An Introduction to islasso

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Abstract

In this short note we present and briefly discuss the R package islasso dealing with regression models having a large number of covariates. Estimation is carried out by penalizing the coefficients via a quasi-lasso penalty, wherein the nonsmooth lasso penalty is replaced by its smooth counterpart determined iteratively by data according to the induced smoothing idea. The package includes functions to estimate the model and to test for linear hypothesis on linear combinations of relevant coefficients. We illustrate R code throughout a worked example, by avoiding intentionally to report details and extended bibliography.

Introduction

Let $\mathbf{y} = \mathbf{X}\beta + \epsilon$ be the linear model of interest with usual zero-means and homoscedastic errors. As usual, $\mathbf{y} = (y_1, \dots, y_n)^T$ is the response vector, \mathbf{X} is the $n \times p$ design matrix (having p quite large) with regression coefficients β . When interest lies in selecting the non-noise covariates and estimating the relevant effect, one assumes the lasso penalized objective function (Tibshirani, 1996),

$$\frac{1}{2}||\mathbf{y} - \mathbf{X}\boldsymbol{\beta}||_2^2 + \lambda||\boldsymbol{\beta}||_1$$

The R functions

The main function of the package is islasso() where the user supplies the model formula as in the usual lm or qlm functions, i.e.

```
islasso(formula, family = gaussian, lambda, alpha = 1, data, weights, subset, offset,
    unpenalized, contrasts = NULL, control = is.control())
```

family accepts specification of family and link function as in Table 1, lambda is the tuning parameter and unpenalized allows to indicate covariates with unpenalized coefficients.

Table 1. Families and link functions allowed in islasso

family	link
gaussian	identity
binomial	logit, probit
poisson	\log
gamma	identity, \log , inverse

The fitter function is is.lasso.fit() which reads as

```
islasso.fit(X, y, family = gaussian, lambda, alpha = 1, intercept = FALSE, weights = NULL,
    offset = NULL, unpenalized = NULL, control = is.control())
```

which actually implements the estimating algorithm as described in the paper. The lambda argument in islasso.fit and islasso specifies the positive tuning parameter in the penalized objective. Any non-negative value can be provided, but if missing, it is computed via K-fold cross validation by the function cv.glmnet() from package glmnet. The number of folds being used can be specified via the argument nfolds of the auxiliary function is.control().

A worked example: the Diabetes data set

We use the well-known diabetes dataset available in the lars package. The data refer to n=442 patients enrolled to investigate a measure of disease progression one year after the baseline. There are ten covariates, (age, sex, bmi (body mass index), map (average blood pressure) and several blood serum measurements (tc, ldl, hdl, tch, ltg, glu). The matrix x2 in the dataframe also includes second-order terms, namely first-order interactions between covariates, and quadratic terms for the continuous variables.

To select the important terms in the regression equation we apply the lasso

```
library(lars)
library(glmnet)
data("diabetes", package = "lars")
a1 <- with(diabetes, cv.glmnet(x2, y))
n <- nrow(diabetes)</pre>
a1$lambda.min * n
> [1] 1344.186
b <- drop(coef(a1, "lambda.min", exact = TRUE))</pre>
length(b[b != 0])
> [1] 15
Ten-fold cross validation "selects" \lambda = 1344.186. corresponding to 15 non null coefficients
names(b[b != 0])
   [1] "(Intercept)" "sex"
                                       "bmi"
                                                                       "hdl"
                                                       "map"
   [6] "ltg"
                                       "age^2"
                                                       "bmi^2"
                                                                       "glu^2"
> [11] "age:sex"
                                       "age:ltg"
                                                                       "bmi:map"
                        "age:map"
                                                       "age:glu"
The last six estimates are
tail(b[b != 0])
       glu<sup>2</sup>
                  age:sex
                              age:map
                                           age:ltg
                                                       age:glu
                                                                   bmi:map
   69.599081 107.479925
                            29.970061
                                         8.506032
                                                    11.675332
                                                                 85.530937
```

A reasonable question is if all the "selected" coefficients are significant in the model. Unfortunately lasso regression does not return standard errors due to nonsmoothness of objective, and some alternative approaches have been proposed., including the (Lockhart et al., 2013). Among the (few) strategies, including the 'covariance test', the 'post-selection inference' and the '(modified) residual bootstrap', here we illustrate the R package **islasso** implementing the recent 'quasi' lasso approach based on the induced smoothing idea (Brown and Wang, 2005) as discussed in Cilluffo et al. (2019)

While the optimal lambda could be selected (without supplying any value to lambda), we use optimal value minimizing the AIC

```
library(islasso)
out <- islasso(y ~ x2, data = diabetes, lambda = a1$lambda.min * n)</pre>
```

The **summary** method quickly returns the main output of the fitted model, including point estimates, standard errors and p-values. Visualizing estimates for all covariates could be somewhat inconvenient, especially when the number of covariates is large, thus we decide to print estimates only if the pvalue is less than a threshold value. We use 0.50

```
summary(out, pval = 0.1)
> Call:
 islasso(formula = y ~ x2, lambda = a1$lambda.min * n, data = diabetes)
>
 Residuals:
     Min
                  Median
>
               1Q
                               3Q
                                      Max
 -126.33 -63.66
                  -11.64
                            58.16
                                   188.83
>
              Estimate Std. Error
                                     Df z value Pr(>|z|)
>
              152.133
                            9.613
                                   1.00
                                          15.83
                                                  <2e-16 ***
 (Intercept)
 Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
 (Dispersion parameter for gaussian family taken to be 5727.516)
>
      Null deviance: 2621009 on 441.0 degrees of freedom
> Residual deviance: 2524119
                             on 440.7
                                        degrees of freedom
 AIC: 5082.3
 Lambda: 1344.2
> Number of Newton-Raphson iterations: 35
```

In addition to the usual information printed by the summary method, the output also includes the column *Df* representing the degrees of freedom of each coefficient. Their sum is used to quantify the model complexity sum(out\$internal\$hi)

> [1] 1.299546

and the corresponding residual degrees of freedom (440.7004542) as reported above. The Wald test (column z value) and p-values can be used to assess important or significant covariates. Results suggest that the value of the minimum lambda choosn by the cross validation procedure of glmnet was too high, hence as an alternative, it is also possible to select the tuning parameter λ by means the Bayesian or Akaike Information Criterion. The function aic.islasso, requires an islasso fit object and specification of the criterion to be used (AIC/BIC). Hence

```
lmb.bic <- aic.islasso(out, method = "bic", interval = c(1, 100))</pre>
```

```
> Optimization through bic
             38.8146 bic = 4911.32373
> lambda =
> lambda =
             62.1854 bic = 4884.06799
> lambda =
            76.6293 bic = 4866.84943
> lambda =
            85.5561 bic = 4864.21083
> lambda =
             84.9453 bic = 4863.96914
> lambda =
             82.8705 \text{ bic} =
                            4865.82101
> lambda =
            84.8383 bic = 4864.50618
> lambda =
             85.2245 bic = 4863.11256
> lambda =
             85.2318 bic = 4861.80825
> lambda =
             85.3557 bic = 4863.60867
> lambda =
             85.2888 bic = 4862.68798
> lambda =
             85.2577 \text{ bic} = 4862.22825
> lambda =
             85.2417 bic = 4863.60663
> lambda =
             85.2356 bic = 4861.80777
> lambda =
             85.2338 bic = 4861.80800
> lambda =
             85.2379 bic = 4861.80746
> lambda =
             85.2393 \text{ bic} = 4861.80727
> lambda =
            85.2402 bic = 4864.61415
> lambda =
            85.2388 bic = 4861.80734
> lambda =
            85.2397 \text{ bic} = 4864.61373
> lambda =
            85.2391 bic =
                            4861.80730
> lambda =
            85.2395 bic = 4861.80726
> lambda =
             85.2396 bic = 4861.80725
> lambda =
             85.2396 bic = 4861.80724
            85.2396 bic = 4861.80724
> lambda =
out1 <- update(out, lambda = lmb.bic)</pre>
summary(out1, pval = 0.05)
>
> Call:
> islasso(formula = y ~ x2, lambda = lmb.bic, data = diabetes)
> Residuals:
       Min
                 1Q
                      Median
                                   3Q
                                           Max
> -135.110 -40.849
                      -5.068
                               35.406 144.718
              Estimate Std. Error
                                      Df z value Pr(>|z|)
> (Intercept) 152.133
                            2.566 1.000 59.278 < 2e-16 ***
                           68.122 0.983
                                          7.196 6.18e-13 ***
> x2bmi
               490.232
> x2map
               231.871
                           63.226 0.985
                                           3.667 0.000245 ***
> x2hd1
              -153.967
                           65.868 0.926 -2.338 0.019413 *
                                           6.353 2.11e-10 ***
               445.526
                           70.126 0.984
> x21tg
> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
> (Dispersion parameter for gaussian family taken to be 2911.317)
      Null deviance: 2621009 on 441.0 degrees of freedom
> Residual deviance: 1240941 on 426.2 degrees of freedom
> AIC: 4797.4
> Lambda: 85.24
```

> Number of Newton-Raphson iterations: 42

Comparisons between methods to select the tuning parameter and further discussions We conclude this short note by emphasizing that **islasso** also accepts the so-called elastic-net penalty, such that

$$\frac{1}{2}||\mathbf{y} - \mathbf{X}\boldsymbol{\beta}||_2^2 + \lambda \{\alpha||\boldsymbol{\beta}||_1 + \frac{1}{2}(1-\alpha)||\boldsymbol{\beta}||_2^2\}$$

where $0 \le \alpha \le 1$ is the mixing parameter to be specified in islasso() and islasso.fit() via the argument alpha, e.g.

```
out2 <- update(out, alpha = 0.5)</pre>
lmb.bic <- aic.islasso(out2, method = "bic", interval = c(1, 100))</pre>
> Optimization through bic
               38.8146 \text{ bic} =
> lambda =
                                 5052.27556
> lambda =
               62.1854 \text{ bic} =
                                 5069.14549
               24.3707 \text{ bic} =
> lambda =
                                 5030.54552
> lambda =
               15.4439 \text{ bic} =
                                 5005.21015
                9.9268 \text{ bic} =
> lambda =
                                 4980.02217
> lambda =
                6.5171 \text{ bic} =
                                 4956.43776
> lambda =
                4.4097 \text{ bic} =
                                 4938.04911
> lambda =
                3.1073 \text{ bic} =
                                 4929.80041
> lambda =
                2.3024 \text{ bic} =
                                 4924.67504
> lambda =
                1.8049 \, \text{bic} =
                                4923.18374
> lambda =
                1.4744 \text{ bic} =
                                 4923.61784
> lambda =
                1.7658 \text{ bic} =
                                 4923.15119
> lambda =
                1.7289 \text{ bic} =
                                 4923.15297
> lambda =
                1.7494 \text{ bic} =
                                4923.15145
> lambda =
                1.7625 \, \text{bic} =
                                 4923.15082
> lambda =
                1.7584 \text{ bic} =
                                 4923.15076
> lambda =
                1.7599 \text{ bic} =
                                 4923.15074
> lambda =
                1.7598 \, \mathrm{bic} =
                                4923.15074
> lambda =
                1.7599 \text{ bic} =
                                 4923.15074
                1.7599 \text{ bic} = 4923.15074
> lambda =
> lambda =
                1.7599 \text{ bic} = 4923.15074
out3 <- update(out, lambda = lmb.bic)
summary(out3, pval = 0.05)
> Call:
  islasso(formula = y ~ x2, lambda = lmb.bic, data = diabetes)
 Residuals:
        Min
                    1Q
                          Median
                                         30
  -152.747
                          -1.882
             -29.406
                                    31.545
                                              150.128
                Estimate Std. Error
                                            Df z value Pr(>|z|)
                 152.133
                                 2.514
                                         1.000
                                                 60.511 < 2e-16 ***
> (Intercept)
> x2sex
                -249.540
                                         1.000
                                                 -3.915 9.06e-05 ***
                               63.747
> x2bmi
                 457.109
                               82.150
                                         1.000
                                                  5.564 2.63e-08 ***
                 336.222
                               70.346
                                                  4.780 1.76e-06 ***
> x2map
                                         1.000
> x2ltg
                 649.340
                               99.634
                                        0.998
                                                  6.517 7.16e-11 ***
```

```
> x2age:sex
              166.531
                          70.542 0.999
                                          2.361
                                                   0.0182 *
                                                  0.0370 *
> x2ldl:ltg
              591.348
                         283.521 0.984
                                          2.086
> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
 (Dispersion parameter for gaussian family taken to be 2793.808)
>
      Null deviance: 2621009 on 441.0 degrees of freedom
> Residual deviance: 1080400 on 386.7 degrees of freedom
> AIC: 4815.2
> Lambda: 1.7599
> Number of Newton-Raphson iterations: 30
```

References

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- Brown B and Wang Y. Standard errors and covariance matrices for smoothed rank estimators. Biometrika 2005; 92: 149–158.