PRIMAL: An Linear Programming-based Sparse Learning Library in R and Python

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1 Introduction

Linear Programming (LP) based sparse learning methods, such as the Dantzig selector (for linear regression) (Candes et al., 2007), sparse quantile regression (Belloni et al., 2011), sparse support vector machines (Wang, 2013), have been widely used in machine learning for high dimensional data analysis (Belloni et al., 2011; Hudson et al., 2009; Bandyopadhyay et al., 2010). Despite of their popularity, their software implementations are quite limited. We describe a new library in both R and Python – PaRametric sImplex Method for spArse Learning (PRIMAL), for the aforementioned LP-based sparse learning methods with the following two key features: 1) It provides a highly efficient optimization engine based on the parametric simplex method, which can efficiently solve large scale sparse learning problems; 2) Besides the estimation procedures, it provides additional functional modules such as data-dependent model selection and model visualization.

2 Background

Many popular sparse learning methods involve solving an optimization problem. Here we are interested in those that can be cast into the following linear programs,

$$\max(c + \lambda \mathbf{1})^T x$$
, subject to $Ax \le b + \lambda \mathbf{1}, x \ge 0$. (1)

where A, b, and c are known variables, λ is a tuning parameter. There are several general methods for solving Linear Programs (LP). In particular, Interior point method (Mehrotra, 1992) is proven to solve LPs in polynomial time, however, its total computational cost is cubically dependent on the scale of the problem, which is not scalable in high dimensions. Moreover, the computation at every iteration cannot take the advantage of the sparsity in the underlying models to boost the computation. The reason behind is that interior point method uses the log barrier to handle the constraints, thus, cannot yield sparse iterates. On the other hand, simplex method has stood the

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test of various practical problems by efficiently finding optimal solutions, though its worst-case iteration complexity has been shown to scale exponentially with the problem scale in existing literature.

These methods, though popular, are usually designed for solving (1) for one single tuning parameter. This is not satisfactory, since an appropriate choice of is usually unknown. Thus, one usually expects an algorithm to obtain multiple solutions tuned over a reasonable range of values for λ . For each value of λ , we need to solve a linear program from scratch, and it is therefore often very inefficient for high dimensional problems.

To overcome the above drawbacks, we adopt a variant of Simplex method — Parametric Simplex Method (PSM) (Pang et al., 2017) as the core engine to efficiently solve LP-based sparse learning problems. PSM parametrizes (1) using the unknown regularization factor as a "parameter". This eventually yields a piecewise linear solution path for a sequence of regularization factors. PSM relies some special rules to iteratively choose the pair of variables to swap, which algebraically calculates the solution path during each pivoting. PSM terminates at a value of parameter, where we have successfully solved the full solution path to the original problem. Although in the worst-case scenario, PSM can take an exponential number of pivots to find an optimal solution path. The results (Pang et al., 2017) suggest that the number of iterations is roughly linear in the number of nonzero variables for large regularization factors with sparse optima. This means that the desired sparse solutions can often be found using very few pivots.

3 Algorithm and Package Design

The PSM engine is the core optimization engine that aims to solve the LP problem with regularization parameters efficiently. Unlike the traditional generic LP solver, the PSM engine is specially designed for statistical methods that has a regularization parameter that requires tuning. Therefore, the PSM engine handles the tuning process efficiently. Furthermore, in order to achieve efficient and reliable matrices and vectors computation, we exploit Eigen library, an open-source C++ library for linear algebra.

More specifically, the PSM algorithm is briefly described as follows:

- Initialize $\lambda = (|\min_j c_j|, |\min_j b_j|)$, which is large enough such that x = 0 is an optimal solution to (1). Let $\lambda^* = \lambda$, λ_{\min} denote the minimal regularization parameter of our interest.
- While $\lambda^* > \lambda_{\min}$ do
 - Decrease λ until x violates the constraints, i.e., x is infeasible. Denote by λ^* the threshold that breaks the feasibility of x.
 - Update x by swapping a nonzero variable with a zero variable as in the simplex method. More precisely, select the variable that attains λ^* as the entering variable, which becomes nonzero afterwards; select the variable that first violates the constraint as the leaving variable, which becomes zero.

As can be seen, we are essentially keeping the iterates to be the optimal solution corresponding to different values of λ . As a result, PSM naturally takes the advantage of the regularization

parameter tuning and obtains the optimal solution path in one run. Note that the subproblems involved in each iteration of the above algorithm – for determining the decreased value of λ and for determining the update for x– both admit closed-form intermediate solutions. Also, the update in last line only involves part of the variables and yields sparse iterates, and the computational cost can be significantly reduced by using sparse matrix vector multiplication. Therefore, the algorithm has a low per-iteration cost. Furthermore, although in the worst-case scenario, PSM can take an exponential number of operations to find an optimal solution path, empirical results suggest that the number of iterations is roughly linear in the number of nonzero variables for large regularization factors with sparse optima. This implies that the desired sparse solution can be found in very few iterations – roughly speaking O(s), where s is the number of nonzero entries in the optimal solution. For sparse learning problems, s is usually much smaller than the number of variables. As a result, the algorithm is highly efficient.

We have four basic modules in our package design: the core optimization engine using parametric simplex method, visualization engine, and application engine.

PSM engine. The PSM described in the previous section is implemented in a pure C/C++ fashion. The code is supported by Eigen library, which is an efficient headers only C++ linear algebra library.

Application engine. The user interface for Dantzig selector, sparse support vector machine, sparse compressive sensing and quantile regression is implemented. We also include the sample code for the applications in the package.

Visualization engine. Visualization functions for visualizing solution path is implemented.

4 Examples

Now we illustrate the user interface using Danzig selector as an example.

4.1 R user Interface

The following example solves

$$\min_{\beta} \|\beta\|_{1} \quad \text{subject to } \|X^{T}(y - X\beta)\|_{\infty} \le \lambda$$
 (2)

for any λ .

```
> library(PRIMAL)
> # Generate a synthetic dataset: the design matrix and coefficient vector
> n = 100 # number of samples
> d = 250 # sample dimension
> c = 0.5 # correlation parameter
> s = 20 # support size of coefficient
> set.seed(1024)
> X = scale(matrix(rnorm(n*d),n,d)+c*rnorm(n))/sqrt(n-1)*sqrt(n)
> flag = runif(s,-1,1)
```

```
> beta1 <- c()
> for(i in 1:s){
+ if(flag[i]>=0) beta1[i]=rnorm(1,1,1)
+ if(flag[i]<0) beta1[i]=rnorm(1,-1,1)
+ }
> beta = c(beta1, rep(0, d-s))
> # Generate response using Gaussian noise
> noise = rnorm(n)
> Y = X%*%beta + noise
>
> # Dantzig selection solved with parametric simplex method
fit.dantzig = Dantzig_solver(X, Y, max_it = 100, lambda_threshold = 0.01)
This line will return a list containing the information of the solution. From fit.dantz
```

This line will return a list containing the information of the solution. From fit.dantzig, we can recover solutions corresponds to any regularization parameter λ .

```
> # Now let's see the regularization parameters along the path
> print(fit.dantzig$lambda)
  [1] 195.22461 176.80544 135.96638 116.72507 104.60908 102.70803
      89.74968 82.94148
                          78.33078
                                   75.14407
                                              74.65001
                                                        64.26936
 [13]
      58.49683 53.85248
                          49.89716
                                    49.71839
                                              47.68773
                                                        42.22191
 [19]
      40.88262 40.79184
                                    38.71244
                                              38.07740
                                                        37.72982
                          38.71504
 [25]
      37.72246
               37.51079
                          37.47705
                                    37.31156
                                              37.09234
                                                        36.65785
 [31]
      36.51983 35.61435
                          34.75857
                                    34.47443
                                              34.22125
                                                        34.16291
                                    33.40794
 [37]
      34.08827 33.72415
                          33.62017
                                              33.33360
                                                        33.10474
 [43]
      32.65156 32.26433
                          29.46930
                                    28.98127
                                              28.01981
                                                        27.95761
 [49]
       27.27683 27.01244
                          26.16928
                                    26.15484
                                              26.14264
                                                        26.05776
 [55]
      25.90875
                25.58693
                          24.42879
                                    24.39550
                                              24.33576
                                                        24.16818
 [61]
      24.06451 23.96691
                          23.70723
                                    23.40995
                                              23.27554
                                                        22.48808
 [67]
       22.08780 21.24536
                          20.62978
                                    20.43703
                                              20.43107
                                                        20.40643
 [73]
       19.56953
                19.39416
                          19.29465
                                    18.91443
                                              18.81399
                                                        18.41681
 [79]
                          18.04614
       18.37950
                18.30446
                                    17.93162
                                              17.79798
                                                        17.70347
 [85]
       17.68071
                17.66381
                          16.97636
                                    16.67124
                                              16.55067
                                                        16.41163
 [91]
       15.36333
                15.25158
                          15.10068
                                    15.06699
                                              15.05869
                                                        14.92861
 [97]
       14.82495
                14.39728
                          14.32287
                                     14.05490
```

For λ larger than the largest value listed above, i.e., 195.22, the optimal solution of (2) is $\beta = 0$. For λ smaller than 195.22 larger than lambda_threshold, the optimal solution is the same as the smallest value listed above that is larger than λ . For example, consider $\lambda = 24$, its solution is the same as $\lambda = 24.06$ in entry 61. We can get the corresponding β by calling fit.dantzig\$beta[,61], and the corresponding value of objective function by calling fit.dantzig\$value[61].

Moreover, we can visualize the regularization path by calling plot.primal(fit.dantzig). The plots are shown in Figure 1.

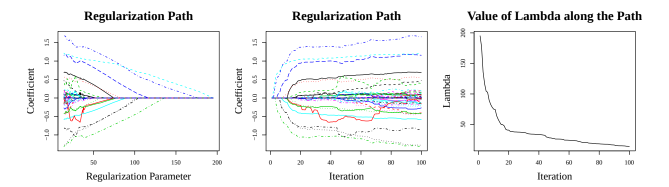


Figure 1: Visualizations of the solution path.

4.2 Python User Interface

Now we use the same example to illustrate the Python user interface.

```
> # load packages
> import numpy as np
> from sklearn.preprocessing import scale
> import pypsm
> from pypsm import Dantzig
> # Generate the design matrix and regression coefficient vector
> n = 100 # sample number
> d = 80 \# sample dimension
> c = 0.5 # correlation parameter
> s = 20  # support size of coefficient
> X = scale(np.random.randn(n,d)+c* np.tile(np.random.randn(n),[d,1]).T)/(n*(n-1))**0.5
> beta = np.append(np.random.rand(s), np.zeros(d-s))
> # Generate response using Gaussian noise, and fit sparse linear models
> noise = np.random.randn(n)
> y = np.matmul(X,beta) + noise
> # fit the model
> solver = Dantzig(X, y)
> solver.train()
result = solver.coef()
solver.plot()
```

The result can be similarly obtained by calling solver.coef()['lambda_list'], solver.coef()['theta_list'] and solver.coef()['target_list']

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