

C. Contribution to Science

- 1. Histone demethylases: Histone demethylases are a class of epigenetic eraser proteins. These enzymes antagonize lysine methylation in chromatin. Our work is centered on understanding the regulation of these enzymes, elucidating their roles in disease and developing pharmacological tools to probe the cellular functions of demethylases. We have uncovered previously unknown mechanistic links by which chromatin context impact demethylation catalysis. Our most significant contribution in this area is the discovery of allosteric regulation in KDM5A, an oncogenic demethylase. We determined that this protein is regulation by a positive feedback mechanism, where binding of the demethylation product to a reader domain within the demethylase stimulates demethylation activity. I served as a primary investigator on the following studies:
 - a. Shiau C, Trnka MJ, Bozicevic A, Ortiz Torres I, Al-Sady B, Burlingame AL, Narlikar GJ, Fujimori DG. Reconstitution of Nucleosome Demethylation and Catalytic Properties of a Jumonji Histone Demethylase. Chem Biol 20: 494-9, 2013. PMCID: PMC3704229
 - b. Ortiz Torres I, Kuchenbecker KM, Nnadi CI, Fletterick RJ, Kelly MJS and **Fujimori DG**. Histone Demethylase KDM5A is Regulated by its Reader Domain Through a Positive-Feedback Mechanism. *Nature Commun* 6: 6204. 2015. PMID 25686748, PMCID: PMC5080983.
 - c. Pack LR, Yamamoto KR, **Fujimori DG**. Opposing chromatin signals direct and regulate the demethylase activity of KDM4C. *J Biol Chem*, 291: 6060-6070, 2016. PMCID: PMC4813556

- d. Longbotham JE, Chio CM, Dharmarajan V, Trnka MJ, Torres I, Goswami D, Ruiz K, Burlingame AL, Griffin PR, **Fujimori DG**. Histone H3 binding to the PHD1 domain of histone demethylase KDM5A enables active site remodeling, Nature Communications, 10: 94, 2019.
- 2. Tools for epigenetics and chromatin: Our lab has developed and/or advanced several methods to study chromatin modifications by relying on chemoselective protein modification strategies. In addition, we have advanced the use of existing methods to elucidate the functions of unknown domains that interact with chromatin. Furthermore, we have developed highly potent small molecule inhibitors for demethylases. I served as a primary investigator or as co-investigator on the following studies:
 - a. Le DD, Cortesi A, Myers SA, Burlingame AL, **Fujimori DG**. Site- and Regiospecific Installation of Methylarginine Analogs into Recombinant Histones and Insights into Effector Protein Binding. *J Am Chem Soc* 135: 2879-2882, 2013. PMCID: PMC4260808
 - b. Dumesic PA, Homer CM, Moresco JJ, Pack LR, Coyle SM, Strahl BD, **Fujimori DG**, Yates III JR, Madhani HD. Product binding enforces the genomic specificity of a yeast polycomb repressive complex. *Cell* 160: 204-218, 2015. PMCID: PMC4303595
 - c. Korczynska M, Le DD, Younger N, Gregori-Puigjané E, Tumber A, Krojer T, Velupillai S, Gileadi C, Nowak RP, Iwasa E, Pollock SB, Ortiz Torres I, Oppermann U, Shoichet BK, **Fujimori DG**. Docking and Linking of Fragments to Discover Jumonji Histone Demethylase Inhibitors. *J Med Chem*, 59: 1580-98, 2016. PMCID: PMC5080985.
- 3. RNA methylation and its role in antibiotic resistance: Modification of the peptidyltransferase center of the bacterial ribosome by Radical SAM enzymes RlmN and Cfr is mechanistically unique. In contrast to a majority of biological methylation substrates, which are electron rich, methylation substrates of these enzymes are electron poor. My lab has contributed to the discovery of a novel mechanism that these enzymes use to carry out methylation. Our work demonstrated that these enzymes, so-called Radical SAM methyl synthases, have several key mechanistic features, such as the formation of a unique covalent intermediate between the enzyme and the substrate as well as the ability to utilize S-adenosylmethionine both in homolyitc and in heterolytic fashion. Enzymatic methylations performed by these enzymes have important roles in the regulation of antibiotic susceptibility. I served as a primary investigator on the following studies:
 - a. Yan, F, Fujimori, DG. RNA Methylation by Radical SAM Enzymes RlmN and Cfr Proceeds via Methylene Transfer and Hydride Shift. Proc Natl Acad Sci USA 108: 3930-34, 2011. PMCID: PMC3054002
 - b. McCusker KP, Medzihradszky KF, Shiver AL, Nichols RJ, Yan F, Maltby DA, Gross CA, Fujimori DG. Covalent Intermediate in the Catalytic Mechanism of the Radical SAM Methyl Synthase RlmN Trapped by Mutagenesis. *J Am Chem Soc* 134: 18074-81, 2012. PMCID: PMC3499099
 - c. Stojkovic V, Noda-Garcia L, Tawfik DS, **Fujimori DG**. Antibiotic Resistance Evolved via Inactivation of a Ribosomal RNA Methylating Enzyme, *Nucl Acids Res*, Aug 5. 2016. PMCID: PMC5062987.
 - d. Stojkovic V, Chu T, Therizols G, Weinberg DE, **Fujimori DG**. miCLIP-MaPseq, a substrate identification approach for radical SAM RNA methylating enzymes. *J Am Chem Soc* 140: 7135-7143, 2018.

Complete List of Published Work in MyBibliography:

http://www.ncbi.nlm.nih.gov/sites/myncbi/danica.fujimori.1/bibliography/41758140/public/?sort=date&direction=ascending

D. Research Support

Ongoing Research Support

R01 GM114044 Fujimori (PI) 09/15/2015 - 08/31/2019 NIH/NIGMS

Allosteric Regulation in the KDM5 Family of Histone Demethylases

In this proposal are aiming to elucidate the mechanistic basis and the functional significance of allosteric regulation of histone demethylation catalyzed by the members of the KDM5 subfamily of jumonji histone demethylases.

New Frontiers Research Award

Fujimori (PI)

06/15/2018 - 06/14/2019

UCSF Program for Breakthrough Biomedical Research

Disrupting Transcription through Targeting of Chromatin Methylation Readers

This award supports development of chemical probes that target epigenetic reader domains.

R01 Al137270 Fujimori (PI)

09/14/2018 - 08/31/2022

NIH/NIGMS

Radical SAM-dependent Methylation in Antibiotic Resistance

In this application we aim to investigate, using biochemical and structural studies, how aberrant methylation of the peptidyl transferase center of the ribosome impacts antibiotic susceptibility.

OVERLAP:

None.

Completed Research Support:

R01 Al095393 (Fujimori PI) NIH/NIAID 06/15/2011 - 05/31/2017

(NCE)

Radical SAM Methyltransferases

The proposed research aims to elucidate the mechanism of modification of ribosomal RNA by methyltransferases RlmN and Cfr. This modification renders bacteria resistant to several important classes of clinically used antibiotics, and its mechanistic understanding could lead to the development of new treatments for multi-drug resistant pathogens.

Catalyst Award Fujimori (PI) 07/01/2015 - 06/30/2016

UCSF Clinical & Translational Science Institute

Targeting Oncogenic KDM4 Demethylases

In this application we propose to expand our efforts to develop potent, selective and cell active inhibitors of histone demethylases through a combination of computational docking, chemical synthesis and activity assays.

Early Career Development Award

Fujimori (PI)

01/15/2011 - 12/31/2015

National Science Foundation

Award ID: 1056143

A Chemical Approach to Elucidate the Mechanism of Radical SAM Methyltransferases

This early career development award offered general support of our research and teaching efforts, and provides funds for helping students from underrepresented groups spend a summer doing research.

New Frontier Research Award

Fujimori (PI)

05/01/2014 - 04/30/2015

UCSF Program for Breakthrough Biomedical Research Regulation of Propagation in Chromatin Demethylation

07/31/2015 (NCE)

The aim of this project was to define a molecular mechanism of spread of chemical signals that determine whether a gene is active or silent, determine the molecular mechanism of cross-talk between the reader and the catalytic domain in the histone demethylase KDM5A and elucidate the role of this cross-talk role in the propagation of chromatin demethylation.

Searle Scholar Award

Fujimori (PI)

07/01/2011 - 06/30/2014

Kinship Foundation Grant No. 11-SSP-157

Histone Demethylases in Cellular Regulation

The aim in this work was to identify substrates and develop chemical tools to help decipher the histone code as it related to dynamic changes in methylation caused by Jumonji C domain-containing histone demethylases.

BIOGRAPHICAL SKETCH

Provide the following information for the Senior/key personnel and other significant contributors. Follow this format for each person. **DO NOT EXCEED FIVE PAGES.**

NAME: Fraser, James Solomon

eRA COMMONS USER NAME (credential, e.g., agency login): FRASERJA

POSITION TITLE: Associate Professor of Bioengineering and Therapeutic Sciences

EDUCATION/TRAINING (Begin with baccalaureate or other initial professional education, such as nursing, include postdoctoral training and residency training if applicable. Add/delete rows as necessary.)

INSTITUTION AND LOCATION	DEGREE (if applicable)	Completion Date MM/YYYY	FIELD OF STUDY
McGill University, Montreal, QC, Canada	B.Sc.	05/2005	Biology
University of California, Berkeley, CA	Ph.D.	12/2010	Molecular and Cell Biology

Personal Statement

The long-term goals of our research are to understand how protein conformational ensembles are reshaped by perturbations and to quantify how these perturbations impact protein function and organismal fitness. To accomplish these goals, we create new computational and biophysical approaches to study how proteins move between different conformational states. A primary guiding principle of our research is that physical perturbations (such as temperature or pressure) can often reveal the same "hidden" conformations exploited by biochemical perturbations (such as ligand binding or mutation). In 2011, I started my independent research career as a QB3 at UCSF Fellow and in 2013 was appointed as an Assistant Professor of Bioengineering and Therapeutic Sciences, with promotion to Associate Professor with tenure in 2016. We are best known for creating multitemperature X-ray data collection approaches, which are especially powerful when paired with multiconformer computational modeling to reveal otherwise inaccessible attributes of conformational ensembles. My group has also pioneered methods to model and evaluate the data emerging from the "resolution revolution" in cryo-electron microscopy. I took advantage of a Sackler Sabbatical Fellowship at UC Berkeley (host: Eva Nogales) to immerse myself in the practical aspects of electron microscopy data collection and processing. I have brought this new perspective back to UCSF, as my group integrates high resolution EM, X-ray, NMR, and computation to improve protein engineering and small molecule discovery. I care deeply about mentorship: my first two postdoctoral fellows are on their desired career trajectories (Daniel Keedy, now Assistant Professor at CCNY; Brandi Hudson, now Scientist at Relay Therapeutics); my first three graduate students have moved onto postdoctoral training (Rahel Woldeyes, with Wah Chiu at Stanford; David Mavor, with Dan Bolon at UMass) or launched successful careers (Andrew van Benschoten is a data scientist at Oracle). In addition, I have incorporated new technologies in teaching, establishing a deep sequencing-based project course that published multiple papers with student co-authors and a new "Methods in Structural Biology" class, with EM, X-ray, and NMR data collection at UCSF or the Advanced Light Source at LBNL.

Key Citations

- 1. Keedy DA*, Hill ZB*, Biel JT, Kang E, Rettenmaier TJ, Brandao-Neto J, Pearce NM, von Delft F, Wells JA, **Fraser JS**. An expanded allosteric network in PTP1B by multitemperature crystallography, fragment screening, and covalent tethering. *eLife*. 2018. PMCID: PMC6039181.
- 2. Otten R*, Liu L*, Kenner LR, Clarkson MW, Mavor D, Tawfik DS, Kern D, **Fraser JS.** Rescue of conformational dynamics in enzyme catalysis by directed evolution. *Nature Communications*. 2018. PMCID: PMC5883053.
- 3. Barad BA, Echols N, Wang RY, Cheng Y, DiMaio F, Adams PD, **Fraser JS**. EMRinger: Side-chain-directed model and map validation for 3D Electron Cryomicroscopy. *Nature Methods*. 2015. PMCID: PMC4589481.
- 4. Fischer M, Coleman RG, **Fraser JS**, Shoichet BK. Incorporation of protein flexibility and conformational energy penalties in docking screens to improve ligand discovery. *Nature Chemistry*. 2014. PMCID: PMC4144196.

A. Positions and Honors

Positions and Employment

2011-2012	QB3 at UCSF Faculty Fellow (P	rincipal Investigator)

Department of Cellular and Molecular Pharmacology, UCSF

California Institute of Quantitative Biosciences (QB3)

2013-2016 Assistant Professor

Department of Bioengineering and Therapeutic Sciences, UCSF

California Institute of Quantitative Biosciences (QB3)

2016 - Consulting Professor

Department of Photon Science

SLAC National Accelerator Laboratory

2016 - Associate Professor

Department of Bioengineering and Therapeutic Sciences, UCSF

California Institute of Quantitative Biosciences (QB3)

2018 - Faculty Scientist

Molecular Biophysics and Integrated Bioimaging Division

Lawrence Berkeley National Lab

Other Experience

Other Expen	ence
2007 -	Author of problems/solutions manual for physical biochemistry textbook "The Molecules of Life"
	(Garland Science, Authors: John Kuriyan, Boyana Konforti, David Wemmer)
2008-2009	Assistant to Professor Howard Schachman for NIH Ethics Training (MCB 293C)
2013-2015	Advanced Light Source Proposal Review (Structural Biology), Panel Member
2015-2018	Linac Coherent Light Source (XFEL) Proposal Review Panel (BIO-C), Chair
2016-	Beamline 8.3.1. at the Advanced Light Source, Head of Participating Research Team
2016-	ASAPbio (Accelerating Science and Publication in biology) Board of Directors, Treasurer
2016-	Relay Therapeutics, Consultant
2017-	Quantitative Biosciences Institute of UCSF, Associate Director
2017-	ALS-ENABLE P30 Resource, Deputy Director
2017-	Collaboration for Structural Simulations and Scattering, Project Director
2018	Protein Society Annual Symposium, Co-Chair
2018-	PHENIX (Python-based Hierarchical Environment for Integrated Xtallography), Advisory Board
Honors	
2001-2005	Canadian Millennium Excellence Undergraduate Scholarship
2004	NSERC Undergraduate Summer Research Award (Mentor: Alan Davidson)
2006-2007	Natural Sciences and Engineering Research Council (Canada) Postgraduate Fellowship
2007-2010	Natural Sciences and Engineering Research Council (Canada) Doctoral Fellowship
2007-2010	National Science Foundation Graduate Research Fellowship
2010	EMBO Short Term Fellowship (Host: Dan Tawfik, Weizmann Institute, Israel)
2010	Warren DeLano Award for Structural Bioinformatics and Computational Biophysics
2011	Nicholas Cozzarelli Prize for Best Dissertation in Molecular and Cell Biology (UCB)
2011	Forbes 30 under 30 Science
2014	Searle Scholar, Kinship Foundation
2014	Pew Scholar, Pew Charitable Trusts
2014	Packard Fellow, The David and Lucile Packard Foundation

B. Contributions to Science

2017-2018

1. Identifying hidden alternative conformations of proteins in biophysical data. We study proteins as conformational ensembles. Although X-ray crystallography is an ensemble experiment, the results are typically summarized with a single static structure. As a graduate student, and now in my own lab, we have developed software to discover the structural ensembles present in the crystal. The ensemble nature of proteins highlighted by this work feeds into all of our mechanistic studies that interpret the functional effects of mutations, that characterize designed and artificially-evolved proteins, or that seek to modulate protein function with small molecules. We are expanding this direction to include modeling and validating protein structural data generated by cryoelectron microscopy (using EMRinger and ensemble modeling) and through integrative approaches to discover cryptic sites.

UCSF/Berkeley Sabbatical Exchange (Host: Eva Nogales)

- a. van den Bedem H, Bhabha G, Yang K, Wright PE, Fraser JS. Automated identification of functional dynamic contact networks from X-ray crystallography. *Nature Methods*. 2013. PMCID: PMC3760795.
- b. Eshun-Wilson L, Zhang R, Portran D, Toso D, Lohr T, Vendruscolo M, Bonomi M, **Fraser JS**, Nogales E. Effects of α-tubulin acetylation on microtubule structure and stability. Submitted Preprint on BioRxiv. 2019. http://dx.doi.org/10.1101/516591
- c. van Zundert GCP*, Hudson BM*, Oliveira SHP, Keedy DA, Fonseca R, Heliou A, Suresh P, Borrelli K, Day T, **Fraser JS**, van den Bedem H. qFit-ligand reveals widespread conformational heterogeneity of drug-like molecules in X-ray electron density maps. J Med Chem. 2018. PMCID: In process by journal.
- d. **Fraser JS**, Clarkson MW, Degnan SC, Erion R, Kern D, Alber T. Hidden alternative structures of proline isomerase essential for catalysis. *Nature*. 2009; 462(7273):669-73. PMCID: PMC2805857.
- 2. Creating multi-temperature X-ray data collection methods to inform mechanistic studies. We recognized that the standard practice of cryocooling crystals could distort protein conformations. In both larger surveys and isolated mechanistic studies, we have demonstrated the value of room temperature data collection for revealing the structural basis of protein conformational dynamics, leading to new insights into the enzymes PTP1B, CypA, H-Ras, and DHFR, and increasing connections to dynamics studies from NMR and simulations. Additionally, we have identified how temperature can bias small molecule discovery, leading some fragment sites inaccessible at cryogenic temperatures, and the positioning of crucial water molecules in the flu ion channel M2.
 - a. **Fraser JS**, van den Bedem H, Samelson AJ, Lang PT, Holton JM, Echols N, Alber T. Accessing protein conformational ensembles by room-temperature X-ray crystallography. *Proceedings of the National Academy of Sciences*. 2011. PMCID: PMC3182744.
 - b. Thomaston JL, Alfonso-Prieto M, Woldeyes RA, **Fraser JS**, Klein ML, Fiorin G, DeGrado WF. High-resolution structures of the M2 channel from influenza A virus reveal dynamic pathways for proton stabilization and transduction. *Proceedings of the National Academy of Sciences*. 2015. PMCID: PMC4655559.
 - c. Biel JT, Thompson MC, Cunningham CN, Corn JE, **Fraser JS.** Flexibility and design: conformational heterogeneity along the evolutionary trajectory of a redesigned ubiquitin. *Structure*. 2017. PMCID: PMC5415430.
 - d. Keedy DA*, Kenner LR*, Warkentin M*, Woldeyes RA*, Thompson MC, Brewster AS, Van Benschoten AH, Baxter EL, Hopkins JB, Uervirojnangkoorn M, McPhillips SE, Song J, Alonso-Mori R, Holton JM, Weis WI, Brunger AT, Soltis SM, Lemke H, Gonzalez A, Sauter NK, Cohen AE, van den Bedem H, Thorne RE, **Fraser JS**. Mapping the Conformational Landscape of a Dynamic Enzyme by XFEL and Multitemperature Crystallography. *eLife*. 2015. PMCID: PMC4721965.
- 3. **Developing new X-ray diffuse scattering and X-FEL experiments to probe correlated motions in proteins**. A major limitation of most biophysical techniques is the inability to directly reveal correlations in motions between distinct regions of macromolecules. Diffuse scattering has the potential to reveal these motions; however, we currently lack the ability to collect, integrate, and refine diffuse scattering data. We are tackling each of these problems directly with collaborators: Michael Wall, Nicholas Sauter, Tom Terwilliger, and Paul Adams. Our long-term goal is to increase the information content of every X-ray diffraction experiment to reveal atomic level coupling at high resolution and improved models of grouped flexibility at low resolution. We are also taking advantage of the new capabilities of next-generation X-ray free electron laser (X-FEL) light sources to perform radiation damage-free imaging of proteins and to watch how protein ensembles respond when perturbed by rapid temperature jumps using the X-FEL.
 - a. Thompson MC, Barad BA, Wolff AM, Cho HS, Schotte F, Schwarz DMC, Anfinrud P, **Fraser JS**. Temperature-Jump Solution X-ray Scattering Reveals Distinct Motions in a Dynamic Enzyme. Submitted Preprint on BioRxiv. 2018. http://dx.doi.org/10.1101/476432
 - b. Van Benschoten AH, Liu L, Gonzalez A, Brewster AS, Sauter NK, **Fraser JS**, Wall ME. Measuring and modeling diffuse scattering in protein X-ray crystallography. *Proceedings of the National Academy of Sciences*. 2016. PMCID: PMC4839442.
 - c. Wall ME, Van Benschoten AH, Sauter NK, Adams PD, **Fraser JS**, Terwilliger TC. Conformational dynamics of a crystalline protein from microsecond-scale molecular dynamics simulations and diffuse X-ray scattering. *Proceedings of the National Academy of Sciences*. 2014. PMCID: PMC4273327.

- d. Thomaston JL, Woldeyes RA, Nakane T, Yamashita A, Tanaka T, Koiwai K, Brewster AS, Barad BA, Chen Y, Lemmin T, Uervirojnangkoorn M, Arima T, Kobayashi J, Masuda T, Suzuki M, Sugahara M, Sauter NK, Tanaka R, Nureki O, Tono K, Joti Y, Nango E, Iwata S, Yumoto F, Fraser JS, DeGrado WF. XFEL structures of the influenza M2 proton channel: Room temperature water networks and insights into proton conduction. *Proceedings of the National Academy of Sciences*. 2017. PMCID: PMC5754760
- 4. Identifying unifying concepts between systems and structural biology. With Nevan Krogan, we have articulated the similarities in genetic epistasis and thermodynamic measurements and applied these insights to large-scale studies of point mutants and posttranslational modifications. This framework forms the basis for the UCSF graduate course that I direct, PUBS (Physical Underpinnings of Biological Systems), which uses deep sequencing to determine the context dependence of fitness effects of mutations. The class is taught through project-based learning where incoming students perform all library preparations, load samples directly on the MiSeq, and write all their own code to process sequencing data.
 - Beltrao P, Albanèse V, Kenner LR, Swaney DL, Burlingame A, Villén J, Lim WA, Fraser JS, Frydman J, Krogan NJ. Systematic functional prioritization of protein posttranslational modifications. Cell. 2012. PMCID: PMC3404735
 - b. Braberg H, Jin H, Moehle EA, Chan YA, Wang S, Shales M, Benschop JJ, Morris JH, Qiu C, Hu F, Tang LK, Fraser JS, Holstege FC, Hieter P, Guthrie C, Kaplan CD, Krogan NJ. From structure to systems: high-resolution, quantitative genetic analysis of RNA polymerase II. *Cell.* 2013. PMCID: PMC3932829
 - c. **Fraser JS**, Gross JD, Krogan NJ. From systems to structure: bridging networks and mechanism. *Mol Cell*. 2013. PMCID: PMC3558917
 - d. Mavor D, Barlow KA, Thompson S, Barad BA, Bonny AR, Cario CL, Gaskins G, Liu Z, Deming L, Axen SD, Caceres E, Chen W, Cuesta A, Gate R, Green EM, Hulce KR, Ji W, Kenner LR, Mensa B, Morinishi LS, Moss SM, Mravic M, Muir RK, Niekamp S, Nnadi CI, Palovcak E, Poss EM, Ross TD, Salcedo E, See S, Subramaniam M, Wong AW, Li J, Thorn KS, Conchúir SÓ, Roscoe BP, Chow ED, DeRisi JL, Kortemme T, Bolon DN, Fraser JS. Determination of Ubiquitin Fitness Landscapes Under Different Chemical Stresses in a Classroom Setting. *eLife*. 2016. PMCID: PMC4862753
- 5. Determining structures of protein mediating microbial-host interactions. I have a longstanding interest in microbiology, beginning from my undergraduate work with Alan Davidson (Toronto) on bacteriophage structure prediction that lead to the surprising discovery of a class of mobile immunoglobulin domains. I have collaborated with the Zusman lab (UC Berkeley) to determine the structure of FrzS, a key signaling regulator of Myxococcus xanthus, with the Fischbach lab (Stanford) to determine how the gut microbiome produces the neurotransmitter tryptamine, and with the Tawfik lab (Weizmann Institute, Israel) to determine the role of epistasis in restricting antibiotic resistance mutations. We are expanding this interest to include the interaction of human enzymes in degrading chitin molecules that can cause inflammation in the context of allergy and asthma, the hijacking of the proline isomerase CypA in lentiviral evolution, and structure-based antibiotic design using cryoEM.
 - a. **Fraser JS**, Yu Z, Maxwell KL, Davidson AR. Ig-like domains on bacteriophages: a tale of promiscuity and deceit. *J Mol Biol.* 2006. PMID: 16631788.
 - b. Williams BB, Van Benschoten AH, Cimermancic P, Donia MS, Zimmermann M, Taketani M, Ishihara A, Kashyap PC, **Fraser JS**, Fischbach MA. Discovery and characterization of gut microbiota decarboxylases that can produce the neurotransmitter tryptamine. *Cell Host Microbe*. 2014. PMCID: PMC4260654
 - c. Kane JR, Stanley DJ, Hultquist JF, Johnson JR, Mietrach N, Binning JM, Jónsson SR, Barelier S, Newton BW, Johnson TL, Franks-Skiba KE, Li M, Brown WL, Gunnarsson HI, Adalbjornsdóttir A, Fraser JS, Harris RS, Andrésdóttir V, Gross JD, Krogan NJ. Lineage-Specific Viral Hijacking of Noncanonical E3 Ubiquitin Ligase Cofactors in the Evolution of Vif Anti-APOBEC3 Activity. *Cell Reports*. 2015. PMCID: PMC4613747.
 - d. Dellus-Gur E, Elias M, Caselli E, Prati F, Salverda ML, de Visser JA, **Fraser JS**, Tawfik DS. Negative epistasis and evolvability in TEM-1 β-lactamase The thin line between an enzyme's conformational freedom and disorder. *J Mol Biol*. 2015. PMCID: PMC4718737.

Complete List of 53 Publications in MyBibliography:

http://www.ncbi.nlm.nih.gov/myncbi/browse/collection/40891283/?sort=date&direction=descending

C. Research Support

Ongoing Research Support

R01 GM123159 Fraser (MPI)/van den Bedem 12/01/17 – 11/31/21

NIH/NIGMS

Resolving ensemble averaged conformations by multi-temperature x-ray crystallography

The objective of this research program is to experimentally access and computationally model multi-scale heterogeneity in allosteric protein-ligand complexes.

Packard Fellowship for Science and Engineering Fraser (PI)

11/01/14 - 10/31/19

The David and Lucile Packard Foundation

The major goal of this project is to create and apply methods to examine non-Bragg (diffuse) scattering to define and study the importance of conformational dynamics in protein function.

NSF 11-522 Snell (PI) 09/01/13 – 09/01/23

NSF - OIA - SCI & TECH CTRS

Biology with X-ray Lasers

The major goal of this center is to encourage the development of methods for biophysics using the newly developed x-ray free electron lasers (X-FEL). We participate by generating samples for X-FEL diffraction and comparing the resulting data to room temperature synchrotron datasets.

LFR-17-476732 Fraser (PI) 03/01/17 – 02/29/20

UC Lab Fees Research Program

Macromolecular movements by simulation and diffuse scatter

The goal of this project is to validate X-ray diffuse scattering data with molecular dynamics simulations. Fraser is the overall project director, overseeing coordination between sites (UCSD, UCI, UCR, LANL).

MCB 1714915 Herschlag (PI) 08/01/17 – 07/31/21

NSF

Collaborative Research: Systematic Investigation of the Structure, Dynamics, and Energetics of Hydrogen Bonds and the Protein Interior Using Ketosteroid Isomerase and Model Systems

The goal of this project is to determine the biophysical and mechanistic basis for enzyme catalysis.

R01 GM0517315 Holton (PI) 07/01/17 – 06/30/22

NIH/NIGMS

Eliminating Critical Systematic Errors In Structural Biology With Next-Generation Simulation

The goal of the project is to use simulations to explore systematic errors to enable improved modeling.

P30 GM0519206 Adams (PI) 07/01/17 – 06/30/22

NIH/NIGMS

ALS Efficiently Networking Advanced Beam Line Experiments (ALS-ENABLE)

Fraser administers the project as Deputy Director of Macromolecular Crystallography and performs outreach. Fraser is the deputy project director, overseeing the crystallography component of the project.

Completed Research Support

R21 GM110580 Fraser (PI) 04/01/14-03/31/17

NIH/NIGMS

Model Comparison in Structural Biology

This project created new metrics for determining the precision and accuracy of protein conformations.

DP5 OD009180 Fraser (PI) 09/01/11 – 08/31/17

NIH/OSC

The Impact of Mutation on the Conformations and Recognition of Ubiquitin

This project used deep mutational scanning and biophysical characterization to study variants of Ubiquitin.