­­Curriculum Vitae: **Christopher L. Baker, Ph.D.**

The Jackson Laboratory

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**EDUCATION**

2004-2010 Ph.D. Department of Genetics, Dartmouth Medical School, Hanover, NH

 Molecular and Cellular Biology Program

2001-2004 B.S., *Summa Cum Laude,* College of Agriculture and Life Sciences, University of Vermont,

**Professional Experience**

2019 Resident Supervisor, The Jackson Laboratory’s Summer Student Program

2017-present Assistant Professor, Sackler School of Graduate Biomedical Sciences, Department of Medicine, Tufts School of Medicine, Boston, MA

2016-present Faculty, Graduate School of Biomedical Sciences and Engineering, The University of Maine, Orono, ME

2016-present Assistant Professor, The Jackson Laboratory, Bar Harbor, ME

2015-2016 Associate Research Scientist, The Jackson Laboratory, Bar Harbor, ME

2010-2015 Postdoctoral Fellow, The Jackson Laboratory, Bar Harbor, ME

**Publications**

Google citation indices: <https://scholar.google.com/citations?user=DpPqyOUAAAAJ&hl=en>

1. Aydin S, Pham DT, Zhang T, Keele GR, Skelly DA, Pankratz M, Choi T, Gygi SP, Reinholdt LG\*, **Baker CL\***, Churchill GA\*, Munger SC\*. Genetic dissection of the pluripotent proteome through multi-omics data integration. bioRxiv. doi: https://doi.org/10.1101/2022.04.22.489216 ***\*corresponding authors***
2. Kuffler L, Skelly DA, Czechanski A, Munger SC, **Baker CL**, Reinholdt LG, Carter GW. Imputation of 3D genome structure by genetic-epigenetic interaction modeling in mice. bioRxiv. doi: https://doi.org/10.1101/2022.02.07.479436
3. Byers C, Spruce C, Fortin H, Hartig E, Czechanski A, Munger SC,Reinholdt RG, Skelly DA, **Baker CL**. **2022**. Genetic control of pluripotency epigenome determines differentiation bias in mouse embryonic stem cells. *The* *EMBO Journal*. 41(2):e109445, doi: 10.15252/embj.2021109445
4. Kelliher CM, Lambreghts R, Xiang Q, **Baker CL**, Loros JJ, Dunlap JC. **2020**. PRD-2 directly regulates casein kinase I and counteracts nonsense mediated decay in the *Neurospora* circadian clock. *eLife,* 9:e64007 doi: 10.7554/eLife.64007
5. Skelly DA, Czechanski A, Byers C, Aydin S, Spruce C, Olivier C, Choi K, Gatti DM, Raghupathy NM, Keele GR, Stanton A, Vincent M, Dion S, Greenstein I, Pankratz M, Porter DK, Martin W, O’Conner C, Qin W, Harrill AH, Choi T, Churchill GA\*, Munger SC\*, **Baker CL\*,** Reinholdt RG\*. **2020**. Mapping the effects of genetic variation on chromatin state and gene expression reveals loci that control ground state pluripotency. *Cell Stem Cell*, 27(3):459-469*.* ***\*corresponding authors***

Preview: D’Antonio M, D-Antonio-Chronowska A, Frazer KA. Revealing Instability: Genetic variation underlies variability in mESC pluripotency. *Cell Stem Cell*, 27(3): 347-349*.*

1. Ortmann D, Brown S, Czechanski A, Aydin S, Muraro D, Huang Y, Tomaz RA, Osnato A, Canu G, Wesley BT, Skelly DA, Stegle O, Choi T, Churchill G, **Baker CL**, Munger SC, Reinholdt LG, Vallier L. **2020**. Naïve pluripotent stem cells exhibit phenotypic variability that is driven by genetic variation. *Cell Stem Cell*, 27(3):470-481.
2. Lau K, Mason EA, Kie J, De Souza DP, Kloehn J, Tull D, McConville MJ, Keniry A, Beck T, Blewitt ME, Ritchie ME, Naik SH, Zalcenstein D, Korn O, Su S, Romero IG, Spruce C, **Baker CL**, McGarr TC, Wells CA, Pera MF. **2020**. Unique properties of a subset of human pluripotent stem cells with high capacity for self-renewal. *Nat Commun.,* 11(1):2420. doi: 10.1038/s41467-020-16214-8.
3. Spruce C, Dlamini S, Ananda G, Bronkema N, Tian H, Paigen K, Carter GW, **Baker CL**. **2020.** HELLS and PRDM9 form a pioneer complex to open chromatin at meiotic recombination hotspots. *Genes Dev.,* 34: 398-412.doi:10.1101/gad.333542.119

Article highlighted in: Alavattam KG, Abe H, Namekawa SH. Pioneering meiotic recombination. 2020. *Genes Dev*., 34:395-397.

1. Mihola O, Pratto F, Brick K, Linhartova E, Flachs P, **Baker CL**, Sedlacek R, Paigen K, Petkov PM, Camerini-Otero D, Trachtulec Z. **2019**. Histone methyltransferase PRDM9 is partially dispensable for meiosis in male mice. *Genome Research,* doi:10.1101/gr.244426.118
2. **Baker CL\*,** Walker M, Arat S, Ananda G, Petkova P, Powers N, Tian H, Spruce C, Ji B, Rausch D, Choi K, Petkov PM, Carter GW, Paigen K\*. **2019.** Tissue-specific *trans* regulation of the mouse epigenome. *Genetics,* https://doi.org/10.1534/genetics.118.301697*.* ***\*corresponding authors***
3. **Baker CL** and Pera MF. **2018**. Capturing Totipotent Stem Cells. *Cell Stem Cell*, 22(1):25-34.
4. Powers NR, Parvanov ED, **Baker CL**, Walker M, Petkov PM, Paigen K. **2016**. The meiotic recombination activator PRDM9 trimethylates both H3K36 and H3K4 at recombination hotspots in vivo. *PLoS Genetics*, 12(6):e1006146. doi:10.1371/journal.pgen.1006146
5. Narasimhan V, Hunt K\*, Mason D\*, **Baker CL\***, *et al.* [35 authors]. **2016**. Health and population effects of rare gene knockouts in adult humans with related parents. *Science,* 352(6284):474-7. doi: 10.1126/science.aac8624 \*equal contribution
6. **Baker CL**, Petkova P, Walker M, Flachs P, Mihola O, Trachtulec Z, Petkov PM, Paigen K. **2015**. Multimer formation explains allelic suppression at PRDM9 hotspots. *PLoS Genetics,* 11(9): e1005512. doi:10.1371/journal.pgen.1005512
7. Walker M, Billings T, **Baker CL**, Powers N, Tian H, Saxl RL, Choi K, Hibbs MA, Carter GW, Handel MA, Paigen K, Petkov PM. **2015**. Affinity-seq detects genome-wide PRDM9 binding sites and reveals the impact of prior chromatin modifications on mammalian recombination hotspot usage. *Epigenetics & Chromatin*, 8:31: doi: 10.1186/s13072-015-0024-6
8. Sun F, Fujiwara Y, Reinholdt R, Hu, J, Saxl RL, **Baker CL**, Petkov PM, Paigen K, Handel MA. **2015**. Nuclear localization of PRDM9 and its role in meiotic chromatin modifications and homologous synapsis. *Chromosoma*: 1-19.
9. **Baker CL**, Kajita S, Walker M, Saxl RL, Raghupathy N, Choi K, Petkov PM, Paigen K. **2015**. PRDM9 drives evolutionary erosion of hotspots through haplotype-specific initiation of meiotic recombination. *PLoS Genetics*, 11(1): e1004916. doi:10.1371/journal.pgen.1004916
10. Larrondo LF, Olivares-Yanez C, **Baker CL**, Loros JL, Dunlap JC. **2015**. Decoupling circadian clock protein turnover from circadian period determination. *Science*, 347(6221):1257277.
11. Bubier JA, Jay JJ, **Baker CL**, Bergeson SE, Ohno H, Metten P, Crabbe JC, Chesler EJ. **2014**. Identification of a QTL in *Mus musculus* for alcohol preference, withdrawal, and Ap3m2 expression using integrative functional genomics and precision genetics. *Genetics,* 197(4):1377-93.
12. **Baker CL**, Walker M, Kajita S, Petkov PM, Paigen K. **2014.** PRDM9 binding organizes hotspot nucleosomes and limits Holliday junction migration. *Genome Research*, 24(5):724-732.

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1. Billings T, Parvanov ED, **Baker CL**, Walker M, Paigen K, Petkov PM. **2013**. DNA binding specificities of the long zing finger recombination protein PRDM9. *Genome Biology*, 14:R35
2. **Baker CL**, Loros JJ, and Dunlap JC. **2012**. The circadian clock *Neurospora crassa*. *FEMS Microbiology Reviews*, 36(1):95-110.
3. **Baker CL** and Dunlap JC. **2009**. Circadian Rhythms: Phosphorylating the CLOCK. *Cell Cycle*, 9(2):231-2.
4. Mehra A, **Baker CL**, Loros JJ, and Dunlap JC. **2009**. Post translational modifications in circadian rhythms. *TiBS*,34(10):483-90.
5. Mehra A, Shi M, **Baker CL**, Colot HV, Loros JJ, Dunlap JC. **2009**. CK2 and temperature compensation in *Neurospora*. *Sleep and Biological Rhythms*, 7(3)162-171.
6. **Baker CL**, Kettenbach AN, Loros JJ, Gerber SA, and Dunlap JC. **2009**. Quantitative proteomics reveals a dynamic interactome and phase-specific phosphorylation in the *Neurospora* circadian clock. *Molecular Cell*, 34(3):354-63.
7. Mehra A, Shi M, **Baker CL**, Colot HV, Loros JJ, and Dunlap JC. **2009**. A role for Casein Kinase 2 in the mechanism underlying circadian temperature compensation. *Cell*, 137(4):749-60.
8. Larrondo LF, Colot HV, **Baker CL**, Loros JJ, and Dunlap, JC. **2009**. Fungal Functional Genomics: Tunable Knockout-Knockin-expression and tagging strategies. *Eukaryotic Cell*, 8(5)800-4.
9. Loros JJ, Dunlap JC, Larrondo LF, Shi M, Belden WJ, Gooch VD, Chen CH, **Baker CL**, Mehra A, Colot HV, Schwerdtfeger C, Lambreghts R, Collopy PD, Gamsby JJ, Hong CI. **2007**. Circadian output, input, and intracellular oscillators: insights into the circadian systems of single cells. *Cold Spring Harb Symp Quant Biol,* 72:201-14.
10. Dunlap JC, Loros JJ, Colot HV, Mehra A, Belden WJ, Shi M, Hong CI, Larrondo LF, **Baker CL**, Chen-Hui C, Schwerdtfeger C, Collopy PD, Gamsby JJ, Lambreghts R. **2007**. A circadian clock in *Neurospora*: How genes and proteins cooperate to produce a sustained, entrainable, and compensated biological oscillator with a period of about a day. *Cold Spring Harb Symp Quant Biol,* 72:57-68.
11. Pregueiro AM, Liu Q, **Baker CL**, Dunlap JC, Loros JJ. **2006**. The *Neurospora* checkpoint kinase 2: a regulatory link between the circadian and cell cycles. *Science,* 313(5787):644-9.

**FuNding**

R35GM133724 Effort: 51% 07/01/2019 ‑ 06/30/2024

National Institute of General Medical Sciences, NIH Annual Direct Costs: $250,000

Cellular Systems Genetic Approaches to Understanding Regulatory Variation Total Direct Costs: $1,250,000

In this proposal we will quickly and efficiently define the pathways and mechanisms that modulate regulatory variation and function during early mammalian development by leveraging natural variation intrinsic in a unique mouse embryonic stem cell system.

Role: PI

R24OD030037 (Reinholdt, Baker, Munger) Effort: 15% 02/26/21 – 01/31/2025

Office of the Director, National Institutes of Health Annual Direct Costs: $495,000

 Total Direct Costs: $1,917,165

Genetically Diverse Mouse Embryonic Stem Cells: A Platform for Cellular Systems Genetics

The objective of this application is to generate a thoroughly-validated panel of genetically diverse mouse embryonic stem cells (mESC) that will enable widespread adoption of cellular systems genetics.

Role: Multi-PI

**PENDING**

R13 Effort: N/A

Office of the Director & National Human Genome Research Institute, NIH

Diversity in a Dish: Pluripotent Stem Cells in Genetic Analysis and Disease Modeling

Our vision is to bring together a unique combination of pluripotent stem cell biologists, systems geneticists, and investigators using these approaches for precision disease modeling. Pluripotent stem cells hold great promise for deciphering disease mechanisms and providing new cell therapies for intractable medical conditions. Only through developing and studying genetically diverse panels of stem cells can we ensure that the benefits of this research are distributed equitably. The goal of this symposium is to help bridge these groups to coalesce into an integrated, pluripotent stem cell-based genetics into a focused discipline.

Role: PI

IMPACT Score: **20**

**COMPLETED**

JAX‑DIF‑FY20‑CLB-Neurogenetics (Baker, Chesler) Effort: N/A 01/31/2020 ‑ 12/31/2021

The Jackson Laboratory Director's Innovation Fund

Investigating Epigenomic Mechanisms underlying the Neurogenetics of Addiction Total Direct Costs: $133,672

The major goals of this study are to determine the effect of genetic background and sex on chromatin accessibility at cis-regulatory elements in the striatum, a critical brain structure implicated in multiple substance abuse disorders; to identify role of genetic background on proportion of cell-types in the striatum; and to integrate expression and behavior QTL from models of addiction with variation in chromatin accessibility.

Role: PI

JAX‑DIF‑FY19‑CB‑SCM (Baker, Pera, Munger, Skelly, Reinholdt) Effort: N/A 04/19/2019 ‑ 04/18/2020

The Jackson Laboratory Director's Innovation Fund

Genetic control of cellular heterogeneity in pluripotent stem cells Total Direct Costs: $220,000

The goal of this proposal is to profile transcript abundance in mouse and human PSCs to delineate the regulatory networks that underlie cell heterogeneity across a panel of diverse genetic backgrounds.We will use 1) single cell (sc)RNA-Seq to capture population dynamics and developmental trajectories, and 2) FACS isolation of specific subpopulations at different stages of pluripotency using cell-surface markers followed by standard RNA-Seq*.*

Role: PI

JAX‑DIF‑FY17‑CB‑SCM (Baker, Churchill, Munger, Reinholdt) Effort: N/A 04/01/2017 ‑ 10/31/2018

The Jackson Laboratory Director's Innovation Fund Annual Direct Costs: $178,000

DO and CC RIX mESCs. An Advanced Platform for Cellular Systems Genetics Total Direct Costs: $178,000

The goal of this proposal is to demonstrate the power of the founder strain, DO, and CC-RIX mESC lines as a platform for cellular systems genetics. To that end, we will profile the chromatin landscape and cellular proteomes of undifferentiated mESC lines comprising the eight founder strains of the DO/CC, 150 individual outbred DO lines, and 10 F1 progeny (CC-RIX) from crosses of CC strains.

Role: PI

JAX‑DIF‑FY17‑KP (Paigen, Baker, Petkov) 03/01/2017 ‑ 12/31/2017

The Jackson Laboratory Director's Innovation Fund Annual Direct Costs: $43,200

Understanding the Chromatin Regulatory System Total Direct Costs: $43,200

The goal of this project is to expand our understanding of a newly discovered system of trans‑acting genes comprising a Chromatin Regulatory System (CRS) that controls the epigenetic landscape.

Role: Co‑Investigator

F32GM101736-01 (Baker, PI) 2012-2014 (salary support)

National Institute of General Medical Sciences, NIH

Ruth I. Kirschstein National Research Service Award

Title: Genetic Dissection of Quantitative Control of Recombination

**Awards and Honors**

2016 PALM (Promoting Active Learning and Mentoring) Fellowship. [*see press*](http://genestogenomes.org/putting-active-learning-into-practice-an-interview-with-palm-fellow-christopher-baker-and-palm-mentor-michelle-smith/)

 *NSF funded fellowship to promote long-term mentorship to improve undergraduate education.*

2014 Outstanding Oral Presentation, 28th International Mammalian Genome Conference

2013 Outstanding Oral Presentation, 27th International Mammalian Genome Conference

2012-2013 Ruth I. Kirschstein National Research Service Award

 *Individual postdoctoral fellowship*

2010-2012 T32 Postdoctoral Fellowship, The Jackson Laboratory

2010 John W. Strohbehn Award for Excellence in Biomedical Research, Dartmouth Medical School *awarded to a single graduating Ph.D.*

2009 Rosaline Borison Memorial Fellowship

2008 Society for Research on Biological Rhythms (SRBR) Excellence Award

 s*tudent travel award*

2007 Albert J. Ryan Fellow, Albert J. Ryan Foundation

2006-2008 National Institute for Health Pre-doctoral Training Grant, Dartmouth Medical School

2003 & 2004 Hughes Endeavor for Life Science Excellence Grant (HELiX), University of Vermont

 *competitive undergraduate research award*

2003 Class of 1939 Scholarship, a merit based award, University of Vermont

2003 James E. Ludlow Endowed Scholarship Award, University of Vermont

2002 & 2003 Holzer Memorial Scholarship, a merit based award, University of Vermont

**Invited Presentations**

1. Texas A&M, College Station, TX. February 20th, 2023.
2. Cold Spring Harbor Laboratory, The Genome Access Course. November 11th, 2019.
3. Sloan Kettering Cancer Center, Manhattan, NY. November 10th, 2019.
4. Tufts University Sackler Medical School, Boston, MA. Department of Genetics Seminar. May 9th, 2018. Natural genetic variation shapes the epigenetic landscape and patterns of inheritance.
5. Time of Our Life Symposium. July 13-14th, 2017. Hanover, NH. Genetic control of the epigenetic landscape.
6. 44th Annual Maine Biological and Medical Science Symposium. April 28-29th, 2017. Mount Desert Biological Laboratory. Non-Mendelian inheritance, meiotic drive, and genetic recombination.

**Conference Presentations**

1. The 35th International Mammalian Genome Conference. July 17-20th, 2022. Vancouver, British Columbia, Canada. Long-read sequencing from genetically diverse mouse embryonic stem cells reveals abundant transcript novelty and usage.
2. Complex Trait Community. September 1-3rd, 2021. Virtual. Genetic control of the pluripotency epigenome determines differentiation bias in mouse embryonic stem cells.
3. Stem Cell Biology. September 17-21st, 2019. Cold Spring Harbor, NY. Genetic variation influences ground state pluripotency in embryonic stem cells through a hierarchy of molecular phenotypes.
4. The 32nd International Mammalian Genome Conference. November 11-14th, 2018. Rio Mar, PR. A nucleosome remodeling factor is required for PRDM9-dependent meiotic recombination.
5. Mammalian Genetic and Genomics: From Molecular Mechanisms to Translational Applications. October 24-27th, 2017. EMBL Heidelberg, Germany. Genetic control of the epigenetic landscape.
6. The Allied Genetics Conference. July 13-17th, 2016. Orlando, FL. Natural genetic variation controls chromatin state in male germ cells.
7. The 28th International Mammalian Genome Conference.October 25-29th, 2014. Bar Harbor, ME. PRDM9 drives evolutionary erosion of hotspots. **Selected as outstanding presentation**
8. The 27th International Mammalian Genome Conference**.** September 15, 2013. Salamanca, Spain. Genome-wide analysis of PRDM9-dependent chromatin modification. **Selected as outstanding presentation**
9. The Center for Genome Dynamics Advisory Board Meeting.June 25, 2013. Bar Harbor, ME. Genome-wide analysis of PRDM9-dependent chromatin modification.

**Poster Presentations**

1. International Society for Stem Cell Research Annual Meeting. June 15-18th, 2022. Distal regulation of heterochromatin impacts the 3D genome and differentiation propensity of mouse embryonic stem cells.
2. The Identity and Evolution of Cell Types. May 4-7th, 2021. Virtual. Genetic control of the pluripotency epigenome determines differentiation bias in mouse embryonic stem cells.
3. 3D Genome: Gene Regulation and Disease. March 17-21st, 2019. Banff, Alberta, Canada. PRDM9-dependent recruitment of HELLS is required for activation of meiotic recombination and fertility.
4. Population, Evolutionary and Quantitative Genetics Conference. May 13-16th, 2018. Madison, WI. Tissue-specific *trans* regulation of the chromatin landscape.
5. Chromosome Architecture and Chromosome Organization. March 23-27th, 2018. Whistler, Canada. Tissue-specific *trans* regulation of the chromatin landscape.
6. Chromatin and Epigenetics. May 3-6th, 2017. EMBL Heidelberg, Germany. Genetic control of epigenetic landscape in germ cells.
7. The 29th International Mammalian Genome Conference. November 8-11th, 2015. Yokohama, Japan. Multimer formation explains allelic suppression of PRDM9 recombination hotspots.
8. The Biology of Genomes. May 5-9th, 2015. Cold Spring Harbor, NY. Poster. Multimer formation explains allelic suppression of PRDM9 recombination hotspots.
9. Gordon Research Conference: Meiosis. June 1-6th, 2014. New London, NH. PRDM9 drives evolutionary erosion of hotspots through haplotype-specific initiation of meiotic recombination.
10. Gordon Research Seminar: Meiosis. June 2-3, 2012. New London, NH. Poster presentation: PRDM9 dependent Histone H3 Lysine 4 trimethylation and DNA binding at human hotspots.
11. The 10th International Conference on Systems Biology 2009. August 30 – September 4, 2009. Stanford, CA. Quantitative proteomics investigation of the *Neurospora* circadian system.
12. Society for Research on Biological Rhythms 20th Anniversary Meeting. May 17-21, 2008. Sandestin, FL. Analysis of protein interactions in the *Neurospora crassa* circadian clock.
13. Neurospora 2008 Asilomar Meeting. March 27-30, 2008. Asilomar, CA. Characterization of the *Neurospora crassa* circadian clock interactome.
14. Albert J. Ryan Foundation Annual Meeting. May 2007. Holderness, NH. Multisite phosphorylation of a *Neurospora* circadian clock protein.

**Training and Mentorship**

*Predoctoral students*

2013-2014 Shimpei Kajita (1 year exchange programs for predoctoral students from Japan)

2017-2021 Candice Byers, Tufts University, Genetics Program

2020-current Haley Fortin, Tufts University, Genetics Program

*Rotations*

2017 winter Candice Byers, Tufts University

2018 spring Uma Aurora, Tufts University

2019 winter Ben Clauss, Tufts University

2020 winter Haley Fortin, Tufts University

2020 spring Elli Hartig, Tufts University

2022 spring Jaycee Choi, Tufts University

2022 fall Hilda Frempong, University of Maine

*Master’s thesis students*

2018-2021 Catrina Spruce, Molecular and Biomedical Sciences Program, University of Maine

*Post-baccalaureate Interns*

2017-2018 Sbonga Dlamini

2021- Anna Struba

*Summer student/interns*

2017, 2018 Naomi Bronkema, Swarthmore College

2019 Elizabeth Raruk

2021 Alex Mora

2022 Adel Misherghi, College of the Atlantic

Brooke Wilson, Southern Maine Community College

*Thesis Committee Membership*

2017-2019 Eraj Khokhar, GSBSE, University of Maine

2018-2019 Jessie Rochester, GSBSE, University of Maine

2018- Uma Aurora, School of Graduate Biomedical Sciences, Tufts University

2018-2022 Alex Stanton, School of Graduate Biomedical Sciences, Tufts University

2019- Ben Clauss, School of Graduate Biomedical Sciences, Tufts University \*committee chair

2022- Caryl Young, GSBSE, University of Maine

*Qualifying Exam Committee*

2018 Uma Aurora, Tufts University, Salwa Mostafa, Tufts University

2019 Ben Clauss, Tufts University \*committee chair

2022 Madison Armstrong, Tufts University, Sherrea Brown, Tufts University, Alexis Garretson, Tufts University, Xinru (Chelsy) Chen, Tufts University

**Teaching Experience**

2021-2022 Course Director, Mammalian Genetics, Tufts/GSBSE/The Jackson Laboratory, Bar Harbor, ME

2020-present Qualifying Exam Advisor, Tufts Medical School GSBS/The Jackson Laboratory, Bar Harbor, ME

2018-2019 Faculty advisor for Tufts Genetics journal club. Course #: GENE-0295-101

2016-2020 Instructor, Mammalian Genetics II, Tufts/GSBSE/The Jackson Laboratory, Bar Harbor, ME (*seminar course for first year graduate students. I teach one lecture and workshop on epigenetics and chromatin*)

2016 Instructor, Colby College Genomics Course, The Jackson Laboratory, Bar Harbor, ME (*two week course focusing on learning genomics, both laboratory and computational modules*)

2014-2017 Teaching Assistant and Instructor, Genetics I and Genetics II, The Jackson Laboratory, Bar Harbor, ME (*college level introductory genetics course for JAX employee’s*)

2013 Instructor, Topics in Biomedical Research, College of the Atlantic, Bar Harbor, ME

2013 Guest Lecturer, Genomics and Bioinformatics, Middlebury College, Middlebury VT

 course instructor Jeremy Ward, Ph.D.

2012 Instructor, Cutting Edge Techniques, The Jackson Laboratory summer student program, Bar Harbor, ME

2006 Teaching Assistant, Molecular Genetics of Prokaryotes and Lower Eukaryotes, Dartmouth College, Hanover, NH

2002 Teaching Assistant, CDAE Department course on computer applications, University of Vermont, Burlington, VT

**Service Committees**

2020-current Tufts Qualifying Exam Advisor for all first-year students

2019-2021 Tufts Graduate School JAX Genetics Program Admissions Committee

2018-current Faculty partner for Protein Purification and Production (PPP) core service at The Jackson Laboratory

2018-2020 Bioinformatics Training Working Group, The Jackson Laboratory. Committee to develop a plan to provide training for JAX researchers and trainees in computational and statistics skills.

2017-2019 Faculty retreat planning committee. Two-year term. \*committee chair 2019

2017-2020 Research Animal Facility Advisory Committee, The Jackson Laboratory

**Professional Membership**

American Association for the Advancement of Science

Genetic Society of America

International Mammalian Genome Society

 2020-present Nomination and Elections committee, 2022 \*committee chair

International Society for Stem Cell Research

**Scientific Community Service**

2017 University of Maine Student Symposium Judge, Cross Insurance Center, Bangor, ME

2015 Science Fair Judge, Conners Emmerson Elementary School, Bar Harbor, ME

2011 Maine State Science Fair Judge, grades 9-12, Bar Harbor, ME

2012-current Guest Speaker for science lessons at local elementary and middle schools

2009 Vermont State Science Fair Judge, grades 6-12, Norwich University, VT

2006 School-to-Career Mentor, service for high school students, NH

Ad hoc reviewer: Nature Communications, PLOS One, Philosophical Transactions B, Mammalian Genome, Genome Research, Stem Cell Reports, NPJ Regenerative Medicine, Development, Genetics, PLOS Genetics, Genome Biology

Ad hoc grant reviewer: Agence nationale de la recherche (The French National Research Agency, ANR), United Kingdom Research and Innovation: Medical Research Council, NSF CAREER Award