

Developing a parsimonious predictor for binary traits in sugar beet (*Beta vulgaris*)

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1 Introduction

The primary goal of breeding schemes in farm animals and crops is generally to increase the agricultural output. Production traits are typically quantitative continuous variables (e.g. milk yield in dairy cattle, or per hectare yield in maize and rice). Many traits of importance in plant and animal breeding follow nonetheless a discrete categorical distribution, both ordered (e.g. calving ease in cattle, grain texture in rice) and unordered (e.g. grain pigmentation in rice, coat colour in cattle). A special case is that of binomial traits, which can take up only two different values, like disease resistance/susceptibility or

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presence/absence of a morphological characteristic. Annual bolting (flowering) behaviour and root vigor are examples of binomial traits of agronomic importance in sugar beet (*Beta vulgaris*).

Advances in biotechnology and genomics, and the advent of high-density molecular markers (especially single-nucleotide polymorphisms, SNP) genotyping have led to the emergence of molecular breeding [8].

One exciting application of molecular breeding are genomic predictions ([7]): something on genomic predictions in general. The recent publication of the reference genome for *Beta vulgaris* genome [4] facilitated molecular breeding also in this crop species. Genomic predictions in sugar beet already done both for continuous ([6,12]) and binary ([2]) traits.

The concept of parsimony: when many possible predictors are available, it is useful to select a subset to limit analysis cost and time. Moreover: use the minimum necessary information set, Occam razor ([3]), and so forth. A model need to be simpler than the data the it fits/explains (e.g. knn with k=1)

Given two models that fit the data, the simplest has to be chosen (Occam's razor)

As the technology advances, and available predictors grow, not only the prediction precision becomes important, but also the actual cost must be considered.

Sugar beets in particular: we work on root vigor [2].

In this paper we propose statistical methods to highlight and select the most useful predictors given a set. We started on real world data and validated our approach on a XXX dataset. We found that it is possible to strongly reduce the dimension of the predictors set and still achieve high performance.

2 Material and methods

2.1 Plant material and SNP genotypes

Root vigor. Available data. SNP technology used, imputation.

Copy-paste from other articles. Dataset description. Text with citations [11] and [10].

2.2 Predictor development procedure

A two-step approach was adopted for the construction of a parsimonious predictor for root vigor.

- a ranker to rank the various available predictors (SNPs in our case). We used the BOSS algorithm - this is an iterative step. we progressively reduced the predictors set, taking away the least useful predictor and applying to the resulting subset a ridge logistic regression approach. Thus, we obtained as many performances estimation as the number of original predictors.

2.2.1 Rank of predictors

This explain the BOSS algorithm [9]

2.2.2 Selection of predictors and classification method

We take one predictor out at each iteration You put the model formula for ridge logistic regression

2.2.3 Predictive ability

Cross validation: how many times, what fractions. Explanation of error rate and other parameters (ROC?)

2.3 Comparison with another method to rank predictors

Another ranker: why use one, and its description. P value and SNP effect (as it is done in GWAS)

SNP variance [5]

2.4 Software

R, weka, perl.

3 Results

Possible charts: - Precision as a function of the number of predictors. - Break-down of two types of error.

If possible: probability of assignment as a function of predictors, maybe with ROC curve? Maybe in discussion?

Plot AUC vs # predictors?

Something on the ranking of predictors (variable importance vs SNP effects)?

Plot coefficients from BOSS?

4 Discussion

General overview why error rates are not evenly distributed? Reminder: it works very well because of LD and H2

Unstable below 30/40 SNPs; little “bump” around 20 SNPs: more marked with BOSS, but also visible with GWAS. Why there? SNPs with large effect on the trait, but low significance? SNPs with large effect but low LD (with

the QTL)? In the latter case, the marker might sometimes be in the opposite phase

Based on results, a panel of 30-35-40 SNPs is recommended for accurate prediction of root vigor (move to breeding applications? Together with development of a custom-chip?)

4.1 SNP effects

Manhattan plot with BOSS weights and weights from the other articles, somehow compared (same chart? two charts? only ten best?).

Do the peaks make sense from the biological perspective?

Variance of SNPs vs genetic variance: → missing heritability? (cite Brachi 2011, Manolio 2009?).

BOSS probability: 1 big peak + smaller peaks. Compare against SNP density? Maybe the big peak corresponds to a physically isolated SNP, whereas smaller peaks correspond to a cluster of SNPs in LD which individually account for a smaller part of the variability, but together play an important predictive role.

4.2 Relative performance of rankers

why using P-values and not other standard rankers (e.g. backward stepwise selection)? Because of the specific nature of the problem

4.3 Genotyping strategies and applications to breeding

genotyping strategies: Costs, possible technologies (gbs, snp chip, microarrays), implications

applications to breeding: why is it important root vigor early detection. Other binomial traits (e.g. disease resistance) May be applied to bolting (another trait which exhibits binomial distribution), which has been shown to be controlled by multiple genes and influenced by environmental factors ([1]).

sugar beet: 30% of world's sugar production (cite Dohm? FAO?). Root vigor linked to yield.

Sugar beet: sugar + energy (citation?)

Other binomial traits: resistance to viral and fungal diseases, bolting (cite Dohm? Someone else?)

Breeding has shaped the genome of sugar beet (comparison with *Beta maritima*, [4]).

Extensions to multinomial traits? Examples?

5 Conclusions

Concluding remarks

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