Kelsey E. Grinde

CONTACT	Mathematics, Statistics, & Computer Science Macalester College 1600 Grand Avenue Saint Paul, MN 55105	kgrinde@macalester.edu (651)-696-6976 kegrinde.github.io
EDUCATION	Ph.D. in Biostatistics University of Washington, Seattle, WA	2019
	Dissertation: Statistical inference in admixed populations. Advisor: Sharon Browning, Ph.D.	s
	B.A. in Mathematics , Concentration in Statistics St. Olaf College, Northfield, MN	2014
	Graduated summa cum laude with Distinction in Statist Advisor: Paul Roback, Ph.D.	ics
WORK EXPERIENCE	Assistant Professor Department of Mathematics, Statistics, & Computer Science Macalester College, Saint Paul, MN	2020-present
	Postdoctoral Teaching Fellow Department of Mathematics, Statistics, & Computer Science Macalester College, Saint Paul, MN	2019–2020
	Graduate Research Assistant Browning Statistical Genetics Lab University of Washington, Seattle, WA	2014–2019
	Graduate Research Assistant Genetic Analysis Center University of Washington, Seattle, WA	2015–2016
	Undergraduate Research Assistant Summer Research Program in Statistical Genetics & Biostatis Dordt College, Sioux Center, IA	2013, 2014 tics
	Undergraduate Research Fellow Center for Interdisciplinary Research St. Olaf College, Northfield, MN	2013–2014
TEACHING EXPERIENCE	 Macalester College STAT 253: Statistical Machine Learning (3 sections) STAT 155: Introduction to Statistical Modeling (13 sect MATH/STAT 455: Mathematical Statistics (4 sections) STAT 494: Statistical Genetics (1 section) 	(upcoming: Fall 2024) ions) 2019–2023 2020–2023 2022
	University of WashingtonBIOST 311: Regression Methods in the Health Sciences,	Co-Instructor 2018

 \bullet BIOST 310: Biostatistics for the Health Sciences, Teaching Assistant

2017

•	BIOST 561: Computational Skills for Biostatistics, Guest Lecturer	2017
•	BIOST 550: Statistical Genetics I, Guest Lecturer	2017
•	BIOST 570: Regression Methods for Independent Data, Teaching Asst.	2016
•	First Year Statistical Theory Exam Review Sessions, Co-Instructor	2016

St. Olaf College

• STAT 322: Statistical Theory, Grader

2013

2011 - 2012

• Academic Support Center, Tutor & Academic Assistant (Intermediate Spanish I & II, Calculus I, Abstract Algebra I)

• Urban Schools and Communities Program, Participant 2012

• Department of Mathematics, Statistics, and Computer Science, Tutor (Calculus I & II, Multivariable Calculus)

PUBLICATIONS

- * denotes an undergraduate student
- + denotes joint first authors

Refereed Journal Articles

- Horimoto, A., Boyken, L., Blue, E., Grinde, K., Nafikov, R., Sohi, H., Nato, A., Bis, J., Brusco, L., Morelli, L., Ramirez, A., Dalmasso, M., Temple, S., Satizabal, C., Browning, S., Seshadri, S., Wijsman, E., & Thornton, T. "Admixture mapping implicates LIG4, MYO16, and FAM155A at 13q33.3 as ancestry-of-origin loci for Alzheimer disease in Hispanic and Latino populations." HGG Advances 4.3 (2023): 1000207. [link]
- 13. Barragan, F.*, Mills, L., Raduski, A., Marcotte, E., **Grinde, K.**, Spector, L., & Williams, A. "Genetic ancestry, differential gene expression, and survival in pediatric b-cell acute lymphoblastic leukemia." *Cancer Medicine* 12.4 (2023): 4761–4772. [link]
- 12. Zucko, D., Hayir, A.*, **Grinde, K.**, & Boris-Lawrie, K. "Circular RNA Profiles in Viremia and ART Suppression Predict Competing circRNA– miRNA–mRNA Networks Exclusive to HIV-1 Viremic Patients." *Viruses* 14.4 (2022): 683. [link]
- Lin, B.+, Grinde, K.+, Brody, J., Breeze, C., Raffield, L., Mychaleckyj, J., Thornton, T., Perry, J., Baier, L., de Las Fuentes, L., Guo, X., Heavner, B., Hanson, R., Hung, Y.-J., Qian, H., Hsiung, C., Hwang, S.-J., Irvin, M., Jain, D., Kelly, T., Kobes, S., Lange, L., Lash, J., Li, Y., Liu, X., Mi, X., Musani, X., Papanicolaou, G., Parsa, A., Reiner, A., Salimi, S., Sheu, W., Shuldiner, A., Taylor, K., Smith, A., Smith, J., Tin, A., Vaidya, D., Wallace, R., Yamamoto, K., Sakaue, S., Matsuda, K., Kamatani, Y., Momozawa, Y., Yanek, L., Young, B., Zhao, W., Okada, Y., Abecasis, G., Psaty, B., Arnett, D., Boerwinkle, E., Cai, J., Chen, I., Correa, A., Cupples, L.A., He, J., Kardia, S., Kooperberg, C., Mathias, R., Mitchell, B., Nickerson, D., Turner, S., Ramachandran, V., Rotter, J., Levy, D., Kramer, H., Köttgen, A., Rich, S., Lin, D.-Y., Browning, S., Franceschini, N., & TOPMed Kidney Working Group. "Whole genome sequence analyses of eGFR in 23,732 people representing multiple ancestries in the NHLBI Trans-Omics for Precision Medicine (TOPMed) consortium." eBioMedicine 63 (2021): 103157. [link]
- 10. Raffield, L., Lu, A., Szeto, M., Little, A., Grinde, K., Shaw, J., Auer, P., Cushman, M., Horvath, S., Irvin, M., Lange, E., Lange, L., Nickerson, D., Thornton, T., Wilson, J., Wheeler, M., NHLBI TOPMed Consortium, TOPMed Hematology & Hemostasis Working Group, Zakai, N., & Reiner, A. "Coagulation factor VIII: Relationship to cardiovascular disease risk and whole genome sequence

- and epigenome-wide analysis in African Americans." *Journal of Thrombosis and Haemostasis* 18.6 (2020): 1335–1347. [link]
- Shungin, D., Haworth, S., Divaris, K., Agler, C., Kamatani, Y., Lee, M.K., Grinde, K., Hindy, G., Alaraudanjoki, V., Pesonen, P., Temuer, A., Holtfreter, B., Sakaue, S., Hirata, J., Yu, Y.H., Ridker, P., Giulianini, F., Chasman, D., Magnusson, P., Sudo, T., Okada, Y., Voelker, U., Kocher, T., Anttonen, V., Laitala, M.L., Orho-Melander, M., Sofer, T., Shaffer, J., Vieira, A., Marazita, M., Kubo, M., Furuichi, Y., North, K., Offenbacher, S., Ingelsson, E., Franks, P., Timpson, N., Johansson, I. "Genome-wide analysis of dental caries and periodontal disease combining clinical and self-reported data." Nature Communications 10.1 (2019): 2773. [link]
- 8. Sofer, T., Zheng, X., Gogarten, S.M., Laurie, C.A., **Grinde, K.**, Shaffer, J.R., Shungin, D., O'Connell, J.R., Durazo-Arvizo, R.A., Raffield, L., Lange, L., Musani, S., Vasan, R.S., Cupples, L.A., Reiner, A.P., Laurie, C.C., Rice, K.M. "A fully-adjusted two-stage procedure for rank normalization in genetic association studies." *Genetic Epidemiology* 43.3 (2019): 263–275. [link]
- 7. **Grinde, K.**, Brown, L., Reiner, A., Thornton, T., Browning, S. "Genome-wide significance thresholds for admixture mapping studies." *American Journal of Human Genetics* 104 (2019): 454–465. [link]
- Grinde, K., Qi, Q., Thornton, T., Liu, S., Shadyab, A.H., Chan, K.H.K., Reiner, A.P., & Sofer, T. "Generalizing polygenic risk scores from Europeans to Hispanics/Latinos." Genetic Epidemiology 43.1 (2019): 50–62. [link]
- 5. **Grinde, K.**, Green, A., Arbet, J., O'Connell, M., Valcarcel, A., Westra, J., & Tintle, N. "Illustrating, quantifying and correcting for bias in post-hoc analysis of gene-based rare variant tests of association." Frontiers in Genetics 8.117 (2017): 1–11. [link]
- 4. Browning, S.R., **Grinde, K.**, Plantinga, A., Gogarten, S.M., Stilp, A.M., Kaplan, R.C., Avilés-Santa, L., Browning, B.L., & Laurie, C.C. "Local ancestry inference in a large US-based Hispanic/Latino study: Hispanic Community Health Study/Study of Latinos (HCHS/SOL)." *G3: Genes|Genomes|Genetics* 6.6 (2016): 1525–1534. [link]
- 3. Greco, B., Hainline, A., Arbet, J., **Grinde, K.**, Benitez, A., & Tintle, N. "A general approach for combining diverse rare variant association tests provides improved robustness across a wider range of genetic architectures." *European Journal of Human Genetics* 24 (2016): 767–773. [link]
- 2. Green, A., Cook, K., **Grinde, K.**, Valcarcel, A., & Tintle, N. "A general method for combining different family-based rare-variant tests of association to improve power and robustness of a wide range of genetic architectures." *BioMed Central Proceedings* 10.7.23 (2016): 165–170. [link]
- Valcarcel, A., Grinde, K., Cook, K., Green, A., & Tintle, N. "A multistep approach to single nucleotide polymorphism—set analysis: An evaluation of power and type I error of gene-based tests of association after pathway-based association tests." BioMed Central Proceedings 10.7.16 (2016): 349–355. [link]

Refereed Abstracts

 Jensen-Otsu, E., Grinde, K., Baxi, A., Harms, M., Teng, B., Strate, L.L., & Ko, C.W. "Anesthesia professional-delivered sedation is associated with similar outcomes compared to nurse administered sedation in patients admitted with acute upper gastrointenstinal bleeding." Gastrointenstinal Endoscopy 87.6S (2018): AB418-AB419. [link]

Open Education Resources

2. **Grinde**, **K**. "Rethinking grading systems in introductory and advanced statistics courses." Online resource page (2023): [link].

Submitted to Consortium for the Advancement of Undergraduate Statistics Education Resources for JEDI-Informed Teaching of Statistics in 2024.

1. Heggeseth, B., Myint, L., & **Grinde**, **K.** "Stat 155 Notes." Online textbook (2021): https://bcheggeseth.github.io/Stat155Notes/.

Other Writing

- 2. **Grinde, K.**⁺, Theobold, A.⁺, & Myint, L⁺. "Beyond Achievement: Access, Identity, and Power in Alternative Grading." *Grading for Growth* blog guest post (2024): [link].
- 1. **Grinde**, **K.** "Statistical Inference in Admixed Populations." Doctoral dissertation, University of Washington. 2019. [link].

SUBMITTED MANUSCRIPTS

1. **Grinde, K.**, Browning, B., Reiner, A., Thornton, T., & Browning, S. "Adjusting for principal components can induce spurious associations in genome-wide association studies in admixed populations." Under Review at *PLOS Genetics*. Preprint available [here].

SOFTWARE & APPLICATIONS

- 3. Hayir, A.*, & Grinde, K. "Interactive Circos Tool." R shiny application (2022): https://kblcircosgraph.shinyapps.io/circos/.
- 2. Huang, Z.*, & **Grinde**, **K.** "Significance Threshold Estimation for Admixture Mapping using Rcpp." R package (2020): https://github.com/GrindeLab/STEAMcpp.
- 1. **Grinde**, **K.** "STEAM: Significance Threshold Estimation for Admixture Mapping." R package (2019): https://github.com/kegrinde/STEAM.

RESEARCH TALKS

Presentations at International or National Venues

- 10. Adjusting for principal components can induce spurious associations in genomewide association studies in admixed populations. International Genetic Epidemiology Society Annual Meeting. Virtual. 2021. (Presentation Award Winner)
- 9. Deriving significance thresholds for genome-wide admixture mapping studies. International Genetic Epidemiology Society Annual Meeting. San Diego, CA. 2018.
- 8. Controlling for multiple testing in genome-wide admixture mapping studies. Western North American Region of the International Biometric Society Meeting. Edmonton, Canada. 2018. (Presentation Award Winner)
- 7. Admixture mapping: controlling for false positives in the presence of population structure. American Society of Human Genetics Annual Meeting. Orlando, FL. 2017. (Poster)
- 6. Generalizing genetic risk scores from Europeans to Hispanics/Latinos. International Genetic Epidemiology Society Annual Meeting. Cambridge, United Kingdom. 2017. (Poster)
- 5. Illustrating, quantifying, and correcting for bias in post-hoc analysis of gene-based rare variant tests of association. Joint Statistical Meetings. Seattle, WA. 2015. (Poster)
- 4. A hierarchical approach to SNP-set analysis: an evaluation of power and type I error of gene-based tests of association after pathway-based analysis. Genetic Analysis Workshop 19. Vienna, Austria. 2014.
- 3. Accounting for variability in paleoecological mixing models. National Conference for Undergraduate Research. Lexington, KY. 2014.

- 2. What now? Post-hoc approaches for gene-based, rare variant tests of association. American Society of Human Genetics Annual Meeting. Boston, MA. 2013. (Poster)
- 1. General approaches for combining multiple rare variant association tests provide improved power across a wider range of genetic architectures. American Society of Human Genetics Annual Meeting. Boston, MA. 2013. (Poster)

Presentations at Regional or Local Venues

- 24. Using PCA to infer and adjust for population structure: what can go wrong? Twin Cities Pop/EvoGen Group, University of Minnesota. Minneapolis, MN. 2024. (Invited)
- 23. Statistical methods for genetic studies in admixed populations. Carleton College Math/Stats Colloquium. Northfield, MN. 2023. (Invited)
- 22. Statistical genetics in populations with mixed ancestry. Creighton University Department of Mathematics. Omaha, NE. 2022. (Invited)
- 21. What's our work: statistical genetics. Macalester College Mathematics, Statistics, and Computer Science Seminar. Saint Paul, MN. 2021.
- 20. Genome-wide significance thresholds for admixture mapping studies. University of Minnesota Interdisciplinary Biostatistics Training in Genetics and Genomics Journal Club. Virtual. 2021. (Invited)
- 19. Statistical genetics in populations with mixed ancestry. Augsburg University Mathematics Colloquium. Virtual. 2020. (Invited)
- 18. Statistical methods for genome-wide admixture mapping studies. University of Minnesota Division of Pediatric Epidemiology and Clinical Research. Virtual. 2020. (Invited)
- 17. Statistical genetics in populations with mixed ancestry. Macalester College Department of Mathematics, Statistics, and Computer Science. Saint Paul, MN. 2019. (Invited)
- 16. Statistial inference in populations with mixed ancestry. Department of Mathematics, Statistics, and Computer Science, St. Olaf College. Northfield, MN. 2019. (Invited)
- 15. Adjusting for principal components can induce spurious associations in genome-wide association studies. Genetic Analysis Center. Seattle, WA. 2019. (Invited)
- 14. Adjusting for population structure in genetic association studies: new insights and the potential pitfalls of using PCs. University of Washington Popgen Lunch. Seattle, WA. 2019. (Invited)
- 13. Statistical inference in populations with mixed ancestry. University of Washington Biostatistics Colloquium. Seattle, WA. 2018. (Invited)
- 12. Admixture mapping in TOPMed. NHLBI Trans-Omics for Precision Medicine (TOPMed) Kidney Working Group. Virtual. 2018.
- 11. Admixture mapping: controlling for false positives in the presence of population structure. Biostatistics Department Retreat, University of Washington. Seattle, WA. 2017. (Poster)
- 10. Issues in implementation of local ancestry inference on the X chromosome. Omics in Latinos Genetic Analysis Center Meeting. Seattle, WA. 2015.
- 9. Estimating genetic maps with large data sets. Biostatistics Department Retreat, University of Washington. Blaine, WA. 2015. (Poster)
- 8. Identifying and correcting for bias in post-hoc ranking strategies: an application to gene-based rare variant tests of association. Dordt College Summer Seminar. Sioux Center, IA. 2014.

- A hierarchical approach to SNP-set analysis: evaluation of power and type I error
 of gene-based tests of association after pathway-based analysis. Dordt College
 Summer Seminar. Sioux Center, IA. 2014.
- Identifying and correcting for bias in post-hoc ranking strategies: an application
 to gene-based rare variant tests of association. University of Michigan Department of Biostatistics. Ann Arbor, MI. 2014.
- A hierarchical approach to SNP-set analysis: evaluation of power and type I error of gene-based tests of association after pathway-based analysis. University of Michigan Department of Biostatistics. Ann Arbor, MI. 2014.
- 4. What now? Post-hoc approaches for gene-based, rare variant tests of association. Great Plains R-Users Group Conference. Sioux Center, IA. 2014. (Poster)
- 3. Accounting for variability in paleoecological mixing models. St. Olaf Natural Sciences and Mathematics Honors' Day Poster Session. Northfield, MN. 2014. (Poster)
- 2. Predicting donors at Red Cross blood drives. St. Olaf Mathematics, Statistics, and Computer Science Colloquium. Northfield, MN. 2014.
- Predicting donors at Red Cross blood drives. American Red Cross. Saint Paul, MN. 2014.

Student Presentations of Joint/Supervised Work

- Barragan, F. Genetic ancestry, gene expression, and survival in children with B-ALL. Pediatric Research, Education, & Scholarship Symposium. Minneapolis, MN. 2022. (Poster)
- Barragan, F. Gene expression differences by race and genetic ancestry in B-cell acute lymphoblastic leukemia. American Society of Human Genetics Annual Meeting. Virtual. 2021. (Poster)
- 5. Barragan, F. Characterizing racial disparities in pediatric cancer: ancestry, gene expression, and survival disparities in B-cell acute lymphoblastic leukemia. Underrepresented Students in STEM Symposium. Minneapolis, MN. 2021. (Poster)
- 4. Barragan, F. Statistical methods for pediatric leukemia: gene expression & ancestry in B-cell acute lymphoblastic leukemia. Macalester Summer Research Showcase. Saint Paul, MN. 2021. (Poster)
- 3. Huang, Z. Statistical methods for genetic association studies in populations with mixed ancestry. Midstates Consortium Research Symposium. Virtual. 2020.
- 2. Huang, Z. Using Rcpp to speed up tool for controlling for multiple testing in genetic studies. Electronic Undergraduate Statistics Research Conference. Virtual. 2020.
- 1. Huang, Z.* "Statistical methods for genetic association studies in populations with mixed ancestry." Macalester Summer Research Showcase. Virtual. 2020. (Poster)

TEACHING, OUTREACH, & MENTORING TALKS

Presentations at International or National Venues

- 4. Panel discussion on academic careers and job search. American Statistical Association Section on Statistics in Genomics and Genetics. Virtual. 2023. (Invited)
- 3. Rethinking (and then rethinking some more) grading systems in introductory and advanced statistics courses. Joint Statistical Meetings. Toronto, Canada. 2023. (Invited)
- 2. Time management, research strategy, and healthy habits for graduate students. American Statistical Association Section on Statistics in Genomics and Genetics. Virtual. 2021. (Invited)

1. Graduate programs in (bio)statistics. Electronic Undergraduate Statistics Research Conference. Virtual. 2020. (Invited)

Presentations at Regional or Local Venues

- 21. Teaching careers roundtable. BIOS 834: Pedagogical Methods for Biostatistics Courses, University of Michigan. Virtual. 2024. (Invited)
- Career discussion. Gender Minorities in Math and Statistics (GeMMS), Carleton College. Northfield, MN. 2024. (Invited)
- 19. Alternative grading strategies. MSCS Inclusive Pedagogy Summit, Macalester College. St. Paul, MN. 2023. (Invited)
- 18. Faculty panel. Preparing Future Faculty Practicum, University of Minnesota. Virtual. 2023. (Invited)
- 17. Tips and tricks with R/RStudio. MSCS Student Advisory Board Skill-Building Sessions, Macalester College. St. Paul, MN. 2023. (Invited)
- Open Educational Resources and textbook affordability: Macalester environmental scan and survey results. Jan Serie Center for Scholarship and Teaching, Macalester College. St. Paul, MN. 2023.
- 15. Inclusivity in teaching panel. Radical MacACCESS, Macalester College. Virtual. 2021. (Invited)
- 14. Pathways into science outreach panel. Fred Hutchinson Cancer Research Center Hutch United Outreach Committee & Wallin Education Partners Program. Virtual. 2021. (Invited)
- 13. Genetic testing: how does it work? (a statistician's perspective). Department of Mathematics, Statistics, and Computer Science, St. Olaf College. Northfield, MN. 2019. (Invited)
- 12. (Bio)statistics PhD programs: application tips and research opportunities. St. Olaf College Biostatistics Class. Northfield, MN. 2019. (Invited)
- 11. Fellowships, scholarships, and grants. University of Washington Biostatistics Student Seminar. Seattle, WA. 2018.
- 10. Admixture mapping: controlling for false positives in the presence of population structure. StatNorthwest. Seattle, WA. 2018. (Poster)
- 9. Graduate student panel. StatNorthwest. Seattle, WA. 2018. (Invited)
- 8. Travel grants and conference funding. University of Washington Department of Biostatistics. Seattle, WA. 2017.
- What is Biostatistics? Forest Ridge School of the Sacred Heart Science Research Class. Bellevue, WA. 2017.
- NSF Graduate Research Fellowship Program information session. University of Washington Department of Biostatistics. Seattle, WA. 2017.
- 5. What is Biostatistics? 7th and 8th Grade STEM PREP Project. Seattle, WA. 2017.
- 4. Applying for outside funding opportunities. University of Washington Biostatistics Student Seminar. Seattle, WA. 2016.
- 3. Graduate and professional student panel. Healthcare Exploration for Youth Program. Seattle, WA. 2016. (Invited)
- 2. Graduate and professional student panel. Healthcare Exploration for Youth Program. Seattle, WA. 2015. (Invited)
- 1. What now? Post-hoc approaches for gene-based, rare variant tests of association. Inter-Disciplinary Explorations Across the Sciences. Sioux Center, IA. 2014. (Poster)

HONORS &	Professional Awards and Recognition	
AWARDS	• Poster/Lightning Talk Award, 2nd Place (for International Research Talk [10]) International Genetic Epidemiology Society Annual Meeting	2021
	• Top Cited Article (for Refereed Journal Article [6]) Genetic Epidemiology Journal	2021
	• Thomas R. Fleming Excellence in Biostatistics Award University of Washington Department of Biostatistics	2019
	• Gertrude M. Cox Scholarship American Statistical Association	2018
	• Dorothy L. Simpson Leadership Award Achievement Rewards for College Scientists Foundation, Seattle Chapter	2018
	• Excellence in Teaching Award University of Washington Department of Biostatistics	2018
	• Distinguished Oral Presentation Award (for International Research Talk [8]) Western North American Region of the International Biometric Society	2018
	• Achievement Rewards for College Scientists (ARCS) Fellowship 201 ARCS Foundation, Seattle Chapter	4-2017
	• Donovan J. Thompson Award (for best score on Ph.D. qualifying exams) University of Washington Department of Biostatistics	2016
	Grants and Other Research Funding	
	• Collaborative Summer Research Award Amount: \$13,461 Funder: Macalester College	2024
	• Article Processing Charge Grant (for Refereed Journal Article [12]) Amount: \$1,466 Funder: Macalester College Dewitt Wallace Library Open Access Fund	2022
	• Collaborative Summer Research Award, Amount: \$5,500 Funder: Macalester College	2020
	<u> </u>	6-2019
	• Travel Grant Amount: \$300 Funder: University of Washington Graduate and Professional Student Se	2018
	• Conference Travel Award Amount: \$1,000 Funder: University of Washington Department of Biostatistics	2018
	 Travel Award Amount: \$500 Funder: University of Washington Graduate School Fund for Excellen Innovation 	2017 ce and
		5-2016
	Undergraduate Awards • Undergraduate Research Project Competition Honorable Mention Concertium for Advancement of Undergraduate Statistics Education	2014
	Consortium for Advancement of Undergraduate Statistics Education • Statistically Significant Award St. Olaf College	2014

St. Olaf College

	Buntrock Scholarship Charles College	2010-2014
	St. Olaf College • Service Leadership Scholarship St. Olaf College	2010-2014
	Phi Beta Kappa National Honor Society	2013
	• Pi Mu Epsilon National Honor Society	2013
	V 11 Ma Eponon Mational Honor Society	2019
SERVICE	Macalester College	
	• Team Member, AAC&U Open Educational Resources Institute	2022 - 2023
	• Faculty Liaison, Admissions	2022 - 2023
	• MSCS Academic Planning Committee	2022 - 2023
	• Co-Creator and Co-Coordinator, MSCS Honors Seminar	2021 - 2023
	• Mentor, DataFest 2021	, 2022, 2023
	• Statistics Tenure Track Search Committee (hired Taylor Okonek)	2022
	• Statistics Visiting/Postdoc Search Committee	2020-2022
	(hired Bryan Martin [2021], James Normington and Laura Lyman [2022])	
	• Scribe, Mid-Course Interview	2021
	University of Washington Department of Biostatistics	
	• Member, Diversity Committee	2017 - 2019
	• Leadership Team, Women in Biostatistics and Statistics	2017-2018
	• Member, Admissions Committee	2017-2018
	• Founding Member, Peer Mentoring Program	2016-2018
	• Member, Educational Policy and Teaching Evaluation Committee	2016-2017
	• Member, Biostatistics Outreach Working Group	2015
	St. Olaf College	
	• President, Spanish Honor House	2013-2014
	• Volunteer Teaching Assistant & Tutor, Northfield Public Schools	2011-2014
	• Volunteer Teaching Assistant, Wayzata High School	2011
PROFESSIONAL ACTIVITIES	Journal Editorial Board Positions and Peer Review • Review Editor for the Statistical Genetics and Methodology section of Frontiers in Genetics	2021–present
	• Peer-Reviewer for Cell Genomics, GENETICS, PLOS Computational Biology, Scientific Reports, and SIAM Undergraduate Research	2018–present rch Online
	Membership in Professional Societies • Caucus for Women in Statistics (CWS)	2018–present
	· · · ·	2016–present
	,	2013–present
	• • • •	2013–present
	Western North American Region (WNAR) of the International	2015–2019
	Biometric Society (IBS)	2010 2010
	Working Croups	

Working Groups

• Kidney Working Group

Trans-Omics for Precision Medicine Whole Genome Sequencing Program

ADVISING Honors Thesis Advisor

• Tina Chen. Title TBD.

2024-2025

• Freddy Barragan. Statistical genetics for pediatric leukemia: characterizing racial disparities in pediatric acute lymphoblastic leukemia.

(funded by NIH Research Supplement to Promote Diversity in Health-Related Research)

• Zuofu Huang. Estimating significance thresholds and the number 2020–2021 of generations since admixture in admixture mapping studies.

Honors Thesis Committee Member

•	Paige Tomer. An investigation into the causes of home field advantage	2024
	in professional soccer.	

- Erin Franke. Gentrification and crime in the Twin Cities: insights and challenges through a statistical lens.
- Zhaoheng Li. A comparison of stacking methods to estimate survival using residual lifetime data from prevalent cohort studies.

Summer Research Supervisor

- Tina Chen, Sydney Ohr, Katelyn McClure

 (funded by Macalester Collaborative Summer Research Award [TC, SO]

 and start-up funds [KM])
- Freddy Barragan 2021 (funded by Mann-Hill Fellowship for Student-Faculty Research)
- Zuofu Huang 2020

(funded by Macalester Collaborative Summer Research Award)

LAST UPDATE May 13, 2024