Regularization and shrinkage estimation

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Large p small n problem

Need regularization methods

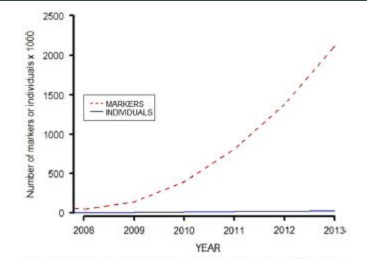


Fig. 1. Relationship between the number of marker effects to be estimated in regression models and the number of genotyped individuals with phenotype in livestock populations.





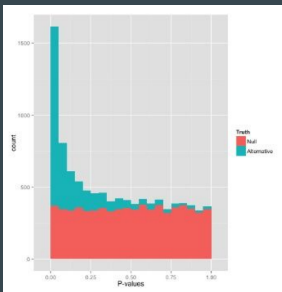
- many tests \rightarrow many false positives
 - e.g. 2000 (independent) tests, α =0.05 → How many expected false positives?

GUESS

- many tests \rightarrow many false positives
 - e.g. 2000 (independent) tests, α =0.05 \rightarrow How many expected false positives?

100 false positives by chance alone

- multiple testing problem
- many SNPs, many statistical tests, many p-values
 - \circ Some *p*-values are lower than the α significance level just by chance







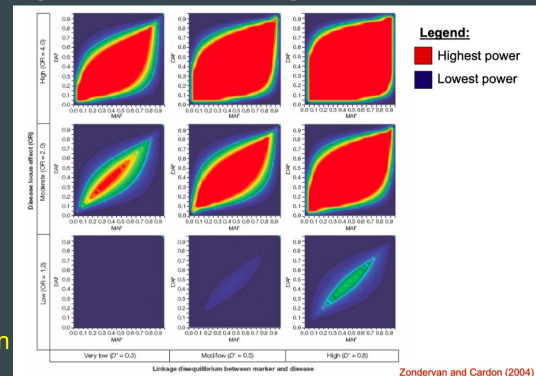
Main difficulties to capture genetic marker signal

- Low effect size (impairs statistical power)
- LD between markers (markers share QTL information)
- Epistasis and interaction (models assume linearity)
- Many markers with (probably) null effect (sparsity)

Main difficulties to capture genetic marker signal

- Sample size
- Magnitude of effect
- MAF marker
- MAF atl
- Range of LD
- Likelihood of the model
- Experimental design

Prediction not that much interest on real effect of markers





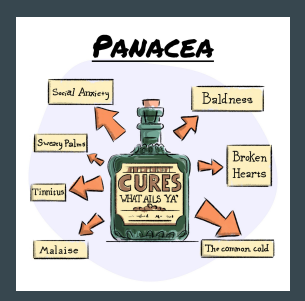
How to cope with the problem

- Increase the sample size (e.g. Bio Banks, large reference populations)
- Reduce the number of tests
 - o Based on LD
 - Choose relevant regions (functional analysis)
- Decrease the significance threshold
 - Bonferroni correction
 - False discovery rate
 - o q-values
- Go Bayesian...



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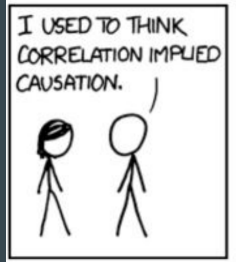


Find a causal mutation

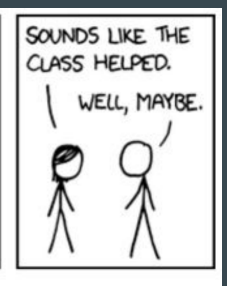
A needle in a haystack



Remember







https://xkcd.com/552/

Prediction aims at good correlation or classification performance!



Luckily:

Predict genomic values is different from finding causal mutations





Inference vs Prediction

Inference

Determine the effect of a covariate on the response

- Determine the <u>causal</u>
 relationship between a
 covariate and the response
- More difficult (in general)

Prediction

- Educated <u>quess</u> of the outcome
- Expected behaviour in the future
- Based on proxies/markers

Inference vs Prediction

GWAS goal

- Detect genomic markers/regions associated to phenotypes (traits) of interest
- Find biological pathways of interest
- Interaction between treatments/drugs and genes

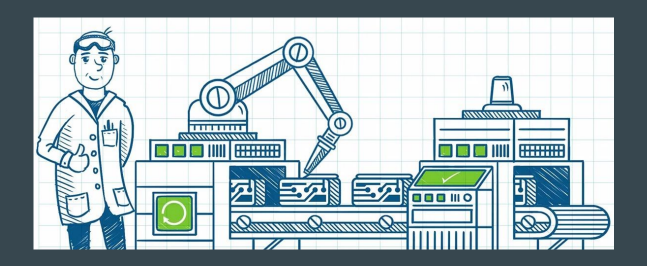
GWP goal

- Calculate the genomic risk score
 /predisposition of a disease or trait
- Calculate the genomic merit of individuals
- Predict future performance
- Sire/dam selection in animal breeding
- Don't need to know the real marker effect



Prediction

Aim: construct a predictive (statistical) machine that provides an accurate guess of a yet to be observed phenotype



Large dimensionality problem

$$y = X\beta + \epsilon,$$

Minimize a loss function (usually minimize MSE)

$$\hat{\beta} = \arg\min_{\beta} \sum_{i} (y_i - \beta_1 - \beta_2 x_i)^2,$$

- p>>>>n
 - Run out of degrees of freedom (cannot estimate so many unknowns).
- $\hat{\beta}_{R} = \arg\min_{\beta} \{ \|y X\beta\|_{2}^{2} + \lambda_{2} \|\beta\|_{2}^{2} \},$ Need a more restrictive penalty function
- Or set a number of unknowns to zero (feature selection e.g. LASSO or elastic net)

$$\|\beta\|_2^2 \le c_2(\lambda_2)$$

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 $\|\beta\|_0 \le c_0(\lambda_0)$

$$\|m{eta}\|_1 = \sum_{j=1}^p |m{eta}_j|,$$

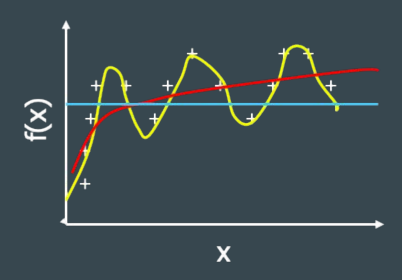


Need to modelize the data

Caution with overfitting with so many variables (SNPs)

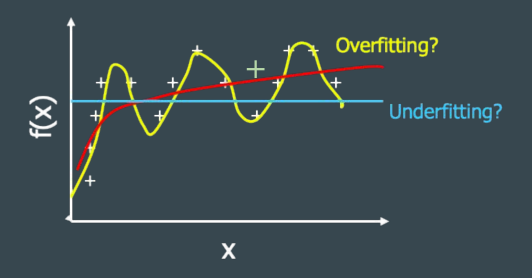


Which model does fit data the best?



Need to modelize the data

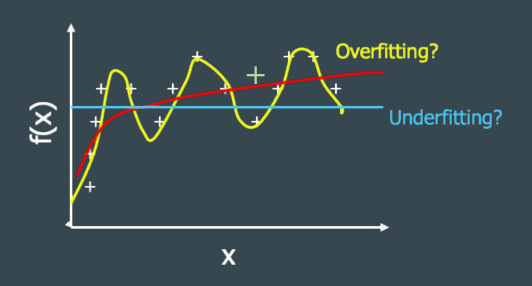
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Need to modelize the data

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Important to control the variance-bias trade-off



Bias

a systematic distortion of a statistical result due to a factor not allowed for in its derivation





Variance

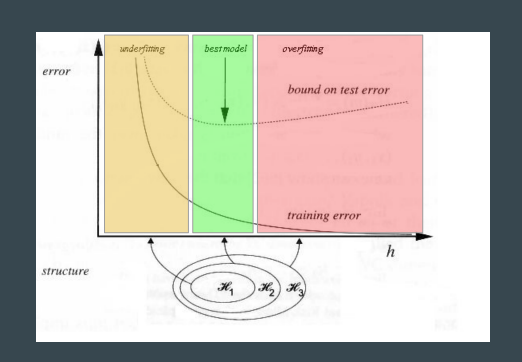
It measures how far a set of numbers is spread out from their average value



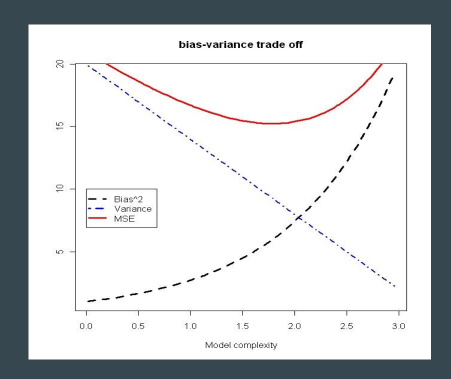
Bias - Variance trade off



Increase model complexity might overfit the data, but impair predictive ability



Bias - Variance trade off



Parameters that control the variance-bias trade off

- Variances in the mixed models
- Lambda parameter in the LASSO
- Hyperparameters in Machine Learning methods

How to control large dimensionality

Applied more restrictive loss functions

$$\hat{\beta} = \arg\min_{\beta} \sum_{i} (y_i - \beta_1 - \beta_2 x_i)^2,$$

$$\hat{\beta}_{R} = \arg\min_{\beta} \{ \|y - X\beta\|_{2}^{2} + \lambda_{2} \|\beta\|_{2}^{2} \},$$

$$J[g(x_{\cdot})|\lambda] = \frac{1}{2}[y - W\theta - g(x_{\cdot})]'R^{-1}[y - W\theta - g(x_{\cdot})] + \frac{\lambda}{2}||g(x_{\cdot})||_{H}^{2},$$



Lasso, Bayesian LASSO, RKHS

How to control large dimensionality

- Apply simple models repeatedly over the data or residuals from previous iteration
 - Neural Networks, LASSO, Boosting, Random Forest
- Resample data during inference to increase variability of results
 - o Random Forest, Bagging
- Variable selection
 - o Random Forest, Neural Networks
- **(**
- Use **cross validation** to train predictive ability



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- Use **cross validation** to train predictive ability

More than one option



Considerations



 If a model have very high goodness of fit, it will not generalize the data (low predictive ability)



• Check predictive ability tuning hyperparameters and using internal and external cross validation.