ANDREW PARKER MORGAN

PERSONAL INFORMATION

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	EDUCATION
2010–2019	MD · University of North Carolina at Chapel Hill Dual-degree student in Medical Scientist Training Program (MSTP)
2012-2017	PhD · University of North Carolina at Chapel Hill Curriculum in Bioinformatics & Computational Biology · Department of Genetics
2005–2009	BS · University of North Carolina at Chapel Hill Biostatistics; biology · School of Public Health
	HONORS AND AWARDS
2019	Isaac Hall Manning Award, for medical student scholarship and leadership

	Decays Distinguished Discontation Assemble (subset discontation in high-rish states)
2017	Dean's Distinguished Dissertation Award, for best dissertation in biological sciences

- 2016 Terry Magnuson Award, for a senior student in Bioinformatics & Computational Biology curriculum
- Excellent student talk, 29th International Mammalian Genome Conference, Yokohama, Japan 2015
- Pillsbury Award, best oral presentation in basic sciences, UNC Medical Student Research Day 2014 Excellent student talk, 28th International Mammalian Genome Conference, Bar Harbor, Maine
- Best student talk, UNC Department of Genetics Scientific Retreat 2013 Excellent student talk, 27th International Mammalian Genome Conference, Salamanca, Spain

Oxford Summer School in Computational Biology, invited participant (top 5% of applicants globally)

2009 Phi Beta Kappa, inductee

> J.N. Couch Award, for excellence in plant biology research Delta Omega Award, top undergraduate student in biostatistics

GRANTS AND FELLOWSHIPS

Ruth L. Kirschstein NRSA Inidividual Predoctoral Fellowship 2014-2018 National Institute of Mental Health F30MH103925

Sponsor: Fernando Pardo-Manuel de Villena, PhD

Bioinformatics & Computational Biology Training Grant 2012-2013

National Institute of General Medical Sciences T₃₂GMo₆₇₅₅₃

PI: Timothy Elston, PhD

2010-2011 Medical Scientist Training Program Training Grant

National Institute of General Medical Sciences T32GMoo8719

PI: Eugene Orringer, MD

SERVICE

2015-	Ad hoc reviewer: Genetics, G ₃ , Scientific Data
2014–2016	Admissions committee, UNC Medical Scientist Training Program
2015–2016	Chief operating officer, UNC Student Health Action Coalition
2013-2015	Data and privacy officer, UNC Student Health Action Coalition
2012-2013	Medical clinic director, UNC Student Health Action Coalition

TEACHING

2015	Systems Genetics Workshop, 29th IMGC, Yokohama, Japan Guest lecture, graduate-level population genomics course (BCB 722)
2014	Academic coach (bioinformatics), UNC Initiative for Maximizing Student Diversity
2013	Teaching assistant for BCB 720: Principles of statistical modelling

PUBLICATIONS

Morgan AP, Bell TA, Crowley JJ, Pardo-Manuel de Villena F (2019) Instability of the pseudoautosomal 2019 region in house mice. Genetics 212: 469-487. 10.1534/genetics.119.302232

Morgan AP, Pardo-Manuel de Villena F (2017) Sequence and structural diversity of mouse Y chromosomes. Mol Biol Evol 34: 3186–3204. 10.1093/molbev/msx250

Rosshart SP, Vassallo BG, Angeletti D, Hutchinson DS, **Morgan AP**, Hickman HD, Ajami NJ, Petrosino JF, Pardo-Manuel de Villena F, Yewdell JW, Rehermann B (2017) Wild mouse gut microbiota promotes host fitness and improves disease resistance. *Cell* 171: 1015–1028. 10.1016/j.cell.2017.09.016

Makhanova NA, **Morgan AP**, Kayashima Y, Makhanov M, Hiller S, Zhilicheva S, Xu L, Pardo-Manuel de Villena F, Maeda N (2017) Genetic architecture of atherosclerosis dissected by QTL analyses in three F2 intercrosses of apolipoprotein E-null mice on C57BL6/J, DBA/2J and 129S6/SvEvTac backgrounds. *PLoS One* 12: e0182882. 10.1371/journal.pone.0182882

Gralinski LE, Menachery VD, **Morgan AP**, Totura A, Beall A, Kocher J, Plante J, Harrison-Shostak CD, Schäfer A, Pardo Manuel de Villena F, Ferris MT, Baric RS (2017) Allelic variation in mouse Ticam2 contributes to SARS-CoV pathogenesis. *G*₃ 7: 1653–1663. 10.1534/g3.117.041434

Morgan AP, Gatti DM, Keane TM, Galante RJ, Pack AI, Mott R, Churchill GA, Pardo-Manuel de Villena F (2017) Structural variation shapes the landscape of meiotic recombination in mouse. *Genetics* **206**: 603–619. 10.1534/genetics.116.197988

Srivastava A*, **Morgan AP***, Najarian M*, Sarsani VK, Sigmon JS, Shorter JR, Kashfeen A, Giusti-Rodgriguez P, Ferris MT, Sullivan PF, Miller DR, Bell TA, McMillan L, Churchill GA, Pardo-Manuel de Villena F (2017) The genomes of the mouse Collaborative Cross. *Genetics* **206**: 537–556. 10.1534/genetics.116.198838

Shorter JR, Odet F, Aylor DL, Pan W, Kao CY, Fu CP, **Morgan AP**, Greenstein S, Bell TA, Stevans AM, Feathers RW, Patel S, Cates SE, Shaw GD, Ahmed S, Miller DR, Chesler EJ, McMillian L, OBrien DA, Pardo-Manuel de Villena F (2017) Male infertility is responsible for nearly half of the strain extinction observed in the Collaborative Cross. *Genetics***206**: 537–556. 10.1534/genetics.116.199596

Morgan AP*, Didion JP*, Doran AG, Holt JM, McMillan L, Keane TM, Pardo-Manuel de Villena F (2016) Whole genome sequence of two wild-derived *Mus musculus domesticus* inbred strains, LEWES/EiJ and ZALENDE/EiJ, with different diploid numbers. *G*₃ 6: 4211–4216. 10.1534/g3.116.034751

Chesler EJ, Gatti DM, Morgan AP, Strobel M, Trepanier L, Oberbeck D, McWeeney S, Hitzemann R, Ferris M, McMullan R, Clayshulte A, Bell TA, Pardo-Manuel de Villena F, Churchill GA (2016) Diversity Outbred Mice at 21: Maintaining allelic variation in the face of selection. *G*3 6:3893–3902. 10.1534/g3.116.035527

Morgan AP, Holt JM, McMullan RC, Bell TA, Clayshulte AMF, Didion JP, Yadgary L, Thybert D, Odom DT, Flicek PW, McMillan L, Pardo-Manuel de Villena F (2016) The evolutionary fates of a large segmental duplication in mouse. *Genetics* 204: 267–2885. 10.1534/genetics.116.191007

Didion JP*, **Morgan AP***, ...(39 *others*)..., Pardo-Manuel de Villena F (2016) *R2d2* drives selfish sweeps in the house mouse. *Mol Biol Evol* **33**: 1381–1395. 10.1093/molbev/msw036

2015 **Morgan AP** (2015) argyle: an R package for analysis of Illumina genotyping arrays. *G*₃ **6**: 281–286. 10.1534/g3.115.023739

Morgan AP*, Fu CP*, Kao CY, Welsh CE, Didion JP, Yadgary L, Hyacinth L, Ferris MT, Bell TA, Miller DR, Giusti-Rodriguez P, Nonneman RJ, Cook KD, Whitmire JK, Gralinski LE, Keller M, Attie AD, Churchill GA, Petkov P, Sullivan PF, Brennan JR, McMillan L, Pardo-Manuel de Villena F (2015) The Mouse Universal Genotyping Array: from substrains to subspecies. *G*3 **6**: 263–279. 10.1534/g3.115.022087

Morgan AP, Welsh CE (2015) Informatics resources for the Collaborative Cross and related mouse populations [review]. *Mamm Genome* **26**: 521–539. 10.1007/s00335-015-9581-z

Didion JP, **Morgan AP**, ... (33 *others*)..., Pardo-Manuel de Villena F. (2015) A multi-megabase copy number gain causes maternal transmission ratio distortion on mouse chromosome 2. *PLoS Genet* 11: e1004850. 10.1371/journal.pgen.1004850

Crowley JJ, Zhabotynsky V, Sun W, ... (35 others)..., **Morgan AP**, McMillan L, Sullivan PF, Pardo-Manuel de Villena F. (2015) Analyses of allele-specific gene expression in highly divergent mouse crosses identifies pervasive allelic imbalance. *Nat Genet* **47**: 353–360. 10.1038/ng.3222

Morgan AP*, Crowley JJ*, Nonneman RJ, Quackenbush CR, Miller CN, Ryan AK, Bogue MA, Paredes SH, Yourstone S, Carroll IM, Kawula TH, Bower MA, Sartor RB, Sullivan PF (2014) The antipsychotic olanzapine interacts with the gut microbiome to cause weight gain in mouse. *PLoS One* 9: e115225.

10.1371/journal.pone.0115225

Liu EY*, **Morgan AP***, Chesler EJ, Wang W, Churchill GA, Pardo-Manuel de Villena F (2014) High-resolution sex-specific linkage maps of the mouse reveal polarized distribution of crossovers in male germline. *Genetics* **197**: 91–106. 10.1534/genetics.114.161653

Rogala AR, **Morgan AP**, Christensen AM, Gooch TJ, Bell TA, Miller DR, Godfrey VL, Pardo-Manuel de Villena F (2014) The Collaborative Cross as a resource for modeling human disease: CC011/Unc, a new mouse model for spontaneous colitis. *Mamm Genome* **25**: 95-108. 10.1007/s00335-013-9499-2