## Kelsey E. Grinde

| CONTACT                | Mathematics, Statistics, and Computer Science<br>Macalester College<br>Saint Paul, MN 55105  | kgrinde@macalester.edu<br>(651)-696-6976<br>kegrinde.github.io |
|------------------------|--|--|
| EDUCATION              | Ph.D. in Biostatistics University of Washington, Seattle, WA Dissertation: Statistical inference in admixed population Advisor: Sharon Browning                        | 2019<br>ns   |
|                        | <b>B.A. in Mathematics</b> , Concentration in Statistics St. Olaf College, Northfield, MN  | 2014   |
|                        | Graduated $summa\ cum\ laude$ with Distinction in Statis Advisor: Paul Roback  | stics  |
| WORK<br>EXPERIENCE     | Assistant Professor Department of Mathematics, Statistics, & Computer Science Macalester College, Saint Paul, MN   | 2020-present   |
|                        | Postdoctoral Teaching Fellow<br>Department of Mathematics, Statistics, & Computer Science<br>Macalester College, Saint Paul, MN  | 2019–2020  |
|                        | Graduate Research Assistant Browning Statistical Genetics Lab University of Washington, Seattle, WA  | 2014–2015, 2016–2019   |
|                        | Graduate Research Assistant<br>Genetic Analysis Center<br>University of Washington, Seattle, WA  | 2015–2016  |
|                        | Undergraduate Research Assistant<br>Summer Research Program in Statistical Genetics & Biostati<br>Dordt College, Sioux Center, IA                                      | 2013, 2014<br>astics   |
|                        | Undergraduate Research Fellow<br>Center for Interdisciplinary Research<br>St. Olaf College, Northfield, MN   | 2013–2014  |
| TEACHING<br>EXPERIENCE | Macalester College  • STAT 253: Statistical Machine Learning Sections Taught: Fall 2024 (×3), Spring 2025, Fall 2025   | 2024–present 5 ( $\times$ 3)                                   |
|                        | • STAT 494: Statistical Genetics<br>Sections Taught: Fall 2022, Spring 2025  | 2022, 2025   |
|                        | <ul> <li>MATH/STAT 455: Mathematical Statistics<br/>Sections Taught: Spring 2020, Spring 2021, Spring 2022</li> </ul>  | 2020–2023<br>2, Spring 2023                                    |
|                        | • STAT 155: Introduction to Statistical Modeling<br>Sections Taught: Fall 2019 (×2), Spring 2020, Fall 2020<br>Fall 2021 (×2), Spring 2022 (×2), Fall 2022, Spring 202 |  |

## University of Washington

| • | BIOST 311:   | Regression Methods in the Health Sciences, Co-Instructor  | 2018 |
|---|--------------|---|------|
| • | 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 S | tatistical 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 (Anderson Elementary School [1st Grade], Waite House Afterschool Programs [K–8th Grade])
- 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

- 15. **Grinde, K.**, Browning, B., Reiner, A., Thornton, T., & Browning, S. "Adjusting for principal components can induce collider bias in genome-wide association studies." *PLOS Genetics* 20.12 (2024): e1011242. [link]
- 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]
- 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]
- 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]
- 6. **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**

- 3. **Grinde, K.**, Heggeseth, B., Johnson, A., & Myint, L. "STAT 253: Statistical Machine Learning Course Notes." Online course text (2025): [link].
- 2. **Grinde, K.** "Rethinking grading systems in introductory and advanced statistics courses." Consortium for the Advancement of Undergraduate Statistics Education Resources for JEDI-Informed Teaching of Statistics (2025): [link].
- 1. Heggeseth, B., Myint, L., & **Grinde**, **K.** "Stat 155 Notes." Online textbook (2021): [link].

#### Other Writing

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

# SOFTWARE & APPLICATIONS

- 4. Chen, T.\*+, McClure, K.\*+, Ohr, S.\*+, Huang, Z., & **Grinde, K.** "STEAM: Significance Threshold Estimation for Admixture Mapping." R package version 0.2.0 (2024): [link].
- 3. Hayir, A.\*, & Grinde, K. "Interactive Circos Tool." R shiny application (2022): [link].
- 2. Huang, Z.\*, & **Grinde**, **K.** "Significance Threshold Estimation for Admixture Mapping using Rcpp." R package (2020): [link].
- 1. **Grinde, K.** "STEAM: Significance Threshold Estimation for Admixture Mapping." R package (2019): [link].

## RESEARCH TALKS

## Presentations at International or National Venues

- 11. Opportunities and challenges of working with (big) genetic data. Big Data Summer Institute Concluding Symposium, University of Michigan Department of Biostatistics, Ann Arbor, MI. 2025. (Invited, Keynote)
- Adjusting for principal components can induce spurious associations in genomewide association studies in admixed populations. International Genetic Epidemiology Society Annual Meeting. Virtual. 2021.
- 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.
- 7. Admixture mapping: controlling for false positives in the presence of population structure. American Society of Human Genetics Annual Meeting. Orlando, FL. 2017. (Poster)
- 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

- 25. UPCOMING: MSCS Colloquium, Department of Mathematics, Statistics, and Computer Science, St. Olaf College. (September 2025.) (Invited)
- 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. Department of Mathematics and Statistics, Carleton College. Northfield, MN. 2023. (Invited)
- 22. Statistical genetics in populations with mixed ancestry. Department of Mathematics, Creighton University. Omaha, NE. 2022. (Invited)
- 21. What's our work: statistical genetics. Department of Mathematics, Statistics, and Computer Science, Macalester College. Saint Paul, MN. 2021.
- 20. Genome-wide significance thresholds for admixture mapping studies. Interdisciplinary Biostatistics Training in Genetics and Genomics Journal Club, University of Minnesota. Virtual. 2021. (Invited)
- 19. Statistical genetics in populations with mixed ancestry. Mathematics Colloquium, Augsburg University. Virtual. 2020. (Invited)
- 18. Statistical methods for genome-wide admixture mapping studies. Division of Pediatric Epidemiology and Clinical Research, University of Minnesota. Virtual. 2020. (Invited)
- 17. Statistical genetics in populations with mixed ancestry. Department of Mathematics, Statistics, and Computer Science, Macalester College. 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 genomewide association studies. Genetic Analysis Center, University of Washington. Seattle, WA. 2019. (Invited)
- 14. Adjusting for population structure in genetic association studies: new insights and the potential pitfalls of using PCs. Population Genetics (PopGen) Lunch, University of Washington. Seattle, WA. 2019. (Invited)
- 13. Statistical inference in populations with mixed ancestry. Biostatistics Colloquium, University of Washington. 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. Department of Biostatistics, University of Michigan. 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. Department of Biostatistics, University of Michigan. 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. Natural Sciences and Mathematics Honors' Day Poster Session, St. Olaf College. Northfield, MN. 2014. (Poster)
- 2. Predicting donors at Red Cross blood drives. Mathematics, Statistics, and Computer Science Colloquium, St. Olaf College. Northfield, MN. 2014.
- Predicting donors at Red Cross blood drives. American Red Cross. Saint Paul, MN. 2014.

## Student Presentations of Joint/Supervised Work

- 11. Ohr, S. Significance threshold estimation for admixture mapping (STEAM), an R package. Midstates Consortium Undergraduate Research Symposium. St. Louis, MO. 2024. (Poster)
- 10. Chen, T. Evaluating the power of admixture mapping: a literature review and simulation study. StatFest. New York, NY. 2024. (Poster)
- 9. McClure, K. and Ohr, S. Significance threshold estimation for admixture mapping (STEAM), an R package. Summer Research Showcase, Macalester College. Saint Paul, MN. 2024. (Poster)
- 8. Chen, T. Evaluating the power of admixture mapping: a literature review and simulation study. Summer Research Showcase, Macalester College. Saint Paul, MN. 2024. (Poster)
- 7. Barragan, S. Genetic ancestry, gene expression, and survival in children with B-ALL. Pediatric Research, Education, & Scholarship Symposium. Minneapolis, MN. 2022. (Poster)
- 6. Barragan, S. Gene expression differences by race and genetic ancestry in B-cell acute lymphoblastic leukemia. American Society of Human Genetics Annual Meeting. Virtual. 2021. (Poster)
- Barragan, S. 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, S. Statistical methods for pediatric leukemia: gene expression & ancestry in B-cell acute lymphoblastic leukemia. Summer Research Showcase, Macalester College. Saint Paul, MN. 2021. (Poster)
- Huang, Z. Statistical methods for genetic association studies in populations with mixed ancestry. Midstates Consortium Undergraduate Research Symposium. Virtual. 2020.
- 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. Summer Research Showcase, Macalester College. Virtual. 2020. (Poster)

## TEACHING, OUTREACH, & MENTORING TALKS

#### Presentations at International or National Venues

- 5. Career Q&A with 2025 Big Data Summer Institute students. University of Michigan Department of Biostatistics, Ann Arbor, MI. 2025. (Invited)
- 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

- 23. UPCOMING: Center for Interdisciplinary Research, St. Olaf College. (September 2025.) (Invited)
- 22. LATEXAdvanced Workshop. Macalester College. St. Paul, MN. 2024.
- 21. Teaching careers roundtable. BIOS 834: Pedagogical Methods for Biostatistics Courses, University of Michigan. Virtual. 2024. (Invited)
- 20. 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. Hutch United Outreach Committee & Wallin Education Partners Program, Fred Hutchinson Cancer Research Center. 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. STAT 284: Biostatistics, St. Olaf College. Northfield, MN. 2019. (Invited)
- 11. Fellowships, scholarships, and grants. Biostatistics Student Seminar, University of Washington. 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. Department of Biostatistics, University of Washington. Seattle, WA. 2017.
- 7. What is Biostatistics? Forest Ridge School of the Sacred Heart Science Research Class. Bellevue, WA. 2017.
- 6. NSF Graduate Research Fellowship Program information session. Department of Biostatistics, University of Washington. Seattle, WA. 2017.
- 5. What is Biostatistics? 7th and 8th Grade STEM PREP Project, University of Washington. Seattle, WA. 2017.
- 4. Applying for outside funding opportunities. Biostatistics Student Seminar, University of Washington. Seattle, WA. 2016.
- 3. Graduate and professional student panel. Healthcare Exploration for Youth Program, University of Washington. Seattle, WA. 2016. (Invited)
- 2. Graduate and professional student panel. Healthcare Exploration for Youth Program, University of Washington. 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**)

#### **GRANTS** Funded by National Organizations

3. Safo, S. and Grinde, K. "Conference: STATGEN25."

2025

Amount: \$19,104

Funder: National Science Foundation (Program No. 21-541)

2. Graduate Research Fellowship

2016-2019

Amount: \$138,000

Funder: National Science Foundation (Program No. 24-591)

1. Statistical Genetics Training Grant

2015 - 2016

Amount: \$22,476

Funder: National Institutes of Health (T32 Training Grant)

## Funded by Local Organizations

6. Collaborative Summer Research Award

2024

Amount: \$13,461

Funder: Macalester College

5. Article Processing Charge Grant

2022

Amount: \$1,466

Funder: Macalester College Dewitt Wallace Library Open Access Fund

(for Refereed Journal Article [12])

4. Collaborative Summer Research Award

2020

Amount: \$5,500

Funder: Macalester College

|                 | 3. Travel Grant<br>Amount: \$300<br>Funder: University of Washington Graduate and Professional Student S  | 2018 enate |
|-----------------|---|------------|
|                 | 2. Conference Travel Award Amount: \$1,000 Funder: University of Washington (UW) Department of Biostatistics  | 2018       |
|                 | <ol> <li>Travel Award         Amount: \$500         Funder: UW Graduate School Fund for Excellence and Innovation     </li> </ol>                                     | 2017       |
| HONORS & AWARDS | <ul> <li>Professional Awards and Recognition</li> <li>Poster/Lightning Talk Award, 2nd Place<br/>International Genetic Epidemiology Society Annual Meeting</li> </ul> | 2021       |
|                 | (for International Research Talk [10])  |            |
|                 | • Top Cited Article<br>Genetic Epidemiology Journal   | 2021       |
|                 | (for Refereed Journal Article [6])  |            |
|                 | • Thomas R. Fleming Excellence in Biostatistics Award<br>University of Washington Department of Biostatistics   | 2019       |
|                 | • Gertrude M. Cox Scholarship<br>American Statistical Association   | 2018       |
|                 | • Dorothy L. Simpson Leadership Award<br>Achievement Rewards for College Scientists Foundation, Seattle Chapte  | 2018<br>r  |
|                 | • Excellence in Teaching Award<br>University of Washington Department of Biostatistics  | 2018       |
|                 | • Distinguished Oral Presentation Award<br>Western North American Region of the International Biometric Society   | 2018       |
|                 | (for International Research Talk [8])   |            |
|                 | • Achievement Rewards for College Scientists (ARCS) Fellowship ARCS Foundation, Seattle Chapter   | 14-2017    |
|                 | • Donovan J. Thompson Award<br>University of Washington Department of Biostatistics   | 2016       |
|                 | (for top performance on PhD qualifying exams)   |            |
|                 | Undergraduate Awards  |            |
|                 | Honorable Mention, Undergraduate Research Project Competition<br>Consortium for Advancement of Undergraduate Statistics Education                                     | 2014       |
|                 | <ul> <li>Honorable Mention, Graduate Research Fellowship Program<br/>National Science Foundation</li> </ul>   | 2014       |
|                 | • Statistically Significant Award<br>St. Olaf Department of Mathematics, Statistics, and Computer Science   | 2014       |
|                 | • Buntrock Scholarship  St. Olaf College  (academic scholarship)  | 010-2014   |
|                 | •   | 010-2014   |
|                 | • Phi Beta Kappa National Honor Society   | 2013       |
|                 | Pi Mu Epsilon National Honor Society  | 2013       |
|                 | (mathematics honor society)   |            |

| SERVICE | Professional Service |
|---------|----------------------|

| Professional Service  • American Statistical Association Section on Statistics in Genome  | ics and Genetics        |  |
|---|-------------------------|--|
| - Proposal Review Committee, STATGEN 2026 20  |                         |  |
| - Co-Chair, STATGEN 2025 Local Organizing Committee   | 2024-2025               |  |
| <ul> <li>Invited Panelist, ASA SSGG Webinar Series</li> </ul>   | 2021 & 2023             |  |
| (see National Teaching/Outreach/Mentoring Talks [2] and [4])  |                         |  |
| <ul> <li>Contributor, ASA SSGG Quarterly Newsletter [link]</li> <li>("Reflections and Tips from Recent Grads on the Job Search Experies</li> </ul>                  | 2021 ence")             |  |
| • Frontiers in Genetics Statistical Genetics and Methodology Sec  | etion                   |  |
| - Community Reviewer  | 2025-present            |  |
| - Review Editor   | 2021 - 2025             |  |
| • Ad Hoc Peer Reviewer for Cell Genomics, GENETICS, Nature<br>Communications, PLOS Computational Biology, Scientific Repo<br>and SIAM Undergraduate Research Online | 2018-present $orts$ ,   |  |
| Macalester College  • Service to the College  |                         |  |
| <ul> <li>Opening Convocation Committee</li> </ul>   | 2025-present            |  |
| <ul> <li>Co-Coordinator, Serie Center Reading Group</li> <li>(Book: Grading for Growth by D. Clark and R. Talbert)</li> </ul>                                       | 2025                    |  |
| <ul> <li>Faculty Liaison to Admissions</li> </ul>   | 2022 – 2024             |  |
| <ul> <li>AAC&amp;U Open Educational Resources Institute Team</li> </ul>   | 2022 – 2023             |  |
| <ul><li>Mid-Course Interview Scribe<br/>(*canceled due to COVID-19)</li></ul>   | <del>2020</del> *, 2021 |  |
| • Service to the Department of Mathematics, Statistics, and Com   | puter Science           |  |
| - Academic Planning Committee 2022–20   | 23, 2024–present        |  |
| - MSCS Honors Seminar Coordinator 2021–20   | 23, 2024–present        |  |
| <ul> <li>Statistics Tenure Track Search Committee</li> </ul>  | 2022, 2025              |  |
| - Statistics Visiting/Postdoc Search Committee 2020-  | -2022, 2024-2025        |  |
| <ul> <li>DataFest Judge</li> </ul>  | 2025                    |  |
| - DataFest Mentor   | 2021, 2022, 2023        |  |
| University of Washington Department of Biostatistics  • Diversity Committee   | 2017–2019               |  |
| Women in Biostatistics and Statistics, Leadership Team  | 2017–2018               |  |
| Admissions Committee  | 2017–2018               |  |
| • Peer Mentoring Program, Founding Member & Mentor  | 2016–2018               |  |
| • Educational Policy and Teaching Evaluation Committee  | 2016–2017               |  |
| Biostatistics Outreach Working Group  | 2015                    |  |
|   |                         |  |
| St. Olaf College • President, Spanish Honor House   | 2013-2014               |  |
| • Volunteer Teaching Assistant & Tutor, Northfield Public School  | ls 2011–2014            |  |
| • Orientation Leader, Liberal Arts Program in Seville   | 2013                    |  |
| • Volunteer Teaching Assistant, Wayzata High School   | 2011                    |  |

| OTHER<br>PROFESSIONAL<br>ACTIVITIES | <ul> <li>International Genetic Epidemiology Society (IGES)</li> <li>American Society of Human Genetics (ASHG)</li> <li>American Statistical Association (ASA)</li> </ul>   | 2018-present<br>2016-present<br>2013-present<br>2013-present |
|-------------------------------------|--|--|
|                                     | <ul> <li>Western North American Region (WNAR) of the International Biometric Society (IBS)</li> <li>Working Groups</li> <li>Kidney Working Group         Trans-Omics for Precision Medicine Whole Genome Sequencing F     </li> <li>Dental Genetics Working Group         Hispanic Community Health Study/Study of Latinos     </li> </ul> | 2015–2019<br>2018–2021<br>Program<br>2016                    |
| ADVISING                            | Summer Research Supervisor  • Tina Chen  (Funded by Local Grant [6])   | 2024   |
|                                     | • Sydney Ohr (Funded by Local Grant [6])   | 2024   |
|                                     | • Katelyn McClure (Funded by start-up funds)   | 2024   |
|                                     | • Sofia Barragan (Funded by Macalester Mann-Hill Fellowship for Student-Faculty Research)  | 2021   |
|                                     | • Zuofu Huang (Funded by Local Grant [4])  | 2020   |
|                                     | <ul> <li>Honors Thesis Advisor</li> <li>Tina Chen. Evaluating the power of admixture mapping.</li> <li>Sofia Barragan. Statistical genetics for pediatric leukemia: characterizing racial disparities in pediatric acute lymphoblastic leuke (Funded by NIH Research Supplement to Promote Diversity in Health-Related</li> </ul>          |  |
|                                     | • Zuofu Huang. Estimating significance thresholds and the number of generations since admixture in admixture mapping studies.  | 2020-2021  |
|                                     | <ul> <li>Honors Thesis Committee Member</li> <li>Paige Tomer. An investigation into the causes of home field advantage in professional soccer.</li> <li>Erin Franke. Gentrification and crime in the Twin Cities: insights and challenges through a statistical lens.</li> </ul>   |  |
|                                     |  |  |
|                                     | • Zhaoheng Li. A comparison of stacking methods to estimate survusing residual lifetime data from prevalent cohort studies.  | ival 2022  |