Gota Morota May 2014

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RESEARCH INTERESTS I am a quantitative geneticist interested in incorporating statistics, machine learning, and bioinformatics to the study of animal genetics in the omics era. The core line of my research is connecting the quantitative genetics and animal breeding theories to currently available molecular information. I am particularly interested in statistical methods for prediction of complex traits using whole-genome molecular markers.

Professional Positions Department of Animal Science

University of Nebraska-Lincoln, Lincoln, Nebraska USA

Assistant Professor (70% Research & 30% Teaching)

Work Experience Department of Animal Sciences

University of Wisconsin-Madison, Madison, Wisconsin USA

Graduate Research Assistant 06/2011 - 05/2014

Animal Genetics Research & Development Group

Zoetis, Inc., Kalamazoo, Michigan USA Quantitative Geneticist (student internship)

06/2013 - 11/2013

Start date: 08/2014

EDUCATION

University of Wisconsin-Madison, Madison, Wisconsin USA

Ph.D., Animal Sciences, May 2014

- Dissertation: "Whole-genome Prediction of Complex Traits Using Kernel Methods."
- Advisor: Prof. Dr. Daniel Gianola
- Committee: Drs. Corinne D. Engelman, Guilherme J. M. Rosa, Grace Wahba and Kent A. Weigel

University of Wisconsin-Madison, Madison, Wisconsin USA

M.S., Dairy Science, December 2011

- Thesis: "Application of Bayesian and Sparse Network Models for Assessing Linkage Disequilibrium in Animals and Plants."
- Advisor: Prof. Dr. Daniel Gianola
- Committee: Drs. Guilherme J. M. Rosa and Kent A. Weigel

Obihiro University of Agriculture and Veterinary Medicine, Obihiro, Hokkaido Japan

B.S., Agricultural Science, March 2008

- Thesis: "Genetic Analysis of Threshold Traits."
- Advisor: Prof. Dr. Mitsuyoshi Suzuki

Professional society memberships

- American Society of Animal Science 2014 Present
- Japanese Society of Animal Breeding and Genetics 2013 Present
- The International Biometric Society (ENAR) 2012 Present

EDITORIAL ACTIVITIES

Ad hoc Reviewer for: Computers and Electronics in Agriculture, DNA Research, Scientia Agricola

Number of reviewed papers: 2012 (1), 2013 (1), 2014 (1)

Manuscripts in Prep

9. <u>Morota G</u> and Gianola D. Kernel-based whole-genome prediction of complex quantitative traits.

Manuscripts in Review

8. Valente BD, <u>Morota G</u>, Peñagaricano F, Rosa GJM, Gianola D, and Weigel KA. The causal meaning of genomic predictors and how it affects construction and comparison of genome-enabled selection models. *BioRxiv doi:* 10.1101/001511, arXiv: http://arxiv.org/abs/1401.1165

PEER REVIEWED JOURNAL PAPERS

As of March 2014, 10 citations total, h-index of 2, i10-index of 0 on Google Scholar and 6 citations total, h-index of 2 according to Web of Science.

2014

- 7. <u>Morota G</u>, Boddhireddy P, Vukasinovic N, Gianola D and DeNise S. Kernel-based variance components estimation and whole-genome prediction of pre-corrected phenotypes and progeny tests for dairy cow health traits. *Frontiers in Genetics*. **5**:56.
- Morota G, Abdollahi-Arpanahi R, Kranis A and Gianola D. (2014) Genome-enabled prediction
 of broiler traits in chickens using genomic annotation. BMC Genomics. 15:109 * Highly
 accessed.
- 5. Abdollahi-Arpanahi R, Pakdel A, Nejati-Javaremi A, Moradi-Shahrbabak M, <u>Morota G</u>, Valente BD, Kranis A, Rosa GJM, and Gianola D. (2014) Dissection of additive genetic variability for quantitative traits in chickens using SNP markers. *Journal of Animal Breeding and Genetics*. doi: 10.1111/jbg.12079
- 4. Abdollahi-Arpanahi R, Nejati-Javaremi A, Pakdel A, Moradi-Shahrbabak M, <u>Morota G</u>, Valente BD, Kranis A, Rosa GJM, and Gianola D (2014). Effect of allele frequencies, effect sizes and number of markers on prediction of quantitative traits in chickens. *Journal of Animal Breeding and Genetics*. doi: 10.1111/jbg.12075

2013

- 3. <u>Morota G</u>, Koyama M, Rosa GJM, Weigel KA, and Gianola D. (2013) Predicting complex traits using a diffusion kernel on genetic markers with an application to dairy cattle and wheat data. *Genetics Selection Evolution*. **45**:17. * **Highly accessed**.
- 2. <u>Morota G</u> and Gianola D. (2013) Evaluation of linkage disequilibrium in wheat with an L1 regularized sparse Markov network. *Theoretical and Applied Genetics.* **126**(8):1991-2002.

2012

1. <u>Morota G</u>, Valente BD, Rosa GJM, Weigel KA, and Gianola D. (2012) An assessment of linkage disequilibrium in Holstein cattle using a Bayesian network. *Journal of Animal Breeding and Genetics*. 129(6):474-487. * Highlighted in the Genomic Selection Virtual Issue.

Papers in Proceedings

2014

- **3.** Gianola D, <u>Morota G</u>, and Crossa J. (2014) Genome-enabled Prediction of Complex Traits with Kernel Methods: What Have We Learned? Proceedings, 10th World Congress of Genetics Applied to Livestock Production. August 17-22, Vancouver, BC, Canada.
- 2. Valente BD, Morota G, Rosa GJM and Gianola D and Weigel KA (2014) The causal meaning of genomic predictors and how it affects the construction and comparison of genome-enabled selection models. Proceedings, 10th World Congress of Genetics Applied to Livestock Production. August 17-22, Vancouver, BC, Canada.

2011

Bueno Filho JS*, <u>Morota G</u>*, Tran Q, Maenner MJ, Vera-Cala LM, Engelman CD, and Meyers KJ. (2011) Analysis of human mini-exome sequencing data from Genetic & Analysis Workshop 17 using a Bayesian hierarchical mixture model. <u>BMC Proceedings</u>, 5(Suppl 9):S93. *equal contribution

INVITED PRESENTATIONS

My PDF presentation slides are available on SlideShare (http://www.slideshare.net/chikudaisei/).

2. Department of Animal Science, University of Nebraska-Lincoln, Nebraska, USA.

2011

2013

1. OBIHIRO GCOE Animal Global Health Seminar. Obihiro University of Agriculture and Veterinary Medicine, Obihiro, Hokkaido Japan

Conference Abstracts

Contributed

2012

• Morota G, Rosa GJM, Weigel KA, and Gianola D. (2012) Application of Bayesian and Sparse Network Models for Assessing Linkage Disequilibrium in Animals and Plants. 26th International Biometric Conference. August 26-31, Kobe, Japan. http://secretariat.ne.jp/ibc2012/30Aug.html#aug-30-14:00-Contributed36. * Second Oral Prize Winners.

<u>Posters</u>

2013

- Morota G, Tsuyuzaki K, and Nikaido I. (2013) MeSHR: R/Bioconductor package for finding statistically overrepresented MeSH terms in a set of genes. Annual Bioconductor Conference BioC 2013. July 18-19, Seattle, USA. https://secure.bioconductor.org/BioC2013/posters.php#8.
- Morota G. (2013) Predicting complex traits using a diffusion kernel on genetic markers with an application to dairy cattle and wheat data. Annual Bioconductor Conference BioC 2013. July 18-19, Seattle, USA. https://secure.bioconductor.org/BioC2013/posters.php#7.

Departmental Seminars	
2014	• Ph.D. Thesis Defense. University of Wisconsin-Madison.
	• Dairy Science Graduate Seminar. University of Wisconsin-Madison.
2013	• Special Seminar. Zoetis, Inc., Kalamazoo, Michigan USA.
2012	• Animal Breeding & Genomics Seminar. University of Wisconsin-Madison.
2011	• Master's Thesis Defense. University of Wisconsin-Madison.
2010	• Animal Breeding & Genomics Seminar. University of Wisconsin-Madison.
2008	\bullet Animal Breeding & Genomics Seminar. University of Wisconsin-Madison.
PARTICIPATION IN CONFERENCES AND WORKSHOPS	
2014	• UC Davis Bioinformatics Training Program: "Using Galaxy for Analysis of High Throughput Sequence Data". University of California, Davis. June 16-20. Taught by the Bioinformatics Core.
	• Short course: "Evolutionary Quantitative Genetics". University of Wisconsin-Madison. May 19-23. Taught by Bruce Walsh.
2013	• Short course: "Statistical methods for prediction of complex traits using whole-genome molecular markers". University of Wisconsin-Madison. May 27-31. Taught by Daniel Gianola and Gustavo de los Campos.
2012	• Short course: "Introduction to genome-enabled selection & Inferring causal phenotype networks using structural equation models". Kyoto University. August 31. Taught by Guilherme J.M. Rosa.
	• Short course: "Identifying Genes for Complex and Mendelian Traits Using Next Generation Sequence Data". 26th International Biometric Conference. August 26. Taught by Suzanne Leal.
	• Short course: "Programming and computer algorithms with focus on genomic selection in animal breeding". University of Georgia. May 15 - June 1. Taught by Ignacy Misztal, Shogo Tsuruta, Ignacio Aguilar, Zulma Vitezica, and Andres Legarra.
2009	• Symposium: "Statistical Genetics of Livestock for the Post-Genomic Era (SGLPGE)". University of Wisconsin-Madison. May 4-6.
2006	• Short course: "Estimation of Variance Conponents in Animal Breeding". Obihiro University of Agriculture and Veterinary Medicine. November. Taught by Shogo Tsuruta.

TEACHING University of Wisconsin-Madison, Madison, Wisconsin USA

Teaching Assistant

ANSCI/DYSCI 363: "Principles of Animal Breeding" Spring, 2011

Teaching Assistant

ANSCI/DYSCI 361: "Introduction to Animal and Veterinary Genetics" Spring, 2011

OSS CONTRIBUTIONS

R packages

• dkDNA: http://cran.r-project.org/web/packages/dkDNA/index.html

Bioconductor packages

- meshr
- MeSH.db
- MeSH.AOR.db
- MeSH.PCR.db
- org.MeSH.XXX.db (120 packages)
- org.MeSH.Aca.db
- \bullet org.MeSH.Aga.PEST.db
- \bullet org.MeSH.Ame.db
- org.MeSH.Aml.db
- \bullet org.MeSH.Ana.db
- org.MeSH.Ani.FGSC.db
- org.MeSH.Ath.db
- org.MeSH.Atu.K84.db
- org.MeSH.Bfl.db
- org.MeSH.Bsu.168.db
- org.MeSH.Bsu.Bsn5.db
- org.MeSH.Bsu.RONN1.db
- org.MeSH.Bsu.TUB10.db
- org.MeSH.Bsu.W23.db
- N. CII. D. II
- \bullet org.MeSH.Bta.db
- \bullet org.MeSH.Cal.SC5314.db
- org.MeSH.Cbr.db
- org.MeSH.Cel.db
- org.MeSH.Cfa.db
- org.MeSH.Cin.db
- org.MeSH.Cja.db
- org.MeSH.Cpo.db
- org.MeSH.Cre.db
- org.MeSH.Dan.db
- \bullet org.MeSH.Dda.3937.db

- \bullet org.MeSH.Ddi.AX4.db
- \bullet org.MeSH.Der.db
- org.MeSH.Dgr.db
- org.MeSH.Dme.db
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- \bullet org.MeSH.Dpe.db
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- org.MeSH.Eco.APEC01.db
- \bullet org.MeSH.Eco.B.REL606.db
- org.MeSH.Eco.BW2952.dborg.MeSH.Eco.CFT073.db
- M CHE DOLOGEA II
- $\bullet \ \ {\rm org.MeSH.Eco.E24377A.db}$
- org.MeSH.Eco.ED1a.dborg.MeSH.Eco.HS.db
- org.MeSH.Eco.IAI1.db
- org.MeSH.Eco.IAI39.db
- org.MeSH.Eco.K12.DH10B.db
- \bullet org.MeSH.Eco.K12.MG1655.db
- org.MeSH.Eco.KO11FL.db
- org.MeSH.Eco.O103.H2.12009.db

- org.MeSH.Eco.O111.H.11128.db
- $\bullet \ \, \mathrm{org.MeSH.Eco.O127.H6.E2348.69.db}$
- org.MeSH.Eco.O157.H7.EC4115.db
- org.MeSH.Eco.O157.H7.EDL933.db
- org.MeSH.Eco.O157.H7.Sakai.db
- org.MeSH.Eco.O157.H7.TW14359.db
- org.MeSH.Eco.O26.H11.11368.db
- org.MeSH.Eco.O26.H7.CB9615.db
- org.MeSH.Eco.S88.db
- org.MeSH.Eco.SE11.db
- org.MeSH.Eco.SMS35.db
- org.MeSH.Eco.UMN026.db
- org.MeSH.Eco.UTI89.db
- org.MeSH.Eqc.db
- org.MeSH.Gga.db
- org.MeSH.Gma.db
- org.MeSH.Hsa.db
- org.MeSH.Laf.db
- org.MeSH.Lma.db
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- org.MeSH.Ocu.db
- org.MeSH.Oni.db
- org.MeSH.Osa.db
- org.MeSH.Pab.db
- org.MeSH.Pae.LESB58.db
- org.MeSH.Pae.PA14.db
- org.MeSH.Pae.PA7.db

- org.MeSH.Pae.PAO1.db
- org.MeSH.Pfa.3D7.db
- org.MeSH.Pto.db
- \bullet org.MeSH.Ptr.db
- org.MeSH.Rno.db
- \bullet org.MeSH.Sau.COL.db
- org.MeSH.Sau.ED98.db
- org.MeSH.Sau.M013.db
- org.MeSH.Sau.MRSA252.db
- org.MeSH.Sau.MSHR1132.db
- org.MeSH.Sau.MSSA476.db
- org.MeSH.Sau.Mu3.db
- org.MeSH.Sau.Mu50.db
- org.MeSH.Sau.MW2.db
- org.MeSH.Sau.N315.db
- org.MeSH.Sau.Newman.db
- org.MeSH.Sau.RF122.db
- \bullet org.MeSH.Sau.USA300FPR3757.db
- $\bullet \ \, {\rm org.MeSH.Sau.USA300TCH1516.db}$
- org.MeSH.Sau.VC40.db
- \bullet org.MeSH.Sce.S288c.db
- org.MeSH.Sco.A32.db
- org.MeSH.Sil.db
- org.MeSH.Spo.972h.db
- org.MeSH.Spu.db
- org.MeSH.Ssc.db
- org.MeSH.Syn.db
- \bullet org.MeSH.Tbr.9274.db
- org.MeSH.Tgo.ME49.db
- org.MeSH.Tgu.db
- org.MeSH.Vvi.db
- \bullet org.MeSH.Xla.db
- org.MeSH.Xtr.db
- \bullet org.MeSH.Zma.db

Github

• https://github.com/morota

Computer Skills

- Programming Languages: Python, Fortran
- Statistical/Numerical Computational Tools: R, Octave

 \bullet Content-description Languages: XML, XHTML, CSS, LATEX

• Operating Systems: Linux and Mac OS X

MISCELLANEOUS • Languages: English and Japanese

REFERENCES References and additional information available upon request.