

# Matheus D. Krause

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I am a motivated Ph.D. Candidate in Plant Breeding with a Statistics minor at Iowa State University. My field of expertise and research interest includes quantitative and statistical genetics, biometry, and biostatistics. I have extensive experience in **R** programming and contributed to many collaborative projects focused on data analysis. In addition, during my academic career, I had hands-on experience in daily breeding operations in different countries. My expected graduation date is the Spring of 2023.

## EDUCATION

### Ph.D. Candidate in Plant Breeding with Statistics minor (2023)

IOWA STATE UNIVERSITY - AMES, IOWA, USA | MAJOR PROFESSOR: DR. WILLIAM D. BEAVIS

**Projects:** I) Unraveling genotype by environment variation to identify mega-environments with genetic and non-genetic factors.  
II) Development of estimation methods to obtain realized rate of genetic gain using routine field trials.

### Master's Degree in Plant Breeding and Genetics (2018)

UNIVERSITY OF SÃO PAULO - PIRACICABA, SP, BRAZIL | MAJOR PROFESSOR: DR. ANTONIO A. F. GARCIA

**Title:** Boosting predictive ability of maize hybrids via genotype by environment interaction under multivariate GBLUP models.

### Bachelor's Degree in Agronomy (2015)

LONDRINA STATE UNIVERSITY - LONDRINA, PR, BRAZIL | MAJOR PROFESSOR: DR. JOSUÉ M. FERREIRA

**Title:** Combining ability of maize inbred lines S<sub>9</sub> derived from ST15 and S709 synthetics.

### Exchange Student in Plant Breeding & Engineering for Mediterranean and Tropical Areas (2014)

SUPAGRO - MONTPELLIER, FRANCE | SUPERVISORS: DRS. RICARDO RALISCH AND JEAN-LUC REGNARD

**Title:** Productibilité semencière de nouveaux géniteurs maïs : faisabilité de l'utilisation de données de production.

## RESEARCH EXPERIENCE

### Graduate Research Assistant | 2018 - PRESENT, DEPARTMENT OF AGRONOMY, IOWA STATE UNIVERSITY

- Development of a metric to estimate realized rate of genetic gain on an annual basis using soybean routine field trials.
- Unraveling genotype by environment variation to identify mega-environments with genetic and non-genetic factors.
- Development of software to correct partially informative markers from backcross one derived double-haploid lines.
- Genome-wide association for complex binary (case-control) and continuous traits.
- Mentored several graduate students (+20) with plant breeding data analysis and **R** programming.

### Graduate Research Assistant | 2016 - 2017, DEPARTMENT OF GENETICS, UNIVERSITY OF SÃO PAULO

- Applying multivariate linear mixed models to predict yield performance of single-cross maize hybrids based on information from genomics and genotype by environment interaction.

### Intern | 2015, TROPICAL BREEDING & GENETICS, SOYBEAN BREEDING

- Plant phenotyping for soybean cyst nematode, *Phytophthora sojae* and *Sclerotinia sclerotiorum*, generation advancement in greenhouses, crossing blocks, seed packing, and general activities in the biotechnology laboratory.

### Undergraduate Research Assistant | 2010 - 2015, DEPARTMENT OF BIOLOGY, LONDRINA STATE UNIVERSITY

- Development and evaluation of inbred lines for common and sweet corn via classical and double-haploid techniques.
- Evaluation and development of single-cross hybrids and synthetic populations for both common and sweet corn.
- Participation in an extension project to teach farmers how to produce their seeds of synthetic populations.

### Intern | 2014, LIMAGRAIN EUROPE, DEPARTMENT OF SEED PRODUCTION TO RESEARCH

- Data mining to support early decision-making on the profitability of female maize inbred lines (seed production).
- Stability analysis to identify stable check cultivars to be used in seed production.

## SELECTED JOURNAL PUBLICATIONS

Krause, MD; Dias, KOG; Singh, AK; Beavis, WD. Using soybean historical field trial data to study genotype by environment variation and identify mega-environments with the integration of genetic and non-genetic factors. Submitted to Field Crops Research, 2022. DOI: 10.1101/2022.04.11.487885. [↗](#)

Dias, KOG; dos Santos, JPR; **Krause, MD**; Piepho, HP; Guimarães, LJM; Pastina, MM; Garcia, AAF. Leveraging probability concepts for cultivar recommendation in multi-environment trials. Theoretical and Applied Genetics, 2022. DOI: 10.1007/s00122-022-04041-y. [↗](#)

Montes, CR; Fox, C; Sanz-Sáez, A; Serbin, SP; Kumagai, E; **Krause, MD**; Xavier, A; Specht, J; Beavis, WD; Bernacchi, CJ; Diers, BW; Ainsworth, EA. High-throughput characterization, correlation, and mapping of leaf photosynthetic and functional traits in the soybean (*Glycine max*) nested association mapping population. Genetics, 2022. DOI: 10.1093/genetics/iyac065. [↗](#)

Verzegnazzi, AL; Santos, I; **Krause, MD**; Hufford, M; Frei, UK; Campbell, J; Almeida, VC; Zuffo, LT; Boerman, N; Lübberstedt, T. Major locus for spontaneous haploid genome doubling detected by a case-control GWAS in exotic maize germplasm. Theoretical and Applied Genetics, 2021. DOI: 10.1002/csc2.20253. [↗](#)

**Krause, MD**; Dias, KOG; dos Santos, JPR; Oliveira, AAO; Guimarães, LJM; Pastina, MM; Margarido, GRA; Garcia, AAF. Boosting predictive ability of tropical maize hybrids via genotype by environment interaction under multivariate GBLUP models. Crop Science, 2020. DOI: 10.1002/csc2.20253. [↗](#)

## SKILL SET

### SPECIFIC SKILLS

- Model building in the framework of linear, bayesian, and generalized mixed models (e.g., *AsremlR*, *lme4*, *rstan*) for field trial analysis, to explore genotype by environment interaction, and for general statistical analyses.
- Theory and application of genomic prediction with frequentist and bayesian approaches, quantitative trait loci mapping, genome-wide association analyses, stochastic simulations, and applications of probability theory.
- Data mining, curation, and acquisition for statistical inference. Cloud / high performance computing.
- I am a constant learner eager to collaborate across diverse groups.

### PROGRAMMING EXPERIENCE

Advanced **R** programming language, basic Python and Julia, Linux,  $\LaTeX$  • Developer of the **R** package SoyURT [↗](#)

### LEADERSHIP

- **2020-2021** Committee member in the 7th and 8th annual R. F. Baker Plant Breeding Symposium at Iowa State University.
- **2011-2013** Student representative of Agronomy in the Regional Council of Engineering and Agronomy in Brazil.

### LANGUAGE

English, Portuguese (native), basic French

## TEACHING, SEMINAR & AWARDS

### TEACHING & SEMINAR

- **2023** Speaker in the “Big data: manage your data before your data kills you” workshop at the Plant and Animal Genome Conference, in San Diego, California - USA.
- **2020 and 2022** Lecture on ‘Simulations in Plant Breeding - An emphasis on *AlphaSimR*’ in Agron 523, Molecular Plant Breeding, graduate-level course under Dr. Thomas Lübberstedt.
- **2017** Instructor of **R** programming at the University of São Paulo.

### AWARDS

- **2022** C. R. Weber for Excellence in Plant Breeding.
- **2020 and 2022** Bayer Travel Scholarship & Mentoring Program.

## ADDITIONAL INFORMATION

### MATHEUS ENJOYS...

Spending time with his family and friends • Reading • Fishing • Being outdoors • Playing guitar • Riding motorcycle.

FOR MORE INFORMATION, PLEASE ACCESS

