



# Using Random Forests As A Prescreening Tool for Genomic Prediction: Impact of Subsets of SNPs on Prediction Accuracy of Total Genetic Values

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# Why Machine Learning Methods Become Popular in Large Genomic Data Analyses ?

- Dealing with “*Large P and Small N*” problem
- Black–box approaches (No prior knowledge required)
- Taking multiple interactions or correlations among predictor variables (e.g. SNP-SNP interactions) into account
- High prediction accuracy (building training and validation procedures into algorithms)

# Knowledge Gap in Genomic Prediction of Total Genetic Values

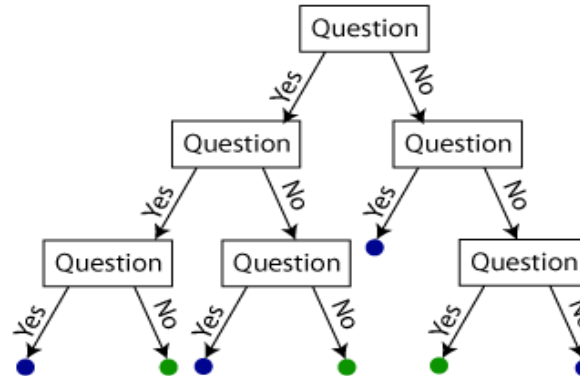
- Do non-additive effects captured by machine learning methods contribute to the prediction accuracy of **total genetic values** ?



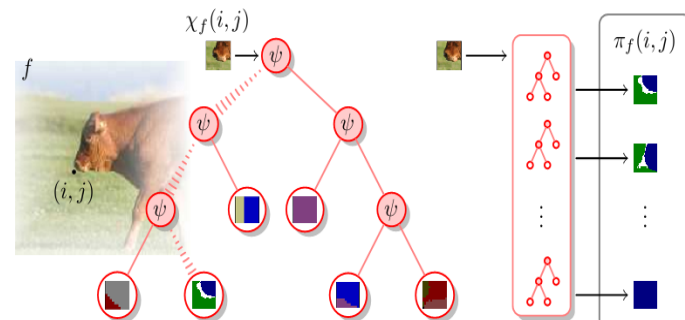
(additive + dominance genetic values)

# Machine Learning Method – Random Forests (RF)

- Leo Breiman, *Random Forests*, Machine Learning, 45, 5-32, 2001.
- A nonparametric tree-based ensemble machine-learning method for classification or regression of multiple variables.



<http://shapeofdata.wordpress.com/2013/07/09/random-forests/>



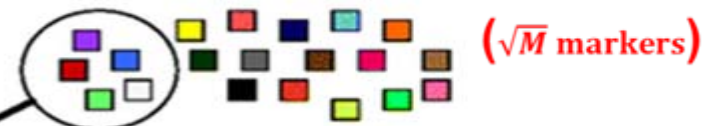
<http://pdollar.wordpress.com/2013/03/08/structured-random-forests/>

# Random Forests – How Does It Work?

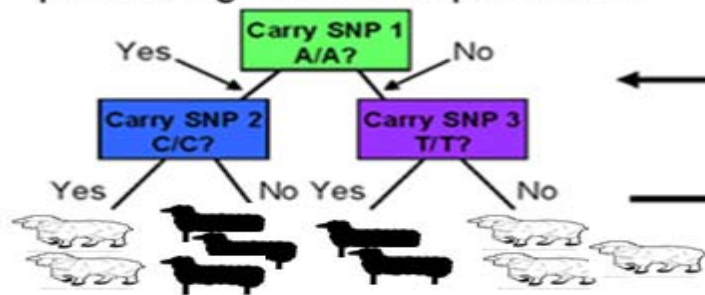
Step 1: select subsample of cases/controls,  
set aside additional samples **(2/3 population)**



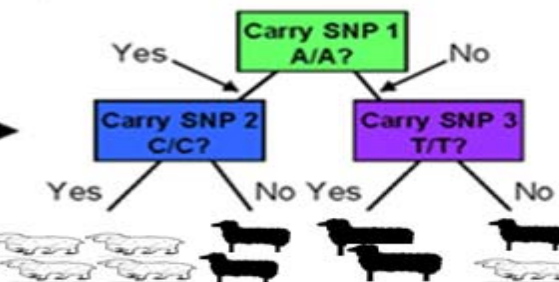
Step 2: select subset of SNPs



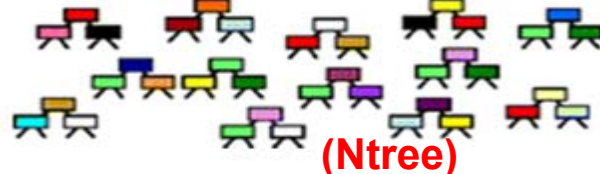
Step 3: Create single tree via recursive  
partitioning on subsample/subset



Step 4: Run additional  
(independent) cases/controls  
through tree to calculate variable  
importances



Step 5: Repeat steps 1-4 to create a  
forest of trees



Step 6: Calculate SNP variable  
importance across all trees in forest

4.54    4.01    2.48

Nicodemus et al. 2010. Hum Genet 127:441-452.

# SNP Variable Importance Value (VIM)

- RF: **%IncMSE** (% Increasing in Mean Squared Error when a SNP is not included)

**Larger the value, the more important the SNP is**

# Beef CRC Cattle Dataset

- 2,109 Brahman Cattle
- 651,253 SNPs (MAF > 0.01)
- Phenotype – Yearling Weight (**pre-adjusted** for average heterozygosity of SNPs, contemporary group and age effects)

# Application of RF in Identifying Subsets of SNP for Genomic Prediction

For each data set, *using 80% Brahman cattle*

Identify Top 500,1000, ... 50,000 SNP Using VIM Values from RF

*Using 20% Brahman cattle*

Predict  $\sigma_a^2$  and  $h^2$  values:

$$\text{GBLUP: } \mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{Z}\mathbf{u} + \mathbf{e}$$
$$V(\mathbf{u}) = \mathbf{G}\sigma_u^2 \text{ and } V(\mathbf{e}) = \mathbf{I}\sigma_e^2 \quad \mathbf{G} = \frac{\mathbf{M}\mathbf{M}^T}{2\sum p_i(1-p_i)}$$

Dominance Model

$$\mathbf{y} = \mathbf{1}_n \mu + \mathbf{X}\mathbf{b} + \mathbf{het}\boldsymbol{\beta} + \mathbf{g} + \mathbf{d} + \mathbf{e},$$

The average heterozygosity of each animal

Genomic breeding values  
 $N \sim (0, \text{GRM}\sigma_g^2)$

Dominance deviations  
 $N \sim (0, \text{DRM}\sigma_d^2)$

Genomic prediction accuracy of total genetic values

# Variance Estimates from Subsets of SNPs

## (RF Selected vs Evenly Spaced vs All SNPs)

	Additive Model			Additive + Dominance Model			
	$h_a^2$	$\sigma_a^2$	ACC	$h_a^2$	$\sigma_a^2$	$\sigma_d^2$	ACC
RF							
500	0.21	140.3	0.45	0.21	140.1	14.2	0.45
1,000	0.26	171.6	0.49	0.26	171.7	24.2	0.49
5,000	0.39	254.9	0.55	0.39	253.4	58.0	0.55
50,000	0.45	299.0	0.60	0.44	294.0	205.1	0.60
Even							
500	0.04	24.8	0.18	0.03	24.8	0.8	0.18
1,000	0.03	25.8	0.21	0.03	23.5	2.2	0.21
5,000	0.09	58.9	0.24	0.08	58.6	15.7	0.24
50,000	0.34	236.5	0.29	0.34	234.2	55.6	0.29
All SNPs	0.38	259.4	0.44	0.38	258.4	49.9	0.44

# Conclusions

- Fitting dominance into the genomic model had **little impact** on the accuracy of genomic prediction of breeding values.
- RF has **potential** to be used as a pre-screening tool for:
  - a) **reduction of high dimensionality** associated with large genomic data;
  - b) identification of **subsets of useful SNPs** for genomic prediction of breeding values.

# Application of RF in Identifying Subsets of SNP for Genomic Prediction of Cattle Live Weight

## Fine-Tuning RF Parameters

$Ntree = 10,000, 12,000, \dots 20,000$

$mtry = \sqrt{M}, 2*\sqrt{M}$  ( $M$ : total no. SNPs)



## Random 5-fold cross-validation scheme:

- Identify Top 500,1000, ... 50,000 SNP Using VIM Values from RF
- Genomic prediction accuracy of total genetic values

# GRM and DRM Calculations

$$G = \frac{Z_a Z'_a}{2 \sum_{k=1}^m p_k q_k}$$

- $Z_a (nxm)$

$$\begin{cases} 2 - 2p_k & (\text{AA}) \\ 1 - 2p_k & (\text{AB}) \text{ (VanRaden et al., 2008)} \\ -2p_k & (\text{BB}) \end{cases}$$

- $p_k$  - menor allele frequency of locus  $k$

$$D^* = \frac{Z_d Z'_d}{4 \sum_{k=1}^m p_k^2 q_k^2}$$

- $Z_d (nxm)$

$$\begin{cases} 2q_k^2 & (\text{AA}) \\ 2p_k(1 - p_k) & (\text{AB}) \text{ (Vitezica et al., 2013)} \\ -2p_k & (\text{BB}) \end{cases}$$

- Matrix  $D^*$  was combined with identity matrix  $I$  as  $D = 0.95D^* + 0.05I$  to improve numerical stability

# Variance Estimates from Subsets of SNPs

## RF Selected vs Evenly Spaced vs All SNPs

	Additive Model			Additive + Dominance Model			
	$h_a^2$	$\sigma_a^2$	% Total $\sigma_a^2$	$h_a^2$	$\sigma_a^2$	$\sigma_d^2$	$\sigma_p^2$
RF							
500	0.21 (0.03)	140.3 (22.8)	667.0 (26.5)	0.21 (0.08)	140.1 (22.7)	14.2 (8.6)	668.1 (26.2)
1,000	0.26 (0.03)	171.6 (25.0)	658.8 (26.1)	0.26 (0.03)	171.7 (25.0)	24.2 (11.4)	660.2 (26.2)
5,000	0.39 (0.04)	254.9 (32.7)	658.2 (26.3)	0.39 (0.04)	253.4 (32.5)	58.0 (21.3)	658.9 (26.4)
50,000	0.45 (0.04)	299.0 (38.7)	669.2 (26.7)	0.44 (0.05)	294.0 (38.3)	205.1 (60.8)	670.3 (27.5)
Even							
500	0.04 (0.02)	24.8 (14.2)	691.0 (26.1)	0.03 (0.02)	24.8 (8.7)	0.8 (1.7)	691.8 (24.9)
1,000	0.03 (0.02)	25.8 (14.4)	691.6 (25.9)	0.03 (0.02)	23.5 (12.6)	2.2 (5.1)	690.3 (25.5)
5,000	0.09 (0.03)	58.9 (21.2)	703.3 (27.8)	0.08 (0.03)	58.6 (21.2)	15.7 (15.6)	705.7 (28.2)
50,000	0.34 (0.05)	236.5 (38.7)	690.0 (26.5)	0.34 (0.05)	234.2 (45.7)	55.6 (51.8)	689.8 (26.9)
All SNPs	0.38 (0.05)	259.4 (38.4)	680.9 (26.5)	0.38 (0.05)	258.4 (38.2)	49.9 (33.5)	680.5 (26.5)

# Genomic Prediction Accuracy of Total Genetic Values

	Additive Model	Additive + Dominance Model
	Acc	Acc
RF		
500	0.45	0.45
1,000	0.49	0.49
5,000	0.55	0.55
50,000	0.60	0.60
Even		
500	0.18	0.18
1,000	0.21	0.21
5,000	0.24	0.24
50,000	0.29	0.29
All SNP	0.44	0.44