The CATE Package for High Dimensional Factor Analysis and Confounder Adjusted Multiple Testing

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In this document, we demonstrate how to use the R package cate to adjust for confounding effects in multiple hypothesis testing. Users who are only interested in high dimensional factor analysis may skip the motivating example and go to Section 4.

1 Introduction

The dataset we will be using is a genome-wide association study of gender (Vawter et al., 2004). In this study, samples were taken postmortem from the brains of 10 individuals, 5 men and 5 women. Three samples were taken from different regions of the brain of each individual, and one aliquot of each sample was sent to each of 3 laboratories for analysis. There were two different microarray platforms used by these labs.

First, let's load the data.

```
library(cate)
data(gender.sm)
names(gender.sm)
   [1] "Y"
                                                            "spikectl"
                                                                              "geneinfo"
   [6] "xchrom"
                         "ychrom"
                                           "xychrom"
                                                                             "autosomal"
## [11] "genecoloring"
                         "samplecoloring"
cbind(X = dim(gender.sm$X), Y = dim(gender.sm$Y), Z = dim(gender.sm$Z)) # matrix dimensions
         Χ
             Y Z
## [1,] 84 84 84
## [2,] 1 500 4
```

There are in total 84 samples and 500 genes in this dataset. There are 12600 genes measured in the original dataset (Vawter et al., 2004; Gagnon-Bartsch and Speed, 2012) and here we look at a 500 subsample. There should have been $10 \times 3 \times 3 = 90$ samples in total, but 6 of them are missing. In the data object gender.sm, X is the gender of each person, Y is the gene expression matrix, and Z includes the batch labels (lab and microarray platform, not individual or brain region).

Since there are several batch variables and unmeasured covariates in this dataset, it should come as no surprise that these confounders can seriously bias the association tests. In other words, the marginal effects of *X* on *Y* may be different from the actual effects of *X* on *Y*. Wang et al. (2015) describe a general framework and a two-step solution for this problem. In the first step, we apply factor analysis to the part of gene expression matrix *Y* that is unrelated to the variable(s) of interest *X*. In the second step, we correct the marginal effects of *X* on *Y* by using the factors obtained in the first step.

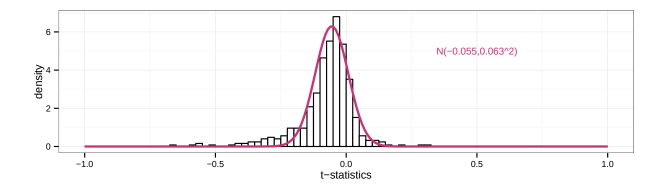


Figure 1: Histogram of t-statistics before confounder adjustment

2 Confounder adjusted multiple testing

To illustrate how the confounders in the <code>gender.sm</code> dataset can bias the association tests, we first run the two-sample t-test for each gene

```
t.stats <- apply(gender.sm$Y, 2, function(y, x) t.test(y~x)$statistic, gender.sm$X)
```

Under the null hypothesis that the gene is unrelated to the gender, the corresponding t-statistic is expected to follow a t-distribution with n-1 degrees of freedom. Since n=84 is fairly large in this study, this t-distribution is very close to the standard normal distribution N(0,1). However, the empirical distribution of t.stats is shown in the next histogram (code not echoed):

It clearly departs from the theoretical null distribution. The median absolute deviation (MAD) of this histogram is only 0.066, much less than the theoretical value 1, i.e. the t-statistics are extremely underdispersed. The histogram is also a little skewed compared to normal or t distribution.

As mentioned earlier, this phenomenon is most likely due to the existence of **confounders**, unmeasured variables that are correlated with both the response *Y* and the variable of interest *X*. To correct for such confounders, we first need to estimate its number. This is implemented in the function est.confounder.num

By default, est.confounder.num uses method = "bcv", which calls the EsaBcv function in the esaBcv package to estimate the number of factors by bi-cross-validation (BCV). To look at the curve of BCV error using different number of factors, the user can turn on the bcv.plot argument. It is recommended to use at least 20 for nRepeat to obtain a more accurate BCV error.

Another method to estimate the number of confounders of is the eigenvalue difference method (Onatski, 2010)

```
est.confounder.num(gender.sm$X, gender.sm$Y, intercept, method = "ed")
## [1] 3
```

The "bcv" method tends to estimate more weak factors and takes longer time than "ed". We recommend to use "bcv" method for most datasets; see Owen and Wang (2015) for more detail.

After finding the number of factors, the user can call the main cate function to adjust for the confounders

```
cate.results <- cate(gender.sm$X, gender.sm$Y, intercept, r = factor.num$r)
names(cate.results)

## [1] "Gamma" "Sigma" "Z" "niter" "converged"
## [6] "alpha" "beta" "beta.cov.row" "beta.cov.col" "beta.t"
## [11] "beta.p.value" "Y.tilde" "alpha.p.value"</pre>
```

For most users, the interesting returned values are

beta: the estimated effects after adjustment;

beta.t: the t-statistics after adjustment;

beta.p.value: the p-values of the estimated effects;

alpha.p.value: the p-value of a χ^2 -test for confounding.

The first thing to look at is perhaps the confounding test, whose null hypothesis is that the estimated factors are not correlated with *X* so the individual association tests are not biased. For the gender dataset, the p-value of this test is

```
cate.results$alpha.p.value
## [1] 0.006312493
```

This is much smaller than 0.05, which indicates, together with the previous histogram, that there are some confounders in the experiment. When this is the case, the user may want to look at the factor analysis results to search for possible sources of confounding (in particular the loadings Gamma and the estimated confounders Z returned by cate).

To discover candidate genes, the user can apply the p.adjust function in the stats package to control certain multiple testing error. For controlling the family-wise error rate (FWER), the user may use the bonferroni or holm option in p.adjust. To increase the number of findings and still control the false discovery rate (FDR), the user may use the BH option. Here are a couple of examples:

```
which(p.adjust(cate.results$beta.p.value, "bonferroni") < 0.05) # control FWER at 0.05
which(p.adjust(cate.results$beta.p.value, "BH") < 0.2) # control FDR at 0.2</pre>
```

Finally, let's review all the available options in cate.

```
args(cate)
## function (X, Y, X.nuis = NULL, r, fa.method = c("ml", "pc", "esa"),
## adj.method = c("rr", "nc", "lqs", "naive"), psi = psi.huber,
## nc = NULL, nc.var.correction = TRUE, calibrate = TRUE)
## NULL
```

Here are some detailed descriptions of all the arguments

X: primary treatment variable(s) whose effects are of interest.

Y: response matrix (e.g. gene expression).

X.nuis: nuisance covariate(s) to include in the regression whose effects are not of interest. Generally X.nuis includes a column of intercept, and can further include the known batch variables (e.g. the matrix Z in the gender.sm dataset) and other demographics variables such as age or gender.

r: number of confounders, usually estimated by est.confounder.num.

fa.method: method used to estimate the confounders. See Section 4 for more detail.

- adj.method: method used to adjust for the confounders. There are two main approaches: robust regression (rr) and negative control (nc). If a fair amount (rule of thumb: ≥ 30) of negative control genes (e.g. spike-in controls) are available, it is recommended to use the nc option. Housekeeping genes can also be used as negative controls, but they are not as reliable as spike-in controls. The robust regression (rr) option assumes the true effects are sparse and can be used when negative controls are not available.
- psi: estimating equation function used when adj.method = "rr". See the rlm function in package MASS
 for more detail.
- nc: positions of the negative controls. Can be a vector of numbers between 1 and p, or a logical vector of length p.
- nc.var.correction: if TRUE (default and recommended), use the variance correction formula in Wang et al. (2015) when adj.method = "nc"; if FALSE, use the oracle variance (same as the RUV4 function in package ruv). See Wang et al. (2015) for more detail.
- calibrate: if TRUE (default), scale the t-statistics beta.t to have median equal to 0 and median absolute deviation (with respect to normal distribution) equal to 1.

For example, to use the nc adjustment method, the user must specify the positions of negative control genes. In the dataset gender.sm, these are given in spikectl (33 spike-in controls) and hkctl (799 housekeeping genes). Here is a sample usage:

The housekeeping genes are usually less reliable than the spike-in controls.

We end this section with Figure 2, two histograms of beta.t after the confounder adjustment using adj.method = "rr" and adj.method = "nc". In both cases, the bulk of the statistics approximately follows the standard normal distribution.

3 Other available confounder adjustment methods on CRAN/Bioconductor

The preprocessed gender.sm dataset was first used by Gagnon-Bartsch and Speed (2012) to demonstrate their confounder correction method "Remove Unwanted Variation" (RUV) using negative controls. See also the R package ruv. The RUV4 function therein is very similar to the adj.method = "nc" option in cate. Two other related packages are sva (Leek and Storey, 2008) and leapp (Sun et al., 2012), which motivate the robust regression method (adj.method = "rr") in cate.

To compare the performance of different methods, the user can use the wrapper functions sva.wrapper, ruv.wrapper, leapp.wrapper in the cate package. They provide a uniform interface and call the corresponding functions in the original packages.

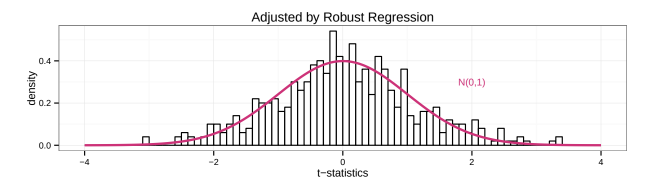
4 Factor analysis

For a data matrix $Y \in \mathbb{R}^{n \times p}$, n being the number of observations and p being the number of variables (e.g. genes), the factor model assumes

$$Y_{n\times p}=Z_{n\times r}\Gamma_{p\times r}^T+E_{n\times p},$$

where Z is a rotation matrix containing latent factors, Γ is a matrix of loadings and E is a noise matrix. The columns of E have covariance matrix Σ .

To perform factor analysis, one simply calls the function factor.analysis with the data matrix and number of factors:



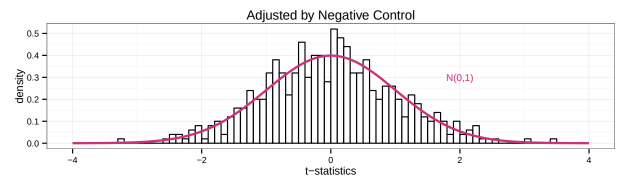


Figure 2: Histograms of test statistics after adjustment

```
mle <- factor.analysis(gender.sm$Y, r = 5)
names(mle)
## [1] "Gamma" "Sigma" "Z" "niter" "converged"</pre>
```

By default, factor.analysis estimates the latent factors by maximum likelihood (method = "ml"). Other available algorithms are principal component analysis (method = "pc") and bi-convex optimization via early stopping alternation (method = "esa"). The default maximum likelihood estimator has good theoretical properties under heteroscedastic noise variance Σ (Bai and Li, 2012) or approximate factor model (Bai and Li, 2015). The esa method tries to minimize the prediction error for Y; for more details we refer the readers to the R package esaBcv. The pc option is usually more desirable when the noise variance is homoscedastic $\Sigma = \sigma^2 I_p$. Finally, using the pc method will always outputs the same results, but the iterative procedures ml and esa may converge to different values depending on the initial point.

The factor analysis function can work for any n and p. In contrasts, both the factanal function in package stats and the fa function in package psych do not support the high dimensional problem where p > n.

References

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