

netReg

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2017-05-05

Introduction

Modelling biological associations or dependencies using linear regression models is often complicated when the analysed data-sets are high-dimensional and less observations than variables are available ($n \ll p$). For these scenarios methods utilizing a priori knowledge, e.g. in the form of biological networks, have been proposed, arguing that this information might provide better estimates for regression coefficients. Recently several network-based regularization techniques have been proposed (C. Li and Li 2008, Kim, Pan, and Shen (2013), Cheng et al. (2014)).

netReg provides a highly-efficient implementation of these graph-penalized regression model. The models introduce a priori generated biological graph information into generalized linear models yielding sparse or smooth solutions for regression coefficients.

netReg computes coefficients using *cyclic coordinate descent* as previously introduced (Fu 1998, Friedman et al. (2007)), (Friedman, Hastie, and Tibshirani 2010). The package is an R-wrapper to an external C++ library that uses **RcppArmadillo** (Eddelbuettel and Sanderson 2014) for fast matrix calculations and **dlib** (King 2009) for gradient-free convex optimization for model selection.

Edgenet tutorial

This section explains how to fit a linear model and do parameter estimation using **edgenet**-regularization. The model is a truncated version from (Cheng et al. 2014) that is able to introduce prior graphs for the design and response matrices for penalization.

At first we generate some toy data randomly:

```
set.seed(23)
X <- matrix(rnorm(1000*5), 1000)
Y <- matrix(rnorm(1000*5), 1000)
```

Then we load the **netReg** library:

```
library(netReg)
```

For **edgenet** we need to create an affinity matrix for the co-variables first. We also can create a graph for the responses, but this is not necessary to demonstrate the method. We could create a random graph like this:

```
aff.mat <- matrix(rbeta(25, 1, 5), 5)
aff.mat <- (t(aff.mat) + aff.mat) / 2
diag(aff.mat) <- 0
```

We created the affinity matrix absolutely random, although in practice a *real* (biological) observed affinity matrix should be used, because in the end the affinity matrix decides the shrinkage of the coefficients.

Model fitting

Fitting a model using edge-based regularization with **netReg** is easy:

```
fit <- edgnet(X=X, Y=Y, G.X=aff.mat, lambda=1, psigx=1, family="gaussian")
print(fit)
```

```
##
## Call: edgnet.default(X = X, Y = Y, G.X = aff.mat, lambda = 1, psigx = 1,
##      family = "gaussian")
##
## Coefficients:
##           [,1]           [,2]           [,3]           [,4]           [,5]
## [1,]  0.01497323  0.015657011  0.0006453391  0.04699207 -0.009302215
## [2,] -0.01076841  0.069246936 -0.0876302568 -0.00405287 -0.007441449
## [3,] -0.02396619 -0.006507272  0.0244506149  0.02685324 -0.014316626
## [4,]  0.00000000 -0.039063322 -0.0301731092 -0.02050031  0.001355670
## [5,]  0.04213578  0.021257125  0.0242694757 -0.02222326 -0.008547463
## Intercept:
##           [,1]
## [1,]  0.019672657
## [2,] -0.047832210
## [3,] -0.004606925
## [4,] -0.025844237
## [5,]  0.032155794
## Parameters:
## lambda psi_gx psi_gy
##      1      1      0
## Family:
## [1] "gaussian"
```

In this case we used a single affinity matrix $G.X$ which represents the relationship of the covariables X . If the design matrix has p variables, $G.X$ has to be an $(p \times p)$ -dimensional symmetric matrix. We can also include a matrix for the response matrix Y with q dependent variables. In that case the affinity matrix $G.Y$ has to be $(q \times q)$ -dimensional (and also symmetric).

The `fit` object contains information about coefficients, intercepts, residuals, etc. Having the coefficients estimated we are able to predict novel data-sets:

```
X.new <- matrix(rnorm(10*5),10)
pred <- predict(fit, X.new)
```

The `pred` objects contains the predicted values for the responses.

Model selection

In most cases we do not have the optimal shrinkage parameters λ , ψ_{gx} and ψ_{gy} . For these settings you can use `netReg`'s included model selection. We use Powell's BOBYQA algorithm ((Powell 2009)) for gradient-free optimization that is included in the C++ library `Dlib`. Doing the model selection only requires calling `cv.edgnet`:

```
cv <- cv.edgnet(X=X, Y=Y, G.X=aff.mat, family="gaussian", maxit=1000)
print(cv)
```

```
##
## Call: cv.edgnet.default(X = X, Y = Y, G.X = aff.mat, maxit = 1000,
##      family = "gaussian")
##
## Parameters:
```

```
## lambda  psigx  psigy
##      0      0      0
## Family:
## [1] "gaussian"
```

You can use the fitted parameters for the normal `edgenet` function. In this scenario λ , ψ_{gx} and ψ_{gy} should be roughly 0 for three reasons:

- we had enough data and a small number of covariables ($n > p$), so we can find the *BLUE* estimator,
- we created X and Y independent of each other,
- our prior graph `aff.mat` had little weight.

Let's do a scenario where we need to shrink some coefficients, i.e. $n \ll p$. We choose a small p , such that the computation does not take too long.

```
p <- 25
X <- matrix(rnorm(10*p), 10)
Y <- matrix(rnorm(10*p), 10)
aff.mat <- matrix(rgamma(p * p, 5, 1), p)
aff.mat <- (t(aff.mat) + aff.mat)
diag(aff.mat) <- 0
cv <- cv.edgenet(X=X, Y=Y, G.X=aff.mat, family="gaussian", maxit=1000)
print(cv)
```

```
##
## Call: cv.edgenet.default(X = X, Y = Y, G.X = aff.mat, maxit = 1000,
##      family = "gaussian")
##
## Parameters:
##   lambda    psigx    psigy
## 6.882653 0.000000 0.000000
## Family:
## [1] "gaussian"
```

In the above example λ should have changed quite a bit, while ψ_{gy} should still be 0. Since we generated `aff.mat` randomly ψ_{gx} should be roughly (or exact) zero as well. This makes sense intuitively since we did not put any biological relationships into the affinity matrices.

References

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