A study of bootstrapping on developing classification model

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Abstract

This report discusses a study of bootstrap to develop classification model based on the proteins expression levels. Our goal is to build a predictive model based on logistic regression to facilicate down syndrome diagnosis, and we compared methods including and Pathwise Coordinate Descent with regularized logistic regression and smoothed bootstrap estimation. Our result shows that

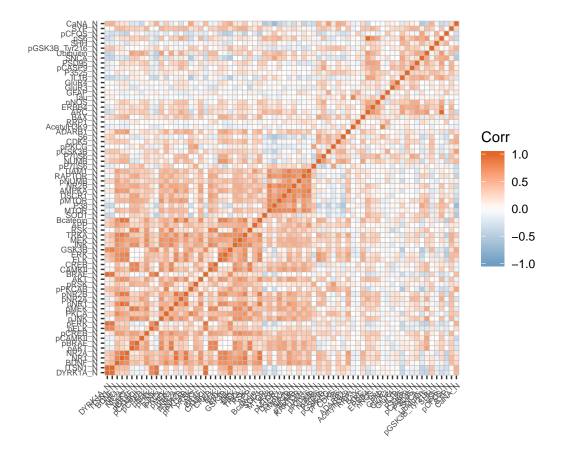
Background

The data *Down.csv* consists of the expression levels of 77 proteins/protein modifications that produced detectable signals in the nuclear fraction of cortex. It has 1080 rows and 79 columns. The first column **MouseID** identifies individual mice; The column **2-78** are values of expression levels of 77 proteins. Column 79 indicates whether the mouse is a control or has Down syndrome. The goal is to develop classification model based on the proteins expression levels.

We found that data is missing for some of the covariates, so we deleted those high missing rate (>20%). For those covariates with missing rate <20%, we assumed them to be missing completely at random(MCAR), so we would analyze the complete cases in the following.

!!!need a missing plot here !!! explain why we need to apply regulized method: singular

Figure 1: missingness plot and multicollinearity plot of the dataset



Method

Logistic Regression

Let y be the vector of n response random variable, X denote the $n \times p$ design matrix $(X_i$ denote the ith row) and β denote the $p \times 1$ coefficient. Let $\eta = E(y) = X\beta$ and given the link function as $g(\eta) = \log \frac{\eta}{1-\eta}$, we have the logistic regression model written as:

$$\log(\frac{\eta}{1-\eta}) = X\beta$$

The likelihood of this logistic regression is:

$$L(\beta; X, y) = \prod_{i=1}^{n} \{ (\frac{\exp(X_i \beta)}{1 + \exp(X_i \beta)})^{y_i} (\frac{1}{1 + \exp(X_i \beta)})^{1 - y_i} \}$$

Maximizing the likelihood is equivalent to maximizing the log likelihood:

$$l(\beta) = \sum_{i=1}^{n} \{y_i(X_i\beta) - \log(1 + \exp(X_i\beta))\}$$

Pathwise Coordinate Descent with regularized logistic regression

Regularization is the common variable selection approaches for high-dimensional covariates. The best known Regularization is called LASSO, which is to add L1-penalty to the objective function. In the context of

logistic regression, we are aiming to maximize the penalized log likelihood:

$$\max_{\beta \in \mathbb{R}^{p+1}} \frac{1}{n} \sum_{i=1}^{n} \{ y_i(X_i \beta) - \log(1 + \exp(X_i \beta)) \} - \lambda \sum_{j=0}^{p} |\beta_j|$$

for some $\lambda \geq 0$. Here the $x_{i,j}$ are standardized so that $\sum_i x_{i,j}/n = 0$ and $\sum_i x_{i,j}^2/n = 0$.

The Newton algorithm for maximizing the log likelihood amounts to interatively reweighted least squares. Hence if the current estimate of the parameter is $\tilde{\beta}$, we can form a quardratic approximation to the negative log likelihood by taylor expansion around the current estimate, which is:

$$f(\beta) = -\frac{1}{2n} \sum_{i=1}^{n} w_i (z_i - \sum_{j=0}^{p} x_{i,j} \beta_j)^2 + C(\tilde{\beta})$$

where

$$z_i = \tilde{\beta}_0 + x_i^T \tilde{\beta} + \frac{y_i - \tilde{p}(x_i)}{\tilde{p}(x_i)(1 - \tilde{p}(x_i))}, \text{ working response}$$

$$w_i = \tilde{p}(x_i)(1 - \tilde{p}(x_i)), \text{ working weights}$$

and $\tilde{p}(x_i)$ is evaluated at the current parameters, the last term is constant. The Newton update is obtained by minimizing the $f(\beta)$

The coordinate descent algorithm to solve the penalized weighted least squares problem

$$\min_{\beta \in \mathbb{R}^{p+1}} \{ -f(\beta) + \lambda P(\beta) \}$$

The above amounts to a sequence of nested loops:

- outer loop: start with λ that all the coefficients are forced to be zerp, then decrement λ ;
- middle loop: update the quardratic $f(\beta)$ using the current estimates of parameters;
- inner loop: run the coordinate descent algorithm on the penalized weighted least square problem.

In our problem, care is taken to avoid coefficients diverging in order to achieve fitted probabilities of 0 or 1 which is the warning message by the R package. When a probability with $\epsilon = 1e - 5$ of 1, we set it to 1, and set the weights to ϵ . 0 is treated similarly.

Smoothing bootstrap and inference

Classical statistical theory ignores model selection in assessing estimation accuracy. Here we consider bootstrap methods for computing standard errors and confidence intervals that take model selection into account. The methodology involves bagging, also known as bootstrap smoothing, to tame the erratic discontinuities of selection-based estimators.

Point estimation:

- First we need to prepare a couple of candidate models
- for each bootstrap in bootstrap with B times, select the best model and get estimates for the coefficient denoted as $t(y^*)$
- smooth $\hat{\mu} = t(y)$ by averaging over the bootstrap replications, defining

$$\tilde{\mu} = s(y) = \frac{1}{B} \sum_{i=1}^{B} t(y^*)$$

Bootstrap smoothing (Efron and Tibshirani 1996), a form of model averaging, is better known as "bagging" in the prediction literature.

smoothed interval In addition to the percentile confidence interval, the nonparametric delta-method estimate of standard deviation for s(y) in the nonideal case

$$\tilde{sd}_B = [\sum_{i=1}^n c\hat{o}v_j^2]^{1/2}$$

where

$$hatcov_j = \sum_{i=1}^{B} (Y_{ij}^* - Y_{.j}^*)(t_i^* - t_.^*)/B$$

with
$$Y_{.j}^* = \sum_{i=1}^B Y_{ij}^*/B$$
 and $t_.^* = \sum_{i=1}^B t_i^*/B = s(y)$.

Results

Pathwise Coordinate Descent Logistic Lasso

Figure 2: Estimation path and cross validation for Pathwise Coordinate Descent Lasso

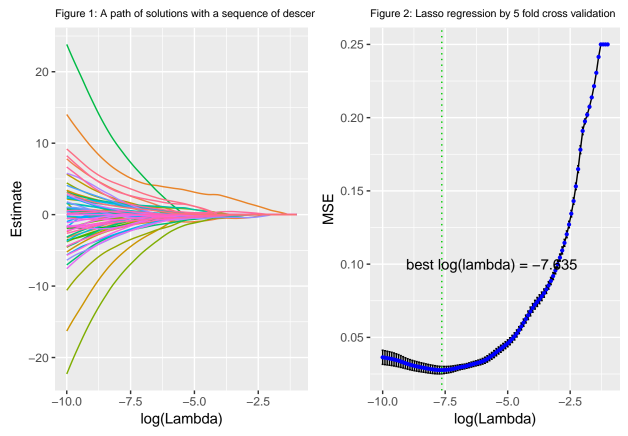


Fig. 2 shows us that as the λ increases, all the variable estimates of parameters shrink accordingly since we penalize all the parameters, though in some. When $\lambda = 0$, the result is the same as least square method and when λ is too large, all the estimates of parameters shrink to 0.

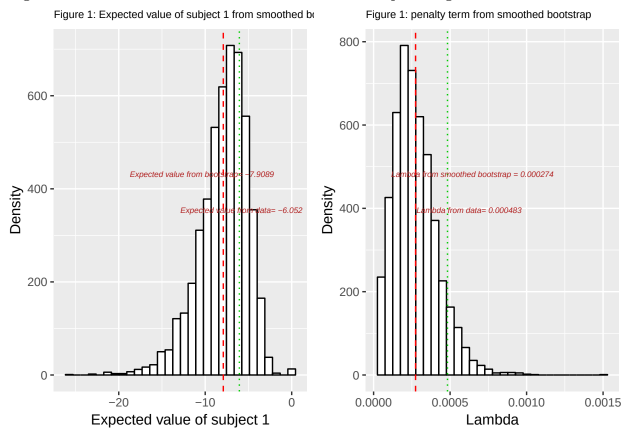
Model selection based on Smoothed Bootstrap for Logistic Lasso

We are still dealing with regularized lasso with this dataset, we wanted to apply smoothed bootstrap in order to get better estimates and inference result.

algorithm:

- bootstrap data from the original dataset
- do cross validation and select the best λ suggest $\lambda = \frac{1}{B} \sum_{i=1}^{B} \lambda_i^{(b)}$ bagging?

Figure 3: Lambda selection based on Smoothed Bootstrap for Logistic Lasso



Cross validation for model prediction comparison

We used 10 fold cross-validation to compare two different models, one is with λ selected from data, the other is selected from the bootstrap.

Table 1: The comparison of performance for two models

| | Misclassification rate | Mean squred error |
|---|------------------------|-------------------|
| Penalty chosen by data Penalty selected from smoothed bootstrap | 0.0322 0.0285 | 0.0239 0.0244 |

^a Dataset: Proteins expression levels of Down syndrome

Significant random variable selection from smoothed bootstrap

for question 4, two method, 1-select covariates based on number of chosen, 2-do hypothesis testing, select significant covariates.

Table 2: Smoothed bootstrap estimation for best model(1)

| | origin | prob | coef | sd | lower | upper | lower.new | upper.new |
|----------------|--------------|------|---------|---------------------|----------|---------|-----------|-----------|
| Intercept | 0.0949 | 1.00 | 0.3143 | 0.3546 | -0.4132 | 1.2144 | -0.3807 | 1.0093 |
| DYRK1A_N | 0.0000 | 0.04 | 0.0310 | 0.1390 | 0.0000 | 0.2384 | -0.2414 | 0.3034 |
| ITSN1_N | 7.9040 | 1.00 | 9.6725 | 2.1862 | 4.4201 | 16.9149 | 5.3875 | 13.9575 |
| BDNF_N | 1.0655 | 0.86 | 1.0811 | 0.7540 | 0.0000 | 3.1864 | -0.3967 | 2.5589 |
| NR1_N | -0.9385 | 0.90 | -2.0857 | 1.2323 | -5.7961 | 0.0000 | -4.5010 | 0.3296 |
| NR2A_N | 0.3742 | 0.54 | 0.4882 | 0.5696 | 0.0000 | 2.5152 | -0.6282 | 1.6046 |
| $pAKT_N$ | 0.6050 | 0.82 | 0.8438 | 0.6632 | 0.0000 | 2.7745 | -0.4561 | 2.1437 |
| $pBRAF_N$ | 0.0000 | 0.62 | 0.1922 | 0.4213 | -0.7271 | 1.5147 | -0.6335 | 1.0179 |
| pCAMKII_N | -0.6756 | 0.93 | -1.0979 | 0.6266 | -2.7897 | 0.0000 | -2.3260 | 0.1302 |
| $pCREB_N$ | 1.2113 | 0.95 | 1.3620 | 0.6984 | 0.0000 | 3.3009 | -0.0069 | 2.7309 |
| pELK_N | -2.1788 | 0.99 | -2.7419 | 1.1203 | -6.1524 | -0.4194 | -4.9377 | -0.5461 |
| $pERK_N$ | -0.2493 | 0.54 | -0.5806 | 0.5946 | -2.7898 | 0.0000 | -1.7460 | 0.5848 |
| $pJNK_N$ | -0.0067 | 0.46 | -0.2080 | 0.3361 | -1.4612 | 0.3040 | -0.8668 | 0.4508 |
| PKCA_N | 1.1962 | 0.82 | 1.2604 | 0.9599 | 0.0000 | 4.0332 | -0.6210 | 3.1418 |
| $pMEK_N$ | 0.0000 | 0.54 | -0.0939 | 0.5630 | -1.8812 | 1.2504 | -1.1974 | 1.0096 |
| pNR1_N | -2.4536 | 0.96 | -2.6193 | 1.2315 | -5.9502 | 0.0000 | -5.0330 | -0.2056 |
| $pNR2A_N$ | 0.0000 | 0.28 | -0.0891 | 0.2675 | -1.1664 | 0.5175 | -0.6134 | 0.4352 |
| $pNR2B_N$ | 1.5568 | 0.90 | 1.8496 | 1.0888 | 0.0000 | 5.1242 | -0.2844 | 3.9836 |
| pPKCAB_N | -1.4553 | 0.94 | -1.4310 | 0.7757 | -3.7448 | 0.0000 | -2.9514 | 0.0894 |
| $pRSK_N$ | -2.4102 | 1.00 | -2.8974 | 0.7656 | -5.2917 | -1.1120 | -4.3980 | -1.3968 |
| AKT_N | 2.7674 | 1.00 | 3.3722 | 0.8799 | 1.2074 | 6.0927 | 1.6476 | 5.0968 |
| BRAF_N | -4.8560 | 1.00 | -5.6899 | 1.7241 | -10.9610 | -1.8681 | -9.0691 | -2.3107 |
| CAMKII_N | -1.5901 | 0.99 | -2.3111 | 0.8889 | -4.9441 | -0.3901 | -4.0533 | -0.5689 |
| CREB_N | -1.2469 | 0.98 | -1.3510 | 0.5539 | -2.9474 | -0.0163 | -2.4366 | -0.2654 |
| ELK_N | -3.6872 | 1.00 | -4.6751 | 1.0545 | -8.0746 | -2.1847 | -6.7419 | -2.6083 |
| ERK_N | -7.4243 | 1.00 | -8.7471 | 1.6856 | -14.5871 | -4.7828 | -12.0509 | -5.4433 |
| $GSK3B_N$ | -0.9853 | 0.88 | -1.5983 | 1.0300 | -4.6946 | 0.0000 | -3.6171 | 0.4205 |
| JNK_N | 0.0000 | 0.56 | -0.1163 | 0.5229 | -1.7227 | 1.2068 | -1.1412 | 0.9086 |
| MEK_N | 1.3308 | 0.98 | 1.6328 | 0.7152 | 0.0194 | 3.6599 | 0.2310 | 3.0346 |
| $TRKA_N$ | 3.7756 | 1.00 | 5.5845 | 2.1337 | 2.0454 | 11.9326 | 1.4024 | 9.7666 |
| RSK_N | -0.6061 | 0.72 | -0.6804 | 0.6776 | -2.6645 | 0.0600 | -2.0085 | 0.6477 |
| APP_N | 5.3514 | 1.00 | 7.8719 | 1.4402 | 5.0187 | 13.0828 | 5.0491 | 10.6947 |
| $Bcatenin_N$ | 0.0000 | 0.21 | 0.1476 | 0.3265 | 0.0000 | 1.6599 | -0.4923 | 0.7875 |
| SOD1_N | 0.4636 | 0.62 | 0.4002 | 0.4175 | 0.0000 | 1.7697 | -0.4181 | 1.2185 |
| $MTOR_N$ | -2.3190 | 0.99 | -2.8751 | 0.9748 | -5.7528 | -0.7263 | -4.7857 | -0.9645 |
| a Poststan tin | F 000 | | | | | | | |

^a Bootstap time=5000

Chosen probability greater than 80% will be considered as important; Chosen probability greater than 96% will be considered as significant(two exceptions); Chosen probability greater than 96% and confidence interval without zero might be a good choice.

Conclusions

References

1 Efron, Bradley. "Estimation and accuracy after model selection." Journal of the American Statistical Association 109.507 (2014): 991-1007.

Table 3: Smoothed bootstrap estimation for best model(2)

| | origin | prob | coef | sd | lower | upper | lower.new | upper.new |
|---------------------|---------|------|---------|---------------------|---------|---------|-----------|-----------|
| P38_N | 0.0000 | 0.44 | -0.1663 | 0.3615 | -1.4422 | 0.6667 | -0.8748 | 0.5422 |
| pMTOR_N | 0.0000 | 0.42 | 0.1135 | 0.3941 | -0.9596 | 1.4826 | -0.6589 | 0.8859 |
| DSCR1_N | 1.2781 | 0.98 | 1.5514 | 0.6235 | 0.0022 | 3.3412 | 0.3293 | 2.7735 |
| AMPKA_N | -0.9954 | 0.93 | -1.4471 | 0.8265 | -3.9602 | 0.0000 | -3.0670 | 0.1728 |
| NR2B_N | -0.1377 | 0.57 | -0.2951 | 0.4053 | -1.6332 | 0.3396 | -1.0895 | 0.4993 |
| pNUMB_N | -0.3484 | 0.74 | -0.7193 | 0.6099 | -2.5185 | 0.0000 | -1.9147 | 0.4761 |
| RAPTOR_N | -1.7634 | 0.96 | -2.1489 | 1.0112 | -4.9061 | 0.0000 | -4.1309 | -0.1669 |
| TIAM1_N | 2.7743 | 1.00 | 3.4095 | 1.0557 | 1.1476 | 6.4593 | 1.3403 | 5.4787 |
| $pP70S6_N$ | 0.4770 | 0.83 | 0.9325 | 0.7488 | 0.0000 | 3.0487 | -0.5351 | 2.4001 |
| NUMB_N | 1.4306 | 0.98 | 1.8409 | 0.8104 | 0.0385 | 4.1741 | 0.2525 | 3.4293 |
| P70S6_N | 0.2829 | 0.67 | 0.4033 | 0.4369 | -0.1974 | 1.8139 | -0.4530 | 1.2596 |
| $pGSK3B_N$ | 0.4776 | 0.71 | 0.3922 | 0.5633 | -0.7969 | 1.9720 | -0.7119 | 1.4963 |
| pPKCG_N | 0.0000 | 0.50 | -0.3093 | 0.4168 | -1.9056 | 0.2988 | -1.1262 | 0.5076 |
| CDK5_N | 0.0000 | 0.51 | 0.0790 | 0.2705 | -0.6702 | 0.9113 | -0.4512 | 0.6092 |
| S6_N | -0.5883 | 0.87 | -0.8153 | 0.5316 | -2.3725 | 0.0000 | -1.8572 | 0.2266 |
| ADARB1_N | 0.1880 | 0.69 | 0.3031 | 0.4307 | -0.5048 | 1.6314 | -0.5411 | 1.1473 |
| AcetylH3K9_N | 0.0000 | 0.33 | -0.1398 | 0.2592 | -1.3143 | 0.2063 | -0.6478 | 0.3682 |
| RRP1_N | -0.8838 | 0.98 | -0.5174 | 1.0229 | -2.2796 | 2.2449 | -2.5223 | 1.4875 |
| BAX_N | 0.0000 | 0.61 | -0.0985 | 0.3247 | -1.1343 | 0.6766 | -0.7349 | 0.5379 |
| ARC_N | -0.5474 | 0.68 | -0.2956 | 0.2802 | -1.3687 | 0.0000 | -0.8448 | 0.2536 |
| ERBB4_N | 1.4902 | 1.00 | 2.0181 | 0.5463 | 0.7995 | 3.6211 | 0.9474 | 3.0888 |
| nNOS_N | -0.6199 | 0.88 | -0.6207 | 0.3817 | -1.7324 | 0.0000 | -1.3688 | 0.1274 |
| Tau_N | 1.5522 | 1.00 | 2.2831 | 0.5607 | 1.0326 | 4.3017 | 1.1841 | 3.3821 |
| GFAP_N | -0.2715 | 0.83 | -0.4706 | 0.4336 | -1.7065 | 0.2579 | -1.3205 | 0.3793 |
| GluR3_N | -1.3348 | 1.00 | -1.7384 | 0.4746 | -3.2161 | -0.7186 | -2.6686 | -0.8082 |
| GluR4_N | 0.1356 | 0.82 | 0.3429 | 0.2973 | -0.0463 | 1.3845 | -0.2398 | 0.9256 |
| IL1B_N | -1.4177 | 0.99 | -1.9549 | 0.6874 | -4.1387 | -0.4795 | -3.3022 | -0.6076 |
| P3525_N | 1.0465 | 0.97 | 1.2122 | 0.5624 | 0.0000 | 2.7394 | 0.1099 | 2.3145 |
| pCASP9_N | -0.1031 | 0.63 | -0.1433 | 0.3663 | -1.2589 | 0.6892 | -0.8612 | 0.5746 |
| PSD95_N | 0.0000 | 0.45 | 0.0777 | 0.3431 | -0.8004 | 1.3061 | -0.5948 | 0.7502 |
| SNCA_N | 0.1480 | 0.75 | 0.4574 | 0.4959 | -0.4088 | 1.9387 | -0.5146 | 1.4294 |
| Ubiquitin_N | 0.9464 | 0.97 | 1.3435 | 0.6419 | 0.0000 | 3.1608 | 0.0854 | 2.6016 |
| $pGSK3B_Tyr216_N$ | 0.0000 | 0.47 | 0.0178 | 0.3412 | -0.9850 | 1.1540 | -0.6510 | 0.6866 |
| SHH_N | -1.5405 | 1.00 | -1.9631 | 0.5122 | -3.6381 | -0.8420 | -2.9670 | -0.9592 |
| $pS6_N$ | -0.1735 | 0.93 | -0.8120 | 0.4846 | -2.3624 | 0.0000 | -1.7618 | 0.1378 |
| pCFOS_N | 0.1882 | 0.76 | 0.3258 | 0.3444 | -0.2652 | 1.3904 | -0.3492 | 1.0008 |
| SYP_N | -0.9364 | 0.99 | -1.2874 | 0.4838 | -2.6677 | -0.1815 | -2.2356 | -0.3392 |
| CaNA_N | 1.7695 | 0.99 | 2.3003 | 0.7909 | 0.4918 | 4.8228 | 0.7501 | 3.8505 |

^a Bootstap time=5000

${\bf Appendix}~{\bf A}$

```
knitr::opts_chunk$set(echo = FALSE, message = FALSE, warning = FALSE, comment = "")
library(tidyverse)
# plot
library(ggcorrplot)
library(ggpubr)
library(grid)
library(gridExtra)
# table
```

```
library(kableExtra)
# lasso
library(glmnet)
# parallel program
library(parallel)
library(doParallel)
library(foreach)
library(iterators)
nCores <- 4
registerDoParallel(nCores)
set.seed(99999)
options(knitr.table.format = "latex")
oridata <- read.csv("Down.csv")</pre>
mydata <- oridata%>%
  dplyr::select(.,-c(BAD_N,BCL2_N,H3AcK18_N,EGR1_N,H3MeK4_N))%>%
  filter(complete.cases(.))
### missing-----
n <- dim(mydata)[1]</pre>
ncol <- dim(mydata)[2]</pre>
list \langle -c(2:(ncol - 1))
namesX <- names(mydata)[-c(1,ncol)]</pre>
# standardize
dataX <- do.call(cbind, lapply(list, function(x) (mydata[,x] - mean(mydata[,x]))/sd(mydata[,x])))</pre>
colnames(dataX) <- namesX</pre>
# design matrix
X <- data.frame(dataX) %>%
 mutate(., intercept = 1)
# response
y <- as.vector(ifelse(mydata[,ncol] == "Down syndrome", 1, 0))
dat <- list(y = y, X = as.matrix(X))</pre>
# check correlation
corplot <- cor(dataX) %>%
  ggcorrplot(.,ggtheme = ggplot2::theme_gray,
             colors = c("#6D9EC1", "white", "#E46726"),
             tl.cex = 6)
grid.arrange(corplot,ncol=1)
### regulized logistic regression: middle loop and inner loop
sfun <- function(beta,lambda) sign(beta) * max(abs(beta)-lambda, 0)
reglogitlasso <- function(lambda, dat, start, tol=1e-10, maxiter = 200,...){
  p <- dim(dat$X)[2]</pre>
  n <- length(dat$y)</pre>
  betavec <- start
  i <- 0
  loglik <- 0
  prevloglik <- Inf
  res <- c(0, loglik, betavec)</pre>
  while (i < maxiter && abs(loglik - prevloglik) > tol && loglik < Inf) {
    i <- i + 1
    prevloglik <- loglik
    for (j in 1:p) {
      u <- dat$X %*% betavec
      expu <- exp(u)
      prob <- expu/(expu+1)</pre>
```

```
w <- prob*(1-prob)</pre>
      w \leftarrow ifelse(abs(w-0) < 1e-5, 1e-5, w)
      z \leftarrow u + (dat\$y-prob)/w
      znoj <- dat$X[,-j] %*% betavec[-j]</pre>
      betavec[j] \leftarrow sfun(mean(w*(dat X[,j])*(z - znoj)), lambda)/(mean(w*dat X[,j]*dat X[,j]))
    loglik <- sum(w*(z-dat$X %*% betavec)^2)/(2*n) + lambda * sum(abs(betavec))
    res <- rbind(res, c(i, loglik, betavec))}</pre>
  return(res)
intial = rep(0, dim(dat$X)[2])
# corres <- reglogitlasso(lambda = exp(-10), dat, start = intial,tol=1e-5)
# start from -1 to -10
## pathwise update: outer loop
path <- function(data,grid){</pre>
  start <- rep(0, dim(data$X)[2])</pre>
  betas <- NULL
  for (x in 1:length(grid)){
    cor.result <- reglogitlasso(lambda = grid[x],dat = data,start= start)</pre>
    lasbeta <- cor.result[nrow(cor.result),3:dim(cor.result)[2]]</pre>
    start <- lasbeta
    betas <- rbind(betas,c(lasbeta))</pre>
  }
  return(data.frame(cbind(grid,betas)))
}
# path.out <- path(dat, grid=exp(seq(-1,-10, length=100)))
# colnames(path.out) <- c("grid", colnames(X))</pre>
path.out <- read.csv("path.out.csv",header = T)%>%
  dplyr::select(.,-c(1))
path.plot <- path.out %>%
  gather(key = par, value = estimate, c(2:dim(path.out)[2])) %>%
  ggplot(aes(x = log(grid), y = estimate, group = par, col = par)) +
  ggtitle("Figure 1: A path of solutions with a sequence of descending lambda's") +
  theme(plot.title = element_text(size = 8),legend.position = "none")+
  xlab("log(Lambda)") +
  vlab("Estimate")
##### 5-fold cross-validation to choose beta lambda
cvresult <- function(dat,grid,K){</pre>
  n <- dim(dat$X)[1]</pre>
  folds <- sample(1:K, n, replace=TRUE)</pre>
  start <- rep(0, dim(dat$X)[2])
  cv.error <- vector()</pre>
  cv.se <- vector()</pre>
  for (x in 1:length(grid)) {
  cv.errors <- vector()</pre>
  for (i in 1:K){
    cor.result <- reglogitlasso(lambda = grid[x],dat = list(X=dat$X[folds!=i,],y=dat$y[folds!=i]),</pre>
                                   start = start)
    lasbeta <- cor.result[nrow(cor.result),3:dim(cor.result)[2]]</pre>
    u <- as.matrix(dat$X[folds == i,]) %*% lasbeta</pre>
    expu <- exp(u)
```

```
prob <- expu / (1 + expu)</pre>
    y <- as.vector(dat$y[folds==i])</pre>
    cv.errors[i] = mean((y-prob)^2)
  start <- lasbeta
  cv.error[x] <- mean(cv.errors)</pre>
  cv.se[x] <- sqrt(var(cv.errors)/K)</pre>
  return(cbind(grid,cv.error,cv.se))
}
\# result \leftarrow cvresult(dat,grid=exp(seq(-1,-10, length=100)),K=5)
# best.lambda <- result[which.min(result[,2]),1]</pre>
best.lambda <- round(0.0004825801 ,6)
result <- read.csv("cvresult.csv",header = T)%>%
  dplyr::select(.,-c(1))
cv.plot <-
    ggplot(result, aes(x=log(result$grid), y=result$cv.error)) +
    geom_errorbar(aes(ymin=result$cv.error-result$cv.se, ymax=result$cv.error+result$cv.se),
                   colour=1) +
    geom_line() +
    geom_point(size=0.8,colour = 4) +
    ggtitle("Figure 2: Lasso regression by 5 fold cross validation")+
    theme(plot.title = element_text(size = 8))+
    xlab("log(Lambda)") + ylab("MSE") +
    geom_vline(xintercept = log(best.lambda),col=3,lty=3) +
    annotate("text", log(best.lambda)+2, 0.1, label = paste("best log(lambda) = ", round(log(best.lambda)+2, 0.1)
grid.arrange(path.plot,cv.plot,ncol=2)
#### final estimation for beta
\# finlasso \leftarrow as.matrix(path(dat,grid = exp(seq(-1,log(best.lambda), length=100))))
# colnames(finlasso) <- c("grid", colnames(X))</pre>
# lasso.beta <- finlasso[nrow(finlasso),2:dim(finlasso)[2]]
temp <- read.csv("coeff.csv",header = T)</pre>
lenb <- dim(temp)[1]</pre>
co.fin.beta <- rbind(temp[lenb,],temp[1:(lenb-1),])</pre>
######### get the estimate and se by smmothing bootstrap
n <- length(y)
p <- dim(dataX)[2]+1</pre>
B <- 5000
taskFun <- function(){</pre>
  # interest in: lambda, u_1(plot), beta, betanum, ynum
  ynum <- vector()</pre>
  betanum <- vector()</pre>
  beta <- vector()</pre>
  bootid <- sample(c(1:n),replace = T)</pre>
  boot.x <- dataX[bootid,]</pre>
  boot.y <- y[bootid]</pre>
  for (j in 1:n) ynum[j] <- sum(j==bootid)</pre>
  min.lambda <- cv.glmnet(x=boot.x, y=boot.y,alpha=1, family="binomial")$lambda.min
  model <- glmnet(x=boot.x, y=boot.y,alpha=1, family="binomial",lambda = min.lambda)</pre>
  beta <- coef(model)</pre>
  for (k in 1:p) betanum[k] <- I(coef(model)[k]!=0)</pre>
  sub1 <- predict(model,dataX)[1] #,type = "response"</pre>
  return(cbind(min.lambda,sub1,t(beta),t(betanum),t(ynum)))
```

```
# out <- foreach(i = 1:B, .combine = rbind) %dopar% taskFun()
# get sparse matrix, transform into data frame
out.df <- read.csv("bootstrap.csv", header = T)%>%
  dplyr::select(.,-c(1))
colnames(out.df) <- c("min.lambda", "subject.1", "intercpt", colnames(dataX),</pre>
                    sprintf("b[%d]",seq(1:p)),sprintf("n[%d]",seq(1:n)))
best.lambda.s <- mean(out.df$min.lambda)</pre>
# plot of expected value of subject 1
model <- glmnet(x=dataX, y=y,alpha=1, family="binomial",lambda = best.lambda)</pre>
sub1 <- predict(model,dataX)[1]#,type = "response"</pre>
s1 <- ggplot(out.df,aes(x=out.df$subject.1)) +</pre>
  geom_histogram(colour="black", fill="white")+
  #qeom_density()
  ggtitle("Figure 1: Expected value of subject 1 from smoothed bootstrap")+
  theme(plot.title = element_text(size = 8))+
  xlab("Expected value of subject 1") + ylab("Density") +
  geom_vline(xintercept = mean(out.df$subject.1),col=2,lty=2) +
  geom_vline(xintercept = sub1,col=3,lty=3)
s2 <- ggplot(out.df,aes(x=out.df$min.lambda)) +</pre>
  geom_histogram(colour="black", fill="white")+
  #qeom_density()
  ggtitle("Figure 1: penalty term from smoothed bootstrap")+
  theme(plot.title = element_text(size = 8))+
  xlab("Lambda") + ylab("Density") +
  geom vline(xintercept = best.lambda.s,col=2,lty=2) +
  geom_vline(xintercept = best.lambda,col=3,lty=3)
grid.arrange(s1+
  annotation_custom(grid.text(paste("Expected value from data= ", round(sub1, 4), sep = ""),
                               x=0.5, y=0.5, hjust=0,
                    gp=gpar(col="firebrick",fontsize=6, fontface="italic")))+
  annotation_custom(grid.text(paste("Expected value from bootstrap= ",
                                     round(mean(out.df$subject.1), 4), sep = ""),
                               x=0.3, y=0.6, hjust=0,
                    gp=gpar(col="firebrick",fontsize=6, fontface="italic"))),
             s2+
  annotation_custom(grid.text(paste("Lambda from data= ", round(best.lambda, 6), sep = ""),
                               x=0.2, y=0.5, hjust=0,
                            gp=gpar(col="firebrick",fontsize=6, fontface="italic")))+
  annotation_custom(grid.text( paste("Lambda from smoothed bootstrap = ",
                                       round(best.lambda.s, 6),sep = ""),
                                 x=0.1, y=0.6, hjust=0,
                            gp=gpar(col="firebrick",fontsize=6, fontface="italic"))),
  ncol=2)
############ cross validation for comparison of prediction error
cvcomp <- function(dat,K){</pre>
  n \leftarrow dim(dat X)[1]
  folds <- sample(1:K, n, replace=TRUE)</pre>
  start \leftarrow rep(0, dim(dat X)[2])
  cv.error.1 <- vector()</pre>
  cv.error.2 <- vector()</pre>
  cv.mse.1 <- vector()</pre>
  cv.mse.2 <- vector()</pre>
```

```
for (i in 1:K){
    cv.x <- dat$X[folds!=i,]</pre>
    cv.y <- dat$y[folds!=i]</pre>
    obs <- glmnet(x=dat$X[folds!=i,],y=dat$y[folds!=i], alpha=1,family="binomial",lambda = best.lambda)
    smooth <- glmnet(x=dat$X[folds!=i,],y=dat$y[folds!=i], alpha=1,family="binomial",lambda = best.lamb</pre>
    y.data <- predict(obs,dat$X[folds == i,],type = "response")>0.5
    y.smooth <- predict(smooth,dat$X[folds == i,],type = "response")>0.5
    y.data2 <- predict(obs,dat$X[folds == i,],type = "response")</pre>
    y.smooth2 <- predict(smooth,dat$X[folds == i,],type = "response")</pre>
    # misclassification rate
    cv.error.1[i] = sum(y.data!=dat$y[folds == i])/length(y[folds == i])
    cv.error.2[i] = sum(y.smooth!=dat$y[folds == i])/length(y[folds == i])
    cv.mse.1[i] = mean((dat$y[folds == i]-y.data2)^2)
    cv.mse.2[i] = mean((dat$y[folds == i]-y.smooth2)^2)
  return(list(error = cbind(mean(cv.error.1), mean(cv.error.2)),
              mse = cbind(mean(cv.mse.1),mean(cv.mse.2))))
}
cvresult <- rbind(round(cvcomp(dat,K=10)$error,4),round(cvcomp(dat,K=10)$mse,4))</pre>
colnames(cvresult) <- c("Penalty chosen by data", "Penalty selected from smoothed bootstrap")</pre>
rownames(cvresult) <- c("Misclassification rate", "Mean squred error")</pre>
kable(t(cvresult), "latex", caption = "The comparison of performance for two models", booktabs = T) %>%
  kable_styling(latex_options = c("hold_position", "scale_down")) %>%
  add_footnote(c("Dataset: Proteins expression levels of Down syndrome"),
               notation = "alphabet")
## select beta based on number of chosen
# colnames(out.df) <- c("min.lambda", "subject.1", "intercpt", colnames(dataX),
                      sprintf("b[\%d]", seq(1:p)), sprintf("n[\%d]", seq(1:n)))
betanum <- out.df[,c((2+p+1):(2+p+p))]
betachosen <- round(t(apply(betanum,2,mean)),2)</pre>
## inference-hypothesis testing
betacoeff <- out.df[,c((2+1):(2+p))]
ynum <- out.df[,c((2+2*p+1):(2+2*p+n))]
meany <- apply(ynum, 2, mean)</pre>
coef <- round(apply(betacoeff, 2, mean),4)</pre>
sd <- vector()
for (k in 1:p) {
  covj <- (t(ynum)-(as.vector(meany)%*%t(rep(1,B))))%*%(betacoeff[,k]-rep(coef[k],B))</pre>
  sd[k] <- round(sqrt((t(covj)%*%covj))/B,4)</pre>
coeff <- cbind(coef,sd)</pre>
newcf <- round(cbind(lower.new = coef -1.96*sd,upper.new = coef +1.96*sd),4)</pre>
cf <- round(cbind(lower = apply(betacoeff,2,quantile,0.025),</pre>
                  upper = apply(betacoeff,2,quantile,0.975)),4)
est.result <- cbind(origin =round(as.vector(co.fin.beta[,2]),4),</pre>
                     prob = as.vector(betachosen),
                     coeff,cf,newcf)
rownames(est.result) <- c("Intercept", colnames(dataX))</pre>
kable(est.result[c(1:35),], "latex", caption = "Smoothed bootstrap estimation for best model(1)", bookt
  kable_styling(latex_options = c("hold_position", "scale_down"),
                 font_size = 6,full_width = F) %>%
  add_footnote(c("Bootstap time=5000"),
```