Study of Oryza sativa genotypes involved in plant height using genome-wide association

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Variable Selection:

After dealing with all the missing values in our datasets we still have too many variables with only few observations, so we need to do some variable selection to have regression models with significant results.

Simple Regression:

Applying a simple linear regression to our data shows us that some estimated coefficients have missing values because of an exact linear relationship between the variables known as perfect multicollinearity. It's mainly because of the pseudo-inverse matrix computation. We decided to drop the variables with missing values for coefficients to avoid having highly correlated variables. The computation of the coefficient follows: $\beta_{OLS} = argmin_{\beta} \sum_{i=0}^{n} (y_i - \beta. x_i)^2$

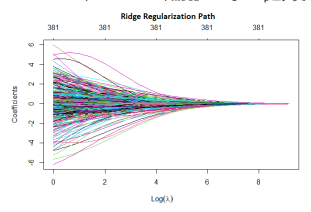
Forward and Stepwise Regression:

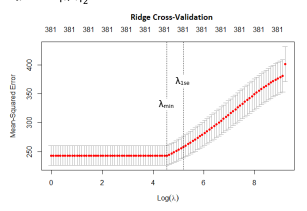
Both methods give us an R² equal to 1 which means that our model is overfitting since he perfectly fit the target data he trained on. So, we need to go further and use penalized regression methods to avoid overfitting and obtain a better model. Knowing that both models give us the same output, we are going to keep only the variables that the forward and stepwise regression has selected. On the other hand, the backward was not possible due to the number of variables.

Penalized Methods:

Ridge Regression:

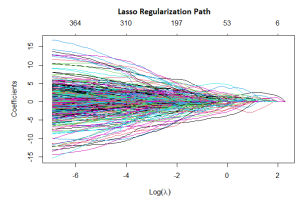
All coefficients decrease until being all equal to zero for the same value λ . Which means the resulting model always includes all the variables and that can be a problem for large datasets specifically in our case where the goal is to identify the genotypes involved in a selected phenotype. The coefficients are computed using the following formula for various values of λ with an L2-norm penalization: $\beta_{RIDGE} = argmin_{\beta} \sum_{i=0}^{n} (y_i - \beta, x_i)^2 + \lambda$. $||\beta||_2^2$

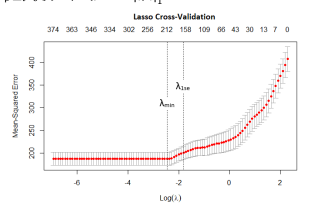




Lasso Regression:

Lasso is mainly useful in high dimensional datasets, where there are more variables than observations, but we only expect a small part of the variables to be truly meaningful. The coefficients become equal to zero one by one. The advantage of Lasso is that it does variable selection, for a given value λ some variable coefficients will be equal to zero. We penalize the function to minimize with an L1-norm such as: $\beta_{LASSO} = argmin_{\beta} \sum_{i=0}^{n} (y_i - \beta_i x_i)^2 + \lambda_i ||\beta_i||_1$

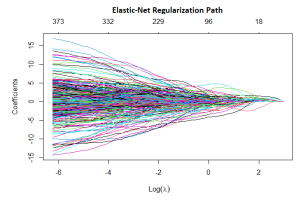


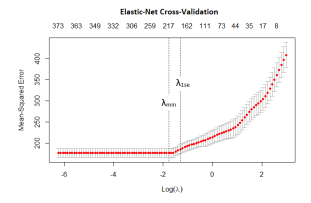


Sophisticated Methods:

Elastic-Net:

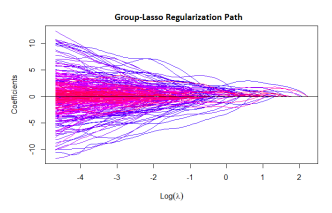
After going through various values for α and λ we found that the couple that minimizes the most the R² value is α = 1 (which corresponds to Lasso) with λ = λ_{min} . However, we have chosen α = 0.5 penalize the likelihood with an L1-term and an L2-term at the same time following the formula: $\beta_{ELASTIC-NET} = argmin_{\beta} \sum_{i=0}^{n} (y_i - \beta. x_i)^2 + \lambda \left(\alpha. \left||\beta|\right|_1 + \frac{1-\alpha}{2}. \left||\beta|\right|_2^2\right)$

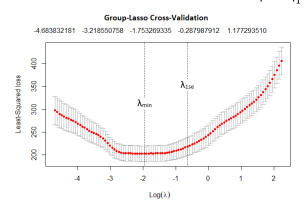




Group-Lasso:

Lasso gives us quite good results. However, there are situations in which the variables have a natural grouped structure. Group-Lasso is built as the sum of squares of coefficients belonging to the same group. This way it considers the possible grouped structure of variables, and it sends to zero whole groups of variables. Where G_k is the k-st group found within our variables and p_k is the number of elements in this group: $\beta_{GROUP-LASSO} = argmin_{\beta} \sum_{i=0}^{n} (y_i - \beta. \, x_i)^2 + \lambda \sum_{k=1}^{K} \sqrt{p_k} . \left| \left| \beta_{G_k} \right| \right|_1$





Support Vector Regression:

SVR finds an appropriate hyperplane to fit the data. We look for the hyperplane that fits the best to our observations by looking for the one with the highest margin. The margin is the distance that separates a hyperplane from the closest observation. In contrast to Ordinary Least Square, the objective function of SVR is to minimize the coefficients, more specifically, the L2-norm of the coefficient vector with respect to a constraint: $min_{\beta} \frac{1}{2} \left| |\beta| \right|_2^2$ with constraint $|y_i - \beta_i x_i| \le \varepsilon$

Summary:

To evaluate the performance of our several models, we use the coefficient of determination: $R^2 = 1 - \frac{\left||y - X\hat{\beta}|\right|_2^2}{\left||y - \bar{y}|\right|_2^2}$

	Ridge	Lasso	Elastic-Net	Group-Lasso	Support Vector Regression
\mathbb{R}^2	0.51	0.92	0.85	0.91	0.81

The R² value tells us how good our model is at making predictions of plant height given the information about genotypes. We clearly see that Ridge Regression is a bad choice for our specific case and that the highest R² is obtained using Lasso or Group-Lasso which is implied by the fact that only few genotypes are really involved in plant height.

List of genotypes mainly involved in plant height obtained using Lasso Regression with λ_{min} "id1000051" "id3011930" 'id1000852" 'id3008445" "ud7001348 "id1000994" id7004166" id3013258" id1002920" "id7004052" "ud1000154" "id1004393" "id3015622" "wd7002954" "wd3001877' "id3012431' id3008476" id3014217 id1000026" id7002916 id7003169 ud7001215 id7003112 d3011218 ud7001352 id1000399" id1002509" id1003248" id7003332" id7003361 id7003377 ud7001201 id3014412 id7002708 id7002717 id1000660 id7002946' id1001692' id3011075" id4000010 id7003201 "id7003704" ud7001582 id3015885 id3009920" id1003840' id7002799" id1000528 id1000529 id3008355' ud7001515 id1003839 id3012912 id3008721 id7002795 id7003442 id3009602 'id1001049 ud7001237 id7002775 id7003947 id3011059' id4000756" vd1000189 id1001664 id1001588 id7004032' id1002087 id3016604" d3010236' id3009276' id3013989' d7002923 id3011813 id3009243 wd3000757 id3009331 id1000007 id3008454 d3008833 id3008397 id3011640 id1001366' id3008682' id3012384 id1001981 id1001915 id3008500 id1001509 id3011423' d1000857 id3011035 ud7001419 id7003190' d4000164 ud7001180 id7003983 id1000980 ud3000950" d3011115 d1001516' i d7003709 id3010511 ud3001031 id3013397 d3012248 id1000661" ud7001238 id7002838 id3017406' wd7002936 dd3000953 id3017627 id3010370 id3017179 id3011960 id1000027 id1001599 ud1000027 ud1000187" id1000731" d1001266" id1001430" d1001224 d7003931' id1003919 id3009037 id3011044 d1005297 d1000030" id3008411 d3015746 id1001638 id1004041 id1002227 id3017002 id7002749 ud1000074 id7003060 wd7002914 id7003064 id3014618 d7003641 id7004040' id3009233 ud7001500 id7003174 id7003039 id7003877 id1004156' id3008674 id1003877 id3017136 id3016668 id7004170 id3009780 wd7002589' id1003465 id3017403 id1001003" id3009824 id1001015 id3008432 id7003272 id7003078 id3009379 id7002999 id3016447 id1001332 id1004872 "id1000841" "id1002622' "id3018096" 'id3009934'' id3008453" 'id1002730 id3012473' "id1000929" "id1000423" ud7001431 "id1000011

Our final goal was to create a model of the problem. Using the results obtained with Lasso Regression we can select the most important genotypes that are mainly involved in the *Oryza sativa* plant height. We finally obtained a list of 207 genotypes selected among our initial 36 901 genotypes using λ_{min} which is the value that gives us the best R². Still, if we want to have fewer genotypes, we can use the model with λ_{1se} which gives a simpler model with only 164 genotypes.