# Milestone 2: Core results 01

#### Anders

## 25/1/2023

#### Data

One clinical trails on breast cancer (advanced HR+/HER2-) using two different drug combination

The data set have mRNA expression of 771 genes at baseline (prior to treatment). This genes are specifically selected based on their potential roles in breast cancer pathology:

The gene set is dived into 25 sets of "signature genes"; which are thought to represent functional unities with respect to cancer biology. Often signaling pathways. Furthermore, 8 immune cells are represented with specific genes. These sets are substantially smaller then the signature genes; which I presume leads to some issue in modeling (as for clinical data too - see next sentence). In the domain knowledge part at the end 5 signature gene sets are use. Make sense to add more...

Additionally, the data-set contains clinical data; which up to now is not used in any models. If included they maybe should have a higher weight or be implemented differently from a sole gene. Maybe in a stacked ensemble model as signature.

#### Respones in study -ish

## Proliferation score

A score based on expression level of some of the genes. Range: -1.1366 to 0.8511

#### Risk of relapse score (ROR)

A combined score based on expression level of genes and some clinical findings. Range: -8.035678 to 75.13174 (only used in combination with proliferation score as described bellow)

#### Risk of relapse score with proliferaton score (ROR\_Prolif)

A combined score of the two above. Range: 1 to 97 (1-100)

The two scores involving ROR also have categorical variants containing: low, medium, high (but not used...)

#### Trail

Two treatments which differ with respect to drug combination - Target: ribociclib and endocrine therapy (letrozole) - Chemotherapy: doxorubicin, cyclophosphamide and paclitaxel. approx. 50 patients in each group. Endpoints: proliferation score, ROR score, combined ROR and prolif

# Major goal

- 1. Find best model to predict outcome of cancer treatment with genetic profile as predictive features
- 2. Features selection in order to understand cancer biology

# Major challanges

Preliminary experiments (on trail 1) showed instability in prediction and feature selection between bootstrap samples of Lasso. I believe this is a classical problem of high-dim data?

# Approch

Test all thinkable models in a search for superior models

### Evaluation of models

Two levels of evaluation is considered:

#### 1. Relative comparison of the different models

1000 bootstrap models are fitted and then evaluated on the original sample. This gives a relative comparison of the various models with respect to data very similar to the given data set. Correlation, MSE and frequency of selected features is compared.

### 2. Expected outcome of future patients

3 strategies are considered:

- 1. Repeated cross-validations (200 rep, 5-fold)
- 2. Bootstrap models with 0.632 (or 0.632?) adjustment (Not done)
- 3. Use the cohort as test data-set (Challenge: This trail have different responses)

# RESULTS

(- mboost)

- xgboost

Features tested:
6 genes
771 genes
node values of mech model
residuals of mech model
Responses tested:
proliferation score
combined score including proliferation and ROR
Models tested:
Lasso
Ridge
Elastic Net
Boosting with stumps as base learner

2

#### PCA on subsets of genes

Stacking using different features in the base models

Sparse group lasso (not done)

Iterative learning (ongoing)

Post Lasso (not done)

# Results of individual modles:

# Lasso - Bootstrap

### 6 genes -> proliferation score

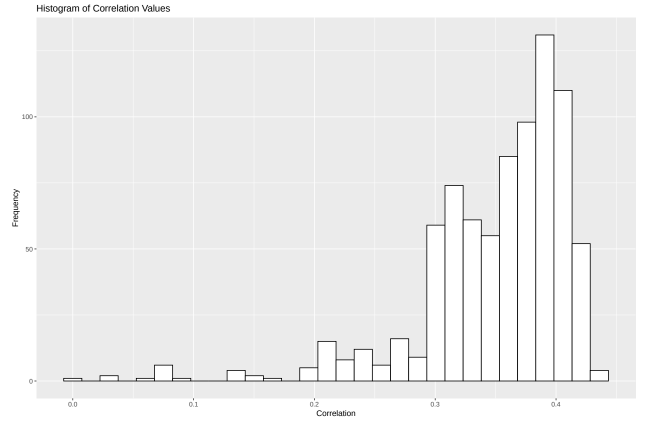
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0.182

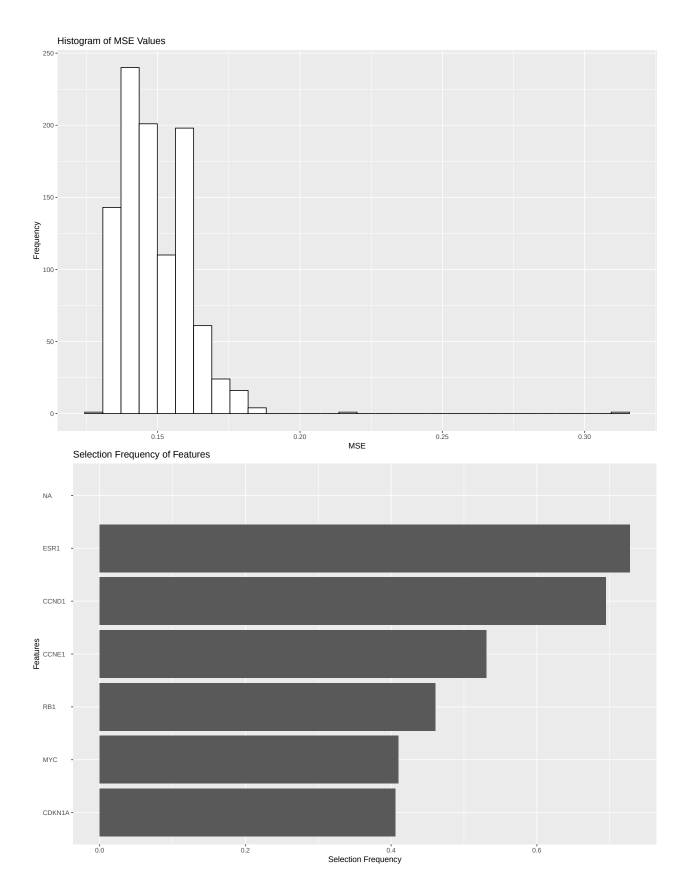
##

## CORRELATIONS RESULTS

## Mean: 0.3498379 ## Median: 0.3672243 ## Variance: 0.003984261 ## st.dev.: 0.063121



## MSE RESULTS
## Mean: 0.1492302
## Median: 0.1469228
## Variance: 0.0001550247
## st.dev.: 0.01245089



##

```
## Features selected 50% or more times:
```

## CCND1 CCNE1 ESR1

## Top 20 featrues:

## [1] "ESR1" "CCND1" "CCNE1" "RB1" "MYC" "CDKN1A" NA NA ## [9] NA NA NA NA NA NA NA NA

## [17] NA NA NA NA

# 6 genes -> ROR\_proliferation score

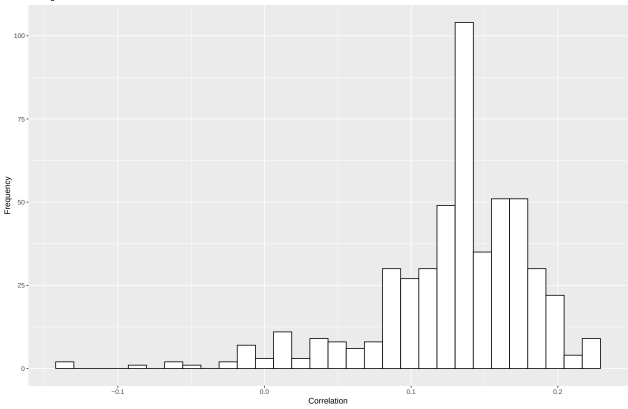
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0.495

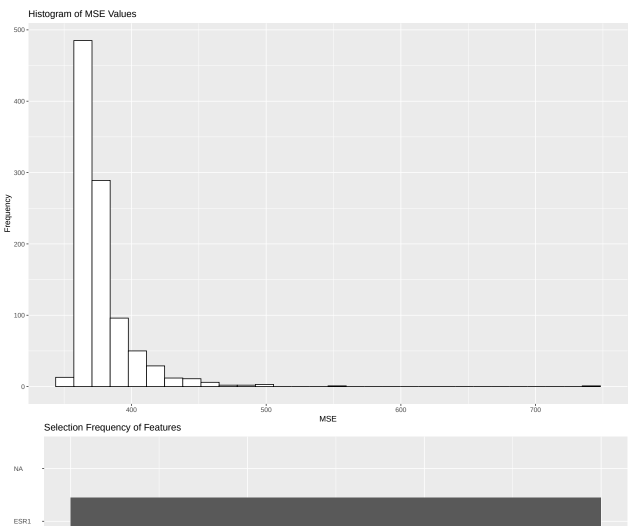
##

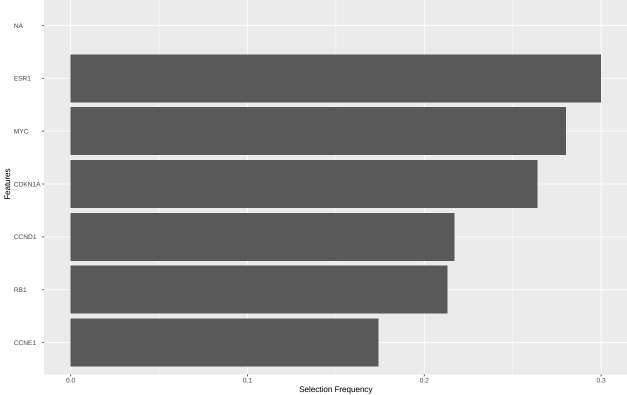
## CORRELATIONS RESULTS
## Mean: 0.1282306
## Median: 0.1311715

## Median: 0.1311715 ## Variance: 0.00285293 ## st.dev.: 0.05341282 Histogram of Correlation Values



## MSE RESULTS
## Mean: 378.094
## Median: 370.7201
## Variance: 549.4448
## st.dev.: 23.44024





```
## Features selected 50% or more times:
##
## Top 20 featrues:
                                                     "CCNE1"
## [1] "ESR1"
                 "MYC"
                          "CDKN1A" "CCND1"
                                            "RB1"
                                                                       NA
                                                              NA
## [9] NA
                 NA
                          NA
                                   NA
                                            NA
                                                     NA
                                                              NA
                                                                       NA
## [17] NA
                 NA
                          NA
                                   NA
```

# 771 genes -> proliferation score

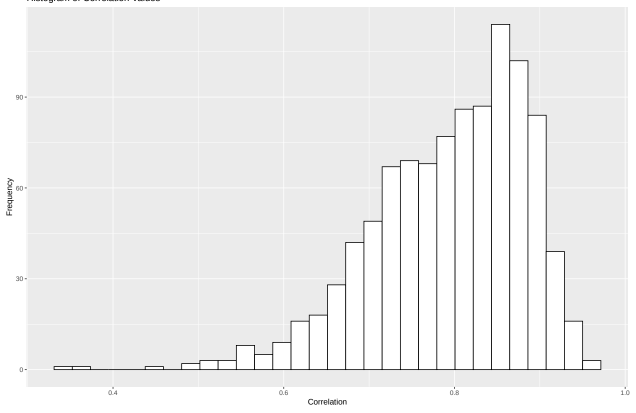
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0.002

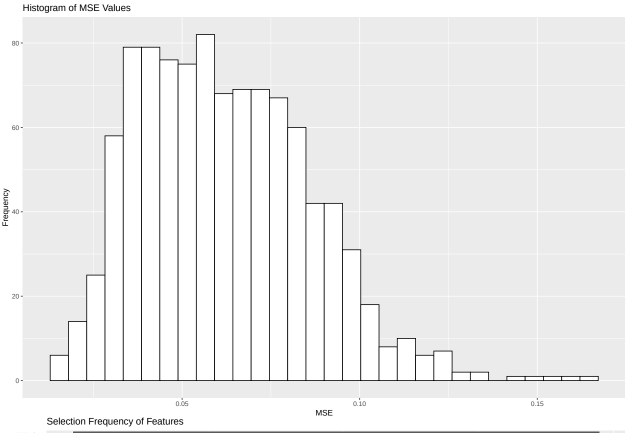
##

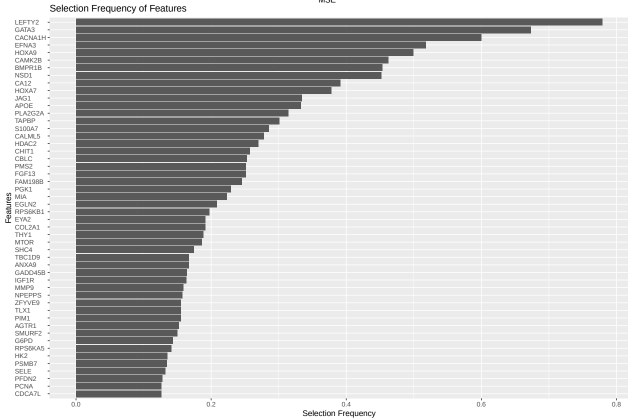
## CORRELATIONS RESULTS
## Mean: 0.7941413
## Median: 0.8101886
## Variance: 0.008119272
## st.dev.: 0.090107

Histogram of Correlation Values



## MSE RESULTS
## Mean: 0.06209131
## Median: 0.0598495
## Variance: 0.0005751012
## st.dev.: 0.02398127





```
## Features selected 50% or more times:
```

## CACNA1H EFNA3 GATA3 LEFTY2

## Top 20 featrues:

## [1] "LEFTY2" "GATA3" "CACNA1H" "EFNA3" "HOXA9" "CAMK2B" "BMPR1B" ## [8] "NSD1" "CA12" "HOXA7" "JAG1" "APOE" "PLA2G2A" "TAPBP"

## [15] "S100A7" "CALML5" "HDAC2" "CHIT1" "CBLC" "FGF13"

### 771 genes -> ROR-proliferation score

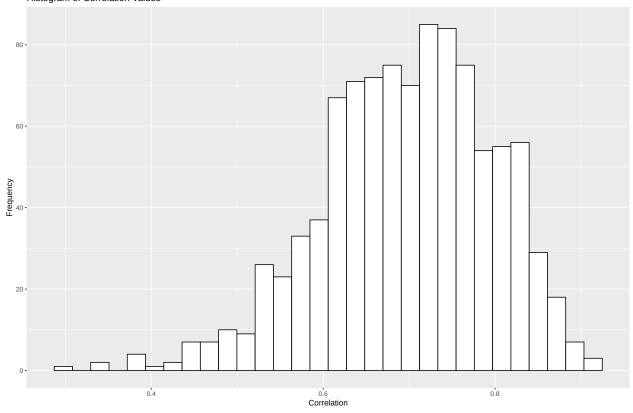
## number of models fitted: 1000

 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0.017

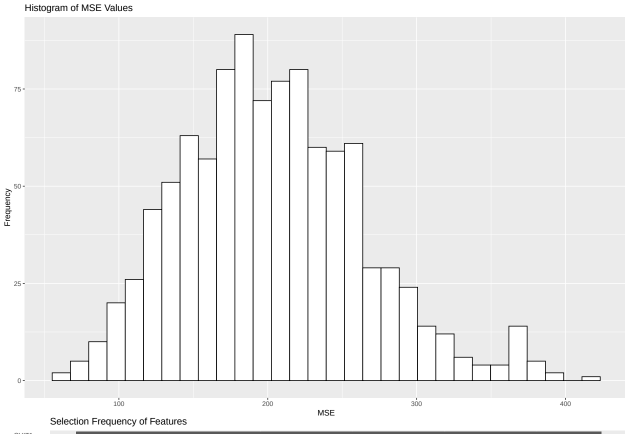
##

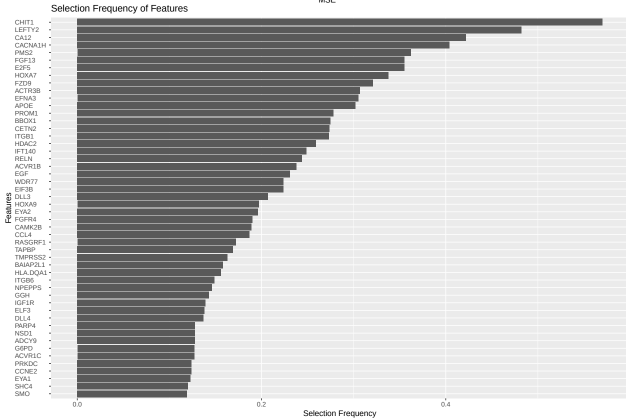
## CORRELATIONS RESULTS
## Mean: 0.6968101
## Median: 0.7035889
## Variance: 0.009901439
## st.dev.: 0.09950598

Histogram of Correlation Values



## MSE RESULTS
## Mean: 203.408
## Median: 198.455
## Variance: 3763.666
## st.dev.: 61.34872



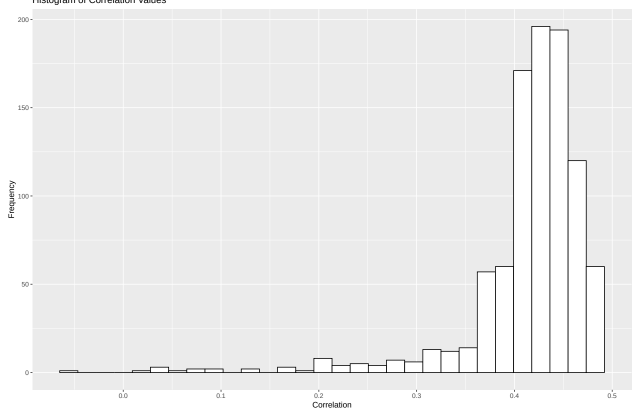


```
## Features selected 50% or more times:
## CHIT1
## Top 20 featrues:
## [1] "CHIT1"
                  "LEFTY2"
                            "CA12"
                                       "CACNA1H" "PMS2"
                                                           "E2F5"
                                                                     "FGF13"
## [8] "HOXA7"
                  "FZD9"
                            "ACTR3B"
                                      "EFNA3"
                                                 "APOE"
                                                           "PROM1"
                                                                     "BBOX1"
## [15] "CETN2"
                  "ITGB1"
                            "HDAC2"
                                      "IFT140" "RELN"
                                                           "ACVR1B"
```

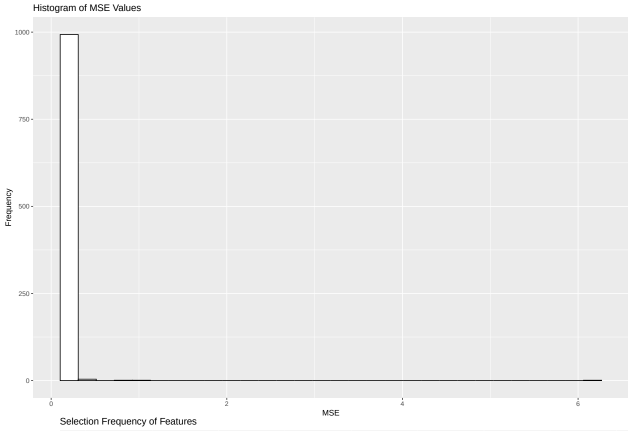
### node values -> proliferation score

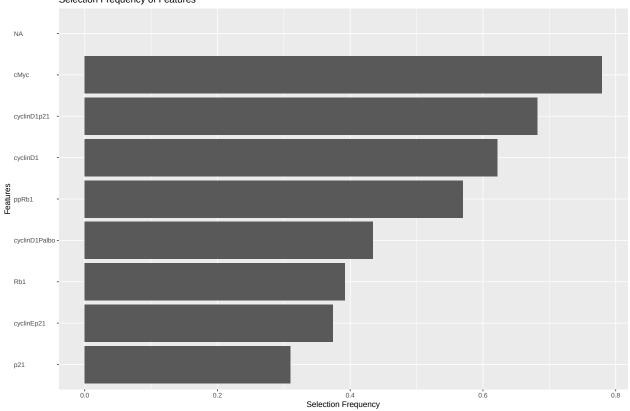
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0.053
##
## CORRELATIONS RESULTS
## Mean: 0.414496

## Mean: 0.414496 ## Median: 0.4275114 ## Variance: 0.00413092 ## st.dev.: 0.06427223 Histogram of Correlation Values



## MSE RESULTS ## Mean: 0.1479731 ## Median: 0.1355805 ## Variance: 0.03673469 ## st.dev.: 0.191663





```
## Features selected 50% or more times:
## cyclinD1 cyclinD1p21 cMyc ppRb1
## Top 20 featrues:
  [1] "cMyc"
                         "cyclinD1p21"
                                         "cyclinD1"
                                                          "ppRb1"
   [5] "cyclinD1Palbo" "Rb1"
                                         "cyclinEp21"
                                                          "p21"
##
   [9] NA
                                                         NA
## [13] NA
                        NA
                                         NA
                                                         NA
## [17] NA
                        NA
                                         NA
                                                         NA
```

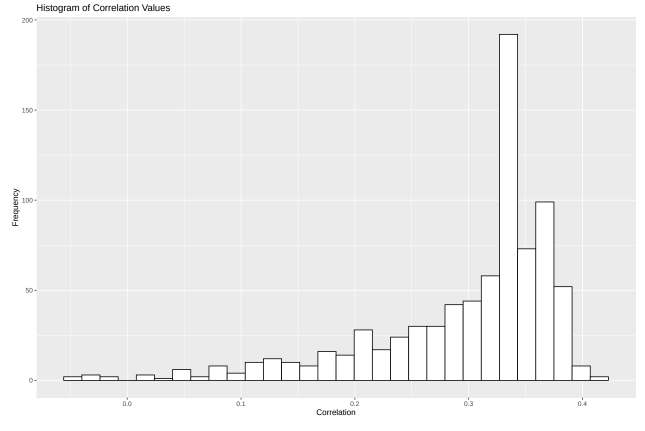
### node values -> ROR-proliferation score

## number of models fitted: 1000

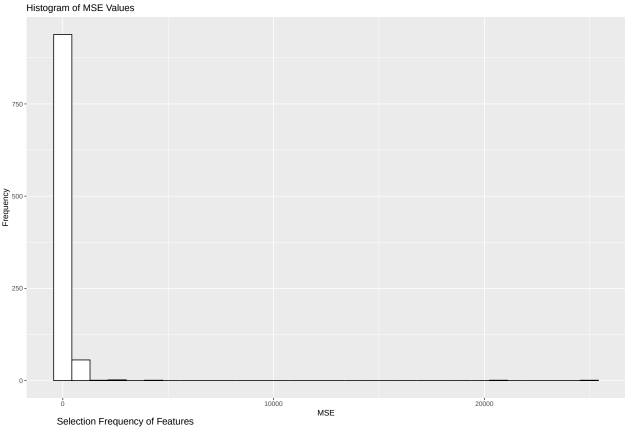
## Fraction of model fits with no selected genes: 0.2

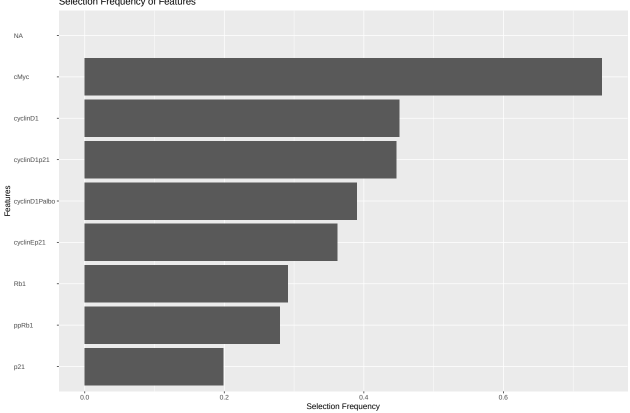
##

## CORRELATIONS RESULTS
## Mean: 0.2964169
## Median: 0.3317433
## Variance: 0.006900552
## st.dev.: 0.08306956



## MSE RESULTS
## Mean: 417.6667
## Median: 353.8176
## Variance: 1054857
## st.dev.: 1027.062





```
## Features selected 50% or more times:
## cMyc
## Top 20 featrues:
   [1] "cMyc"
                         "cyclinD1"
                                          "cyclinD1p21"
                                                          "cyclinD1Palbo"
                                                          "p21"
    [5] "cyclinEp21"
                         "Rb1"
                                          "ppRb1"
##
##
   [9] NA
                         NA
                                         NA
                                                          NA
## [13] NA
                         NA
                                         NA
                                                          NA
## [17] NA
                         NA
                                         NA
                                                          NA
```

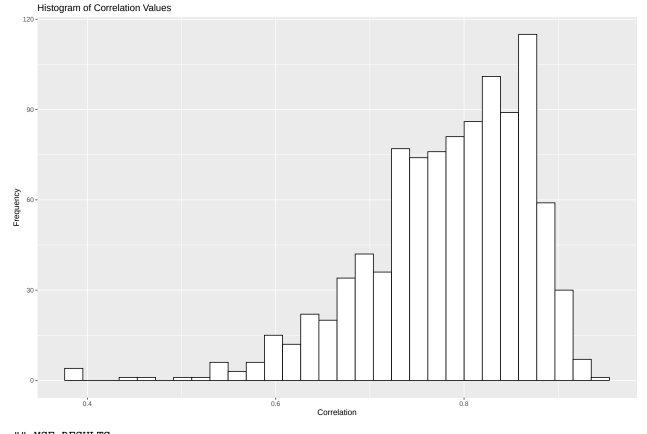
### Mechanistic + Residuals -> proliferation score (additive)

## number of models fitted: 1000

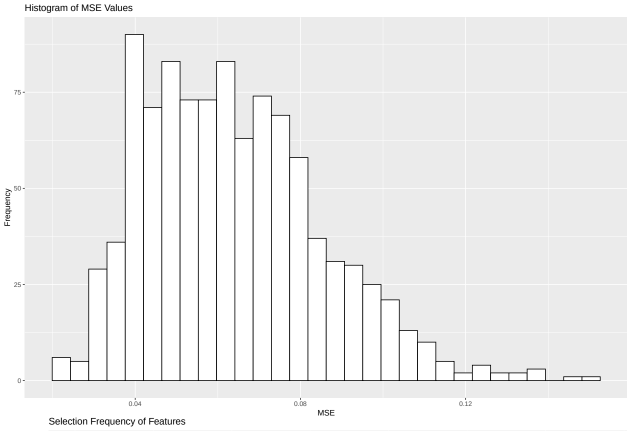
## Fraction of model fits with no selected genes: 0

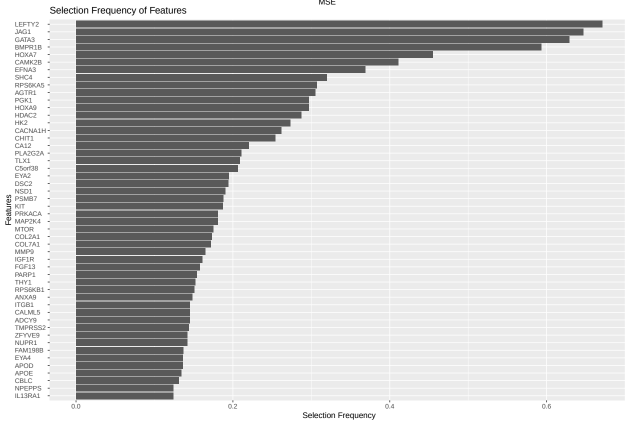
##

## CORRELATIONS RESULTS
## Mean: 0.7835808
## Median: 0.7962129
## Variance: 0.007298891
## st.dev.: 0.08543355



## MSE RESULTS
## Mean: 0.06384841
## Median: 0.06161072
## Variance: 0.0004577716
## st.dev.: 0.0213956

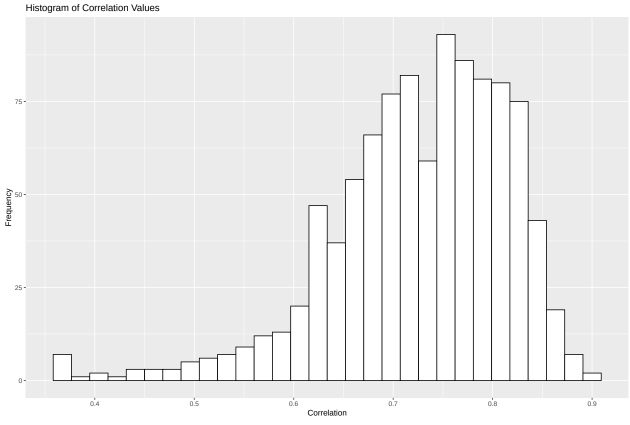




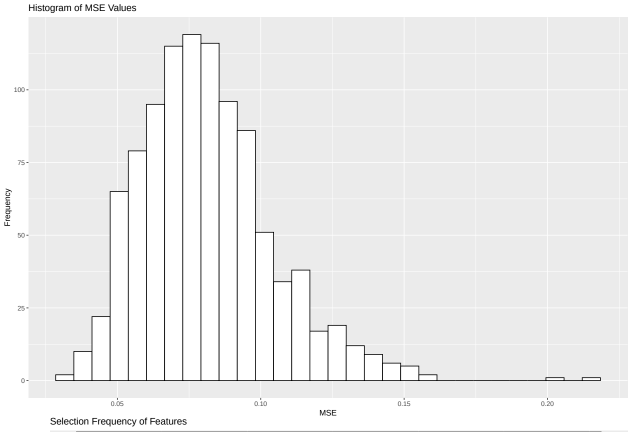
```
## Features selected 50% or more times:
## BMPR1B GATA3 JAG1 LEFTY2
## Top 20 featrues:
  [1] "LEFTY2" "JAG1"
                            "GATA3"
                                      "BMPR1B"
                                                 "HOXA7"
                                                           "CAMK2B"
                                                                     "EFNA3"
   [8] "SHC4"
                  "RPS6KA5" "AGTR1"
                                      "HOXA9"
                                                 "PGK1"
                                                           "HDAC2"
                                                                     "HK2"
## [15] "CACNA1H" "CHIT1"
                            "CA12"
                                      "PLA2G2A" "TLX1"
                                                           "C5orf38"
```

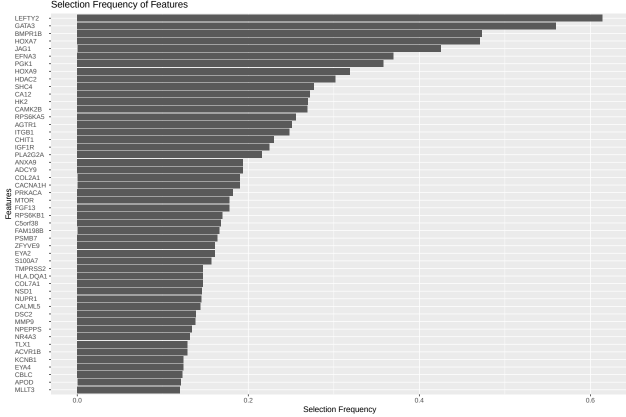
# $Mechanistic + Residuals -> proliferation\ score\ (multiplicative)$

```
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS
## Mean: 0.7266736
## Median: 0.739984
## Variance: 0.008014481
## st.dev.: 0.08952363
```



## MSE RESULTS
## Mean: 0.0813415
## Median: 0.07892236
## Variance: 0.0005355891
## st.dev.: 0.0231428

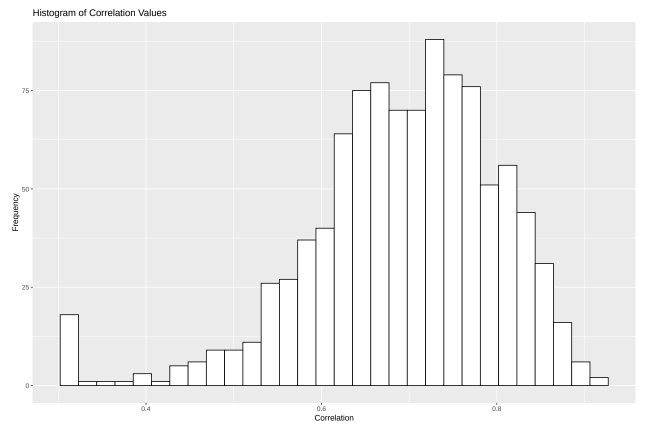




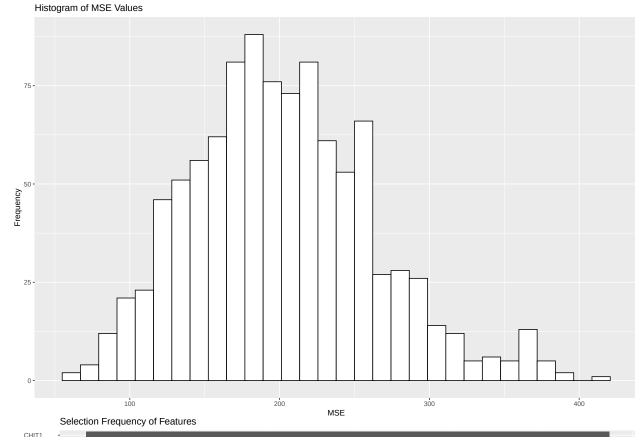
```
## Features selected 50% or more times:
## GATA3 LEFTY2
## Top 20 featrues:
  [1] "LEFTY2"
                  "GATA3"
                            "BMPR1B"
                                       "HOXA7"
                                                 "JAG1"
                                                                      "PGK1"
                                                           "EFNA3"
   [8] "HOXA9"
                             "SHC4"
                                                 "HK2"
                  "HDAC2"
                                       "CA12"
                                                           "CAMK2B"
                                                                      "RPS6KA5"
## [15] "AGTR1"
                  "ITGB1"
                            "CHIT1"
                                       "IGF1R"
                                                 "PLA2G2A" "ADCY9"
```

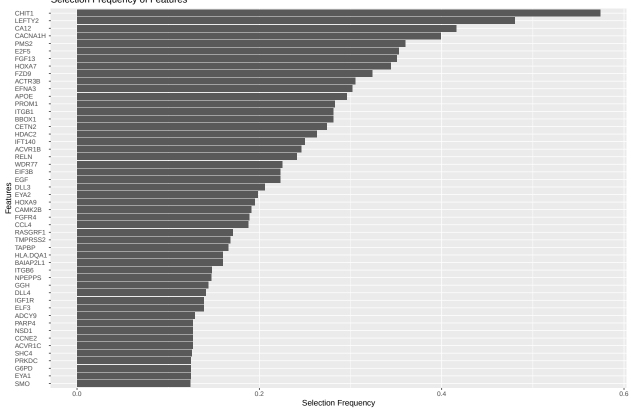
# Mechnaistic + Residuals -> ROR-proliferation score (additive)

```
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS
## Mean: 0.6925953
## Median: 0.7037142
## Variance: 0.01193153
## st.dev.: 0.1092315
```



## MSE RESULTS
## Mean: 202.3235
## Median: 197.4845
## Variance: 3726.169
## st.dev.: 61.04236

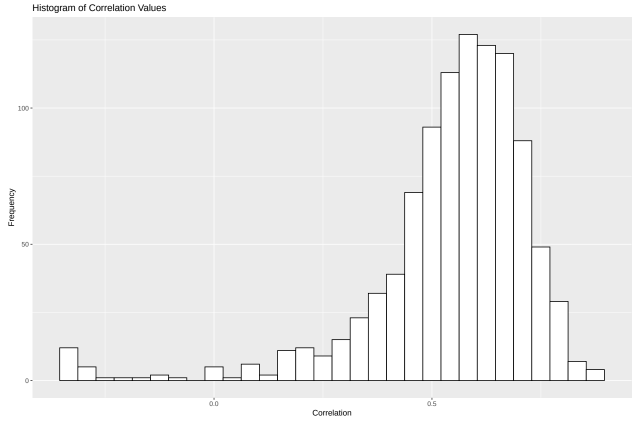




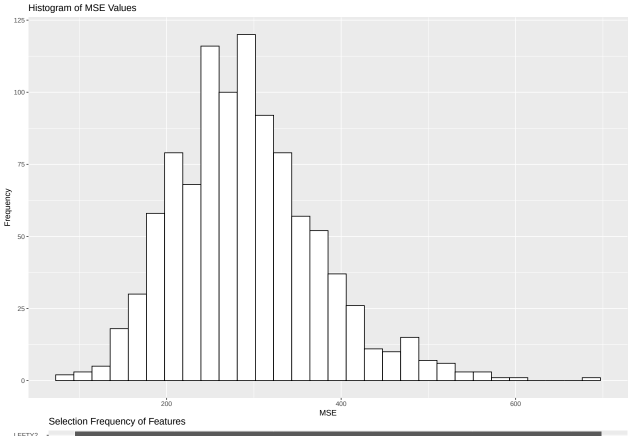
```
## Features selected 50% or more times:
## CHIT1
## Top 20 featrues:
  [1] "CHIT1"
                  "LEFTY2"
                            "CA12"
                                       "CACNA1H" "PMS2"
                                                           "E2F5"
                                                                     "FGF13"
## [8] "HOXA7"
                  "FZD9"
                                                                     "BBOX1"
                            "ACTR3B"
                                      "EFNA3"
                                                 "APOE"
                                                           "PROM1"
## [15] "ITGB1"
                  "CETN2"
                            "HDAC2"
                                       "IFT140"
                                                "ACVR1B"
                                                           "RELN"
```

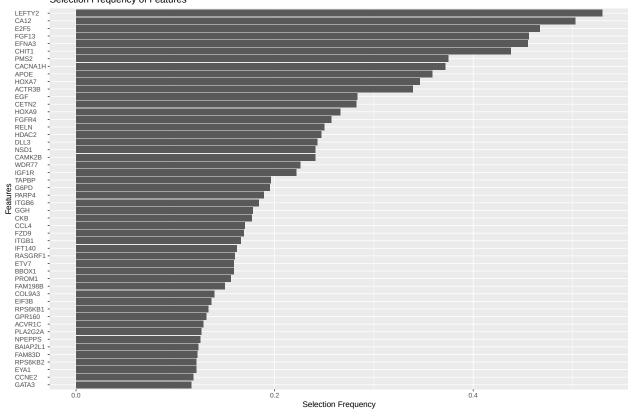
# Mechnaistic + Residuals -> ROR-proliferation score (multiplicative)

```
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS
## Mean: 0.5427757
## Median: 0.5790305
## Variance: 0.03640698
## st.dev.: 0.1908061
```



## MSE RESULTS
## Mean: 291.0186
## Median: 284.2399
## Variance: 6854.567
## st.dev.: 82.79231





```
## Features selected 50% or more times:
## CA12 LEFTY2
## Top 20 featrues:
   [1] "LEFTY2" "CA12"
                             "E2F5"
                                       "FGF13"
                                                 "EFNA3"
                                                            "CHIT1"
                                                                      "PMS2"
   [8] "CACNA1H" "APOE"
                                                 "EGF"
                                                                      "HOXA9"
                             "HOXA7"
                                       "ACTR3B"
                                                            "CETN2"
                  "RELN"
## [15] "FGFR4"
                             "HDAC2"
                                       "DLL3"
                                                 "CAMK2B"
                                                            "NSD1"
```

# Summery results: lasso proliferation score (bootstrap)

Model	cor_mean	$sd\_cor$	MSE_mean	$MSE\_sd$
lasso 6 genes	0.3498379	0.0631210	0.1492302	0.0124509
lasso 771 genes	0.7941413	0.0901070	0.0620913	0.0239813
Nodes	0.4144960	0.0642722	0.1479731	0.1916630
Residual additive	0.7835808	0.0854335	0.0638484	0.0213956
Residual multiplicative	0.7266736	0.0895236	0.0813415	0.0231428

## Summery results: lasso ROR+proliferation score (bootstrap)

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
lasso 6 genes	0.1282306	0.0534128	378.0940	23.44024
lasso 771 genes	0.6968101	0.0995060	203.4080	61.34872
Nodes	0.2964169	0.0830696	417.6667	1027.06231
Residual additive	0.6925953	0.1092315	202.3235	61.04236
Residual multiplicative	0.5427757	0.1908061	291.0186	82.79231

# Lasso - Repeated cross-validation

200 repeats of five fold cross-validation

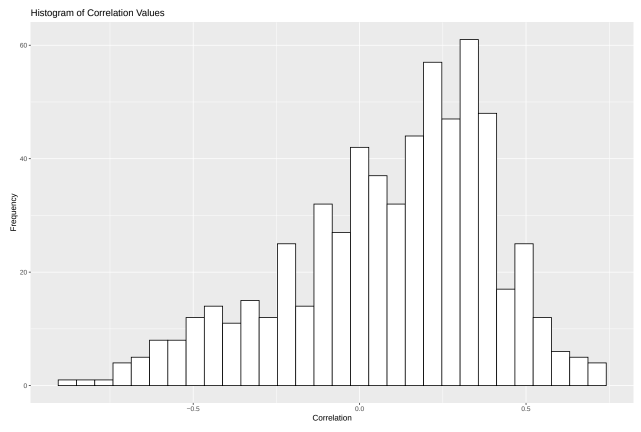
## 6 genes -> proliferation score

```
## number of models fitted: 1000
```

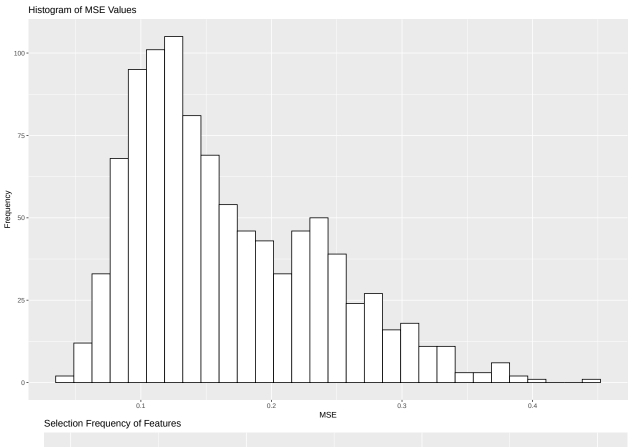
## Fraction of model fits with no selected genes: 0.373

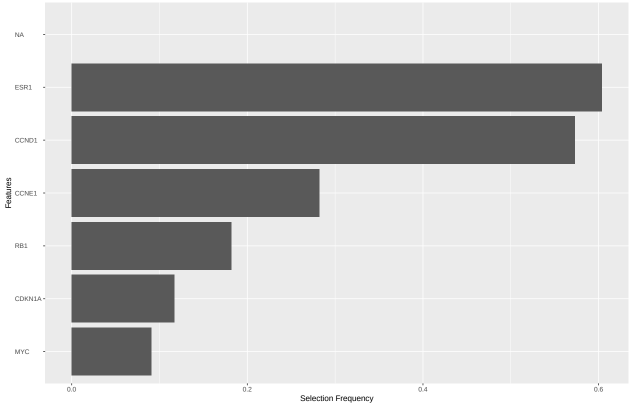
##

## CORRELATIONS RESULTS
## Mean: 0.09196628
## Median: 0.1502755
## Variance: 0.09413904
## st.dev.: 0.3068209



## MSE RESULTS
## Mean: 0.1655931
## Median: 0.1468293
## Variance: 0.005160684
## st.dev.: 0.0718379





```
## Features selected 50% or more times:
## CCND1 ESR1
## Top 20 featrues:
## [1] "ESR1"
                 "CCND1" "CCNE1"
                                   "RB1"
                                            "CDKN1A" "MYC"
                                                                        NA
                                                              NA
## [9] NA
                 NA
                          NA
                                   NA
                                                     NA
                                                              NA
                                                                        NA
```

NA

# 6 genes -> ROR\_proliferation score

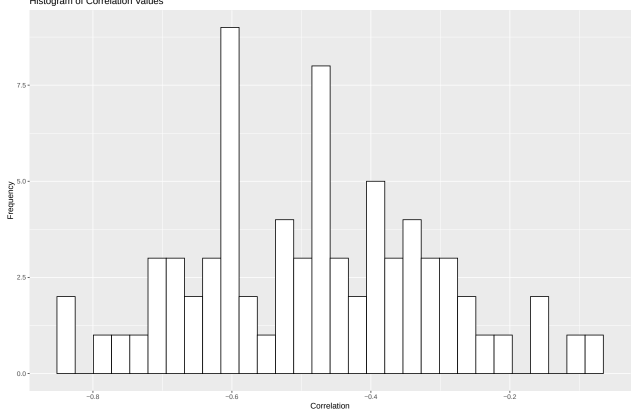
NA

## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0.926
##
## CORRELATIONS RESULTS
## Mean: -0.4822298
## Median: -0.4810641

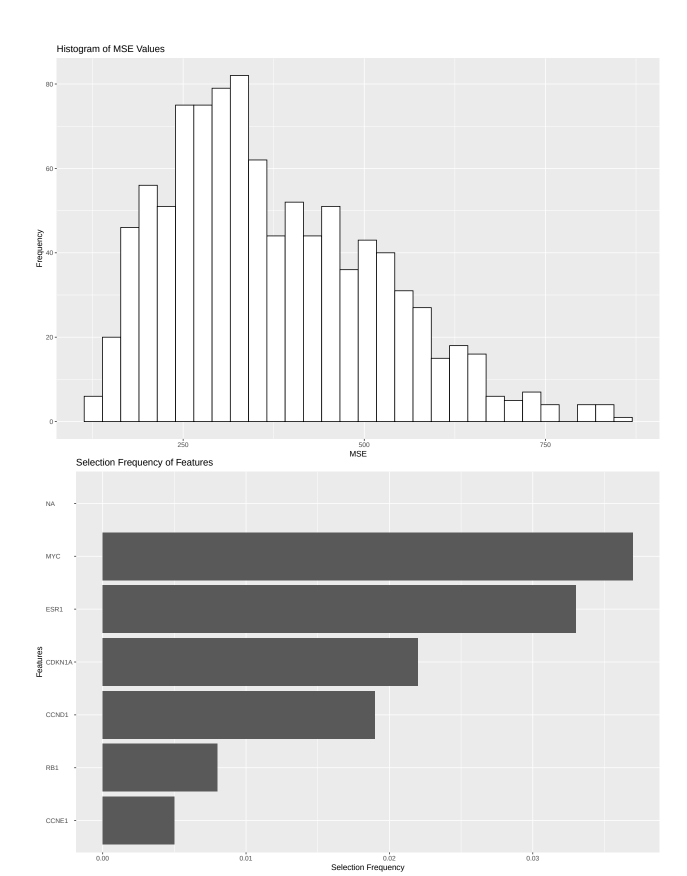
NA

## Variance: 0.02975572
## st.dev.: 0.1724985
Histogram of Correlation Values

## [17] NA



## MSE RESULTS
## Mean: 374.1519
## Median: 343.2105
## Variance: 20780.92
## st.dev.: 144.1559

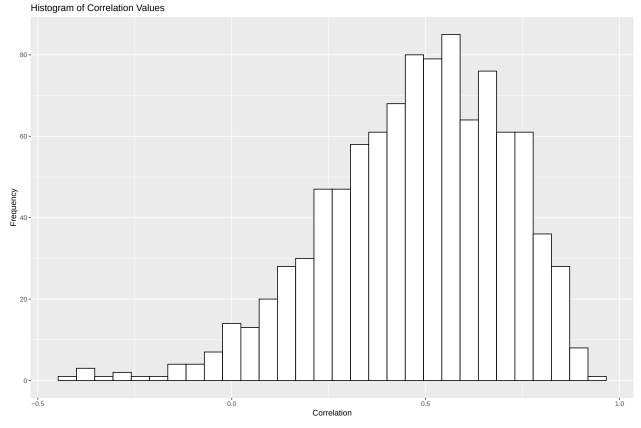


```
## Features selected 50% or more times:
##
## Top 20 featrues:
  [1] "MYC"
                 "ESR1"
                          "CDKN1A" "CCND1"
                                            "RB1"
                                                      "CCNE1"
                                                                        NA
                                                               NA
## [9] NA
                 NA
                          NA
                                   NA
                                            NA
                                                      NA
                                                               NA
                                                                        NA
## [17] NA
                 NA
                          NA
                                   NA
```

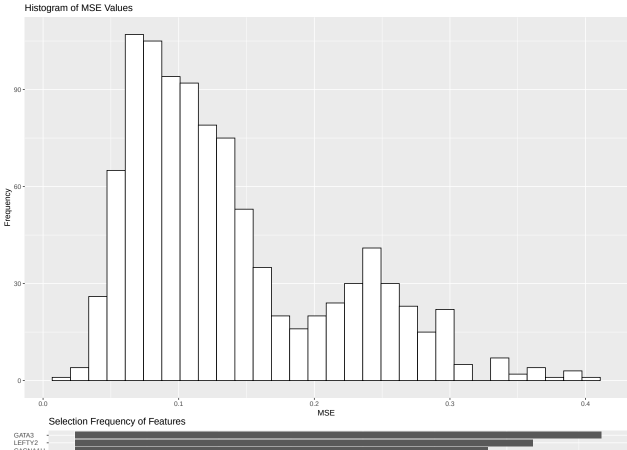
# 771 genes -> proliferation score

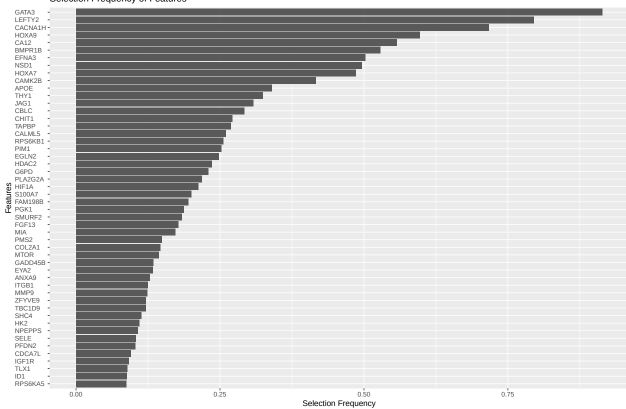
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0.011
##
## CORRELATIONS RESULTS

## Mean: 0.4737037 ## Median: 0.4959203 ## Variance: 0.05337068 ## st.dev.: 0.2310209



## MSE RESULTS
## Mean: 0.1376002
## Median: 0.1154157
## Variance: 0.005670929
## st.dev.: 0.07530557

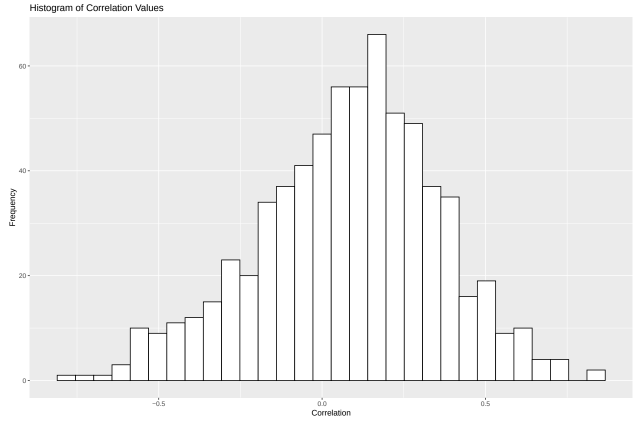




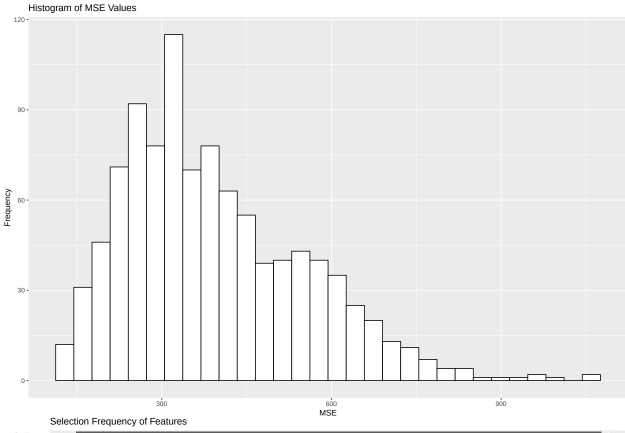
```
## Features selected 50% or more times:
## BMPR1B CA12 CACNA1H EFNA3 GATA3 HOXA9 LEFTY2
## Top 20 featrues:
## [1] "GATA3"
                  "LEFTY2"
                            "CACNA1H" "HOXA9"
                                                 "CA12"
                                                           "BMPR1B"
                                                                     "EFNA3"
## [8] "NSD1"
                            "CAMK2B" "APOE"
                                                 "THY1"
                  "HOXA7"
                                                           "JAG1"
                                                                     "CBLC"
## [15] "CHIT1"
                  "TAPBP"
                            "CALML5" "RPS6KB1" "PIM1"
                                                           "EGLN2"
```

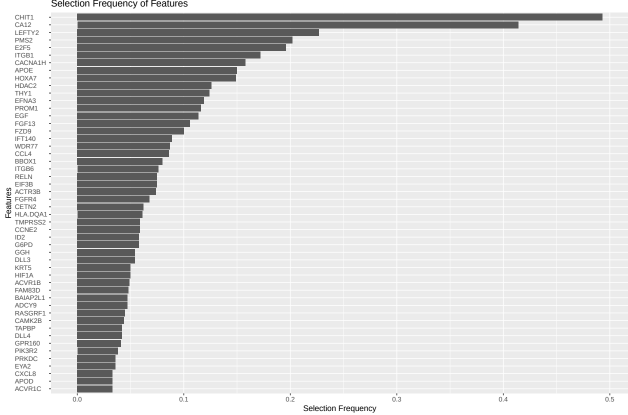
# 771 genes -> ROR-proliferation score

## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0.321
##
## CORRELATIONS RESULTS
## Mean: 0.08062366
## Median: 0.1014264
## Variance: 0.07657135
## st.dev.: 0.2767153



## MSE RESULTS
## Mean: 393.8069
## Median: 360.5105
## Variance: 25486.55
## st.dev.: 159.6451



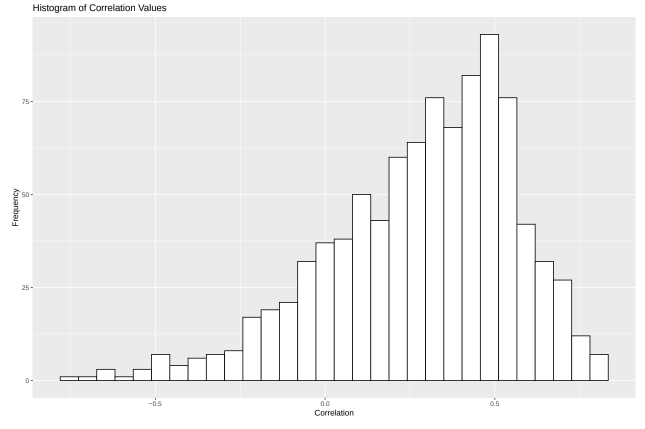


```
## Features selected 50% or more times:
##
## Top 20 featrues:
                  "CA12"
  [1] "CHIT1"
                             "LEFTY2"
                                       "PMS2"
                                                 "E2F5"
                                                            "ITGB1"
                                                                      "CACNA1H"
## [8] "APOE"
                             "HDAC2"
                                                                      "EGF"
                  "HOXA7"
                                       "THY1"
                                                 "EFNA3"
                                                            "PROM1"
## [15] "FGF13"
                  "FZD9"
                             "IFT140" "WDR77"
                                                 "CCL4"
                                                            "BBOX1"
```

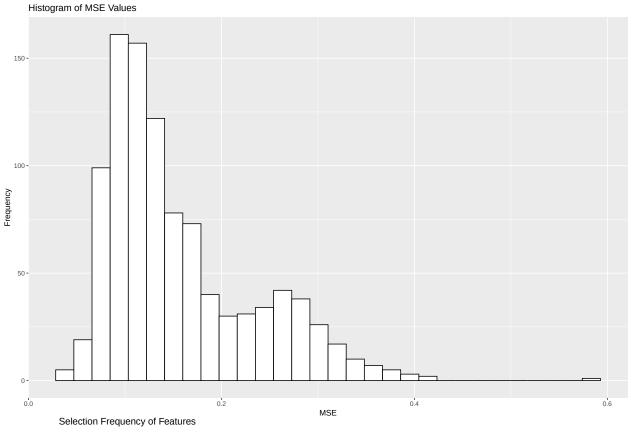
# ${\bf node\ values\ \text{--}>\ proliferation\ score}$

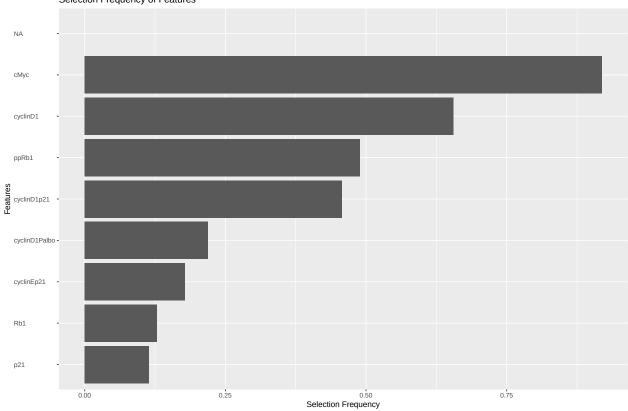
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0.063
##
## CORRELATIONS RESULTS
## Mean: 0.2842257
## Median: 0.3249779

## Variance: 0.07664357 ## st.dev.: 0.2768458



## MSE RESULTS
## Mean: 0.1560308
## Median: 0.1314678
## Variance: 0.005819908
## st.dev.: 0.07628832





```
## Features selected 50% or more times:
## cyclinD1 cMyc
## Top 20 featrues:
   [1] "cMyc"
                         "cyclinD1"
                                         "ppRb1"
                                                          "cyclinD1p21"
                                                          "p21"
    [5] "cyclinD1Palbo" "cyclinEp21"
                                         "Rb1"
##
   [9] NA
                        NA
                                         NA
                                                          NA
## [13] NA
                        NA
                                         NA
                                                          NA
## [17] NA
                        NA
                                                         NA
                                         NA
```

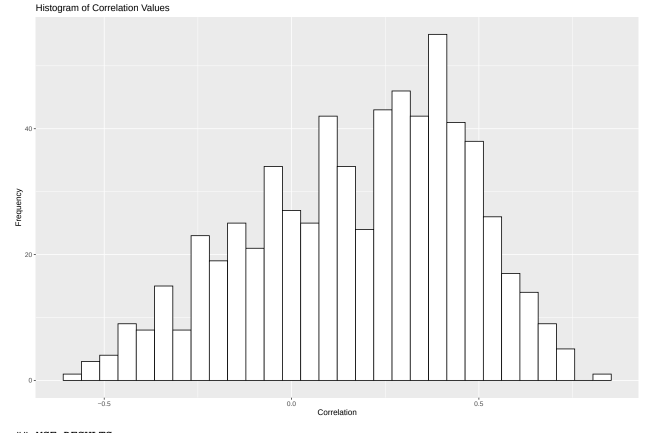
### node values $\rightarrow$ ROR-proliferation score

## number of models fitted: 1000

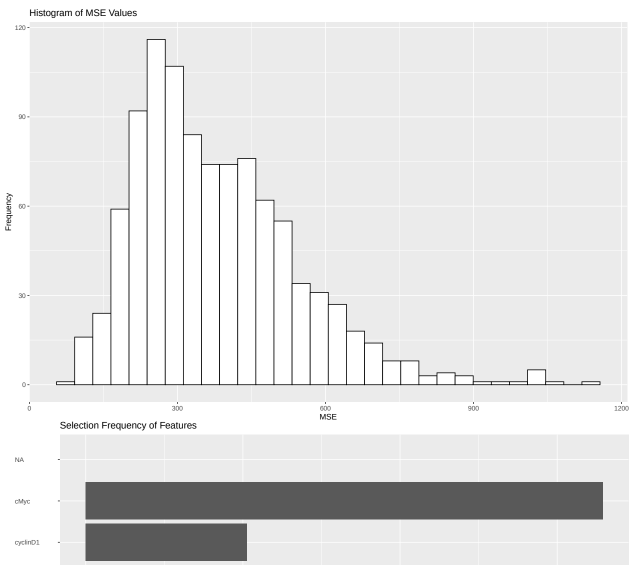
## Fraction of model fits with no selected genes: 0.341

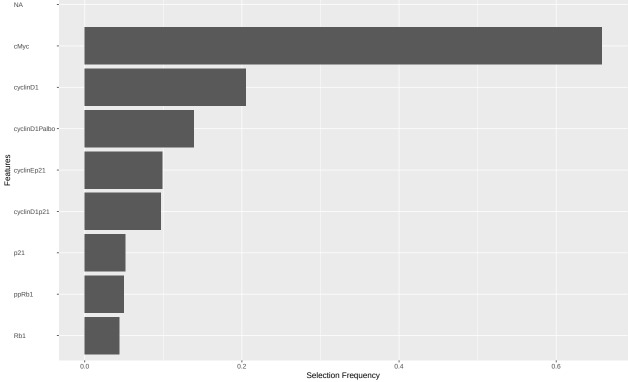
##

## CORRELATIONS RESULTS ## Mean: 0.1806504 ## Median: 0.2237481 ## Variance: 0.08150408 ## st.dev.: 0.2854892



## MSE RESULTS
## Mean: 380.1157
## Median: 349.3312
## Variance: 27088.84
## st.dev.: 164.5869



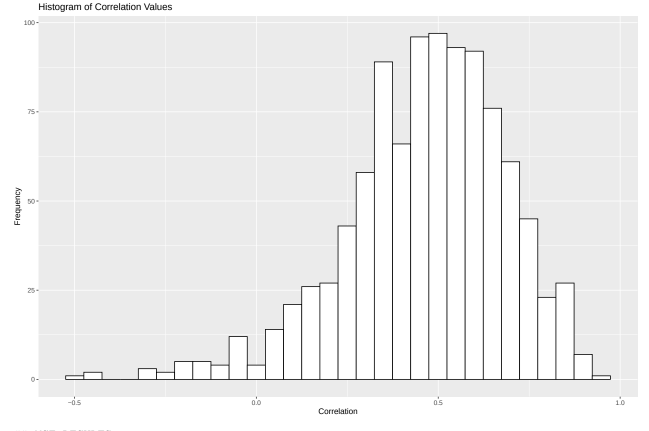


```
## Features selected 50% or more times:
## cMyc
## Top 20 featrues:
   [1] "cMyc"
                         "cyclinD1"
                                         "cyclinD1Palbo" "cyclinEp21"
    [5] "cyclinD1p21"
                         "p21"
                                         "ppRb1"
                                                          "Rb1"
##
##
   [9] NA
                         NA
                                         NA
                                                          NA
## [13] NA
                        NA
                                         NA
                                                          NA
## [17] NA
                                         NA
                        NA
                                                          NA
```

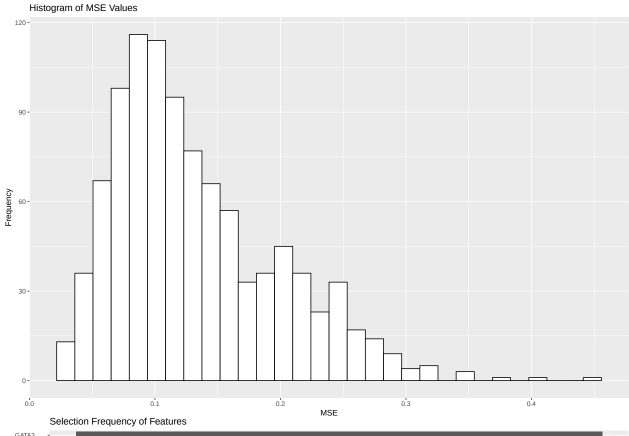
### Mechanistic + Residuals -> proliferation score (additive)

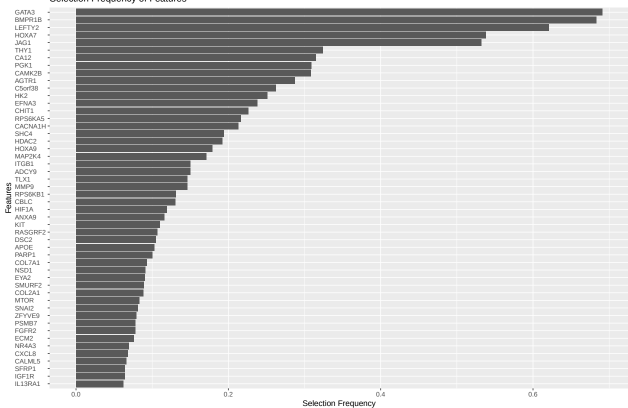
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##

## CORRELATIONS RESULTS
## Mean: 0.4633095
## Median: 0.4870052
## Variance: 0.04959996
## st.dev.: 0.2227105



## MSE RESULTS
## Mean: 0.1331785
## Median: 0.1164927
## Variance: 0.004278014
## st.dev.: 0.06540653



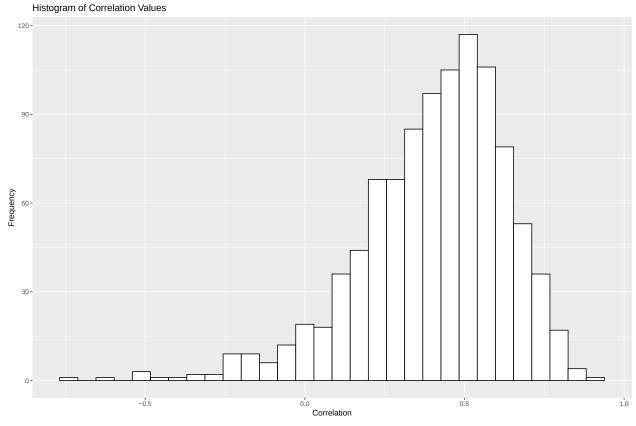


##

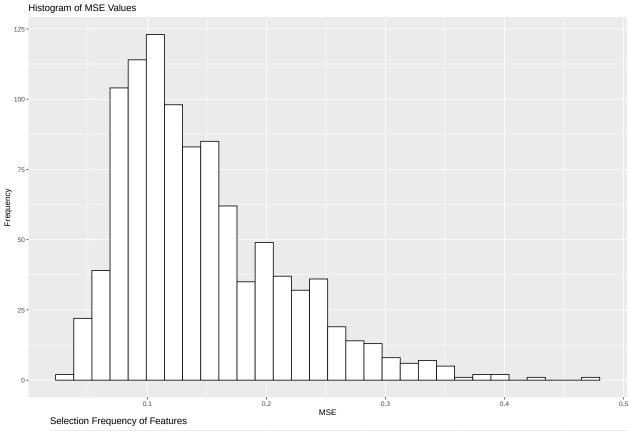
```
## Features selected 50% or more times:
## BMPR1B GATA3 HOXA7 JAG1 LEFTY2
## Top 20 featrues:
## [1] "GATA3"
                  "BMPR1B"
                            "LEFTY2"
                                      "HOXA7"
                                                 "JAG1"
                                                                     "CA12"
                                                           "THY1"
## [8] "PGK1"
                  "CAMK2B" "AGTR1"
                                      "C5orf38" "HK2"
                                                           "EFNA3"
                                                                     "CHIT1"
## [15] "RPS6KA5" "CACNA1H" "SHC4"
                                      "HDAC2"
                                                 "HOXA9"
                                                           "MAP2K4"
```

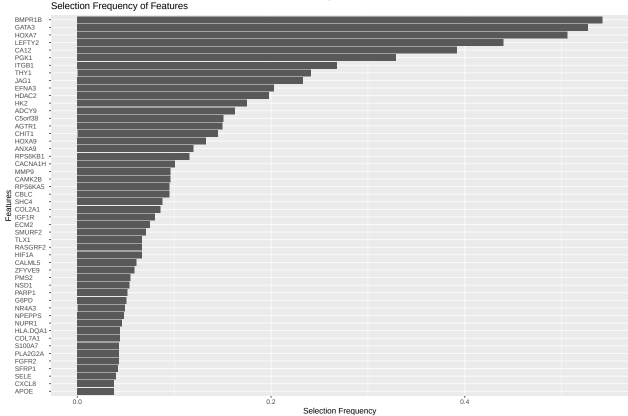
#### Mechanistic + Residuals -> proliferation score (multiplicative)

```
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS
## Mean: 0.4028471
## Median: 0.437445
## Variance: 0.05302116
## st.dev.: 0.2302632
```



## MSE RESULTS
## Mean: 0.1455819
## Median: 0.1286019
## Variance: 0.004632394
## st.dev.: 0.06806169





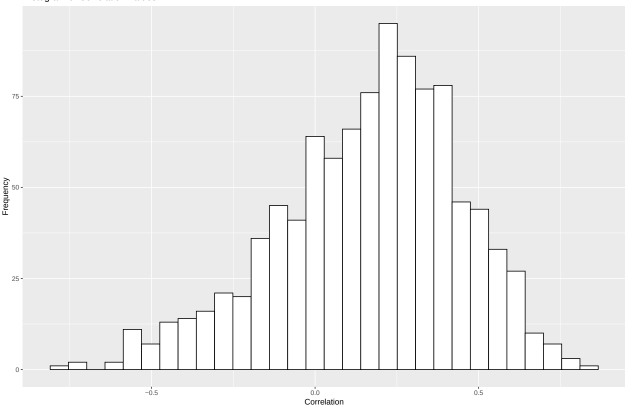
##

```
## Features selected 50% or more times:
## BMPR1B GATA3 HOXA7
## Top 20 featrues:
  [1] "BMPR1B"
                  "GATA3"
                             "HOXA7"
                                       "LEFTY2"
                                                 "CA12"
                                                           "PGK1"
                                                                      "ITGB1"
## [8] "THY1"
                  "JAG1"
                                       "HDAC2"
                             "EFNA3"
                                                 "HK2"
                                                           "ADCY9"
                                                                      "C5orf38"
## [15] "AGTR1"
                  "CHIT1"
                            "HOXA9"
                                       "ANXA9"
                                                 "RPS6KB1" "CACNA1H"
```

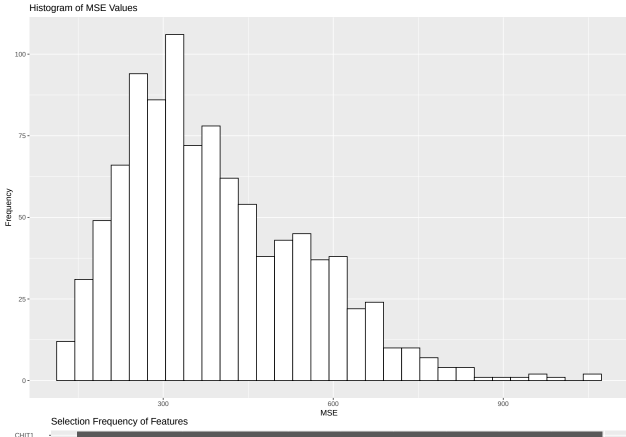
#### Mechnaistic + Residuals -> ROR-proliferation score (additive)

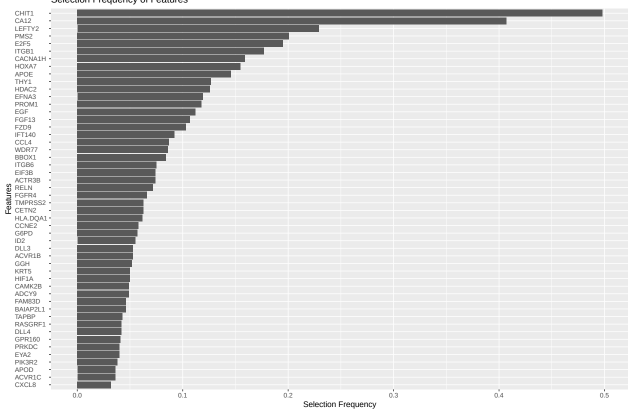
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS
## Mean: 0.16425
## Median: 0.198308
## Variance: 0.07775369

## st.dev.: 0.2788435
Histogram of Correlation Values



## MSE RESULTS
## Mean: 392.5436
## Median: 360.3419
## Variance: 25240.73
## st.dev.: 158.8733



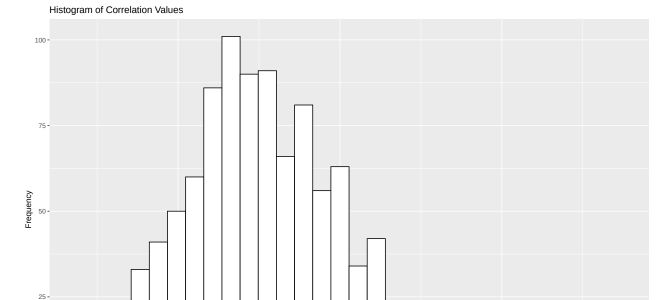


##

```
## Features selected 50% or more times:
##
## Top 20 featrues:
                  "CA12"
   [1] "CHIT1"
                             "LEFTY2"
                                       "PMS2"
                                                  "E2F5"
                                                            "ITGB1"
                                                                       "CACNA1H"
   [8] "HOXA7"
                   "APOE"
                             "THY1"
                                                                       "EGF"
                                        "HDAC2"
                                                  "EFNA3"
                                                            "PROM1"
## [15] "FGF13"
                  "FZD9"
                             "IFT140"
                                       "CCL4"
                                                  "WDR77"
                                                            "BBOX1"
```

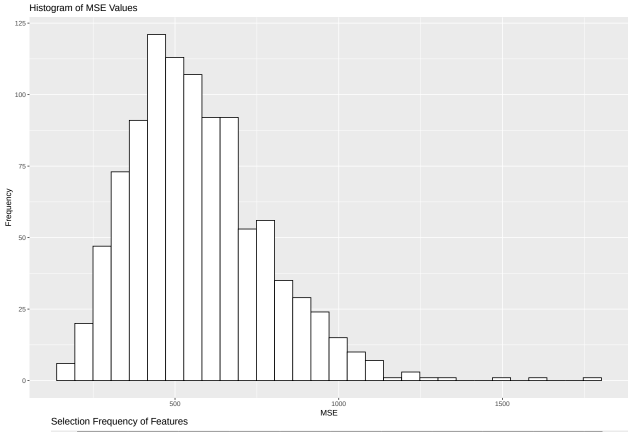
#### Mechnaistic + Residuals -> ROR-proliferation score (multiplicative)

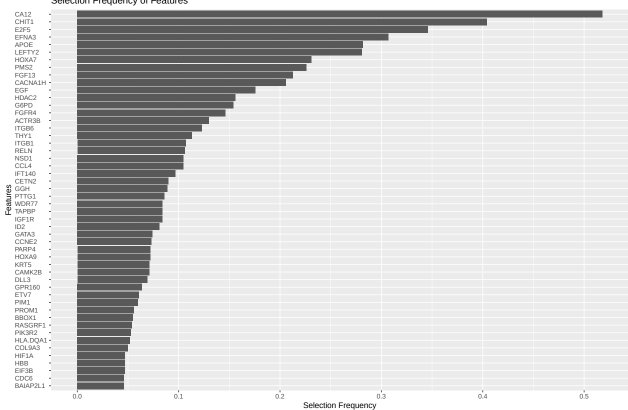
```
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS
## Mean: -0.2145253
## Median: -0.2415987
## Variance: 0.06311304
## st.dev.: 0.2512231
```



## MSE RESULTS
## Mean: 568.8063
## Median: 541.2023
## Variance: 43510.24
## st.dev.: 208.5911

Correlation





##

```
## Features selected 50% or more times:
## CA12
## Top 20 featrues:
  [1] "CA12"
                  "CHIT1"
                            "E2F5"
                                       "EFNA3"
                                                 "APOE"
                                                           "LEFTY2"
                                                                     "HOXA7"
   [8] "PMS2"
                            "CACNA1H" "EGF"
                                                           "G6PD"
                  "FGF13"
                                                 "HDAC2"
                                                                     "FGFR4"
## [15] "ACTR3B" "ITGB6"
                            "THY1"
                                      "ITGB1"
                                                 "RELN"
                                                           "CCL4"
```

#### Summery results: lasso proliferation score (repeated cross-validation)

Model	cor_mean	$sd\_cor$	${\rm MSE\_mean}$	$MSE\_sd$
lasso 6 genes	0.0919663	0.3068209	0.1655931	0.0718379
lasso 771 genes	0.4737037	0.2310209	0.0620913	0.0753056
Nodes	0.2842257	0.2768458	0.1560308	0.0762883
Residual additive	0.4633095	0.2227105	0.1331785	0.0654065
Residual multiplicative	0.4028471	0.2302632	0.1455819	0.0680617

#### Summery results: lasso ROR+proliferation score (repeated cross-validation)

cor_mean	$sd\_cor$	${\rm MSE\_mean}$	$MSE\_sd$
-0.4822298	0.1724985	374.1519	144.1559
0.0806237	0.2767153	393.8069	159.6451
0.1806504	0.2854892	380.1157	164.5869
0.1642500	0.2788435	392.5436	158.8733
-0.2145253	0.2512231	568.8063	208.5911
	-0.4822298 0.0806237 0.1806504 0.1642500	-0.4822298     0.1724985       0.0806237     0.2767153       0.1806504     0.2854892       0.1642500     0.2788435	-0.4822298     0.1724985     374.1519       0.0806237     0.2767153     393.8069       0.1806504     0.2854892     380.1157       0.1642500     0.2788435     392.5436

## Ridge bootstrap

#### 771 genes -> proliferation score

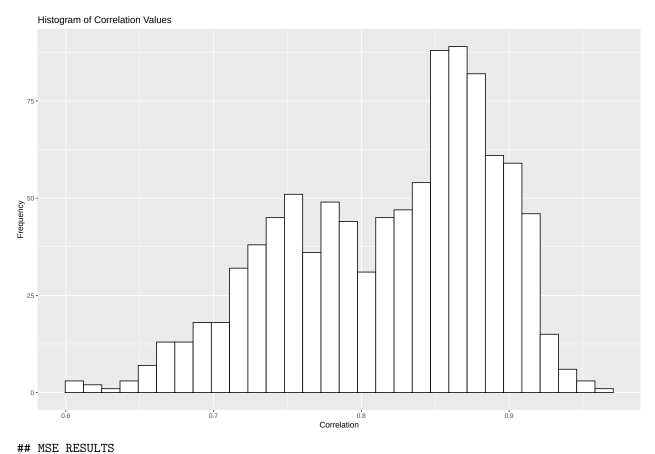
```
## number of models fitted: 1000
```

## Fraction of model fits with no selected genes: 0

##

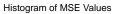
## CORRELATIONS RESULTS

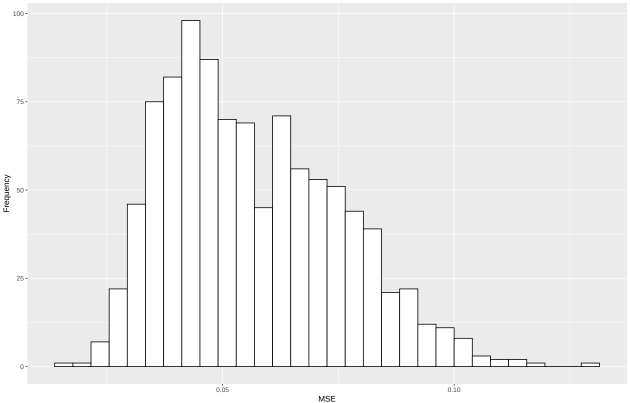
## Mean: 0.8189568 ## Median: 0.8355063 ## Variance: 0.005044925 ## st.dev.: 0.07102764



## Mse Results ## Mean: 0.05665241 ## Median: 0.0532761

## Variance: 0.0003488416 ## st.dev.: 0.0186773





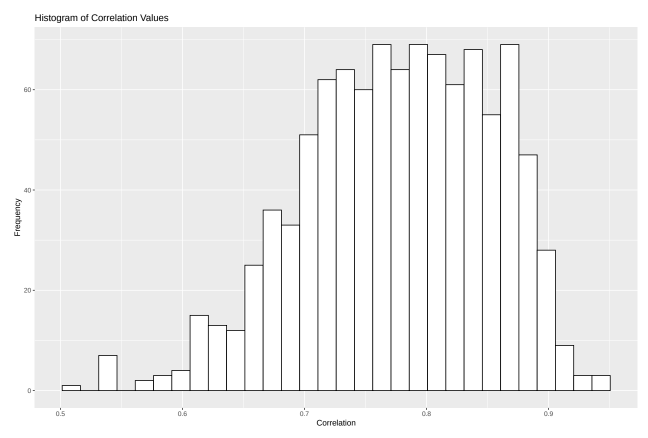
### 771 genes -> ROR-proliferation score

## number of models fitted: 1000

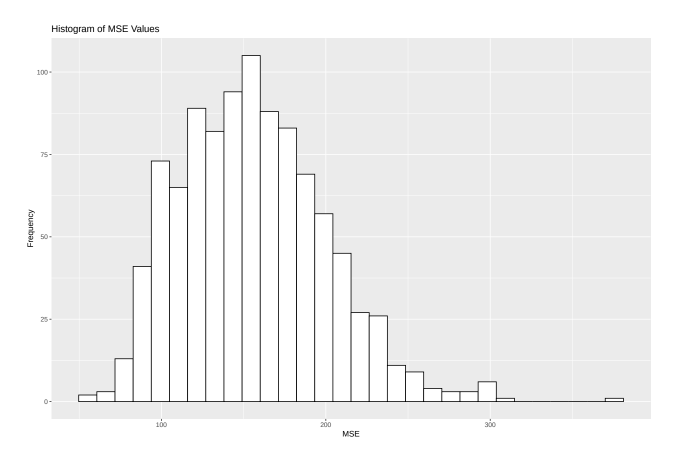
 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0

##

## CORRELATIONS RESULTS
## Mean: 0.7761924
## Median: 0.7811637
## Variance: 0.006001423
## st.dev.: 0.07746885



## MSE RESULTS
## Mean: 156.065
## Median: 154.0679
## Variance: 1952.025
## st.dev.: 44.18173



# Ridge repeated cross-validation

```
771 genes -> proliferation score
```

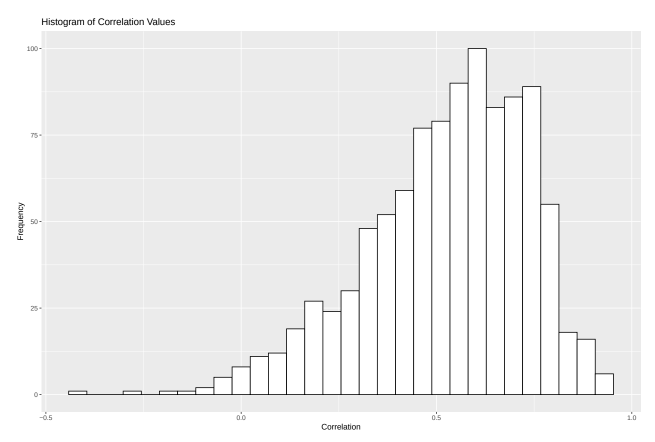
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0

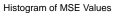
##

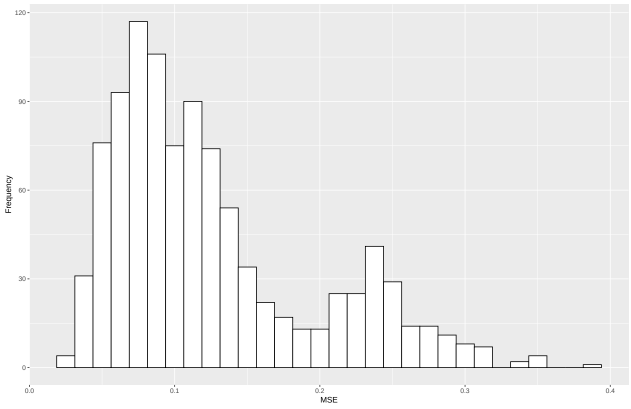
## CORRELATIONS RESULTS
## Mean: 0.5268342
## Median: 0.5562175

## Variance: 0.04291266 ## st.dev.: 0.2071537



## MSE RESULTS
## Mean: 0.1256548
## Median: 0.1059589
## Variance: 0.004890454
## st.dev.: 0.06993178





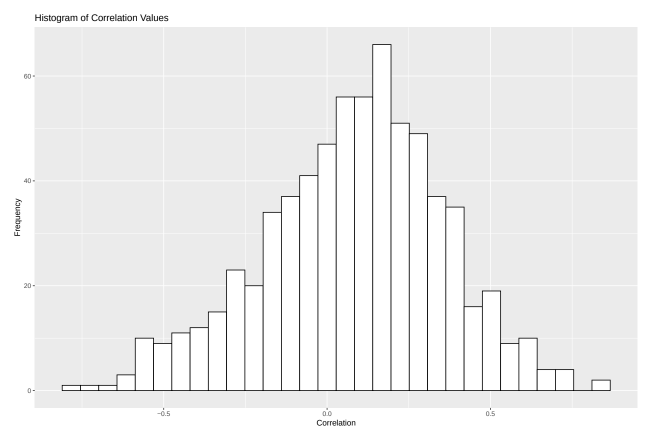
### 771 genes -> ROR-proliferation score

## number of models fitted: 1000

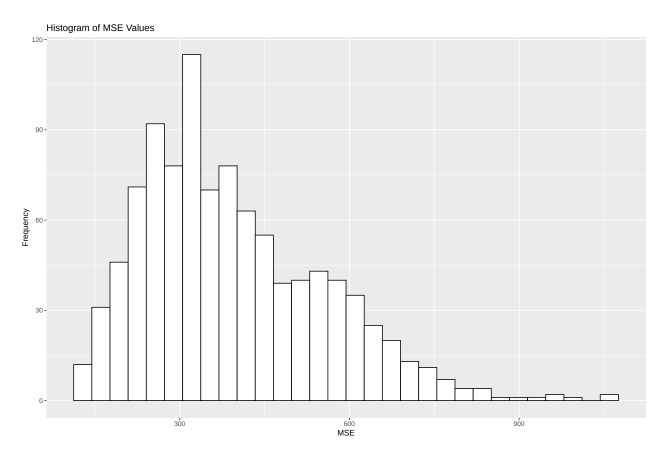
 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0.321

##

## CORRELATIONS RESULTS
## Mean: 0.08062366
## Median: 0.1014264
## Variance: 0.07657135
## st.dev.: 0.2767153



## MSE RESULTS
## Mean: 393.8069
## Median: 360.5105
## Variance: 25486.55
## st.dev.: 159.6451



#### Summery results: ridge 771 genes bootstrap and repeated cross-validation

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
prolif boot	0.8189568	0.0710276	0.0566524	0.0186773
ROR-prolif boot	0.7761924	0.0774688	156.0649552	44.1817261
prolif rep cross-val	0.5268342	0.2071537	0.1256548	0.0699318
ROR-prolif rep cross-val	0.0806237	0.2767153	393.8068910	159.6450765

## Elastic Net - bootstrap

#### 771 genes -> proliferation score

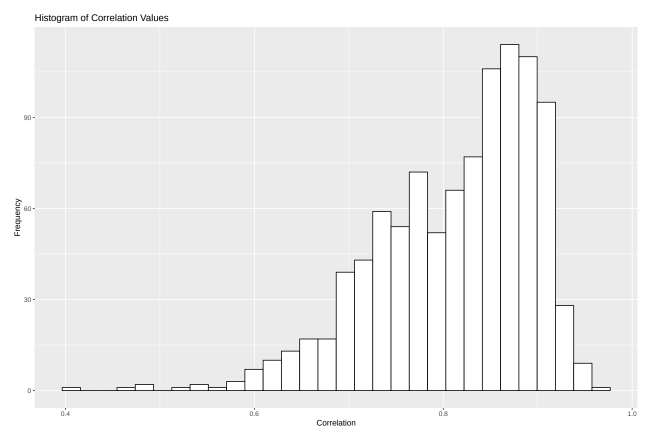
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0

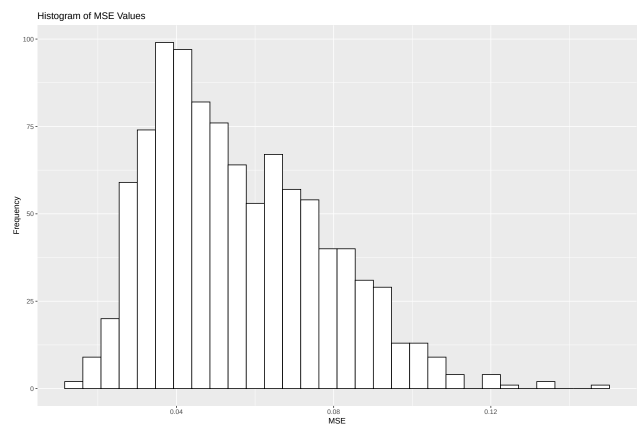
##

## CORRELATIONS RESULTS
## Mean: 0.8125591

## Median: 0.8324809 ## Variance: 0.007038791 ## st.dev.: 0.0838975



## MSE RESULTS
## Mean: 0.0558955
## Median: 0.05152763
## Variance: 0.0004710269
## st.dev.: 0.02170316



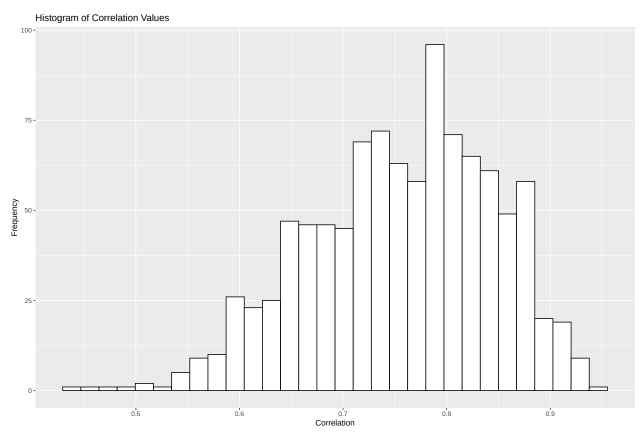
### 771 genes -> ROR-proliferation score

## number of models fitted: 1000

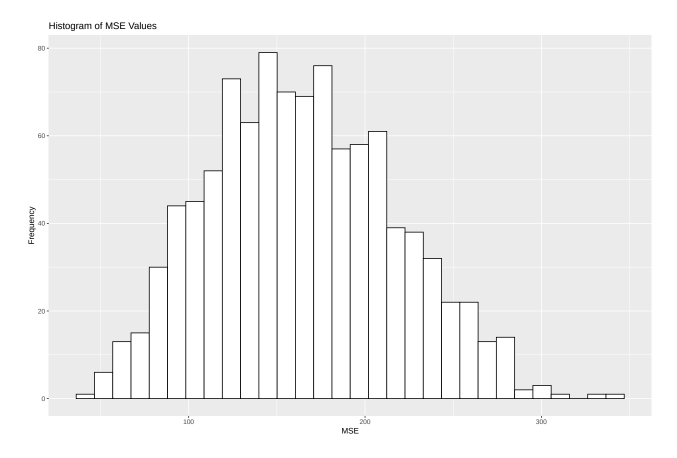
 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0

##

## CORRELATIONS RESULTS
## Mean: 0.7565123
## Median: 0.7644687
## Variance: 0.007887981
## st.dev.: 0.08881431



## MSE RESULTS ## Mean: 164.4116 ## Median: 162.2643 ## Variance: 2785.22 ## st.dev.: 52.77518



## Elastic Net: cross-validation

#### 771 genes -> proliferation score

## number of models fitted: 1000

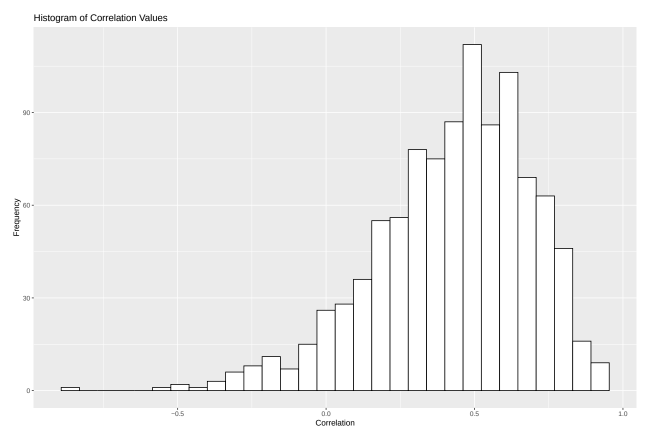
## Fraction of model fits with no selected genes: 0

##

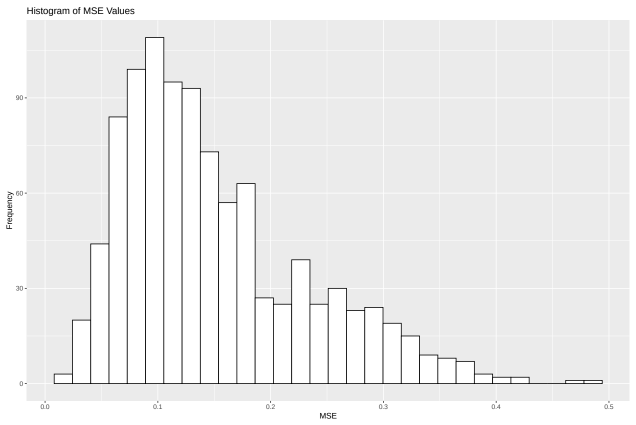
## CORRELATIONS RESULTS

## Mean: 0.427189

## Median: 0.4644365 ## Variance: 0.06953924 ## st.dev.: 0.2637029



## MSE RESULTS
## Mean: 0.1501025
## Median: 0.1284895
## Variance: 0.006763425
## st.dev.: 0.08224004



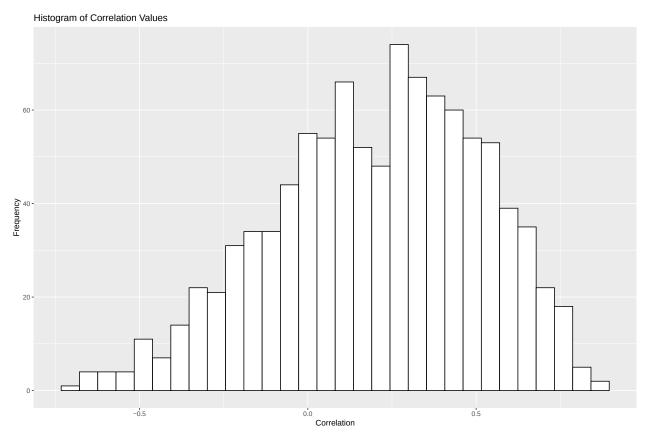
### 771 genes -> ROR-proliferation score

## number of models fitted: 1000

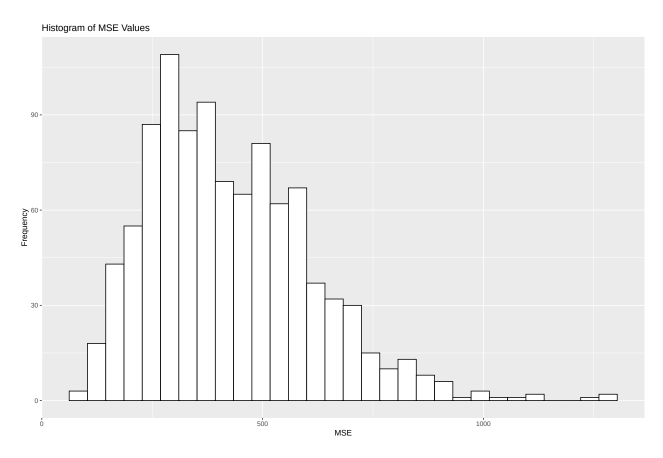
 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0.002

##

## CORRELATIONS RESULTS ## Mean: 0.2049341 ## Median: 0.226514 ## Variance: 0.0968435 ## st.dev.: 0.3111969



## MSE RESULTS
## Mean: 427.3517
## Median: 396.2622
## Variance: 34687.75
## st.dev.: 186.2465



#### Summery results: elastic net 771 genes bootstrap and repeated cross-validation

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
prolif boot	0.8125591	0.0838975	0.0558955	0.0217032
ROR-prolif boot	0.7565123	0.0888143	164.4116160	52.7751849
prolif rep cross-val	0.4271890	0.2637029	0.1501025	0.0822400
ROR-prolif rep cross-val	0.2049341	0.3111969	427.3517412	186.2464725

## Boosting with stumps as base learner - bootstrap

#### 771 genes -> proliferation score

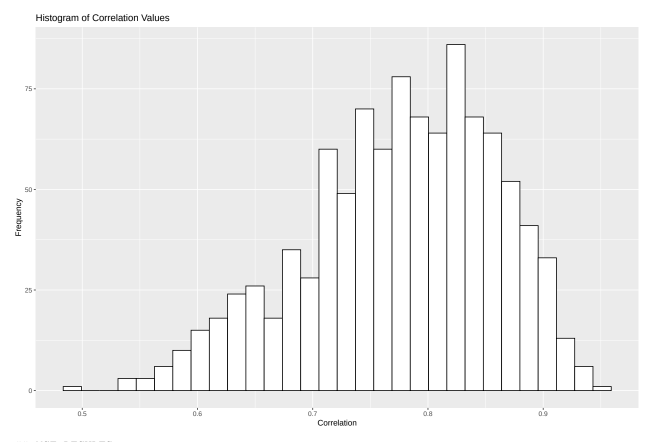
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0

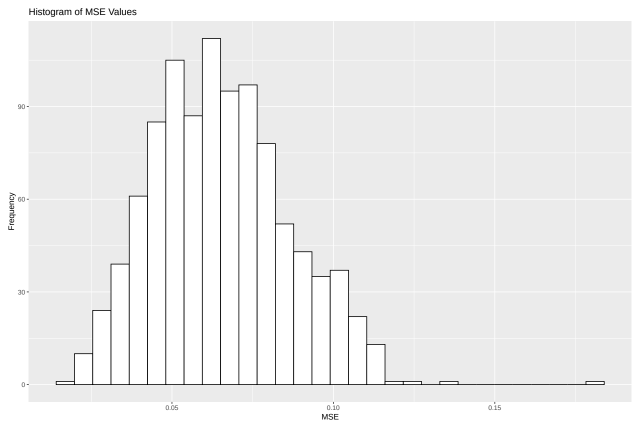
##

## CORRELATIONS RESULTS
## Mean: 0.7760479
## Median: 0.7841718
## Variance: 0.006853002

## st.dev.: 0.08278286



## MSE RESULTS
## Mean: 0.06537103
## Median: 0.06397191
## Variance: 0.0004410638
## st.dev.: 0.02100152



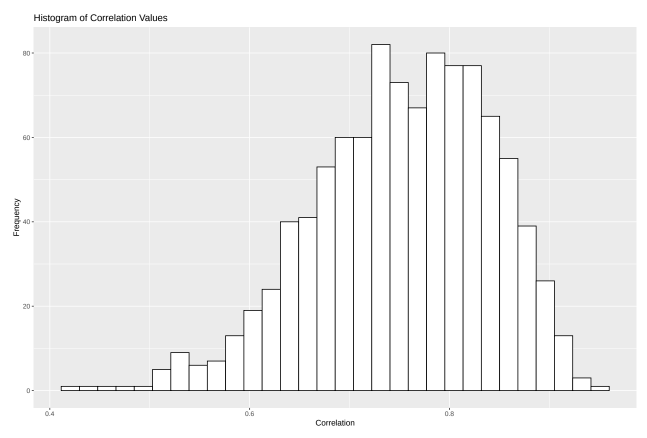
### 771 genes -> ROR-proliferation score

## number of models fitted: 1000

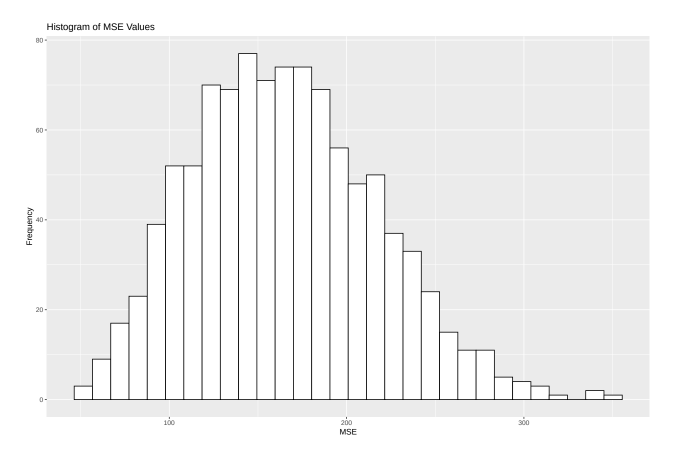
 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0

##

## CORRELATIONS RESULTS
## Mean: 0.7530515
## Median: 0.7593775
## Variance: 0.007786865
## st.dev.: 0.08824321



## MSE RESULTS
## Mean: 165.145
## Median: 162.2361
## Variance: 2690.091
## st.dev.: 51.86608



# Boosting with stumps as base learner cross-validation

#### 771 genes -> proliferation score

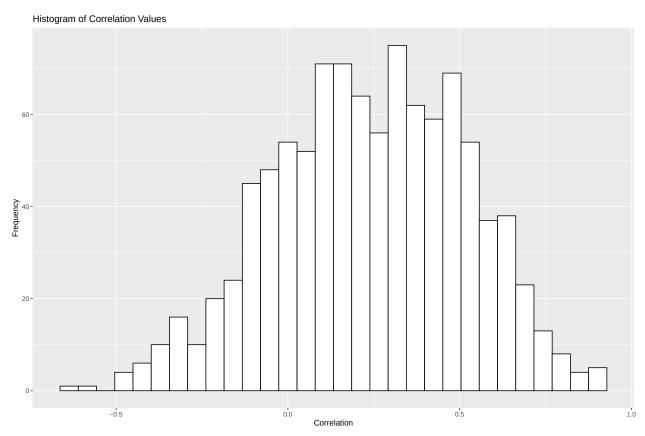
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0

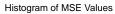
##

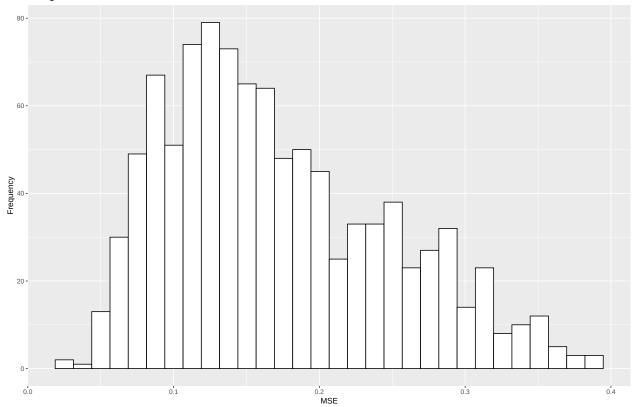
## CORRELATIONS RESULTS ## Mean: 0.2364594

## Median: 0.2426694 ## Variance: 0.07812256 ## st.dev.: 0.2795041



## MSE RESULTS
## Mean: 0.1712618
## Median: 0.1554505
## Variance: 0.005837176
## st.dev.: 0.07640141





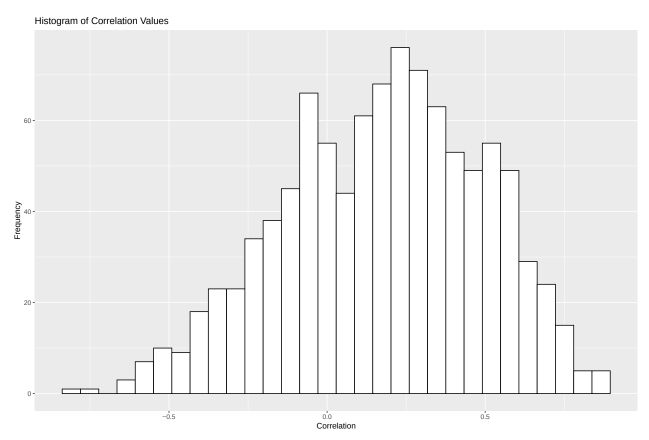
### 771 genes -> ROR-proliferation score

## number of models fitted: 1000

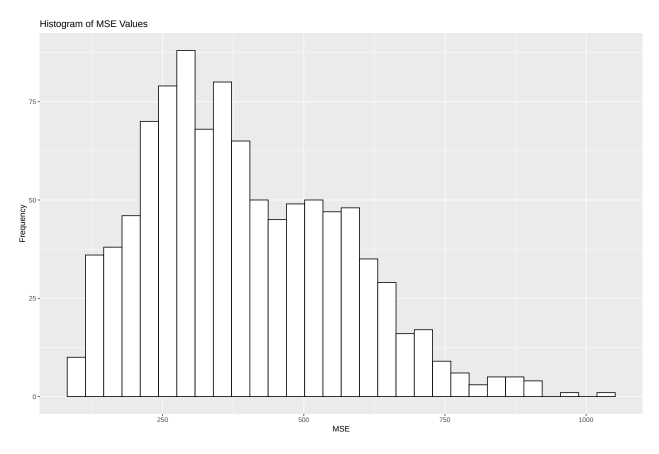
 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0

##

## CORRELATIONS RESULTS
## Mean: 0.1744792
## Median: 0.1948469
## Variance: 0.09932263
## st.dev.: 0.3151549



## MSE RESULTS
## Mean: 394.2634
## Median: 366.3266
## Variance: 29461.83
## st.dev.: 171.6445



#### Summery results: Boosting with stumps 771 genes bootstrap and repeated cross-validation

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
prolif boot	0.7760479	0.0827829	0.0653710	0.0210015
ROR-prolif boot	0.7530515	0.0882432	165.1450271	51.8660843
prolif rep cross-val	0.2364594	0.2795041	0.1712618	0.0764014
ROR-prolif rep cross-val	0.1744792	0.3151549	394.2634498	171.6444924

### START USING DOMAIN KNOWLEDGE

# PCA on signature gene sets using repeated cross-validation

Ridge: 771 genes -> ROR-proliferation score

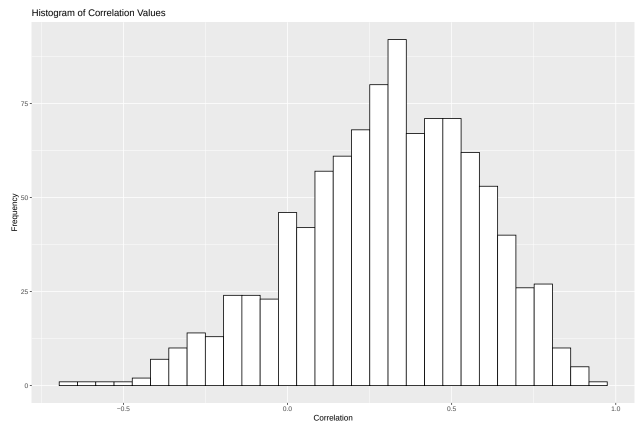
## number of models fitted: 1000

 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0

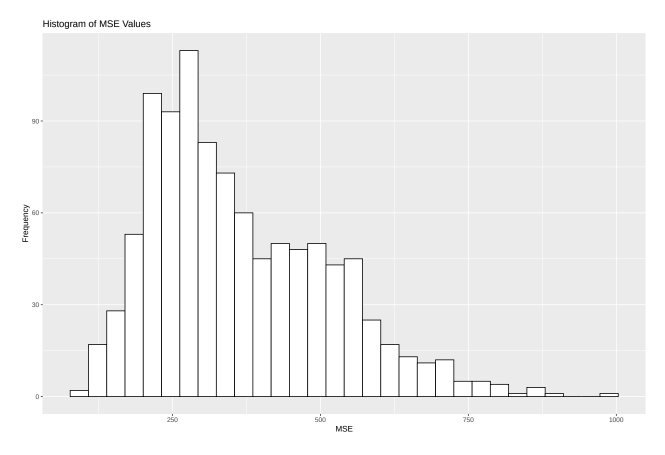
##

## CORRELATIONS RESULTS

## Mean: 0.3028091 ## Median: 0.319312 ## Variance: 0.07725244 ## st.dev.: 0.2779432



## MSE RESULTS
## Mean: 364.5733
## Median: 327.0747
## Variance: 22807.28
## st.dev.: 151.0208



Ridge: 771 genes -> ROR-proliferation score + interactions between PCs

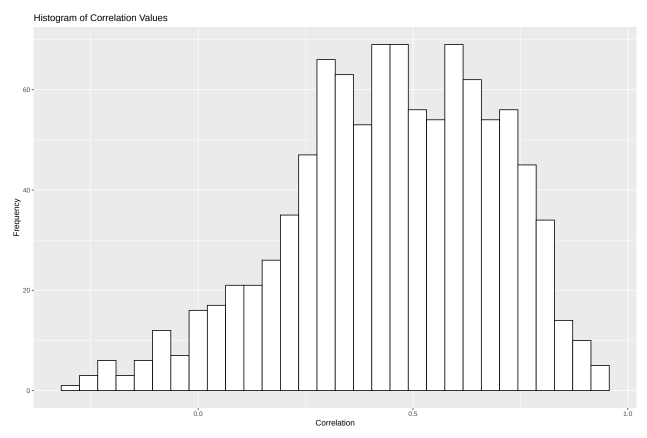
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0

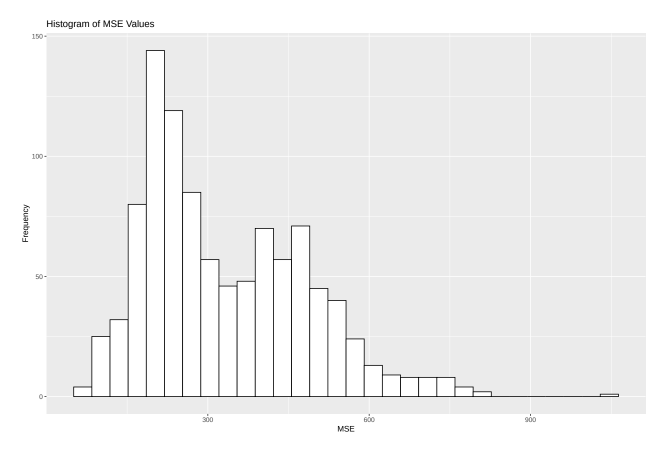
##

## CORRELATIONS RESULTS

## Mean: 0.4505837 ## Median: 0.463892 ## Variance: 0.05781415 ## st.dev.: 0.2404457



## MSE RESULTS
## Mean: 333.3975
## Median: 291.1335
## Variance: 23000.06
## st.dev.: 151.6577



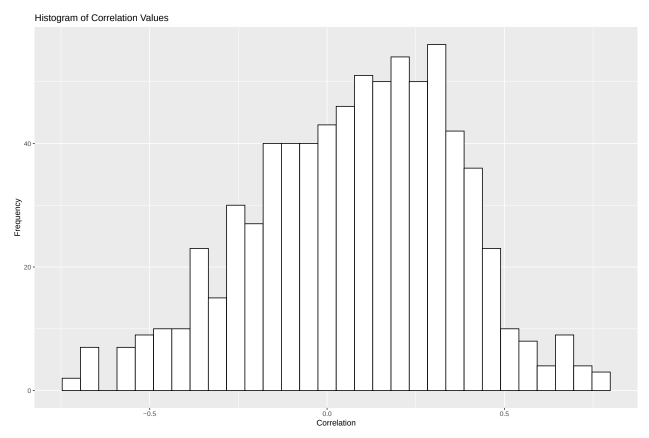
### Lasso: 771 genes -> ROR-proliferation score

## number of models fitted: 1000

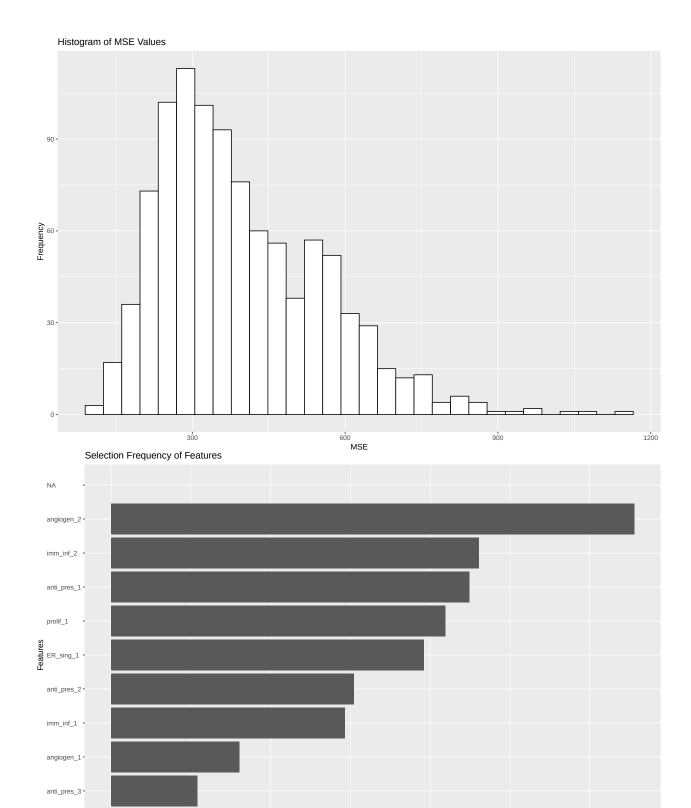
## Fraction of model fits with no selected genes: 0.251

##

## CORRELATIONS RESULTS ## Mean: 0.07849613 ## Median: 0.1028317 ## Variance: 0.08158919 ## st.dev.: 0.2856382



## MSE RESULTS
## Mean: 396.4902
## Median: 361.1201
## Variance: 25641.71
## st.dev.: 160.1303



prolif\_2

0.0

Selection Frequency

0.4

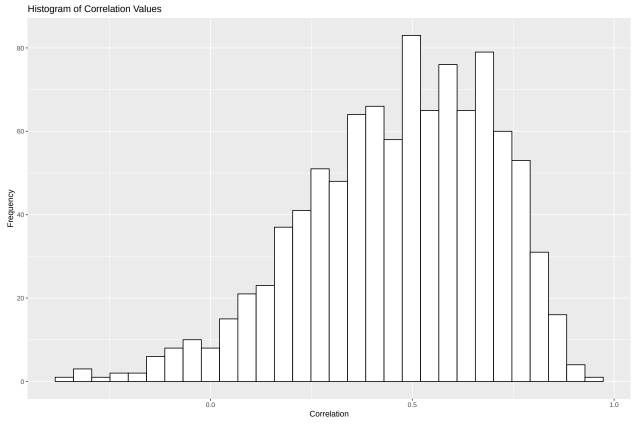
0.6

0.2

```
## Features selected 50% or more times:
## angiogen_2
## Top 20 featrues:
## [1] "angiogen_2" "imm_inf_2" "anti_pres_1" "prolif_1" "ER_sing_1"
## [6] "anti_pres_2" "imm_inf_1" "angiogen_1" "anti_pres_3" "prolif_2"
## [11] NA NA NA NA NA NA
## [16] NA NA NA NA NA
```

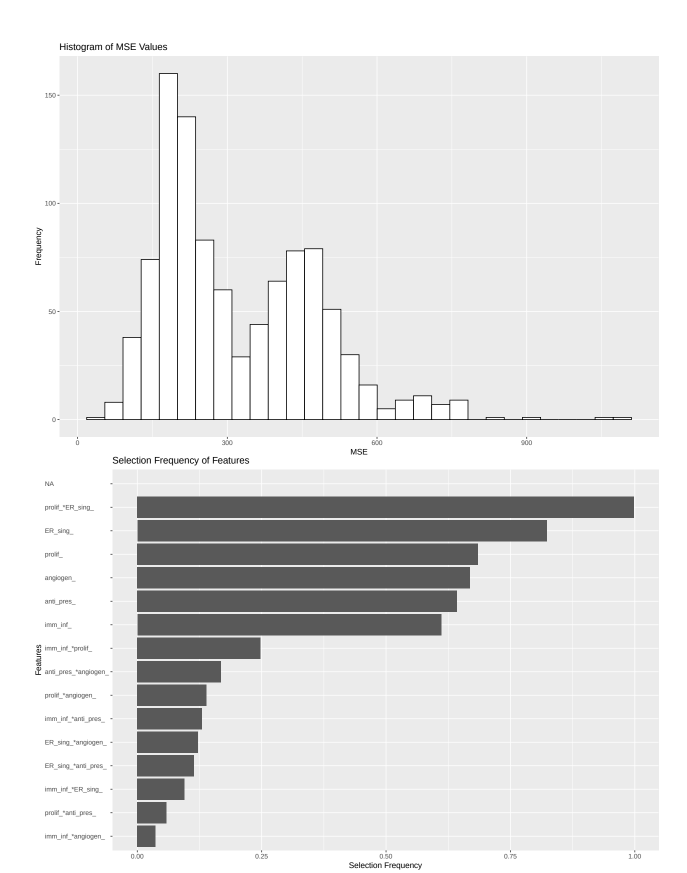
## Lasso: 771 genes -> ROR-proliferation score + interactions between PCs

```
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0.002
##
## CORRELATIONS RESULTS
## Mean: 0.4687056
## Median: 0.4972697
## Variance: 0.05426225
```



## MSE RESULTS
## Mean: 321.0086
## Median: 265.9525
## Variance: 24834.46
## st.dev.: 157.5895

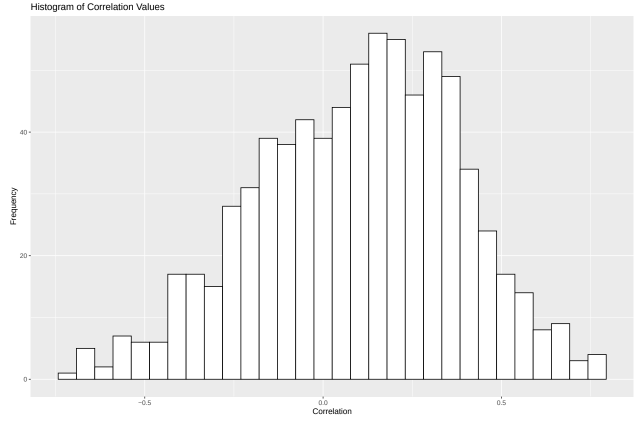
## st.dev.: 0.2329426



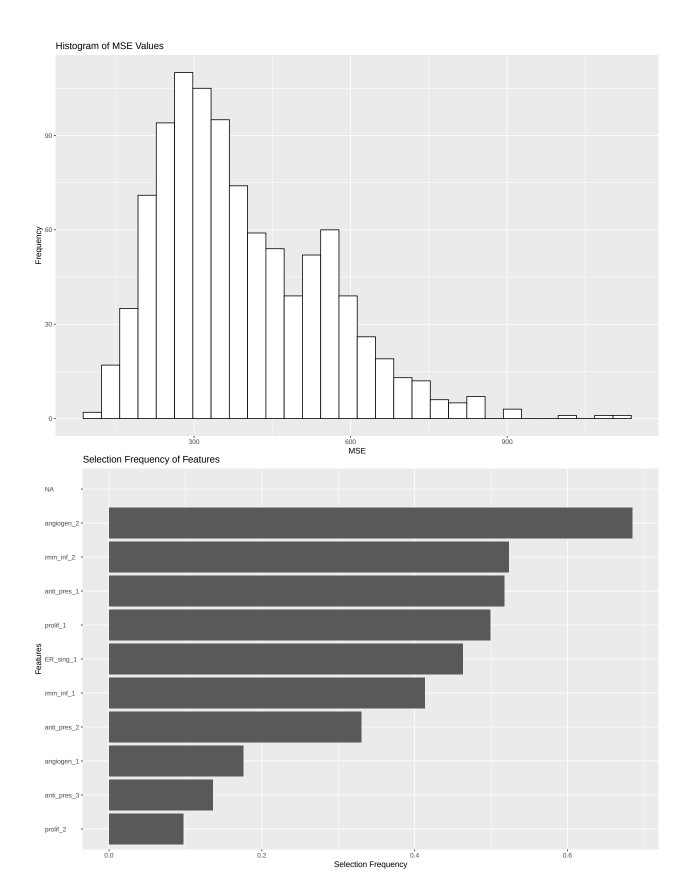
```
## Features selected 50% or more times:
## imm_inf_ prolif_ ER_sing_ anti_pres_ angiogen_ prolif_*ER_sing_
## Top 20 featrues:
  [1] "prolif_*ER_sing_"
                               "ER_sing_"
                                                       "prolif_"
   [4] "angiogen_"
                                                       "imm_inf_"
                               "anti_pres_"
##
  [7] "imm_inf_*prolif_"
                               "anti_pres_*angiogen_" "prolif_*angiogen_"
## [10] "imm_inf_*anti_pres_"
                               "ER_sing_*angiogen_"
                                                       "ER_sing_*anti_pres_"
## [13] "imm_inf_*ER_sing_"
                               "prolif_*anti_pres_"
                                                       "imm_inf_*angiogen_"
## [16] NA
                               NA
## [19] NA
                               NA
```

# ElasticNet: 771 genes -> ROR-proliferation score

## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0.24
##
## CORRELATIONS RESULTS
## Mean: 0.09606929
## Median: 0.1167623
## Variance: 0.08091802
## st.dev.: 0.2844609



## MSE RESULTS ## Mean: 392.3862 ## Median: 358.5637 ## Variance: 24990.96 ## st.dev.: 158.0853

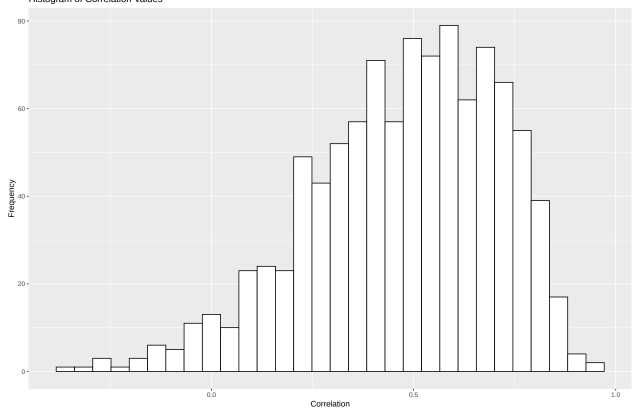


```
## Features selected 50% or more times:
## imm_inf_2 anti_pres_1 angiogen_2
## Top 20 featrues:
  [1] "angiogen_2"
                     "imm_inf_2"
                                   "anti_pres_1" "prolif_1"
                                                               "ER_sing_1"
                      "anti_pres_2" "angiogen_1" "anti_pres_3" "prolif_2"
## [6] "imm_inf_1"
## [11] NA
                                   NA
## [16] NA
                     NA
                                   NA
                                                 NA
                                                               NA
```

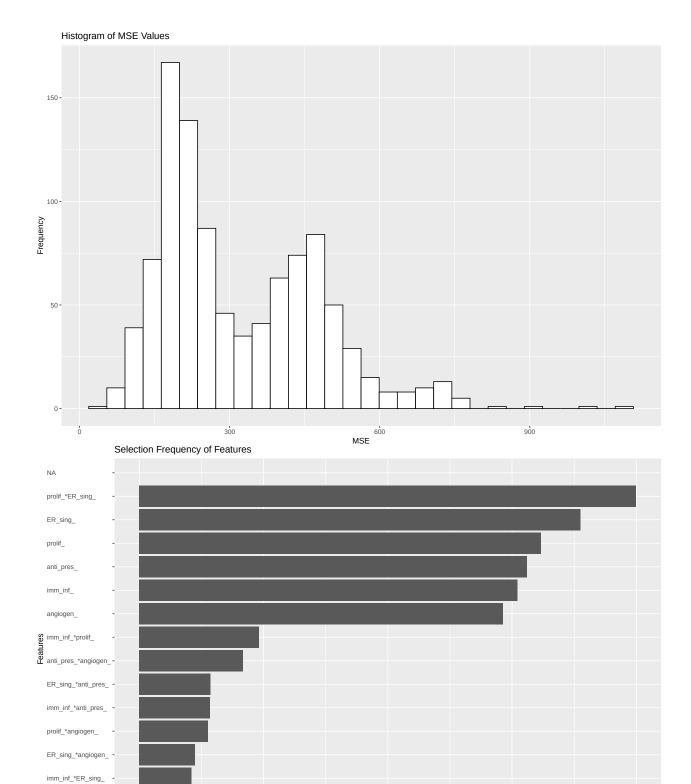
## ElasticNet: 771 genes -> ROR-proliferation score + interactions between PCs

## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0.001
##
## CORRELATIONS RESULTS
## Mean: 0.4749342
## Median: 0.5041917

## Variance: 0.05452847 ## st.dev.: 0.2335133 Histogram of Correlation Values



## MSE RESULTS
## Mean: 319.7269
## Median: 263.5692
## Variance: 24916.52
## st.dev.: 157.8497



prolif\_\*anti\_pres\_ imm\_inf\_\*angiogen\_

0.00

0.25

0.50 Selection Frequency 0.75

```
## Features selected 50% or more times:
## imm_inf_ prolif_ ER_sing_ anti_pres_ angiogen_ prolif_*ER_sing_
## Top 20 featrues:
## [1] "prolif_*ER_sing_"
                               "ER_sing_"
                                                      "prolif_"
## [4] "anti_pres_"
                               "imm_inf_"
                                                      "angiogen_"
## [7] "imm_inf_*prolif_"
                               "anti_pres_*angiogen_" "ER_sing_*anti_pres_"
## [10] "imm_inf_*anti_pres_"
                              "prolif_*angiogen_"
                                                      "ER_sing_*angiogen_"
## [13] "imm_inf_*ER_sing_"
                               "prolif_*anti_pres_"
                                                      "imm_inf_*angiogen_"
## [16] NA
## [19] NA
                               NA
```

## Summery results: PCA ROR+proliferation score (repeated cross-validation)

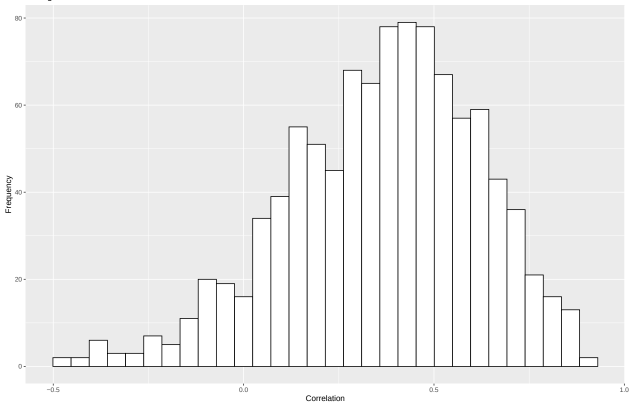
Model	cor_mean	sd_cor	MSE_mean	MSE_sd
ridge	0.3028091	0.2779432	364.5733	151.0208
ridge_interac	0.4505837	0.2404457	333.3975	151.6577
lasso	0.0784961	0.2856382	396.4902	160.1303
$lasso\_interact$	0.4687056	0.2329426	321.0086	157.5895
elastic	0.0960693	0.2844609	392.3862	158.0853
$elastic\_interact$	0.4749342	0.2335133	319.7269	157.8497

# Stacking on signature gene sets using repeated cross-validation

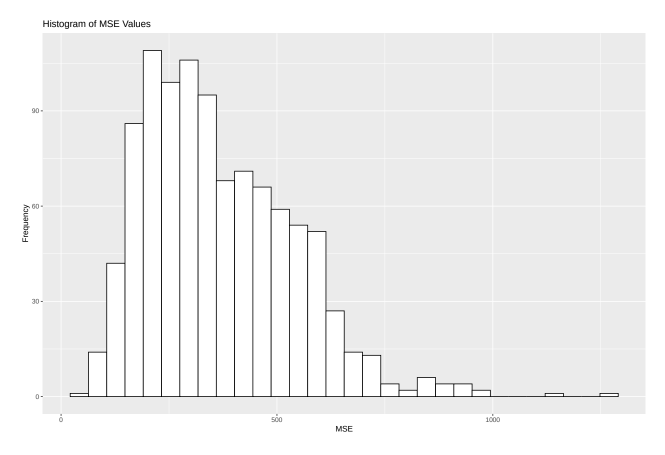
Ridge: 771 genes -> ROR-proliferation score

```
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS
## Mean: 0.363015
## Median: 0.3907333
## Variance: 0.06475058
## st.dev.: 0.2544613
```





## MSE RESULTS
## Mean: 367.3704
## Median: 335.367
## Variance: 29532.74
## st.dev.: 171.8509



Ridge: 771 genes -> ROR-proliferation score + interactions between PCs

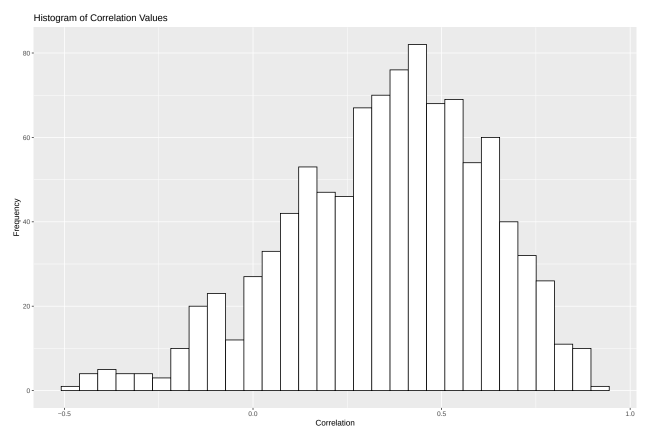
## number of models fitted: 1000

## Fraction of model fits with no selected genes: 0

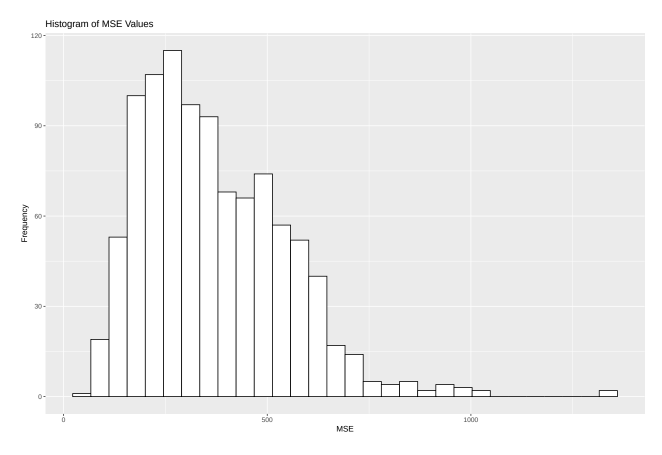
##

## CORRELATIONS RESULTS

## Mean: 0.3555283 ## Median: 0.3864856 ## Variance: 0.06877803 ## st.dev.: 0.2622557



## MSE RESULTS
## Mean: 369.2144
## Median: 337.1152
## Variance: 31306.72
## st.dev.: 176.9371



# Lasso: 771 genes -> ROR-proliferation score

## number of models fitted: 1000

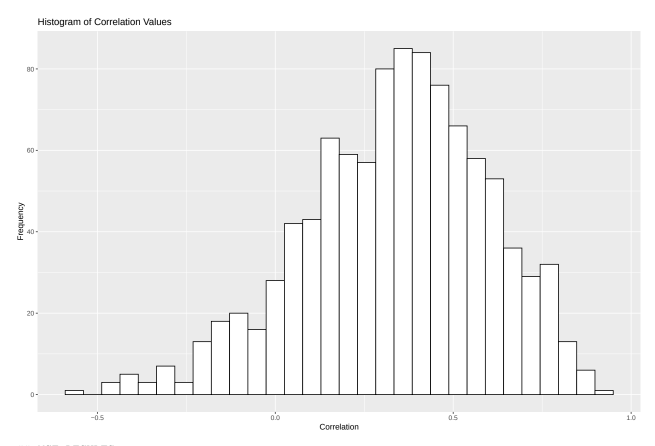
## Fraction of model fits with no selected genes: 0

##

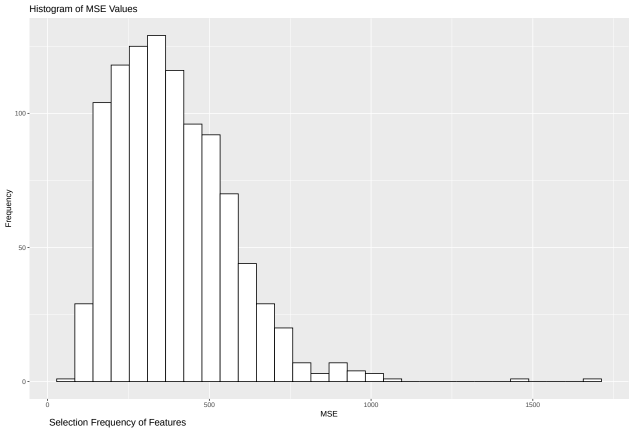
## CORRELATIONS RESULTS

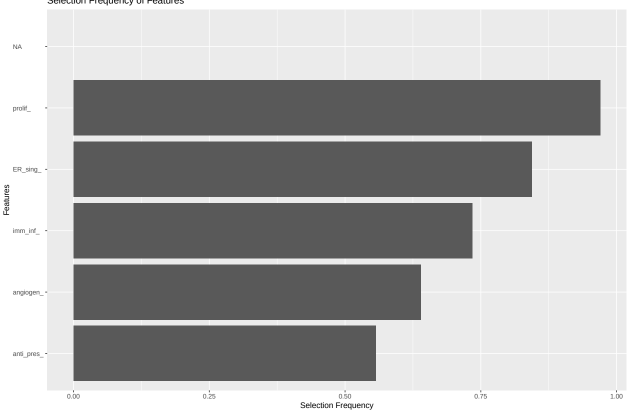
## Mean: 0.3373612 ## Median: 0.3601618 ## Variance: 0.06786039

## st.dev.: 0.2605003



## MSE RESULTS
## Mean: 386.6455
## Median: 360.5423
## Variance: 32369.44
## st.dev.: 179.9151



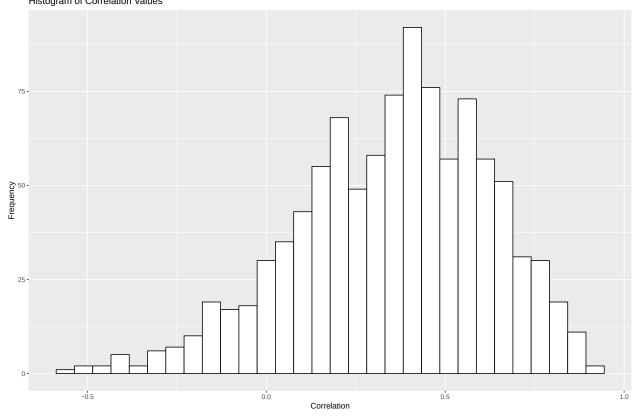


```
## Features selected 50% or more times:
## imm_inf_ prolif_ ER_sing_ anti_pres_ angiogen_
## Top 20 featrues:
   [1] "prolif_"
                     "ER_sing_"
                                   "imm_inf_"
                                                "angiogen_" "anti_pres_"
##
   [6] NA
                     NA
                                   NA
                                                              NA
## [11] NA
                     NA
                                   NA
                                                NA
                                                              NA
## [16] NA
                     NA
                                   NA
                                                NA
                                                              NA
```

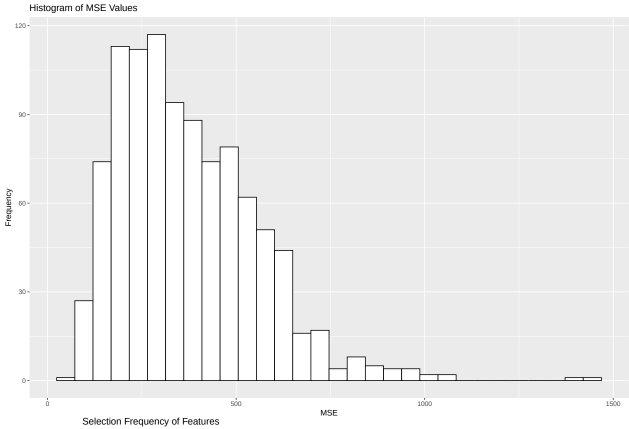
# Lasso: 771 genes -> ROR-proliferation score + interactions between PCs

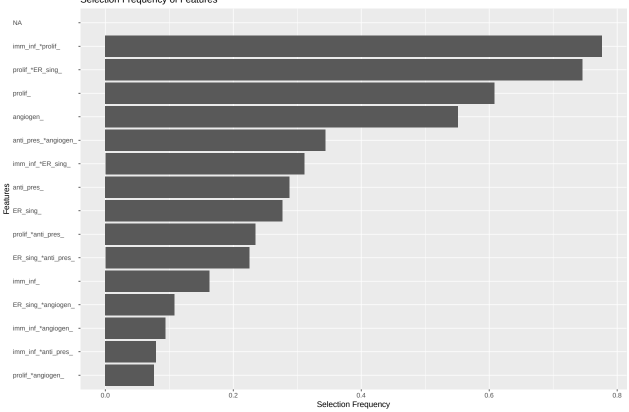
## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS

## Mean: 0.3521323 ## Median: 0.381805 ## Variance: 0.07224413 ## st.dev.: 0.2687827 Histogram of Correlation Values



## MSE RESULTS
## Mean: 373.2299
## Median: 341.6008
## Variance: 33334.83
## st.dev.: 182.5783

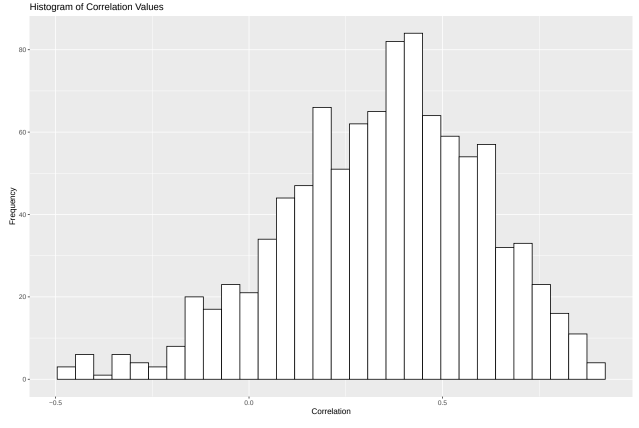




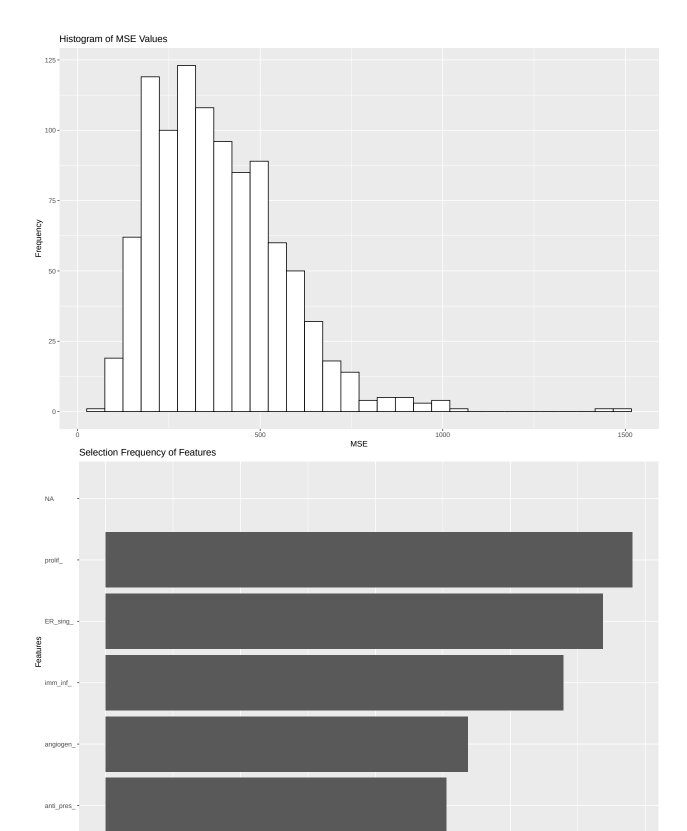
```
## Features selected 50% or more times:
## prolif_ angiogen_ imm_inf_*prolif_ prolif_*ER_sing_
## Top 20 featrues:
  [1] "imm_inf_*prolif_"
                                "prolif_*ER_sing_"
                                                       "prolif_"
                                "anti_pres_*angiogen_" "imm_inf_*ER_sing_"
##
   [4] "angiogen_"
##
  [7] "anti_pres_"
                                "ER_sing_"
                                                       "prolif_*anti_pres_"
## [10] "ER_sing_*anti_pres_"
                                "imm inf "
                                                       "ER_sing_*angiogen_"
## [13] "imm_inf_*angiogen_"
                                "imm_inf_*anti_pres_"
                                                       "prolif_*angiogen_"
## [16] NA
                                                       NA
## [19] NA
                               NA
```

# ElasticNet: 771 genes $\rightarrow$ ROR-proliferation score

## number of models fitted: 1000
## Fraction of model fits with no selected genes: 0
##
## CORRELATIONS RESULTS
## Mean: 0.3417178
## Median: 0.3669518
## Variance: 0.06675737
## st.dev.: 0.2583745



## MSE RESULTS
## Mean: 383.7542
## Median: 356.8975
## Variance: 31468.03
## st.dev.: 177.3923



0.00

0.25

0.50 Selection Frequency

0.75

```
## Features selected 50% or more times:
## imm_inf_ prolif_ ER_sing_ anti_pres_ angiogen_
## Top 20 featrues:
   [1] "prolif_"
                      "ER_sing_"
                                   "imm_inf_"
                                                "angiogen_" "anti_pres_"
##
   [6] NA
                     NA
                                   NA
                                                              NA
## [11] NA
                     NA
                                   NA
                                                NA
                                                              NA
## [16] NA
                     NA
                                   NA
                                                NA
                                                              NA
```

## ElasticNet: 771 genes -> ROR-proliferation score + interactions between PCs

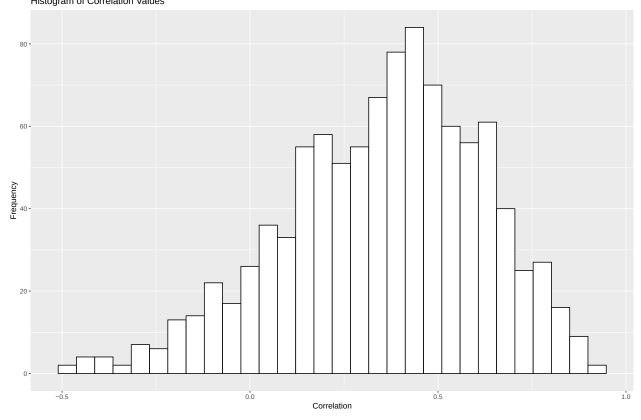
## number of models fitted: 1000

 $\mbox{\tt \#\#}$  Fraction of model fits with no selected genes: 0

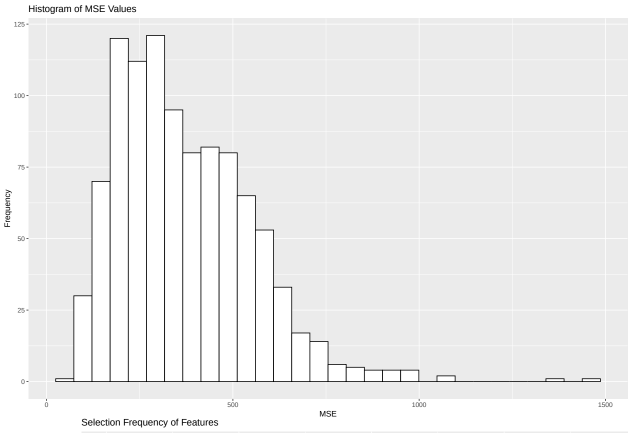
##

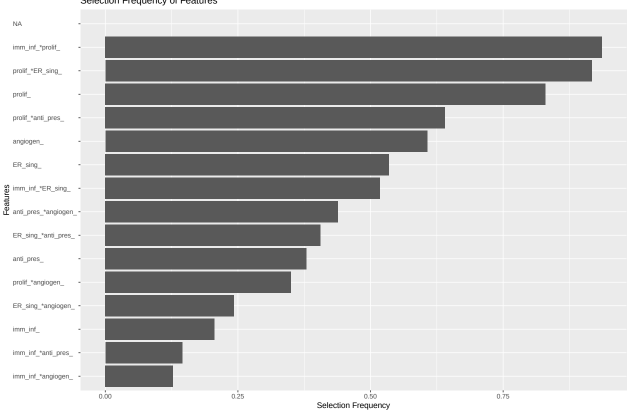
## CORRELATIONS RESULTS

## Mean: 0.353272
## Median: 0.3797595
## Variance: 0.07046161
## st.dev.: 0.2654461
Histogram of Correlation Values



## MSE RESULTS ## Mean: 371.1879 ## Median: 339.6459 ## Variance: 32238 ## st.dev.: 179.5494





```
## Features selected 50% or more times:
## prolif_ ER_sing_ angiogen_ imm_inf_*prolif_ imm_inf_*ER_sing_ prolif_*ER_sing_ prolif_*anti_pres_
## Top 20 featrues:
  [1] "imm_inf_*prolif_"
                               "prolif_*ER_sing_"
                                                      "prolif_"
   [4] "prolif_*anti_pres_"
                               "angiogen_"
                                                      "ER_sing_"
## [7] "imm_inf_*ER_sing_"
                               "anti_pres_*angiogen_" "ER_sing_*anti_pres_"
                               "prolif_*angiogen_"
## [10] "anti_pres_"
                                                     "ER_sing_*angiogen_"
                               "imm_inf_*anti_pres_" "imm_inf_*angiogen_"
## [13] "imm_inf_"
## [16] NA
## [19] NA
                               NA
```

# Summery results: Stacking ROR+proliferation score (repeated cross-validation)

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
ridge	0.3630150	0.2544613	367.3704	171.8509
ridge_interac	0.3555283	0.2622557	369.2144	176.9371
lasso	0.3373612	0.2605003	386.6455	179.9151
$lasso\_interact$	0.3521323	0.2687827	373.2299	182.5783
elastic	0.3417178	0.2583745	383.7542	177.3923
$elastic\_interact$	0.3532720	0.2654461	371.1879	179.5494

# Post Lasso

not done

# Summery of all results

Summery results: lasso proliferation score (bootstrap)

Model	cor_mean	$sd\_cor$	${\rm MSE\_mean}$	$MSE\_sd$
lasso 6 genes	0.3498379	0.0631210	0.1492302	0.0124509
lasso 771 genes	0.7941413	0.0901070	0.0620913	0.0239813
Nodes	0.4144960	0.0642722	0.1479731	0.1916630
Residual additive	0.7835808	0.0854335	0.0638484	0.0213956
Residual multiplicative	0.7266736	0.0895236	0.0813415	0.0231428

# Summery results: lasso ROR+proliferation score (bootstrap)

Model	cor_mean	$sd\_cor$	${\rm MSE\_mean}$	$MSE\_sd$
lasso 6 genes	0.1282306	0.0534128	378.0940	23.44024
lasso 771 genes	0.6968101	0.0995060	203.4080	61.34872
Nodes	0.2964169	0.0830696	417.6667	1027.06231
Residual additive	0.6925953	0.1092315	202.3235	61.04236
Residual multiplicative	0.5427757	0.1908061	291.0186	82.79231

Summery results: lasso proliferation score (repeated cross-validation)

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
lasso 6 genes	0.0919663	0.3068209	0.1655931	0.0718379
lasso 771 genes	0.4737037	0.2310209	0.0620913	0.0753056
Nodes	0.2842257	0.2768458	0.1560308	0.0762883
Residual additive	0.4633095	0.2227105	0.1331785	0.0654065
Residual multiplicative	0.4028471	0.2302632	0.1455819	0.0680617

## Summery results: lasso ROR+proliferation score (repeated cross-validation)

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
lasso 6 genes	-0.4822298	0.1724985	374.1519	144.1559
lasso 771 genes	0.0806237	0.2767153	393.8069	159.6451
Nodes	0.1806504	0.2854892	380.1157	164.5869
Residual additive	0.1642500	0.2788435	392.5436	158.8733
Residual multiplicative	-0.2145253	0.2512231	568.8063	208.5911

# Summery results: ridge 771 genes bootstrap and repeated cross-validation

Model	cor_mean	$sd\_cor$	$MSE\_mean$	$MSE\_sd$
prolif boot	0.8189568	0.0710276	0.0566524	0.0186773
ROR-prolif boot	0.7761924	0.0774688	156.0649552	44.1817261
prolif rep cross-val	0.5268342	0.2071537	0.1256548	0.0699318
ROR-prolif rep cross-val	0.0806237	0.2767153	393.8068910	159.6450765

## Summery results: elastic net 771 genes bootstrap and repeated cross-validation

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
prolif boot	0.8125591	0.0838975	0.0558955	0.0217032
ROR-prolif boot	0.7565123	0.0888143	164.4116160	52.7751849
prolif rep cross-val	0.4271890	0.2637029	0.1501025	0.0822400
ROR-prolif rep cross-val	0.2049341	0.3111969	427.3517412	186.2464725

## Summery results: Boosting with stumps 771 genes bootstrap and repeated cross-validation

Model	cor_mean	sd_cor	MSE_mean	MSE_sd
prolif boot	0.7760479	0.0827829	0.0653710	0.0210015
ROR-prolif boot	0.7530515	0.0882432	165.1450271	51.8660843
prolif rep cross-val	0.2364594	0.2795041	0.1712618	0.0764014
ROR-prolif rep cross-val	0.1744792	0.3151549	394.2634498	171.6444924

# Summary using domain knowledge

Summery results: PCA ROR+proliferation score (repeated cross-validation)

Model	cor_mean	$sd\_cor$	MSE_mean	MSE_sd
ridge	0.3028091	0.2779432	364.5733	151.0208
$ridge\_interac$	0.4505837	0.2404457	333.3975	151.6577
lasso	0.0784961	0.2856382	396.4902	160.1303
lasso_interact	0.4687056	0.2329426	321.0086	157.5895
elastic	0.0960693	0.2844609	392.3862	158.0853
$elastic\_interact$	0.4749342	0.2335133	319.7269	157.8497

# Summery results: Stacking ROR+proliferation score (repeated cross-validation)

Model	cor_mean	$sd\_cor$	$MSE\_mean$	$MSE\_sd$
ridge	0.3630150	0.2544613	367.3704	171.8509
$ridge\_interac$	0.3555283	0.2622557	369.2144	176.9371
lasso	0.3373612	0.2605003	386.6455	179.9151
$lasso\_interact$	0.3521323	0.2687827	373.2299	182.5783
elastic	0.3417178	0.2583745	383.7542	177.3923
$elastic\_interact$	0.3532720	0.2654461	371.1879	179.5494

# Summery most interesting maybe

Response is just ROR+proliferation score, and only used repeated cross-validation.

Model	cor_mean	$sd\_cor$	MSE_mean	MSE_sd
Lasso	0.0806236574665814	0.276715287017282	393.806891045194	159.645076482245
Ridge	0.0806236574665814	0.276715287017282	393.806891045194	159.645076482245
ElasticNet	0.204934145140861	0.311196888316335	427.351741193103	186.246472452805
Boosting	0.174479177155333	0.315154930519824	394.263449789716	171.644492361002
Residual (lasso/additive)	0.164249989278317	0.278843491604259	392.543628848029	158.873314150767
PCA ON GENE SETS				
ridge	0.302809117868443	0.277943223198749	364.573285885524	151.020807845973
ridge_interac	0.450583687359656	0.240445740955287	333.397484377894	151.657700615748
lasso	0.0784961285212545	0.285638208410115	396.490198751861	160.130286797427
lasso_interact	0.468705571862351	0.23294259166303	321.008611919544	157.589534266407
elastic	0.0960692932769097	0.284460933078766	392.386193654839	158.085285268069
elastic_interact	0.474934188616362	0.233513328626198	319.726863673702	157.84966217654
STACKING ON GENE				
SETS				
ridge	0.363015012016748	0.254461343637221	367.370358869747	171.850920863394
ridge_interac	0.355528263391855	0.262255655363312	369.214374412132	176.937052266379
lasso	0.337361194772032	0.260500275299529	386.645536492531	179.915095828192
lasso_interact	0.352132288232874	0.268782684463518	373.22993558893	182.578291674399
elastic	0.341717815852986	0.258374481559797	383.754227227308	177.392304737835
elastic_interact	0.353272035127402	0.265446063898304	371.187896979205	179.549442321733