Implementation of Sparse-Group Lasso Methods on Gene Expression Dataset

A useful R package

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1. THEOREM OVERVIEW

Our data consist of an n response vector y, and an n by p matrix of features X. In many recent applications, we have p >> n: a case where standard linear regression fails. It finds a solution with few nonzero entries in β . In Group Lasso, we suppose our predictor variables were divided into m different groups. We focus on the "sparse-group lasso", we look for beta to minimize

$$\hat{M}(\beta) = \frac{1}{2t} ||\beta - (\beta_0 - t\nabla(r_{(-k)}, \beta_0)) + \frac{1}{2t} ||\beta - \beta_0||_2^2$$
(1)

where $\alpha \in [0,1]$, a convex combination of the lasso and group-lasso penalties, and $r_{(-k)} = y - X^{(l)}\beta^{(l)}$ Combining the subgradient conditions with basic algebra, we get that $\beta = 0$ if

$$||S(\beta_0 - t\nabla l(r_{(-k)}, \beta_0, t\alpha\lambda)||_2 \le t(1 - \alpha)\lambda.$$
(2)

and otherwise β satisfies

$$(1 + \frac{t(1-\alpha)\lambda}{\|\hat{\beta}\|_2})\hat{\beta} = S(\beta_0 - t\nabla l(r_{(-k)}, \beta_0), t\alpha\lambda)$$
(3)

Taking the norm of both sides, we see that our generalized gradient step (i.e., the solution to Equation (10)) is

$$||\hat{\beta}||_2 = (||S(\beta_0 - t\nabla l(r_{(-k)}), \beta_0), t\alpha\lambda)||_2 - t(1 - \alpha)\lambda)_+$$
(4)

If we plug this into Equation (11), we see that our generalized gradient step (i.e., the solution to Equation (10)) is

$$\hat{\beta} = (1 - \frac{t(1-\alpha)\lambda}{\|S(\beta_0 - t\nabla l(r_{(-k)}), \beta_0), t\alpha\lambda)\|_2}) + S(\beta_0 - t\nabla l(r_{(-k)}), \beta_0), t\alpha\lambda)$$
(5)

If we iterate Equation (12), and recenter each pass at $(\beta_0)_{new} = (\beta_0)_{old}$, then we will converge on the optimal solution for $\hat{\beta}^{(k)}$ given fixed values of the other coefficient vectors. If we apply this per block, and cyclically iterating through the blocks we will converge on the overall optimum. For ease of notation in the future, we let $U(\beta_0, t)$ denote our updated formula

$$U(\beta_0, t) = \left(1 - \frac{t(1 - \alpha)\lambda}{||S(\beta_0 - t\nabla l(r_{(-k)}), \beta_0), t\alpha\lambda)||_2}\right) + S(\beta_0 - t\nabla l(r_{(-k)}), \beta_0), t\alpha\lambda)$$
(6)

Note that in our case (linear regression)

$$\nabla l(r_{(-k)}, \beta_0) = -X^{(k)T} r_{(-k)} / n \tag{7}$$

2. ALGORITHM OVERVIEW

- 1. (Outerloop) Cyclically iterate through the groups; at each group(k) execute Step2.
- 2. Check if the group's coefficients are identically 0, by seeing if they obey

$$||S(X^{(k)T}r_{(-k)}, \alpha\lambda)||_2 \le (1-\alpha)\lambda \tag{8}$$

- 3. If not, within the group apply Step 2.
- 4. (Inner loop) Until convergence iterate:

Start with $\beta^{(k,l)} = \theta^{(k,l)} = \beta_0^{(k)}$, step size t=1, and counter l=1. Repeat 0. The following until convergence:

- 4.1 Update gradient g by $g = \nabla l(r_{(-k)}, \beta^{(k,l)});$
- 4.2 Optimize step size by iterating t = 0.8 * t until

$$l(U(\beta^{(k,l)},t)) \le l(\beta^{(k,l)}) + g^T \Delta_{(l,t)} + \frac{1}{2t} ||\Delta_{(l,t)}||_2^2$$
(9)

4.3 Update $\theta^{(k,l)}$ by

$$\theta^{(k,l+1)} \leftarrow U(\beta^{(k,l)},t)$$

4.4 Update the center via a Nesterov step by

$$\beta^{(k,l+1)} \leftarrow \theta(k,l) + \frac{l}{l+3} (\theta(k,l+1) - \theta(k,l)) \tag{10}$$

4.5 Set l = l + 1; where Δ is the change between our old solution and new solution

$$\Delta_{(l,t)} = U(\beta^{(k,l)}, t) - \beta^{(k,l)}.$$

Algorithm 1 OneDim

```
Require: Matrix X, Matrix y
Ensure: \beta
 1: procedure Initialization(X,y)
       Use West algorithm to calculate mean and variance,
       Set y'=y-mean to avoid calculating the intercept.
 3:
       Return scale as a vector.
 4:
 5: end procedure
 6:
 7: function BETTERPATHCALC(X, y)
       Calculate Lambda
 8:
       return Lambda
 9:
10: end function
11:
12:
   function LINNEST(OuterLoop)
       while (do groupChange==1)
                                                              ▶ to check if we need to keep computing beta
13:
           groupChange=0;
14:
       end while
15:
       while (outer limitation: outermostCounter < outerIter[0] && outermostCheck < outerThresh[0]) do
16:
          linSolver;
                                                         \triangleright set constraints to execute linSolver and update \beta
17:
       end while
18:
19: end function
20:
21: function LINSOLVER(inner loop)
22:
       for i \leftarrow 0, np do
                                                                                            ▶ for each group
23:
           compute gradient grad;
           if (zeroCheck \le pow(lambda2, 2) * lp) then
                                                                          ▷ check if this group is zero group
24:
25:
                                                                         ▷ start inner loop to compute beta
26:
           else
              while (innerlimitation : count <= innerIter \&\&check > thresh) do
27:
                 set the constraints to update \beta by innerIter and innerThresh
28:
                 norm=|z|;
29:
30:
                 compute gradient U;
                 diff=-1;
31:
32:
                 check the inequation:
                 diff=RHS-LHS;
33:
34:
                  while diff < 0 do
                     continue the loop
35:
36:
                     t=t*g; get new diff;
                                                                                         ⊳ optimize step size
                 end while
37:
38:
                  update \theta by U;
                  update beta by \theta and U by Nesterov-style momentum
39:
              end while
40:
           end if
41:
       end for
42:
43: end function
```

variable	comment
eta	$r_{(-k)} = y - X^{(l)}\hat{\beta}^{(l)}$
grad	the Gradient
zeroCheck	to check if group is zero group
betaIsZero	indicator for beta is zero
${\rm rangeGroupInd}[i]$	the position for the first element in group i
theta	to store the updated beta
U	updated formula to compute beta
t	step size
1	counter
np	number of groups
lp	length of each group
n	number of observation
${\rm group Change}$	check if beta is ok, otherwise need to do linSolver
isActive	indicator if groupChange is 1;

Table 1: Other Definitions

Algorithm 2 Supplimental Functions

Require: Matrix X, Matrix y

Ensure: β

- 1: **function** LINGRADCALC(X, y)
- 2: compute $ldot = -r_{(-k)}/n$
- 3: end function
- 4: function LinNegLogLikelihoodCalc(X, y)
- 5: compute unpenalized loss function
- 6: end function

3. SIMULATION IN C++

We implemented the algorithm in paper in C++ with Matrix615.h and did simulations with randomly generated data. Our project is written by xcode on macbook pro. It can read data from file or use randomly generated data to calculate $\hat{\beta}$, and predict y. It also output R^2 and RMSE, with which we can test the accuracy of prediction.

Output of C++ Example estimated beta= -8.83592 -1.19675 4.07672 - 8.518446.68252 1.2149 -6.05425 -2.07061 3.66666 -8.41446-8.23652 -1.48738 -7.90107 3.58552 -4.16999 230803 -1.96936 -2.30416 -4.27065 -4.196243.42028 - 1.13221 $3.9326 \quad 3.78405 \quad -8.59944 \quad 4.62309 \quad 3.34582 \quad -2.03059 \quad 6.10709 \quad 1.84812$ 1.14444 4.66506 1.3533 -4.11551 1.42855 2.72132 -4.51502 9.08728 -2.667841.72567-3.81311 -3.65004 1.54093 2.49248 1.25922 4.44533 1.6477 -2.01475 -1.477987.66825 $2.69199 \quad 3.36841 \quad 1.3401 \quad -9.23822 \quad 7.84053 \quad -3.32986 \quad 2.00972 \quad -4.95939 \quad 2.09015 \quad -7.63303 \quad -7.63300 \quad -7.63300 \quad -7.63300 \quad -7.63300 \quad -7.63300 \quad -7.63$ $-1.51501 \quad -1.30075 \quad 1.73774 \quad -1.01883 \quad -1.35253 \quad 1.07117 \quad -4.37588 \quad -3.28805 \quad 1.25195 \quad -4.11551888 \quad -3.28805 \quad 1.25195 \quad -4.11551888 \quad -3.28805 \quad 1.25195 \quad -4.11551888 \quad -3.28805 \quad -3.28$ ****** Predicting value y... fitted y: -0.110665 0.435365-1.454465 -0.7685620.543765R-square = 0.9473rmse = 0.88622total time: 0.000663s

In the SGL R-package from paper, the author did not calculate mean and variance in standardization step. We write a cpp edition which has much better efficiency.

Comparison of standardization code

```
library(microbenchmark)

> cppFunction('int stdize(NumericMatrix x) {
+ double mean=x(0,0), var=0;
+ int count=2;
+ for(int i=0;i<x.ncol();i++) {
+ for(int j=0;j<x.nrow();j++) {
+ if(i! = 0||j! = 0) {
+ var+=1.0*(count-1)count*(x(j,i)-mean)*(x(j,i)-mean);
+ mean=mean+(x(j,i)-mean)/count;
+ }
+ count++;
```

```
+ }
+ \text{var} = \text{sqrt}(\text{var}(\text{x.nrow}()*\text{x.ncol}()-1));
+ \text{ for(int } k=0; k< x.nrow(); k++)
+ x(k,i)=(x(k,i)-mean)/var;
+ }
+ \operatorname{return}(x);
+ }')
> origin=function(x)\{ X < -x \}
+ \text{ means} < - \text{ apply}(X,2,\text{mean})
+ X < - t(t(X) - means)
+ \text{ var } < - \text{ apply}(X, 2, \text{function}(x)(\text{sqrt}(\text{sum}(x^*x))))
+ X < - t(t(X) / var)
+ x < - X
+ X.transform < - list(X.scale = var, X.means = means)
> x = matrix(sample (100), 10)
> microbenchmark(stdize(x), origin(x))
Unit: microseconds
expr
             min
                                                       median
                          lq
                                        mean
                                                                                         neval
                                                                      uq
                                                                               max
stdize(x)
             2.968
                          3.2160
                                       5.23288
                                                      4.0525
                                                                    5.537
                                                                               24.139
                                                                                             100
             101.172
                          107.1185
origin(x)
                                       133.48869
                                                      111.7765
                                                                    139.781
                                                                                375.977
                                                                                            100
```

4. TIMING

The time complexity of our algorithm is $O(n^2p)$ where n is the data number and and p is the number of predictors.

In the timing simulation step, we apply the data generated from the paper, but we compare our model and the original code instead. "New" is the timing of our model while "origin" is the original model in the paper. We can see clear improvement in our model From Table 2.

Short path					
method	1 group	2 groups	3 groups		
n = 150, p = 1500, m = 10					
new	0.0341	0.08	0.0913		
origin	1.02	1.031	1.052		
n = 200, p = 2000, m = 200					
new	0.0543	0.0462	0.0498		
origin	0.4307	0.4367	0.4032		
n = 150, p = 10,000, m = 100					
new	0.5329	0.4625	0.5197		
origin	1.277	1.171	1.263		

Table 2: Short path simulation

Long path						
method	1 group	2 groups	3 groups			
n = 150, p = 1500, m = 10						
new	0.2602	0.9926	1.227			
origin	1.138	1.757	2.325			
n = 200, p = 2000, m = 200						
new	0.0882	0.1384	0.2118			
origin	1.188	1.154	1.534			
n = 150, p = 10,000, m = 100						
new	1.188	1.188	1.188			
origin	2.3	2.74	3.202			

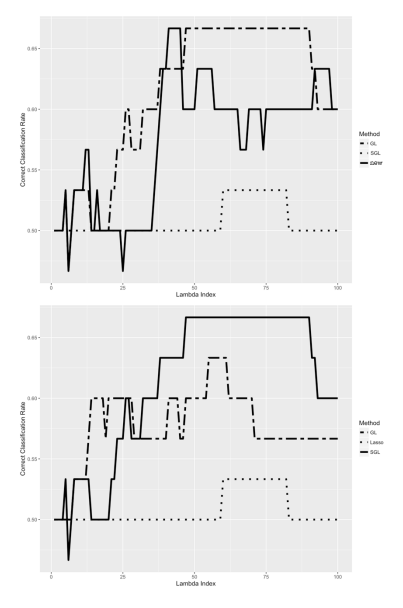
Table 3: Long path simulation

5. ACCURACY

The real dataset we used for comparisonwas the breast cancer data by Ma et al. (2004). This dataset contains gene expression values from 60 patients with estrogen positive breast cancer. The patients were treated with tamoxifen for 5 years and classified according to whether cancer recurred (there were 28 recurrences). Gene expression values were run on a GPL1223: Arcturus 22k human oligonucleotide microarray. Unfortunately, there was significant missing data. As a first pass, all genes with more than 50% missing data were removed. Other missing values were imputed by simple mean imputation. This left us with 12, 071 of our 22, 575 original genes. We again grouped genes together by cytogenetic position data, removing genes that were not recorded in the GSEA C1 dataset. Our final design matrix had 4989 genes in 270 pathways (an average of 18.5 genes per pathway). Thirty patients were chosen at random and used to build the three models. The remaining 30 were used to test their accuracies.

Referring to the second figure below, we see that in this example the sparse-group lasso outperforms the lasso and group lasso. The sparse-group lasso reaches 70% classification accuracy (though this is a narrow peak, so may be slightly biased high), while the group lasso peaks at 60% and the lasso comes in last at 53% accuracy. At its optimum the sparse-group lasso includes 54 genes from 11 bands, while the group lasso selects all 74 genes from 15 bands (again, largely smaller bands for the group lasso), and the lasso selects three genes all from separate bands. This example really highlights the advantage of the sparse-group lasso, it allows us to use group information, but does not force us to use entire groups.

The first figure compare the new method and the original model. We don't have a very stable result but on some interval, we can still have high enough accuracy. These two examples highlight two different possibilities for the sparse-group lasso. In the breast cancer data, the addition of group information is critical for classification, and the grouping may help give insight into the biological mechanisms. However, we should note that , the group whose "information" is largely just increases model variance the sparse-group lasso is certainly not perfect for every scenario with grouped data, but as evidenced in the cancer data it can sometimes be helpful.



REFERENCES

Simon N, Friedman J, Hastie T, et al. A sparse-group lasso[J]. Journal of Computational and Graphical Statistics, 2013, 22(2): 231-245.