BIOGRAPHICAL SKETCH

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NAME: Glass, Karen C.

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

POSITION TITLE: Associate Professor of Pharmacology

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
University of Massachusetts (UMass), Amherst	B.S.	05/99	Microbiology
University of Vermont (UVM)	Ph.D.	10/05	Microbiology & Molecular Genetics, Christopher S. Francklyn
Rapid Data Collection and Structure Solving at the NSLS: A Practical Course in Macromolecular X-Ray Diffraction Measurement, Brookhaven, NY	Training Course	April 6-11, 2003	X-Ray Crystallography
University of California, Santa Cruz	Postdoctoral	11/06	MCD Biology, Harry F. Noller
NMRFAM Protein Structure Determination Workshop, University of Wisconsin-Madison	Training Course	June 6-11, 2010	Nuclear Magnetic Resonance
University of Colorado Denver (UCD)	Postdoctoral	08/10	Pharmacology, Tatiana G. Kutateladze
New York Structural Biology Center, New York, NY	Training Program	2022-2023	Cryo-EM Access and Training (NCCAT)

A. Personal Statement

The focus of my laboratory is to understand the epigenetic basis of disease. In particular, we are interested in how recognition of histone post-translational modifications by bromodomains contributes to their role in normal biological processes and in the development of diseases such as malaria. Interestingly, the histone code is highly dynamic, and changes as parasitic cells differentiate during development, at different stages of the cell life-cycle. and through disease progression. I have a strong background in Biochemistry, Molecular Biology, and Enzymology, with specific training in structural biology techniques including X-ray Crystallography, Nuclear Magnetic Resonance (NMR), and more recently single particle cryo-electron microscopy (cryo-EM). My research group has made significant progress to understand how multiple histone modifications function to collectively regulate the binding activity of bromodomain-containing proteins. An exciting project that has recently developed in my lab is aimed at characterizing the structure and function of the Plasmodium falciparum Bromodomain Protein 1 (PfBDP1). PfBDP1 is critical for the ability of this parasite to infect red blood cells by regulating the expression of invasion related genes. However, the function of this protein at the molecular level is unknown. The C-terminal bromodomain recognizes acetylated lysine residues on histone proteins within the nucleosome, likely bridging PfBDP1 to chromatin. Bromodomains are validated drug targets in cardiovascular disease and cancer, and PfBDP1 presents an exciting new therapeutic opportunity for the treatment of malaria. Our longterm goal is to understand the molecular mechanisms driving the biological activity of the PfBDP1 protein. By delineating how PfBDP1 recognizes specific epigenetic signals to regulate its interaction with the underlying genetic information, ultimately controlling the expression of genes critical for red blood cell entry, we will be able

to leverage this information to prevent malaria parasite proliferation and limit the devastating side effects caused by infection.

Ongoing projects that I would like to highlight include:

- 1. R01 GM129338 Glass/Frietze (MPI) 09/19/2018 08/31/2022 (NCE thru 08/31/2023) Deciphering the molecular mechanisms of histone code recognition by ATAD2/B.
- Stein, G. (Project 2- Frietze/Glass) 04/01/2021 03/31/2026 2. P01 CA240685 Epigenetic Control and Genome Organization, Project 2: Bromodomains as epigenetic modulators of endocrine responsiveness in ER+ breast cancer.

Relevant publications:

- a. Evans CM, Phillips M, Malone KL, Tonelli M, Cornilescu G, Cornilescu C, Holton SJ, Gorjanacz M, Wang L, Carlson S, Gay JC, Nix JC, Demeler B, Markley JL, and Glass KC. (2021). Coordination of di-acetylated histone ligands by the ATAD2 bromodomain. Int. J. Mol. Sci. 2021, 22(17), 9128. PMCID: PMC8430952.
- b. Singh AK, Phillips M, Alkrimi S, Tonelli M, Boyson SP, Malone KL, Nix JC, Glass KC. (2022) Structural insights into acetylated histone ligand recognition by the BDP1 bromodomain of Plasmodium falciparum. Int J Biol Macromol. 2022 Oct 31;S0141-8130(22)02502-8. DOI: 10.1016/j.ijbiomac.2022.10.247. PMID: 36328269

B. Positions, Scientific Appointments, and Honors **Positions**

06/21-present Associate Professor,	Department of Pharmacology, Larner College of Medicine, University of
Vermont (UVM).	

07/19-presen	t Adjunct Associate Professor, Department of Biochemistry, Larner College of Medicine, UVM.
06/16-05/21	Associate Professor, Department of Pharmaceutical Sciences, Albany College of Pharmacy and
	Health Sciences (ACPHS)

08/11-06/19	Adjunct Assistant Profess	sor, Department of Bioch	emistry, Larner Colle	ge of Medicine, UVM.
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08/10-05/16 Assistant Professor, Department of Pharmaceutical Sciences, ACPHS

08/09-12/09 Affiliate Professor, Department of Biology, Metropolitan State College, Denver, CO 11/06-08/10 Postdoctoral Research Fellow, Department of Pharmacology, UCD, Aurora, CO

Postdoctoral Research Associate, Department of Molecular, Cellular and Developmental 11/05-11/06 Biology, UCSC, Santa Cruz, CA

Scientific Appointments	
2022-present	Member, NIGMS study section, Training and Workforce Development-B (TWD-B)
2022-present	Member, NMRFAM User Program External Advisory Board
2021-present	Member, Liaison Committee on Education, American Institute of Physics
2021-present	Member, Pilot grant review committee, UVM Cancer Center
2021-present	Member, The Association for Women in Science (AWIS)
2021-present	Member, Graduate admissions committee, UVM Cellular, Molecular and Biomedical Sciences
2021-present	Member, Education Committee, American Crystallography Association
2021	Ad Hoc Reviewer, NIH Fellowships: Genes, Genomes and Genetics (F08)
2021-present	Member, Biophysical Society
2020-2021	Member, ACPHS Promotion & Tenure committee
2020	Ad Hoc Reviewer, NIH Fellowships: Genes, Genomes and Genetics (F08)
2019-2021	Member, Admissions Committee, Masters in Pharmaceutical Sciences Graduate Program,
	Albany College of Pharmacy and Health Sciences
2018	Ad Hoc Reviewer, NIH Molecular Genetics A Study Section (MGA)
2016	Ad Hoc Reviewer, NIH Molecular Genetics A Study Section (MGA)
2015-present	Member, grant review committee, American Cancer Society Institutional Research Grant (ACS-

IRG) at the University of Vermont 2014-present **Member**, UVM Cellular and Molecular Biology Graduate Program

Member, ACPHS Research Committee 2015-2018

Advisory Board Member: 2nd Epigenomics & Novel Therapeutic Targets Disease Conference. 2015 May 26 - 27, 2016, Boston, MA.

Member, ACPHS Graduate Faculty Curriculum Committee 2014-2020

- 2013 Co-organizer of the 2013 Vermont Cancer Center annual symposium with the theme of Epigenetics and Cancer. 2013 Advisory Board Member: 3rd Epigenetics in Drug Discovery Conference, May 8-10, 2013, Boston, MA. 2013-present **Member**, UVM Graduate Faculty 2012-2014 Founding Chair, ACPHS Graduate Faculty Curriculum Committee
- 2011-present Manuscript referee for journals including: FEBS Letters, Journal of Biological Chemistry.
 - Journal of Medicinal Chemistry, and Nucleic Acids Research (among others).
- 2011-present Member, University of Vermont Cancer Center
- 2011-present **Member**, American Society for Biochemistry and Molecular Biology
- 2010-present **Member**, American Crystallographic Association

Honors

2022	Taylor & Francis Biomolecular Crystallography poster prize, ACA Annual Meeting, Portland, OR
2021-22	National MAVEN Senior Scientist (NIGMS funded leadership program)
2014	ACPHS Researcher of the Year award
2012	ASBMB annual meeting thematic best poster in the Gene Regulation category
2011	Poster presentation award winner in the Faculty/Staff category. Vermont Cancer Center's 2011

- Clinical and Translational Research Symposium: DNA Repair & Cancer. Poster presentation award winner in the Faculty/Staff category. Vermont Cancer Center's 2010 2010
- Clinical and Translational Research Symposium: Inflammation & Cancer.
- 2010 Poster presentation award. University of Colorado Denver Postdoctoral Research Day.
- 2008 Postdoctoral Award for Outstanding Achievement, Department of Pharmacology, UCD, SOM.
- 2008 Keystone Symposia Scholarship, 'Molecular Basis for Chromatin Modifications and Epigenetic Phenomena', Snowmass, CO.
- Postdoctoral Fellowship, National Institutes of Health (NRSA F32GM083462) 2008-10
- 2008 Postdoctoral Fellowship, American Heart Association (declined).
- 2007-08 Postdoctoral Fellowship, American Cancer Society (08-049-01-GMC)
- Travel award from the UVM graduate college for a tRNA Synthetase conference, Seoul, Korea. 2004
- 2001-03 Vermont Department of Energy Experimental Program to Stimulate Competitive Research (DOE EPSCoR) graduate research fellowship.
- 2001 American Crystallography Association student travel grant, ACA meeting, Los Angeles, CA.
- 1999 Graduated with honors, Magna Cum Laude.
- Howard Hughes grant for undergraduate research. 1999
- 1998 Honors Research Grant for undergraduate thesis research.
- 1997 Golden Kev National Honors Society, member.
- 1996 Alpha Lambda Delta, a national academic honors society for freshmen in the top 10% of their class.
- C. Contributions to Science (Please note name change in 2009 from Champagne KS to Glass KC)
- Allosteric regulation of histidine biosynthesis: As a graduate student in the laboratory of Dr. Christopher S. Francklyn at UVM I characterized the structure and function of the N1-5'-phosphoribosyl ATP transferase (ATP-PRTase) in Lactococcus Lactis, which catalyzes the first step of histidine biosynthesis. The L. lactis ATP-PRTase ezyme is unique and contains two subunit types, one of which is a paralogous to histidyl-tRNA synethtase (HisRS). I established that the HisZG ATP-PRTase from L. lactis is a 250 kDa multimeric enzyme complex consisting of four HisG and four HisZ subunits using size exclusion chromatography, and quantitative protein sequencing. Under the quidance of Dr. Sylvie Doublié and Dr. Stephen J. Everse I solved the first structure of a PRPP-bound ATP-PRTase, and provided a structural model for its allosteric activation by comparing inhibited and activated versions of ATP-PRTs from both the hetero-octameric and hexameric families. Mutational analysis followed by kinetic binding assays identified the histidine binding sites in a region highly conserved between HisZ and the functional HisRS, confirming the role of HisZ as a regulatory subunit in the L. lactis ATP-PRTase. My research provided evidence on how a histidyl-tRNA synthetase-like domain evolved through evolution to function as a regulatory domain in amino acid biosynthesis.
 - a. Bovee ML, Champagne KS, Demeler B, Francklyn CS. (2002) The Quaternary Structure of the HisZ-HisG N-1-(5'-Phosphoribosyl)-ATP Transferase from *Lactococcus lactis*. *Biochemistry*. 41(39): 11838-11846. PMID: 12269828

- b. Champagne KS, Sissler M, Larrabee Y, Doublié S, Francklyn CS. (2005) Activation of the hetero-octameric ATP phosphoribosyl transferase through subunit interface rearrangement by a tRNA synthetase paralog. *J Biol Chem.* 280(40): 34096-34104. PMID: 16051603
- **c. Champagne KS**, Piscitelli E, Francklyn CS. (2006) Substrate recognition by the hetero-octameric ATP phosphoribosyltransferase from *L. lactis. Biochemistry*. 45(50): 14933-43. PMID: 17154531
- Molecular mechanisms of histone recognition by the plant homeodomain (PHD): I became interested in the field of Epigenetics and wanted to understand how the 'histone code' might extend and modify our genetic (DNA) information to regulate key cellular processes. Modifications on the histone tail have been shown to be important in altering chromatin structure, and they regulate gene expression by facilitating access of DNA-binding transcription factors. Modifications to the histone tail also act as markers, allowing non-histone proteins to interact with the chromatin. When I began my postdoctoral training with Tatiana G. Kutateladze the molecular basis of histone recognition by chromatin reader domains was poorly understood. I made several seminal discoveries during this period. Using tryptophan fluorescence, I revealed that there is cross-talk between adjacent histone modifications on the histone H3 tail that regulate the functions of proteins interacting with these marks. We showed that di-methylation of arginine 2 on histone H3 (H3R2me2) modulates the interaction of the RAG2 PHD finger with tri-methylated lysine 4 (H3K4me3) on the histone tail. and is essential for V(D)J recombination. I also demonstrated the molecular mechanism of H3K4me3 recognition by the Inhibitor of Growth PHD fingers is conserved within this family, and that histone binding both recruits and activates ING4/5-associated histone acetyltransferase complexes on chromatin. My research also revealed that PHD fingers are divided into sub-families based on their selection of different ligands including unmodified histone H3, and acetylated or methylated lysine.
 - a. Matthews AG, Kuo AJ, Ramón-Maiques S, Han S, Champagne KS, Ivanov D, Gallardo M, Carney D, Cheung P, Ciccone DN, Walter KL, Utz PJ, Shi Y, Kutateladze TG, Yang W, Gozani O, Oettinger MA. (2007) RAG2 PHD finger couples histone H3 lysine 4 trimethylation with V(D)J recombination. *Nature*. Dec 13; 450(7172): 1106-10. PMCID: PMC2988437
 - b. Champagne KS, Saksouk N, Peña PV, Johnson K, Ullah M, Yang XJ, Côté J, Kutateladze TG. (2008) The crystal structure of the ING5 PHD finger in complex with an H3K4me3 histone peptide. *Proteins*. 72(4): 1371-6. PMCID: PMC2756976
 - **c.** Hung T*, Binda O*, **Champagne KS***, Kuo AJ, Johnson K, Chang HY, Simon MD, Kutateladze TG and Gozani O. (2009) ING4-mediated crosstalk between histone H3K4 trimethylation and H3 acetylation attenuates cellular transformation. *Mol Cell*. 33(2): 248-256. PMCID: PMC2650391
 - d. Kim S, Natesan S, Cornilescu G, Carlson S, Tonelli M, McClurg UL, Binda O, Robson CN, Markley JL, Balaz S, Glass KC. (2016) Mechanism of Histone H3K4me3 Recognition by the Plant Homeodomain of Inhibitor of Growth 3. *J Biol Chem*. Aug 26;291(35):18326-41. PMCID: PMC5000080
- 3. Molecular mechanisms of histone recognition by bromodomains: In my independent research program I have continued studying the structure and function of chromatin reader domains, particularly bromodomains, which interact specifically with acetylated histones. The 61 human bromodomain-containing proteins have a wide variety of biological activities. However, while the structure of many of these bromodomain modules are solved, how these protein modules differentiate between multiple acetyllysine modifications to read the histone code is unknown. We recently established the molecular basis of histone acetyllysine recognition by the BRPF1 bromodomain and discovered that the BRPF1 bromodomain interacts with multiple acetylated histone peptides. We also solved the first bromodomain structure in complex with histone H2A acetylated at lysine 5 (H2AK5ac). These structural and mechanistic details of histone recognition by bromodomains is crucial for the development of new therapeutic interventions and molecular tools to study a variety of cancers, and has fundamentally advanced our understanding of how bromodomains recognize and select for acetyllysine marks.
 - **a.** Lee W, Hu K, Tonelli M, Bahrami A, Neuhardt E, **Glass KC**, Markley JL. (2013) Fast automated protein NMR data collection and assignment by ADAPT-NMR on Bruker spectrometers. *J Magn Reson*. *Aug* 30;236C:83-88. PMCID: PMC3858185
 - b. Poplawski A, Hu K, Lee W, Natesan S, Peng D, Carlson S, Shi X, Balaz S, Markley JL, Glass KC. (2014) Molecular Insights into the Recognition of N-Terminal Histone Modifications by the BRPF1 Bromodomain. *J Mol Biol.* 426(8): 1661-1676. PMCID: PMC3969779

- c. Lubula MY, Eckenroth BE, Carlson S, Poplawski A, Chruszcz M, and Glass KC (2014) Structural insights into recognition of acetylated histone ligands by the BRPF1 bromodomain. FEBS Lett. 588(21): 3844-54. PMCID: PMC4252766
- d. Lloyd JT, Glass KC. (2018) Biological function and histone recognition of family IV bromodomain-containing proteins. J Cell Physiol. 2018 Mar;233(3):1877-1886. DOI: 10.1002/jcp.26010. Epub 2017 Jun 13. Review. PMCID: PMC5683942.
- 4. Combinatorial action of chromatin reader domains and histone modifications: As a direct result of my research, we provided new insights into the higher-level regulation of gene expression that is modulated through histone modifications and their readers. The histone code is orders of magnitude more complex than the genetic code, and understanding this process is compounded by the presence and interaction of multiple chromatin reader domains within single enzymatic complexes. My research on the HBO1 and MOZ histone acetyltransferase complexes has highlighted how altering either the subunit composition of an enzymatic complex or the availability of particular combinations of histone modifications can dramatically affect the activity of chromatin remodelers, and epigenetic signaling programs within the cell. For example, we proposed a model describing how the cooperative action of multiple chromatin reader domains within the MOZ histone acetyltransferase regulates its acetylation activity in response to the epigenetic landscape.
 - a. Saksouk N, Avvakumov N*, Champagne KS*, Hung T*, Doyon Y, Cayrou C, Paquet E, Ulla M, Landry AJ, Côté V, Yang XJ, Gozani O, Kutateladze TG and Côté J. (2009) HBO1 HAT complexes target chromatin throughout gene coding regions via multiple PHD finger interactions with histone H3 tail. *Mol Cell*. 33:257-265. PMCID: PMC2677731 *These authors contributed equally to the work.
 - b. Lalonde ME, Avvakumov N, Glass KC, Joncas FH, Saksouk N, Holliday M, Paquet E, Yan K, Tong Q, Klein BJ, Tan S, Yang XJ, Kutateladze TG, Côté J. (2013) Exchange of associated factors directs a switch in HBO1 acetyltransferase histone tail specificity. Genes Dev. 27(18):2009-24. PMCID: PMC3792477
 - c. Avvakumov N, Lalonde ME, Saksouk N, Paquet E, Glass KC, Landry AJ, Doyon Y, Cayrou C, Robitaille GA, Richard DE, Yang XJ, Kutateladze TG and Côté J. (2012) Conserved Molecular Interactions within the HBO1 Acetyltransferase Complexes Regulate Cell Proliferation. *Mol Cell Biol*. Feb;32(3):689-703. PMCID: PMC3266594
 - **d.** Carlson S and **Glass KC**. (2014) The MOZ Histone Acetyltransferase in Epigenetic Signaling and Disease. *J Cell Physiol*. Nov 229(11): 1571-4. PMCID: PMC4750494
- 5. Recognition of multiple histone post-translational modifications by chromatin reader domains: Although the bromodomain structural fold is conserved across all BRD-containing proteins, each BRD module exhibits specific preferences for different histone acetyllysine ligands on core and variant histone proteins. My research group has made significant progress to understand how multiple modifications alters the binding activity of bromodomain containing proteins. These studies have fundamentally advanced our understanding of how bromodomains recognize and select for acetyllysine marks.
 - a. Obi JO, Lubula MY, Cornilescu G, Henrickson A, McGuire K, Evans CM, Phillips M, Demeler B, Markley JL and Glass KC. (2020) The BRPF1 bromodomain is a molecular reader of di-acetyllysine. Curr Res in Struct Biol. 2020, 2:104-115. DOI: 10.1016/j.crstbi.2020.05.001 PMCID: PMC3969779
 - b. Lloyd JT, McLaughlin K, Lubula MY, Gay JC, Dest A, Gao C, Phillips M, Tonelli M, Cornilescu G, Marunde MR, Evans CM, Boyson SP, Carlson S, Keogh MC, Markley JL, Frietze S, and Glass KC. (2020) Structural insights into the recognition of mono- and di-acetylated histones by the ATAD2B bromodomain. *J. Med Chem.* 2020 Nov 12;63(21):12799-12813. PMCID: PMC7884259.
 - **c.** Phillips M, Quinn K, Paculova H, Montgomery C, Joseph FM, Boyson SP, Chang S, Nix JC, Young NL, Frietze SE, **Glass KC**. (2022) Regulation of ATAD2B bromodomain binding activity by the histone code. **BioRxiv** 2022.11.14.516501; DOI: 10.1101/2022.11.14.516501.

Complete List of Published Work in MyBibliography

https://www.ncbi.nlm.nih.gov/myncbi/1J3jpw-PqCEAd/bibliography/public/