Kelsey Erin Grinde

CONTACT	Mathematics, Statistics, & Computer Science Macalester College Saint Paul, MN 55105	763-567-8325 kgrinde@macalester.edu kegrinde.github.io
EDUCATION	Ph.D. in Biostatistics University of Washington, Seattle, WA Dissertation: Statistical inference in admixed population Advisor: Sharon Browning, Ph.D.	2019 ons
	B.A. in Mathematics , Concentration in Statistics St. Olaf College, Northfield, MN	2014
	Graduated summa cum laude with Distinction in Stati Advisor: Paul Roback, Ph.D.	stics
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 & Biostat Dordt Colllege, Sioux Center, IA	2013, 2014 istics
	Undergraduate Research Fellow Center for Interdisciplinary Research St. Olaf College, Northfield, MN	2013–2014
TEACHING EXPERIENCE	Courses Taught	
EAPERIENCE	• MATH/STAT 455: Mathematical Statistics Macalester College, Saint Paul, MN	2020
	• STAT 155: Introduction to Statistical Modeling (3 section Macalester College, Saint Paul, MN	zions) 2019–2020
	• BIOST 311: Regression Methods in the Health Science University of Washington, Seattle, WA (co-taught with Brian Williamson)	s 2018

	 Teaching Assistantships BIOST 310: Biostatistics for the Health Sciences University of Washington, Seattle, WA 	2017
	• BIOST 570: Regression Methods for Independent Data University of Washington, Seattle, WA	2016
	Other Teaching Experience • Guest Lecturer (2 first year graduate courses) University of Washington, Seattle, WA	2017
	• Co-Instructor, First Year Statistical Theory Exam review sessions University of Washington, Seattle, WA	2016
	• Grader, STAT 322: Statistical Theory St. Olaf College, Northfield, MN	2013
	• Academic Assistant & Tutor, math and Spanish (all levels) St. Olaf College, Northfield, MN	2011-2012
	• Urban Schools and Communities Program St. Olaf College Off-Campus Studies, Minneapolis, MN	2012
HONORS & AWARDS	Academic Honors and Awards • Thomas R. Fleming Excellence in Biostatistics Award University of Washington Department of Biostatistics (highest honor given to a graduating Ph.D. student)	2019
	 Donovan J. Thompson Award University of Washington Department of Biostatistics (best combined score on Ph.D. qualifying exams) 	2016
	• Statistically Significant Award St. Olaf College	2014
	• Phi Beta Kappa St. Olaf College	2013
	• Pi Mu Epsilon St. Olaf College	2013
	Fellowships, Scholarships, and Training Grants • Graduate Research Fellowship National Science Foundation	2016–2019
	• Gertrude M. Cox Scholarship American Statistical Association	2018
	• Achievement Rewards for College Scientists Fellowship ARCS Foundation Seattle Chapter	2014-2017
	• Statistical Genetics Training Grant National Institutes of Health	2015-2016
	• Buntrock Scholarship St. Olaf College	2010–2014
	The skine Couries and Leadenskin America	

Teaching, Service, and Leadership Awards
 Dorothy L. Simpson Leadership Award
 Achievement Rewards for College Scientists Foundation Seattle Chapter

•	Excellence in Tea	aching Award	
	University of Was	shington Department of Biostati	istics

USRESP Undergraduate Research Project Competition

• Service Leadership Scholar St. Olaf College

• Honorable Mention

2010-2014

2018

2014

Research Communication and Travel Awards

•	Travel Grant	2018
	University of Washington Graduate and Professional Student Senate	
•	Distinguished Oral Presentation Award	2018
	Western North American Region of the International Biometric Society	
•	Conference Travel Award	2018
	University of Washington Department of Biostatistics	
•	Travel Award	2017
	University of Washington Graduate School Fund for Excellence and Innova	tion

PUBLICATIONS

- 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.
- 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.
- 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.
- 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.
- 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.

(Selected as the International Genetic Epidemiology Society Communications Committee's highlight from this issue of *Genetic Epidemiology*.)

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.

- 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.
- 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.
- 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.
- 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.

SUBMITTED MANUSCRIPTS

3. Lin, B.*, **Grinde, K.***, Brody, J., Raffield, L., Thornton, T., ..., & Franceschini, N. "Discovery of rare genetic variants from whole genome sequencing analyses of kidney function (eGFR) in 23,732 participants from multi-ethnic populations: the Trans-Omics for Precision Medicine (TOPMed) program."

*joint first authors

- 2. Jensen-Otsu, E., **Grinde, K.**, Teng, B.J., Baxi, A.C., Harms, M.A., 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."
- Snyder, J., Iwata, T., Patil, K., Tryon, V., Grinde, K., Mizumori, S., Treuting, & Treuting, P. "Comparison of end-of-life pathology in aged male Fischer 344 and Long Evans rats."

RESEARCH PRESENTATIONS

- 22. Statistical genetics in populations with mixed ancestry. Mathematics, Statistics, and Computer Science Colloquium, Augsburg University. Minneapolis, MN, 2020. (*Invited Speaker*) (postponed due to COVID-19)
- 21. Statistical genetics in populations with mixed ancestry. Department of Mathematics, Statistics, and Computer Science, Macalester College. Saint Paul, MN, 2019. (*Invited Speaker*)
- Adjusting for principal components can induce spurious associations in genomewide association studies. Genetic Analysis Center, University of Washington. Seattle, WA, 2019. (*Invited Speaker*)
- 19. Adjusting for population structure in genetic association studies: new insights and the potential pitfalls of using PCs. Popgen Lunch, University of Washington. Seattle, WA, 2019. (*Invited Speaker*)
- 18. Statistical inference in populations with mixed ancestry. Biostatistics Colloquium, University of Washington. Seattle, WA, 2018. (*Invited Speaker*)
- 17. Deriving significance thresholds for genome-wide admixture mapping studies. International Genetic Epidemiology Society Annual Meeting. San Diego, CA, 2018.
- 16. Controlling for multiple testing in genome-wide admixture mapping studies. Western North American Region of the International Biometric Society Meeting. Edmonton, Canada, 2018. (Oral Presentation Award Winner)

- 15. 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)
- 13. 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*)
- 12. 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.
- 11. 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.
- 8. 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.
- 7. What now? Post-hoc approaches for gene-based, rare variant tests of association. Great Plains R-Users Group Conference. Sioux Center, IA, 2014. (*Poster*)
- 6. Accounting for variability in paleoecological mixing models. St. Olaf Natural Sciences and Mathematics Honors Day Poster Session. Northfield, MN, 2014. (*Poster*)
- 5. Accounting for variability in paleoecological mixing models. National Conference for Undergraduate Research. Lexington, KY, 2014.
- 4. 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. St. Paul, MN, 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*)

OUTREACH & MENTORING PRESENTATIONS

- 12. (Bio)statistics PhD programs: application tips and research opportunities. Biostatistics Class, St. Olaf College. Northfield, MN, 2019.
- 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.

- 8. Travel grants and conference funding. University of Washington Department of Biostatistics. Seattle, WA, 2017.
- 7. What is Biostatistics? Science Research Class, Forest Ridge School of the Sacred Heart. 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, Distance Learning Center & 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. Seattle, WA, 2016.
- 2. Graduate and professional student panel. Healthcare Exploration for Youth Program. Seattle, WA, 2015.
- 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*)

SERVICE & LEADERSHIP

ACTIVITIES

Macalester College

• Scribe, Mid-Course Interview (postponed due to COVID-19)

2020

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 Journal Peer Review

ournal Feer Review	
• GENETICS	2020
• SIAM Undergraduate Research Online (SIURO)	2020
PLOS Computational Biology	2019
• Scientific Reports	2018

Working Groups

Kidney Working Group	2018-present
NHLBI Trans-Omics for Precision Medicine Whole Genome Sequen	ncing Program
• Dental Genetics Working Group	2016
Hispanic Community Health Study/Study of Latinos (HCHS/SOL	4)

Professional Organization Membership

- American Society of Human Genetics
- American Statistical Association
- Caucus for Women in Statistics
- International Genetic Epidemiology Society
- Western North American Region of the International Biometric Society

COMPUTING R, highly proficient
EXPERIENCE Unix/Linux, proficient
Python, familiar

SOFTWARE Significance Threshold Estimation for Admixture Mapping (R package)

• Available on GitHub: https://github.com/kegrinde/STEAM

LANGUAGES English, fluent/native

Spanish, highly proficient

RESEARCH Statistical genetics

INTERESTS Biostatistics

Multiple testing

LAST UPDATE June 18, 2020