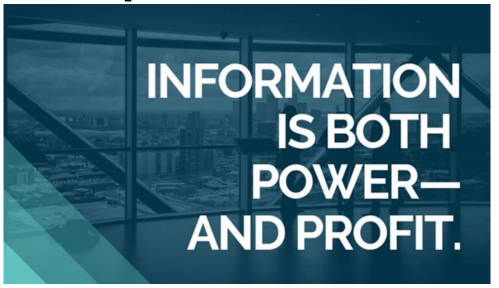


Machine learning-based AI applied to breeding

Alencar Xavier
Breeding Analyst at Corteva
Adjunct professor at Purdue

Adequate use of





Outline

1. Introduction

- More data
- Branching ML

2. Machines

- Filters
- Engines

3. Analytics

- Target G x E x M
- Validation
- Cases of study

4. Conclusion



1. Introduction

- More data
- Branching ML

2. Machines

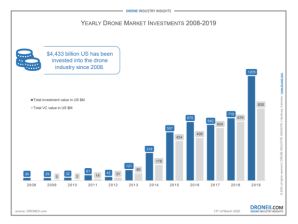
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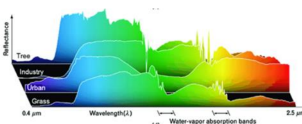
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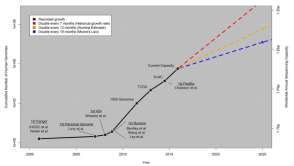
More Pheno





https://www.mdpi.com/2076-3417/12/5/2570

More Geno



The Cost of Sequencing a Human Genome, NIH. https://www.genome.gov/27565109/the-cost-of-sequencing-a-human-genome/



Stephens, Z. D.et al. (2015). Big data: astronomical or genomical? PLoS biology, 13(7), e1002195.

More Env ⁹

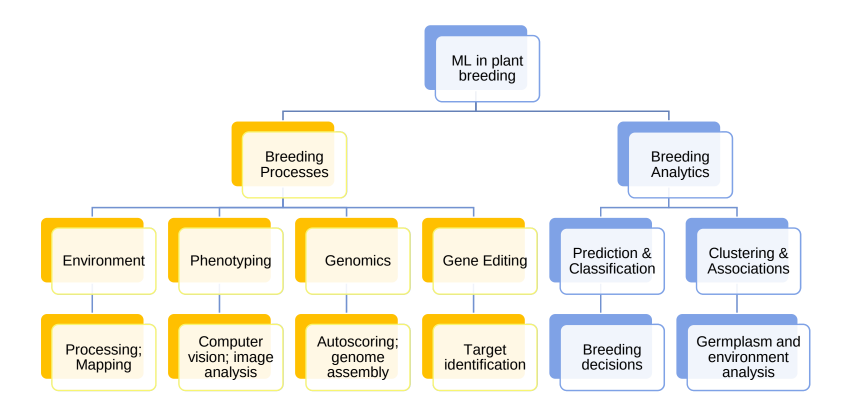
- **UC Merced GridMET**
- **NWS NOAA**
- NASA GISS, NASA power
- **Harmonized SoilDB**
- **USDA SSURGO**

More Computing _____











ML in breeding processes

Enhancing databases, automating lab tasks field work

phenotyping

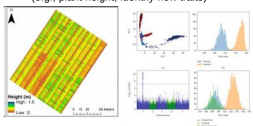
Disease, stress scoring



https://www.mdpi.com/2673-2688/2/3/26 https://www.biomedcentral.com/collections/phenomics

Phenotype automation

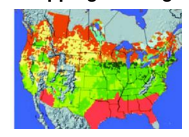
(e.g., plant height, identify new traits)



https://www.mdpi.com/2072-4292/8/12/1031 https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7706325/

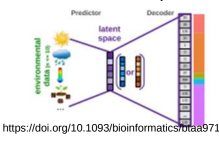
environment

Mapping / zoning



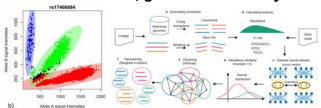
https://www.publish.csiro.au/cp/CP14007

Latent weather, soil



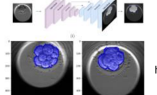
biotech

SNP calls, genome assembly

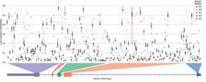


https://doi.org/10.1186/1753-6561-3-s7-s58 https://www.nature.com/articles/s41467-022-29843-y

Embryo rescue DH production



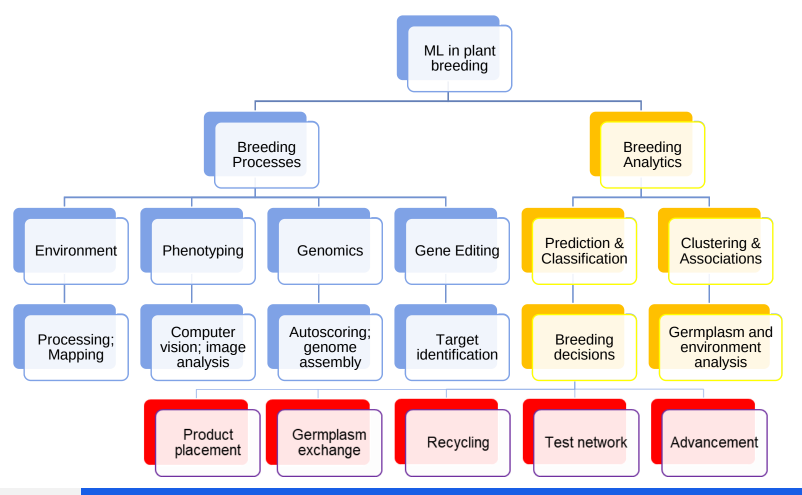




https://doi.org/10.1093/bioinformatics/btab268

https://www.nature.com/articles/s41598-022-06336-v







1. Introduction

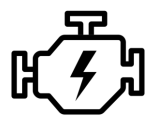
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Machine Learning Engines



A new approach fits multivariate genomic prediction models efficiently

Alencar Xavier^{1,2*†} and David Habier^{1*†}

Walking through the statistical black boxes of plant breeding

Alencar Xavier¹ · William M. Muir² · Bruce Craig³ · Katy Martin Rainey¹

ORCID ID: 0000-0

Plant Breeding

Technical nuances of machine learning: implementation and validation of supervised methods for genomic prediction in plant breeding

Alencar Xavier 1*

Impact of Genomic Prediction Model, Selection Intensity, and Breeding Strategy on the Long-Term Genetic Gain and Genetic Erosion in Soybean Breeding

Using unsupervised learning techniques to assess interactions among complex traits in soybeans

Alencar Xavier · Benjamin Hall · Shaun Casteel · William Muir · Katy Martin Rainey

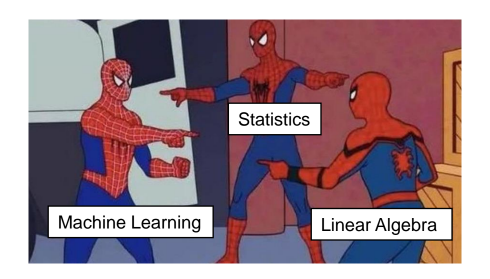
Article

Joint Modeling of Genetics and Field Variation in Plant Breeding Trials Using Relationship and Different Spatial Methods: A Simulation Study of Accuracy and Bias

Efficient Estimation of Marker Effects in

Éder David Borges da Silva 1,2,40, Alencar Xavier 3,40 and Marcos Ventura Faria 20







Key idea of supervised learning: FILTERING

Simple filter

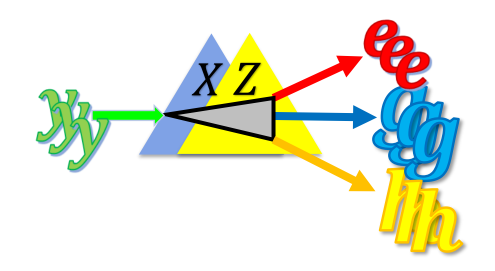
$$y = g + e$$

Multiple filters

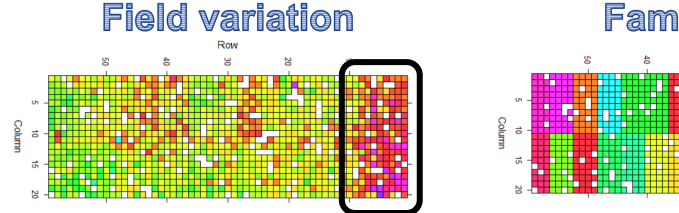
$$y = g + h + e$$

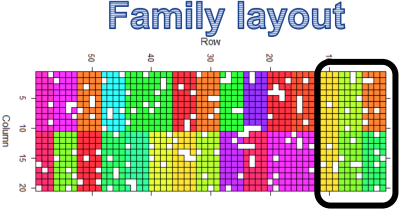
Multi-task filter

$$Y = G + H + E$$



Why bother with multiple filters?

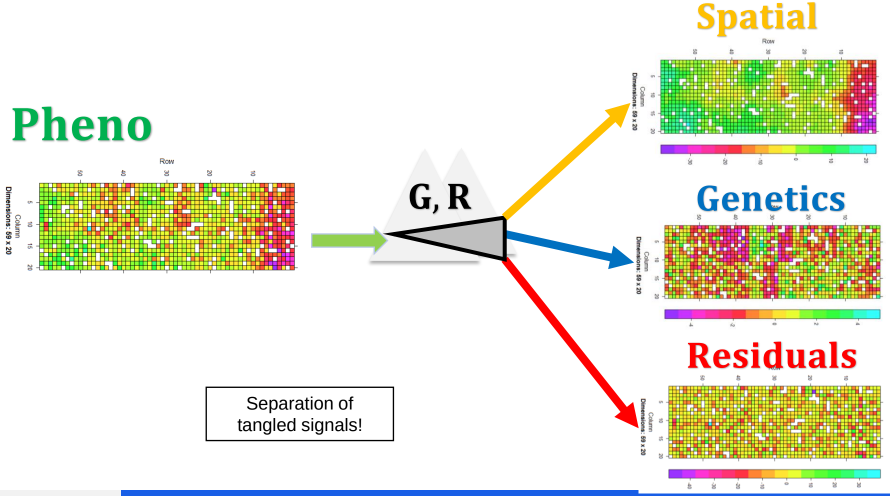




Some families were placed on unfavorable side of the field...

SoyNAM field, Indiana 2014



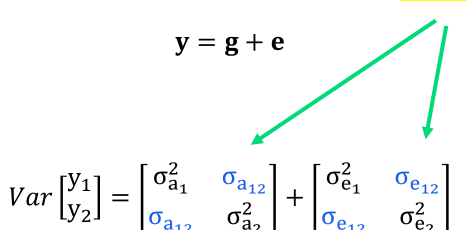




Why bother with multi-task filters?

Simple (bivariate) model:

INFORMATION GAIN



Why bother with multi-task filters?

$$\begin{aligned} \mathbf{y} &= \mathbf{Z}\mathbf{g} + \mathbf{e}, & \mathbf{y} \sim \mathbf{N}(\mathbf{0}, \mathbf{V}) \\ \begin{bmatrix} \mathbf{y}_1 \\ \mathbf{y}_2 \end{bmatrix} &= \begin{bmatrix} \mathbf{Z}_1 & \mathbf{0} \\ \mathbf{0} & \mathbf{Z}_2 \end{bmatrix} \begin{bmatrix} \mathbf{g}_1 \\ \mathbf{g}_2 \end{bmatrix} + \begin{bmatrix} \mathbf{e}_1 \\ \mathbf{e}_2 \end{bmatrix} \end{aligned}$$

Covariance structure

$$V = G \otimes \Sigma_a + I \otimes \Sigma_e = G \otimes \begin{bmatrix} \sigma_{a_1}^2 & \sigma_{a_{12}} \\ \sigma_{a_{12}} & \sigma_{a_{2}}^2 \end{bmatrix} + I \otimes \begin{bmatrix} \sigma_{e_1}^2 & \sigma_{e_{12}} \\ \sigma_{e_{12}} & \sigma_{e_{2}}^2 \end{bmatrix}$$

Model equation

$$\begin{bmatrix} Z_1' \Sigma_e^{11} Z_1 + G^{-1} \Sigma_a^{11} & Z_1' \Sigma_e^{12} Z_2 + G^{-1} \Sigma_a^{12} \\ Z_2' \Sigma_e^{12} Z_1 + G^{-1} \Sigma_a^{12} & Z_2' \Sigma_e^{11} Z_2 + G^{-1} \Sigma_a^{22} \end{bmatrix} \begin{bmatrix} g_1 \\ g_2 \end{bmatrix} = \begin{bmatrix} Z_1' (\Sigma_e^{11} y_1 + \Sigma_e^{12} y_2) \\ Z_2' (\Sigma_e^{22} y_2 + \Sigma_e^{12} y_1) \end{bmatrix}$$

Univariate vs bivariate

$$\begin{split} g_1 &= (Z_1' \Sigma_e^{11} Z_1 + G^{-1} \Sigma_a^{11})^{-1} (Z_1' \Sigma_e^{11} y_1) \\ g_1 | g_2 &= (Z_1' \Sigma_e^{11} Z_1 + G^{-1} \Sigma_a^{11})^{-1} (Z_1' (\Sigma_e^{11} y_1 + \Sigma_e^{12} y_2) - (Z_1' \Sigma_e^{12} Z_2 + G^{-1} \Sigma_a^{12}) g_2) \end{split}$$

GAIN

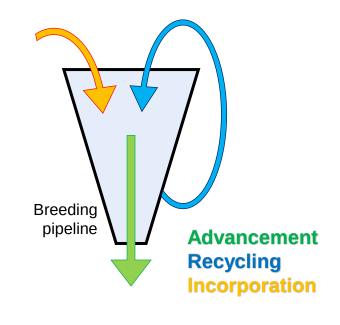
Does the choice of filter matter?

• **ADDITIVE LINEAR FILTERS** (GEBV)

- Pattern: ADDITIVE GENETICS heritable
- Method: GBLUP, RIDGE, LASSO
- Suits: RECYCLING, ADVANCEMENT

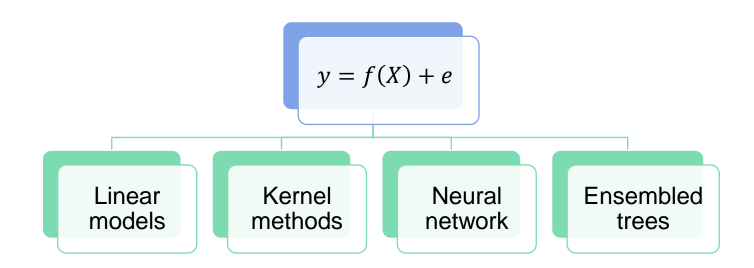
• NON-LINEAR FILTERS (EGV)

- Pattern: ANY GENETIC SIGNAL
- Method: RKHS, DNN, Random Forest
- Suits: ADVANCEMENT, PRODUCT PLACEMENT





Main classes of learners



Solving: y = Xb + e

Finding \rightarrow argmin($e'e + \lambda b'b$)



(Use diagonals of LHS)

$$\hat{b}_{j}^{t+1} = \frac{x_{j}'(y - X_{-j}\hat{b}_{-j})}{x_{j}'x_{j} + \lambda}$$



$$\hat{b}^{t+1} = b^t - \frac{2r}{n} [X'(y - X\hat{b}^t) + \lambda \hat{b}^t]$$

Second order

$$\hat{\mathbf{b}} = (\mathbf{X}'\mathbf{X} + \lambda)^{-1}(\mathbf{X}'\mathbf{y})$$



l've created a monster!!

Used for p>>n solvers

glmnet, BGLR, bWGR, GS3

Used for Deep Neural Nets

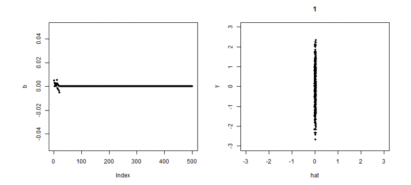
TensorFlow Keras, PyTorch, MXNet

Used for everything else

ASREML, Ime4, SAS

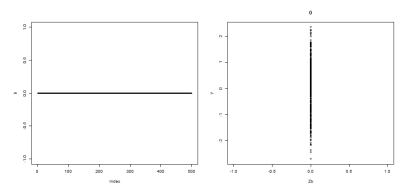


Coordinate descent



$$\hat{b}_{j}^{t+1} = \frac{x_{j}'(y - X_{-j}\hat{b}_{-j})}{x_{i}'x_{i} + \lambda}$$

Gradient descent



$$\hat{\mathbf{b}}^{t+1} = \mathbf{b}^t - \frac{2\mathbf{r}}{\mathbf{n}} [\mathbf{X}'(\mathbf{y} - \mathbf{X}\hat{\mathbf{b}}^t) + \lambda \hat{\mathbf{b}}^t]$$

What about the deep learning? 3

$$y = \alpha(\alpha(XB_1)B_2)b_3 + e$$

i.e., just a "stack of solvers"



Data > Method

Unnecessarily complex analysis should not be used as a foil to disguise lower quality datasets

Kruuk (2004 apud Walsh and Lynch 2018)



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Analytics





"Breeding objective"

Set of traits of interest (TOI)
 bred into a

Target population of genotypes (TPG)
 for a given

Target population of environments (TPE)



TPE, TPG, TPM

Target population of environments (TPE)

- Influences accuracies via GxE correlation
- Which environments should I be able to predict?

Target population of genotypes (TPG)

- Influences accuracies via genetic relationship
- Which genetics should I be able to predict?
- Target population of management (TPM)
 - Herein nested in TPE

From QTLs to Adaptation Landscapes: Using Genotype-To-Phenotype Models to Characterize G×E Over Time

Daniela Bustos-Korts¹*, Marcos Malosetti¹, Karine Chenu², Scott Chapman^{3,4}, Martin P. Boer¹, Bangyou Zheng³ and Fred A. van Eeuwiik¹*

What Should Students in Plant Breeding Know About the Statistical Aspects of Genotype × Environment Interactions?

Fred A. van Eeuwiik,* Daniela V. Bustos-Korts, and Marcos Malosetti

An Equation to Predict the Accuracy of Genomic Values by Combining Data from Multiple Traits, Populations, or Environments

Yvonne C J Wienties . Piter Bijma, Roel F Veerkamp, Mario P L Calus

Genetics, Volume 202, Issue 2, 1 February 2016, Pages 799–823, https://doi.org/10.1534/genetics.115.183269

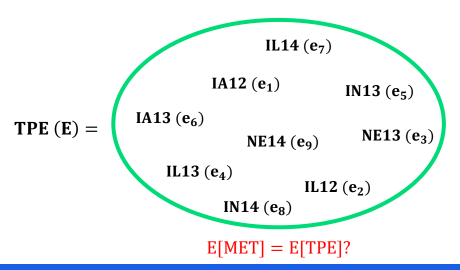


TPE

• Any given trial happens in each environment-management combination, that is sample of much larger population:

$$e_i \in E$$

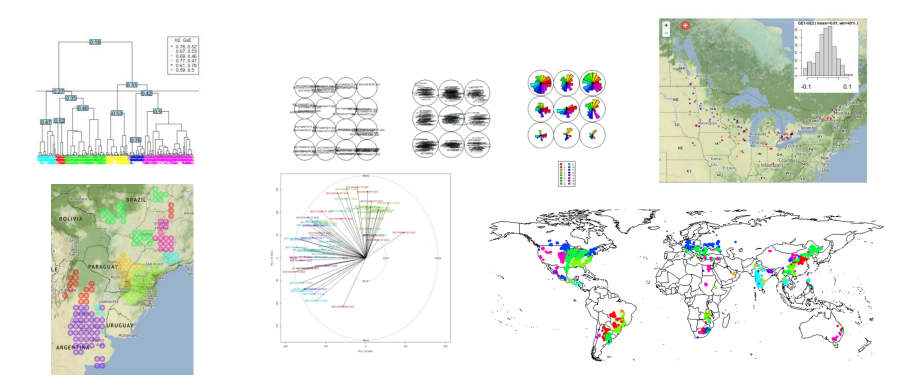
That is:



$$\begin{bmatrix} y_{e_i} \\ y_{e_j} \\ g_E \end{bmatrix} = \begin{bmatrix} \sigma_{g(e_i)}^2 + \sigma_{\varepsilon(e_i)}^2 & \sigma_{g(e_i,e_j)} & \sigma_{g(e_i,E)} \\ \sigma_{g(e_j,e_i)} & \sigma_{g(e_j)}^2 + \sigma_{\varepsilon(e_j)}^2 & \sigma_{g(e_j,E)} \\ \sigma_{g(E,e_j)} & \sigma_{g(E,e_j)} & \sigma_{g(E,e_j)}^2 \end{bmatrix}$$



NOTE: GxExM patterns within TPE are largely assessed using different methods of ML





TPG + TPE

- Accuracy (<u>Wientjes et al 2016</u>) = correlation(true signal, estimated signal),
- It is a function of heritability, GxE, representativeness of the calibration set
- For:

$$y = g + e$$
,
 $var(y) = V$, $var(g) = G$

Then accuracy is

$$a_{i} = cor(g_{i}, \hat{g}_{i}) = \frac{cov(g_{i}, \hat{g}_{i})}{var(g_{i})var(\hat{g}_{i})} = \frac{var(\hat{g}_{i}) r_{GxE}^{2}}{var(g_{i})var(\hat{g}_{i})} = r_{GxE}^{2} \sqrt{\frac{G_{i,y}V^{-1}G_{y,i}}{G_{i,i}}}$$

Thus, we know how much signal to expect in any given prediction

Validation schemes

1) CV type – Test intent

- Random CV = Upper-bound predictive potential
- <u>Leave-one-out</u> = Assess structured scenarios (e.g., geography-out, year-out)
- **Holdout** = Reproduce true applications (e.g., predict individuals from upcoming)

2) TPE/TPG relation

	Genotype	Environment	Difficulty
CV00	New	New	****
CV0	Observed	New	***
CV1	New	Observed	***
CV2	Observed	Observed	*

Adapted from Crossa et al. (2017) doi.org/10.1016/j.tplants.2017.08.011

3) Signal availability

Genetic information available in different cross-validation setups

- Intra-family: Linkage*
- · Within-family: Linkage and LD
- Across-family: Relationships**, Linkage and LD
- Leave-family-out: Relationships and LD
- Untested environments: Same as above x (GxE)



Validation metrics

Correlations

- Most common metrics in breeding (e.g., predictability)
- Pertinent to ranking and selection of complex traits

Prediction error

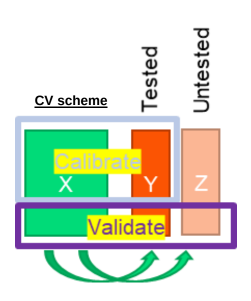
- Utilized when the predicted values must be as close as possible to original scale
- Pertinent to risk prediction (e.g., disease risk)

Success

- Accommodate complex or subjective criteria, independent or otherwise
- Pertinent to decision involving data from multiple sources (e.g., advancement)



Amount of signal that can be captured in different structures

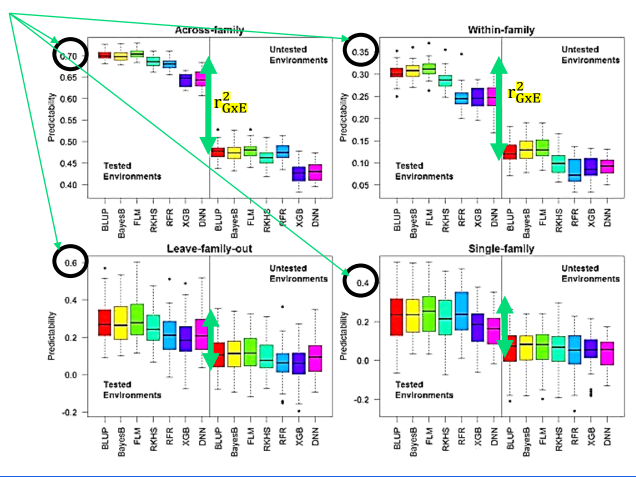


SoyNAM data

ES: 2012 (7 loc) PS: 2013 (4 loc) #Fam = 40

Genos = 5600

SNPs = 4300Obs: 3k-5k obs/loc



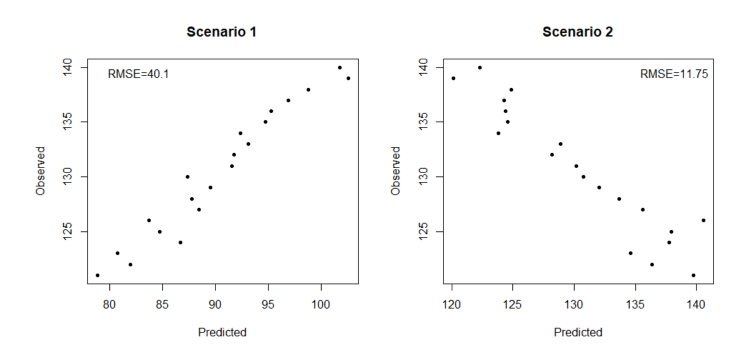


Case of study



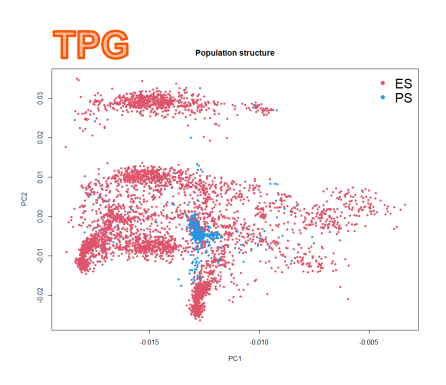


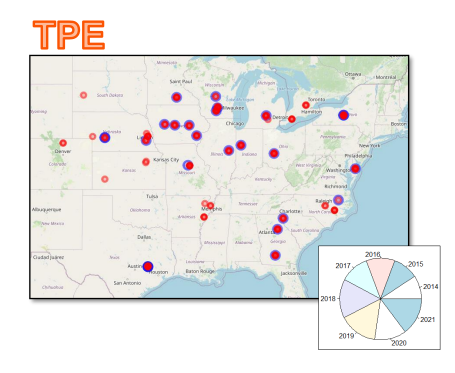
Evaluation criterion





2022 G2F GxE prediction competition







What was modeled?

$$y|E_i = \mu_i + g|E_i$$
 (Two FILTERS)

Phenotype @ ith Loc = ith Loc Mean + Genetic effect @ ith Loc

- The winning approach:
 - Predict location means using mixed model and random forest
 - Predict genetic performance with index from <u>multi-response</u> based on <u>TPE/TPG</u>



2022 G2F GxE prediction competition

Realized results

Ranking with alternative metrics

Team Name	Within RMSE	Toom Name	Cor Within Loc	Team Name	Cor Across Loc
	WILLIIII KIVISE	Team Name	Cor within Loc	Iealii Naille	COI ACTOSS LOC
CLAC	<mark>2.329</mark>	<mark>CLAC</mark>	<mark>0.357</mark>	breedingteam	0.650
igorkf	2.345	CGM	0.353	DataJanitors	0.644
phenomaize	2.374	MPB_Group	0.342	CLAC	<mark>0.631</mark>
UCD_MegaLMM	2.387	UCD_MegaLMN	0.338	Purdue	0.631
CGM	2.391	SmAL	0.285	UCD_MegaLMM	0.628
breedingteam	2.398	DeepCropVision	0.281	phenomaize	0.617
Purdue	2.402	CropEnthusiast	0.279	igorkf	0.600
SmAL	2.425	AllModelsAreWro	ng 0.272	CGM	0.587
ML_APT	2.472	DataJanitors	0.256	SmAL	0.586
MPB_Group	2.544	supermanwasd	0.243	AllModelsAreWrong	0.575

Source: Jacob Washburn, Jose Ignacio Varela, Alencar Xavier



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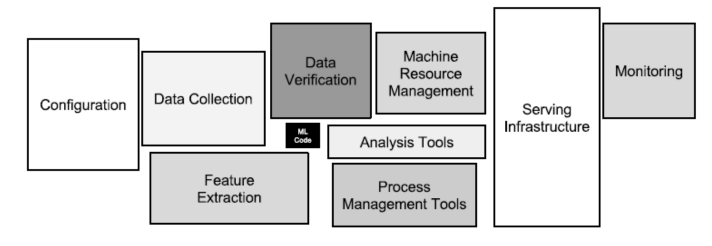
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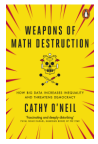
There is more to ML than proof of concepts using cross-validations

Hidden Technical Debt in Machine Learning Systems

doi/10.5555/2969442.2969519



- How easily can an entirely new algorithmic approach be tested at full scale?
- What is the transitive closure of all data dependencies?
- How precisely can the impact of a new change to the system be measured?
- Does improving one model or signal degrade others?
- How quickly can new members of the team be brought up to speed?





Thank you for your attention!

Final remarks:

- 1) Plant breeding uses machine learning for multiple purposes in processes and analytics
- 2) Filter settings are important to maximize signal, but it is less important than data
- 3) Validation metrics and validation schemes matter to design meaningful models

Questions??

Alencar Xavier

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