

Alex Handler Wagner, Ph.D. – Curriculum Vitae

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Personal Profile

My ongoing work is to integrate genomic evidence into knowledge used for interpreting genomic variation. My research program is defined by the development of several precision medicine web tools, for aggregating and disseminating genomic knowledge particularly in the cancer domain. These include the [Drug-Gene Interaction Database](#), the Clinical Interpretations of Variants in Cancer (CIViC) knowledgebase, and the [VICC meta-knowledgebase](#). My community science research program requires active engagement in professional consortia, and I currently hold advisory, membership, or leadership roles in several organizations, including the Variant Interpretation for Cancer Consortium (VICC) (as Director), the Global Alliance for Genomics and Health (GA4GH) (GKS Work Stream co-lead), the Atlas for Variant Effects Alliance (CVI and DCD committees), and ClinGen (Steering Committee, Somatic Cancer, Cancer Variant Interpretation, and Data Platform work groups). I also co-lead the development of the GA4GH [Variation Representation Specification](#).

Research Experience

Aug 2020 - Present	Institute for Genomic Medicine, Nationwide Children's Hospital <i>Principal Investigator</i>
Aug 2020 - Present	Departments of Pediatrics and Biomedical Informatics, the Ohio State University <i>Assistant Professor</i>
Jun 2019 - Jul 2020	Division of Oncology, Washington University in St. Louis <i>Instructor</i> Advisors: Obi L. Griffith, Ph.D.; Timothy J. Ley, M.D.
Jan 2015 - May 2019	McDonnell Genome Institute, Washington University in St. Louis <i>Postdoctoral Research Scholar</i> Advisors: Obi L. Griffith, Ph.D.; Ramaswamy Govindan, M.D.
Aug 2010 - Dec 2014	Coordinated Laboratory for Computational Genomics, University of Iowa <i>Graduate Research Assistant</i> Advisors: Terry A. Braun, Ph.D.; Edwin M. Stone, M.D., Ph.D.

Education

2010-2014	Ph.D, Computational Genetics - University of Iowa GPA: 3.90
2010-2013	Graduate Certificate, Bioinformatics - University of Iowa GPA: 3.96
2003-2008	BS, Biology (Minor: Mathematics) - Iowa State University GPA: 3.51 - Cum Laude

Peer-reviewed Publications

*Denotes Co-First Authorship

1. M Cannon, J Stevenson, K Stahl, R Basu, ..., **AH Wagner** (2024) "DGIdb 5.0: rebuilding the drug-gene interaction database for precision medicine and drug discovery platforms". *Nucleic Acids Research*.
2. M Cannon, J Stevenson, K Kuzma, S Kiwala, JL Warner, ..., **AH Wagner** (2023) "Normalization of drug and therapeutic concepts with Thera-Py". *JAMIA Open*.
3. R Dolin, B Heale, ..., **AH Wagner**, Srikar Chamala (2023) "Sync for Genes Phase 5: Computable artifacts for sharing dynamically annotated FHIR-formatted genomic variants". *Learning Health Systems*.
4. D Danis, JOB Jacobsen, **AH Wagner**, T Groza, ..., PN Robinson (2023) "Phenopacket-tools: Building and validating GA4GH Phenopackets". *PloS one*.
5. MS Ladewig, JOB Jacobsen, **AH Wagner**, D Danis, ..., PN Robinson (2023) "GA4GH Phenopackets: A Practical Introduction". *Advanced Genetics*.
6. K Krysiak, ..., **AH Wagner**, M Griffith, OL Griffith (2023) "CIViCdb 2022: evolution of an open-access cancer variant interpretation knowledgebase". *Nucleic Acids Research*.
7. KP Dalton, ..., **AH Wagner** (2022) "Accessing clinical-grade genomic classification data through the ClinGen Data Platform". *Pacific Symposium on Biocomputing*.
8. W Goar, ..., **AH Wagner** (2022) "Development and application of a computable genotype model in the GA4GH Variation Representation Specification". *Pacific Symposium on Biocomputing*.
9. Y Zhao, M Brush, C Wang, **AH Wagner**, H Liu, RR Freimuth (2022) "Leveraging a pharmacogenomics knowledgebase to formulate a drug response phenotype terminology for genomic medicine". *Bioinformatics*.
10. O Neumann, ..., **AH Wagner**, A Vogel, P Schirmacher, A Stenzinger, D Kazdal (2022) "Genomic architecture of FGFR2 fusions in cholangiocarcinoma and its implication for molecular testing". *British Journal of Cancer*.
11. JOB Jacobsen, ..., **AH Wagner**, JL Warner, C Weiland, MA Haendel, PN Robinson (2022) "The GA4GH Phenopacket schema defines a computable representation of clinical data". *Nature Biotechnology*.
12. K Krysiak, ..., **AH Wagner**, OL Griffith, M Griffith (2022) "A community approach to the cancer-variant-interpretation bottleneck". *Nature Cancer*.
13. N Matentzoglou, ..., **AH Wagner**, CJ Mungall (2022) "A Simple Standard for Sharing Ontological Mappings (SSSOM)". *Database*.
14. P Horak, et al. (2022) "Standards for the classification of pathogenicity of somatic variants in cancer (oncogenicity): Joint recommendations of Clinical Genome Resource (ClinGen), Cancer Genomics Consortium (CGC), and Variant Interpretation for Cancer Consortium (VICC)". *Genetics in Medicine*.
15. S LaHaye, ..., **Alex H Wagner**, AK Eisfeld, K Mrózek, V Magrini, CE Cottrell, ER Mardis, RK Wilson, P White (2021) "Discovery of clinically relevant fusions in pediatric cancer". *BMC Genomics*.
16. **AH Wagner**, Y Akkari, M Li, A Roy, K Tsuchiya, G Raca (2021) "Recommendations for future extensions to the HGNC gene fusion nomenclature". *Leukemia*.
17. HL Rehm, et al. (2021) "GA4GH: International policies and standards for data sharing across genomic research and healthcare". *Cell Genomics*.
18. **AH Wagner**, L Babb, ..., RK Hart (2021) "The GA4GH Variation Representation Specification: A computational framework for variation representation and federated identification". *Cell Genomics*.
19. A Stenzinger, ..., **AH Wagner**, F Tacke, D Capper, KR Müller, F Klauschen (2021) "Artificial intelligence and pathology: From principles to practice and future applications in histomorphology and molecular profiling". *Seminars in cancer biology*.

20. SL Freshour*, S Kiwala*, KC Cotto*, ..., **AH Wagner** (2021) "Integration of the Drug–Gene Interaction Database (DGIdb 4.0) with open crowdsource efforts". *Nucleic Acids Research*.
21. **AH Wagner**, B Walsh, G Mayfield, ..., M Griffith, OL Griffith, A Margolin (2020) "A harmonized meta-knowledgebase of clinical interpretations of somatic cancer genomic variants". *Nature Genetics*.
22. S Rao, B Pitel, **AH Wagner**, ..., M Griffith, OL Griffith, S Madhavan (2020) "Collaborative, Multidisciplinary Evaluation of Cancer Variants Through Virtual Molecular Tumor Boards Informs Local Clinical Practices". *JCO Clinical Cancer Informatics*.
23. **AH Wagner***, S Kiwala*, AC Coffman, JF McMichael, KC Cotto, TB Mooney, EK Barnell, K Krysiak, AM Danos, OL Griffith, M Griffith (2020) "CIViCpy: a Python Software Development and Analysis Toolkit for the CIViC knowledgebase". *JCO Clinical Cancer Informatics*.
24. G Alterovitz, B Heale, J Jones, D Kreda, F Lin, L Liu, X Liu, KD Mandl, DW Poloway, R Ramoni, **AH Wagner**, JL Warner (2020) "FHIR Genomics: enabling standardization for precision medicine use cases". *npj Genom Med*.
25. AM Danos*, K Krysiak*, EK Barnell*, ..., **AH Wagner**, S Madhavan, M Griffith, OL Griffith (2019) "Standard operating procedure for curation and clinical interpretation of variants in cancer". *Genome Med*.
26. EK Barnell, P Ronning, KM Campbell, K Krysiak, BJ Ainscough, C Ramirez, N Spies, J Kunisaki, ZL Skidmore, F Gomez, L Trani, M Matlock, **AH Wagner**, SJ Swamidass, M Griffith, OL Griffith (2019) "Standard operating procedure for somatic variant refinement of tumor sequencing data". *Genetics in Medicine*.
27. **AH Wagner***, S Devarakonda*, ZL Skidmore, K Krysiak, A Ramu, L Trani, J Kunisaki, ..., M Griffith, OL Griffith, R Govindan (2018) "Recurrent WNT Pathway Alterations are Frequent in Relapsed Small Cell Lung Cancer". *Nature Communications*.
28. BJ Ainscough*, EK Barnell*, KM Campbell, **AH Wagner**, TE Rohan, R Govindan, M Griffith, ER Mardis, SJ Swamidass, OL Griffith (2018) "A deep learning approach to automate refinement of somatic variant calling from cancer sequencing data". *Nature Genetics*.
29. AM Danos*, DI Ritter*, **AH Wagner**, K Krysiak, ..., S Kulkarni, M Griffith, S Madhavan, OL Griffith (2018) "Adapting crowdsourced clinical cancer curation in CIViC to the ClinGen minimum variant level data community-driven standards". *Human Mutation*.
30. K Cotto*, **AH Wagner***, YY Feng, S Kiwala, AC Coffman, G Spies, A Wollam, NC Spies, OL Griffith, M Griffith (2017) "DGIdb 3.0: a redesign and expansion of the drug-gene interaction database". *Nucleic Acids Research*.
31. M Griffith, NC Spies, K Krysiak, JF McMichael, AC Coffman, AM Danos, BJ Ainscough, CA Ramirez, DT Rieke, L Kujan, EK Barnell, **AH Wagner**, ..., OL Griffith (2017) "CIViC is a community knowledgebase for expert crowdsourcing the clinical interpretation of variants in cancer". *Nature Genetics*.
32. BJ Ainscough, M Griffith, AC Coffman, **AH Wagner**, J Kunisaki, MNK Choudhary, JF McMichael, RS Fulton, RK Wilson, OL Griffith, ER Mardis (2016) "DoCM: a database of curated mutations in cancer". *Nature methods*.
33. M Griffith, OL Griffith, K Krysiak, ZL Skidmore, MJ Christopher, JM Klco, A Ramu, TL Lamprecht, **AH Wagner**, ..., TJ Ley (2016) "Comprehensive genomic analysis reveals FLT3 activation and a therapeutic strategy for a patient with relapsed adult B-lymphoblastic leukemia". *Experimental hematology*.
34. ZL Skidmore, **AH Wagner**, R Lesurf, KM Campbell, J Kunisaki, OL Griffith, M Griffith (2016) "GenVisR: Genomic Visualizations in R". *Bioinformatics*.
35. **AH Wagner**, AC Coffman, BJ Ainscough, NC Spies, ZL Skidmore, KM Campbell, K Krysiak, D Pan, JF McMichael, JM Eldred, JR Walker, RK Wilson, ER Mardis, M Griffith, OL Griffith (2016) "DGIdb 2.0: mining clinically relevant drug/gene interactions". *Nucleic Acids Research*.
36. SS Whitmore, **AH Wagner**, AP DeLuca, AV Drack, EM Stone, BA Tucker, S Zeng, TA Braun, RF Mullins, TE Scheetz (2014) "Transcriptomic analysis across nasal, temporal, and macular regions of human neural retina and RPE/choroid by RNA-Seq". *Experimental Eye Research*.

37. TP Sharma, CM McDowell, Y Liu, **AH Wagner**, D Thole, BP Faga, RJ Workinger, TA Braun, AF Clark (2014) "Optic nerve crush induces spatial and temporal gene expression patterns in retina and optic nerve of BALB/c mice". *Molecular Neurodegeneration*.
38. TA Braun, RF Mullins, **AH Wagner**, J Andorf, R Johnston, B Bakall, AP DeLuca, G Fisherman, R Weleber, A Cideciyan, S Jacobson, V Sheffield, B Tucker, EM Stone (2013) "Non-exonic and synonymous variants in ABCA4 are an important cause of Stargardt disease". *Human Molecular Genetics*.
39. **AH Wagner**, KR Taylor, AP DeLuca, TL Casavant, RF Mullins, EM Stone, TE Scheetz, TA Braun (2013) "Prioritization of Retinal Disease Genes: An Integrative Approach". *Human Mutation*.
40. **AH Wagner**, VN Anand, W Wang, JE Chatterton, D Sun, AR Shepard, N Jacobson, L Pang, AP DeLuca, TL Casavant, TE Scheetz, RF Mullins, TA Braun, AF Clark (2013) "Exon-level expression profiling of ocular tissues". *Experimental Eye Research*.
41. AP DeLuca, **AH Wagner**, KR Taylor, B Faga, D Thole, VC Sheffield, ..., TA Braun (2011) "Sequencing and disease variation detection tools and techniques". *IEEE/ACS International Conference on Computer Systems and Applications (AICCSA)*.

Other Published Works

1. JA Arbesfeld, EY Da, ..., **AH Wagner** (2023) "Mapping MAVE data for use in human genomics applications". *bioRxiv*. [Manuscript Preprint]
2. **AH Wagner**, et al. (2019) "ga4gh/vr-spec: 1.0 GA4GH Approved". *Global Alliance for Genomics and Health*. [Technical Standard]
3. **AH Wagner**, K Krysiak, KM Campbell, EK Barnell (2019) "Cancer Genomics for the Clinician: Tumor Heterogeneity". *Springer Publishing Group*. [Book Chapter]

Funding: Research and Fellowship Grant Awards

2024 - Present	NIH NHGRI U24 Competitive Subaward - GREGoR Consortium U24HG011746 - Role: Subaward PI
2023 - Present	NIH NCI U24 - ClinGen Somatic Biomedical Knowledgebase U24CA275783 - Role: Co-I
2022 - Present	NIH OD U54 - Bridge2AI BRIDGE Center (Standards) U54HG012513 - Role: Standards Core co-Lead
2022 - 2023	NIH NHGRI R35 Supplement - AI-Ready gnomAD R35HG011949 - Role: PI
2021 - Present	NIH NHGRI R35 - Genomic Innovator R35HG011949 - Role: PI
2019 - Present	NIH NHGRI K99/R00 - Knowledge Harmonization R00HG010157 - Role: PI
2017 - 2019	NIH NCI F32 - Cancer Genomics F32CA206247 - Role: PI

Funding: Training Awards

2016 - 2017	NIH NCI T32 Postdoctoral Training - Cancer Biology T32CA009547 - Role: Trainee
2013 - 2014	NIH NCI T32 Predoctoral Training - Genetics T32GM008629 - Role: Trainee
2011 - 2013	NIH NCI T32 Predoctoral Training - Bioinformatics T32GM082729 - Role: Trainee

Research Recognition Awards

- 2021** **Genomic Innovator Award**
National Human Genome Research Institute, National Institutes of Health
 National recognition awarded for early career success in consortia-driven scientific research.
- 2018** **ICTS Precision Medicine Abstract Award**
Washington University in St. Louis
 The most outstanding research was selected from more than 75 applicants at the Washington University Precision Medicine Symposium to present to Eric Green, the director of the National Human Genome Research Institute (NHGRI).
- 2015** **D.C. Spiestersbach Dissertation Prize Nominee**
University of Iowa, Genetics Program Nomination
 This biennial award recognizes excellence in doctoral research. Each of the twenty biological/life sciences programs at the University of Iowa nominates one dissertation submitted between July 1, 2013 and June 30, 2015 to compete for the award.
- 2013** **Outstanding Student Research Award - Bioinformatics and Computational Biology**
University of Iowa, College of Engineering
 This annual award recognizes a single student in the College of Engineering for exemplary research in the fields of bioinformatics and computational biology.

Travel and Trainee Awards

- 2019** **Trainee Abstract Award**
 Curating the Clinical Genome Annual Conference - *Washington D.C.*
- 2018** **1st Place Student/Trainee Travel Award**
 Cancer Genomics Consortium Annual Conference - *Nashville, TN*
- 2018** **AACR-Bristol Myers Squibb Scholar-in-Training Award**
 American Association for Cancer Research - *Chicago, IL*
- 2013** **NSF Travel Grant**
 Intelligent Systems for Molecular Biology Annual Conference - *Berlin, Germany*
- 2013** **Graduate Student Senate Travel Grant**
 Association for Research in Vision and Ophthalmology Annual Conference - *Seattle, WA*

Presentations

[†]Oral Presentation; [‡]Invited Speaker; *Upcoming

- September 2024** [‡]*Clinical applications of GA4GH Genomic Knowledge Standards for scalable (re)interpretation of pediatric cancer variants*
GA4GH 12th Plenary - Melbourne, Australia
- September 2024** [‡]*Advancing AI-readiness of GREGoR data with GA4GH Genomic Knowledge Standards*
GREGoR Consortium Annual Meeting - Boston, MA
- July 2024** [‡]*Enhancing the AI-readiness of gnomAD with GA4GH Genomic Knowledge Standards*
NIH ODSS Special Session at ISMB - Montreal, Canada
- Mar 2024** [‡]*Advancing AI-readiness of the gnomAD database with GA4GH genomic knowledge standards*
NIH ODSS AI Supplement PI Meeting - Virtual
- Nov 2023** [‡]*Application of the VICC Oncogenicity Framework in a Pediatric Setting (Declined; medical)*
AMP Annual Conference - Salt Lake City, UT

Nov 2023	[‡] <i>Variation Representation (Declined; medical)</i> AMP Annual Conference - Salt Lake City, UT
Jun 2023	[‡] <i>Developing variant representation standards for scalable genomic medicine</i> EMQN Seminar Series - Virtual
Jan 2023	[†] <i>Accessing clinical-grade genomic classification data through the ClinGen Data Platform</i> Pacific Symposium on Biocomputing - Waimea, HI
Nov 2022	[‡] <i>Gene (Fusion) Product Nomenclature</i> AMP Annual Conference - Phoenix, AZ
Sep 2022	[‡] <i>A starter kit for interoperable genomic knowledge</i> GA4GH Plenary - Barcelona, Spain
Sep 2022	[‡] <i>Decoding the standards for genomic data sharing</i> NIH BISTI Seminar - Virtual
Aug 2022	[†] <i>A Unified Framework for Gene Fusion Representation</i> Cancer Genomics Consortium Annual Conference - St. Louis, MO
Jul 2022	[‡] <i>The GA4GH Variation Representation Specification</i> International Graph Genome Symposium 2022 Conference - Ascona, Switzerland
Jun 2022	[†] <i>Cross-consortia Recommendations for a Unifying Framework Supporting Curation and Representation of Gene Fusions</i> ClinGen Curating the Clinical Genome - Virtual
Apr 2022	<i>The GA4GH Variation Representation Specification</i> American Association for Cancer Research Annual Conference - New Orleans, LA
Jan 2022	[‡] <i>Pitfalls of Contemporary Genomic Variation Representation (TERI talk)</i> Pacific Symposium on Biocomputing 2022 - Waimea, HI
Nov 2021	[†] <i>The GA4GH Variation Representation Specification (VRS): a Computational Framework for the Precise Representation of Molecular Variation</i> AMIA Annual Conference - San Diego, CA
Oct 2021	[‡] <i>Genomic Knowledge Standards</i> GA4GH 9th Plenary Connect - Virtual
Sep 2021	[‡] <i>Phenopackets VRS Integration</i> GA4GH 9th Plenary - Virtual
Aug 2021	[†] <i>Multi-consortia initiative to standardize the representation and curation of oncogenic fusions</i> Cancer Genomics Consortium Annual Conference - Virtual
Jun 2021	[†] <i>The ClinGen Data Platform is Driving Interoperability of Expert Evidence through Standards for Computable Genomic Knowledge</i> ClinGen Retreat - Virtual
May 2021	[†] <i>Interoperability of ClinGen Expert Evidence through Standards for Computable Genomic Knowledge</i> ClinGen Curating the Clinical Genome - Virtual
Apr 2021	<i>Advancing Genomic Medicine through Standards for Computable Genomic Knowledge</i> NHGRI Research Conference - Virtual
Apr 2021	<i>A standard operating procedure for the curation of gene fusions</i> AACR Annual Conference - Virtual
Mar 2021	[‡] <i>Clinical Interoperability of Variant Evidence</i> Genomics in Health Implementation Forum - Virtual

Mar 2021	‡ <i>VRS 1.3 Planning and Implementation Guidance</i> GA4GH Implementation Connect - Virtual
Mar 2021	‡ <i>VCF / VRS Alignment</i> GA4GH Implementation Connect - Virtual
Dec 2020	‡ <i>ASH Precision Medicine Working Group: Integrating Variant Interpretations</i> ASH Annual Conference - Virtual
Nov 2020	† <i>Clinical variant harmonization and standards for knowledgebase interoperability (panelist)</i> AMIA Annual Conference - Virtual
Nov 2020	† <i>Clinical variant harmonization and standards for knowledgebase interoperability (panelist)</i> AMIA Annual Conference - Virtual
Sep 2020	‡ <i>VCF / VRS Alignment</i> GA4GH 8th Plenary - Virtual
Sep 2020	‡ <i>VA & VR Implementations: Practical Considerations and Emerging Draft Standards</i> GA4GH 8th Plenary - Virtual
Aug 2020	† <i>Integrating Cytogenomic and Sequence Variants</i> Cancer Genomics Consortium Annual Conference - Virtual
Apr 2020	† <i>Data harmonization standards from the Variant Interpretation for Cancer Consortium</i> AACR Annual Conference - Virtual
Nov 2019	‡ <i>Computational representation of patient data (panelist)</i> AMIA Annual Conference - Washington, D.C.
Aug 2019	‡ <i>Community resources for clinical variant classification in cancers</i> Cancer Genomics Consortium Annual Conference - Nashville, TN
Aug 2019	† <i>A new somatic variation model enables precise search strategies for clinical interpretations of patient tumors</i> Cancer Genomics Consortium Annual Conference - Nashville, TN
May 2019	† <i>Somatic variant curation standards enable improved identification of relevant clinical interpretations for tumor variants</i> Curating the Clinical Genome - Washington, D.C.
Apr 2019	‡ <i>Variant Interpretation for Cancer Consortium: Workstream Goals 2019</i> GA4GH Implementation Connect - Hinxton, United Kingdom
Mar 2019	<i>Cancer genome interpretation with CIViCpy</i> AACR Annual Conference - Atlanta, GA
Mar 2019	‡ <i>The Variant Interpretation for Cancer Consortium, a Genomic Knowledge Workstream Driver Project of the GA4GH</i> GA4GH-AMED Symposium - Sendai, Japan
Oct 2018	‡ <i>The Variant Interpretation for Cancer Consortium</i> GA4GH 6th Annual Plenary - Basel, Switzerland
Aug 2018	† <i>Coordinating variant interpretation knowledgebases improves clinical interpretation of genomic variants in cancers</i> Cancer Genomics Consortium Annual Conference - Nashville, TN
Apr 2018	<i>Standardization and coordination of variant interpretation knowledgebases improves clinical genome actionability</i> American Association for Cancer Research - Chicago, IL
Jun 2016	<i>The Drug Gene Interaction Database</i> Curating the Clinical Genome - Hinxton, United Kingdom

Feb 2016	<i>The Drug Gene Interaction Database</i> AGBT Annual Conference - Orlando, FL
Jul 2013	<i>Positive and Unlabeled Learning for Prioritization (PULP)</i> ISMB Annual Conference - Berlin, Germany
May 2013	<i>Positive and Unlabeled Learning for Prioritizing Candidate Variants in Retinal Degenerative Diseases</i> ARVO Annual Conference - Seattle, WA
May 2013	<i>Positive and Unlabeled Learning for Prioritizing Candidate Variants in Retinal Degenerative Diseases</i> BICB Industry Symposium - Minneapolis, MN
May 2012	<i>RNA Sequencing for Identification of Genetic Factors in Retinal Disease</i> ARVO Annual Conference - Ft. Lauderdale, FL
Aug 2011	<i>Using RNA Sequencing To Identify And Isolate Causative Genetic Factors In Retinal Disease</i> ISU / UIowa Joint Bioinformatics Retreat - Ames, IA

Instruction

Apr 2024	Workshop Organizer. Genomic Knowledge Standards 2-day Secret Agent Workshop. GA4GH Connect - <i>Ascona, Switzerland</i>
Feb 2024	Panelist. CV and Cover Letter Preparation - Professional Development Seminar Series. Cancer Genomics Consortium - <i>Virtual</i>
2023 - Present	Workshop Instructor. IGM Trainee Writing Workshop. Institute for Genomic Medicine - <i>Columbus, OH</i>
Aug 2023	Workshop Organizer. VICC/ClinGen/CIViC Hackathon. CGC Annual Meeting - <i>St. Louis, MO</i>
Oct 2022	Instructor. Functional Variant Interpretation - Advanced Course on Biomarkers in Precision Medicine. ESMO Asia - <i>Singapore</i>
Sep 2022	Workshop Organizer. The GA4GH Variation Representation Specification Hackathon. GA4GH Plenary 2022 - <i>Barcelona, Spain</i>
Aug 2022	Workshop Moderator. CGC Bioinformatics Workshop. Cancer Genomics Consortium Annual Meeting - <i>St. Louis, MO</i>
Apr 2022	Workshop Moderator. Genomic Knowledge Standards 2-day Spaghetti Western Workshop. GA4GH Connect 2022 - <i>Montreal, Canada</i>
Jul 2021	Panelist. Clinical Genomics Career Panel. ClinGen - <i>Virtual</i>
Jun 2021	Workshop Organizer. Variation Representation Specification (VRS) Workshop. Genomics in Health Implementation Forum - <i>Virtual</i>
Jun 2021	Instructor. Harmonizing Variant Interpretation - Advanced Course on Biomarkers in Precision Medicine. ESMO Asia - <i>Singapore</i>
Dec 2019	Guest Lecturer. Variant Interpretation. University of Zurich - <i>Zurich, Switzerland</i>
Aug 2019	Guest Lecturer. Bioinformatics: Understanding what's Underneath the Hood. Cancer Genomics Consortium Annual Conference - <i>Nashville, TN</i>
Mar 2019	Workshop Leader. Genomic Knowledge Standards. AMED-GA4GH Workshop - <i>Sendai, Japan</i>

Nov 2018 **Teaching Assistant & Lecturer.** Advanced Sequencing Technologies and Applications. Cold Spring Harbor Laboratories - *Cold Spring Harbor, NY*

Aug 2018 **Workshop Instructor.** Introduction to bioinformatics learning resources. Cancer Genomics Consortium Annual Conference - *Nashville, TN*

Aug 2017 - Jun 2019 **Workshop Instructor.** Escape from Perlglatory: Developing in Python and Ruby. McDonnell Genome Institute - *Saint Louis, MO*

Nov 2017 **Teaching Assistant & Lecturer.** Advanced Sequencing Technologies and Applications. Cold Spring Harbor Laboratories - *Cold Spring Harbor, NY*

Nov 2016 **Workshop Instructor.** CIViC Hackathon. Netherlands Cancer Institute (NKI) - *Amsterdam, Netherlands*

Nov 2016 **Teaching Assistant & Lecturer.** Advanced Sequencing Technologies and Applications. Cold Spring Harbor Laboratories - *Cold Spring Harbor, NY*

Aug 2014 - Dec 2014 **Teaching Assistant.** Bioinformatics Techniques. Department of Biomedical Engineering, University of Iowa - *Iowa City, IA*

Jan 2014 - May 2014 **Teaching Assistant.** Bioimaging and Bioinformatics. Department of Biomedical Engineering, University of Iowa - *Iowa City, IA*

Jan 2014 - May 2014 **Teaching Assistant.** Computational Genomics. Department of Biomedical Engineering, University of Iowa - *Iowa City, IA*

Aug 2013 - Dec 2013 **Teaching Assistant.** Bioinformatics Techniques. Department of Biomedical Engineering, University of Iowa - *Iowa City, IA*

Aug 2013 - Dec 2013 **Instructor.** Introduction to Bioinformatics Computing with Python. Department of Biomedical Engineering, University of Iowa - *Iowa City, IA*

Service

2024 - Present **Site PI - U24CA275783 (WUSTL/NCH/Glasgow).** ClinGen Steering Committee.
As site PI at Nationwide Children's Hospital, I lead the data dissemination and standardization aims of the ClinGen somatic cancer variant curation grant.

2024 - Present **Work Stream Co-Lead.** GA4GH Genomic Knowledge Standards Work Stream.
The GA4GH Genomic Knowledge Standards Work Stream is responsible for development of standards for the FAIR sharing of genomic knowledge such as variant classifications. Work Stream co-leads are nominated by the community and are selected by the GA4GH executive committee.

2022 - Present **Elected Member.** HGVS Variation Nomenclature Committee (HVNC).
The HGVS Variation Nomenclature Committee (HVNC) is responsible for the development and maintenance of the HGVS nomenclature for describing sequence variation. Members are elected for 4 year terms.

2022 - 2025 **Member.** Cancer Genomics Consortium Annual Meeting Program Committee.
This committee sets the program for the annual meeting of the Cancer Genomics Consortium. I have served on this committee for 3 subsequent years: a member in 2022-23 and 2023-24, and an invited advisor for 2024-25.

2019 - Present **Appointed Lead.** Variation Representation.
The Variation Representation group of the Genomic Knowledge Standards Work Stream is responsible for developing and maintaining standards for the representation of variation as used in genomic knowledgebases. This position is by appointment of the Work Stream leads.

- 2019 - Present** **Director.** Variant Interpretation for Cancer Consortium.
- The Variant Interpretation for Cancer Consortium works to standardize the curation, representation, and interpretation of clinically-relevant evidence associated with genomic variation in cancers. VICC Directors are responsible for guiding the organization through policy, outreach, and oversight. New directors are appointed by sitting directors.*
- 2015 - Present** **Editor.** Clinical Interpretation of Variants in Cancer (CIViC) knowledgebase.
- The CIViC knowledgebase is used to curate information about genetic variants and their role in informing clinical interpretation of cancers. Platform editors are content and platform experts that review and approve submissions by community members. I have moderated over 450 entries on the CIViC platform.*
- 2020 - Present** **Appointed Member.** ClinGen Committees – Data Platform, PGx, and CVI.
- ClinGen committees are groups of experts that are invited to participate in a particular domain. Data Platform is a team focused on genomic knowledge interoperability, Pharmacogenomics (PGx) is a team focused on pharmacogenomic knowledge standardization, and Cancer Variant Interpretation (CVI) focuses on the development of rules for use by Somatic Cancer Variant Curation Expert Panels (SC-VCEPs).*
- 2019 - Present** **Appointed Member.** Atlas of Variant Effects Alliance (AVE) Committees – CVI and DCD.
- AVE committees are groups of experts that are invited to participate in a particular domain. Clinical Variant Interpretation (CVI) is a committee focused on applying MAVE data to clinical interpretation, Data Coordination and Dissemination (DCD) is a committee focused on MAVE data standardization and sharing.*
- 2020 - 2022** **Appointed Member.** ASH Precision Medicine Somatic Working Group.
- The ASH Precision Medicine Working Group focused on the integration of variant interpretation data for hematological malignancies from different institutions.*