

# STK-IN4300 Statistical Learning Methods in Data Science

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#### Outline of the lecture

- Feature Assessment when  $p \gg N$ 
  - Feature Assessment and Multiple Testing Problem
  - The false discovery rate
- Stability Selection
  - Introduction
  - Selection probability
  - Stability path
  - Choice of regularization

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#### In the previous lecture:

- talked about the p >> N framework;
- focused on the construction of prediction models.

# More basic goal:

- assess the significance of the M variables;
  - ightharpoonup in this lecture M is the number of variables (as in the book);
- e.g., identify the genes most related to cancer.

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## Assessing the significance of a variable can be done:

- as a by-product of a multivariate model,
  - selection by a procedure with variable selection property;
  - absolute value of a regression coefficient in lasso;
  - if and how fast a variable enter in a boosting model.
- evaluating the variables one-by-one:
  - univariate tests:

multiple hypothesis testing

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# Consider the data from Rieger et al. (2004):

- study on the sensitivity of cancer patients to ionizing radiation treatment;
- oligo-nucleotide microarray data (M = 12625);
- N = 58:
  - ▶ 44 patients with normal reaction;
  - ▶ 14 patients who had a severe reaction.

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**TABLE 18.4.** Subset of the 12,625 genes from microarray study of radiation sensitivity. There are a total of 44 samples in the normal group and 14 in the radiation sensitive group; we only show three samples from each group.

|             | Normal |       |        |   | Radiation Sensitive |        |         |   |
|-------------|--------|-------|--------|---|---------------------|--------|---------|---|
| Gene 1      | 7.85   | 29.74 | 29.50  |   | 17.20               | -50.75 | -18.89  |   |
| Gene 2      | 15.44  | 2.70  | 19.37  |   | 6.57                | -7.41  | 79.18   |   |
| Gene 3      | -1.79  | 15.52 | -3.13  |   | -8.32               | 12.64  | 4.75    |   |
| Gene 4      | -11.74 | 22.35 | -36.11 |   | -52.17              | 7.24   | -2.32   |   |
| :           | :      | :     | :      | : | :                   | :      | :       | : |
| Gene 12,625 | -14.09 | 32.77 | 57.78  |   | -32.84              | 24.09  | -101.44 |   |

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## The simplest way to identify significative genes:

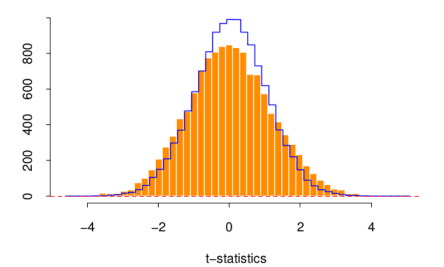
• two-sample t-statistic for each gene,

$$t_j = \frac{\bar{x}_{2j} - \bar{x}_{1j}}{se_j}$$

#### where

- $\bar{x}_{kj} = \sum_{i \in C_k} x_{kj} / N_k;$
- $C_k$  are the indexes of the  $N_k$  observations of group k;
- $se_j = \hat{\sigma}_j \sqrt{\frac{1}{N_1} + \frac{1}{N_2}};$
- $\hat{\sigma}_{j}^{2} = \frac{1}{N_{1} + N_{2} 2} \left( \sum_{i \in C_{1}} (x_{ij} \bar{x}_{1j})^{2} + \sum_{i \in C_{2}} (x_{ij} \bar{x}_{2j})^{2} \right).$

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## From the histogram (12625 t-statistics):

- the values range from -4.7 to 5.0;
- assuming  $t_i \sim N(0,1)$ , significance at 5% when  $|t_i| \ge 2$ ;
- in the example, 1189 genes with  $|t_i| \ge 2$ .

#### However:

- out of 12625 genes, many are significant by chance;
- supposing (it is not true) independence:
  - expected falsely significant genes,  $12625 \cdot 0.05 = 631.25$ ;
  - standard deviation,  $\sqrt{12625 \cdot 0.05 \cdot (1 0.05)} \approx 24.5$ ;
- the actual 1189 is way out of range.

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Without assuming normality, permutation test:

- perform  $K = \binom{58}{14}$  permutations of the sample labels;
- compute the statistic  $t_i^{[k]}$  for each permutation k;
- the p-value for the gene j is

$$p_j = \frac{1}{K} \sum_{k=1}^{K} \mathbb{1}(|t_j^{[k]}| > |t_j|)$$

(not all  $\binom{58}{14}$  are needed, random sample of K=1000)

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For  $j \in 1, ..., M$  test the hypotheses:

 $H_{0j}$ : treatment has no effect on gene j

 $H_{1j}$ : treatment has an effect on gene j

 $H_{0i}$  is rejected at level  $\alpha$  if  $p_i < \alpha$ :

- $\alpha$  is the type-I error;
- we expect a probability of falsely rejecting  $H_{0j}$  of  $\alpha$ .

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#### Feature Assessment when $p \gg N$ : family-wise error rate

Define 
$$A_j = \{H_{0j} \text{ is falsely rejected}\} \longrightarrow Pr(A_j) = \alpha.$$

The **family-wise error rate** (FWER) is the probability of at least one false rejection,

$$Pr(A) = Pr(\bigcup_{j=1}^{M} A_j)$$

- for p large,  $Pr(A) \gg \alpha$ ;
- it depends on the correlation between the test;
- if tests independent,  $Pr(A) = 1 (1 \alpha)^M$ :
- test with positive dependence,  $Pr(A) < 1 (1 \alpha)^M$ ;
  - positive dependence is typical in genomic studies.

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## Feature Assessment when $p \gg N$ : family-wise error rate

The simplest approach to correct the p-value for the multiplicity of the tests is the **Bonferroni method**:

- reject  $H_{0j}$  if  $p_j < \alpha/M$ ;
- it makes the individual test more stringent;
- controls the FWFR
  - it is easy to show that  $FWER \leq \alpha$ ;
- it is very (too) conservative.

# In the example:

- with  $\alpha = 0.05$ ,  $\alpha/M = 0.05/12635 \approx 3.9 \times 10^{-6}$ ;
- no gene has a p-value so small.

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#### Feature Assessment when $p \gg N$ : the false discovery rate

# Instead of FWER, we can control the false discovery rate (FDR):

 expected proportion of genes incorrectly defined significant among those selected as significant,

|             | Called          | Called      |       |
|-------------|-----------------|-------------|-------|
|             | Not Significant | Significant | Total |
| $H_0$ True  | U               | V           | $M_0$ |
| $H_0$ False | T               | S           | $M_1$ |
| Total       | M-R             | R           | M     |

- in formula, FDR = E[V/R];
- procedure to have the FDR smaller than an user-defined  $\alpha$ .

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## Feature Assessment when $p \gg N$ : the false discovery rate

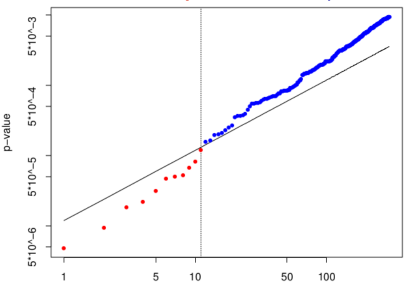
#### Algorithm 18.2 Benjamini-Hochberg (BH) Method.

- 1. Fix the false discovery rate  $\alpha$  and let  $p_{(1)} \leq p_{(2)} \leq \cdots \leq p_{(M)}$  denote the ordered p-values
- 2. Define

$$L = \max\left\{j : p_{(j)} < \alpha \cdot \frac{j}{M}\right\}. \tag{18.44}$$

3. Reject all hypotheses  $H_{0j}$  for which  $p_j \leq p_{(L)}$ , the BH rejection threshold.





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#### Feature Assessment when $p \gg N$ : the false discovery rate

## In the example:

- $\alpha = 0.15$ ;
- the last  $p_i$  under the line  $\alpha \cdot (j/M)$  occurs at j=11;
- the smallest 11 p-values are considered significative;
- in the example,  $p_{(11)} = 0.00012$ ;
- the corresponding t-statistic is  $|t_{(11)}| = 4.101$ ;
- a gene is relevant if the corresponding t-statistics is in absolute value larger than 4.101.

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#### Feature Assessment when $p \gg N$ : the false discovery rate

It can be proved (Benjamini & Hochberg, 1995) that

$$\mathsf{FDR} \leqslant \frac{M_0}{M} \alpha \leqslant \alpha$$

- regardless the number of true null hypotheses;
- regardless the distribution of the p-values under  $H_1$ ;
- suppose independent test statistics;
- in case of dependence, see Benjamini & Yekutieli (2001).

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#### Stability Selection: introduction

## In general:

- the  $L_1$ -penalty is often use to perform model selection;
- no oracle property (strict conditions to have it);
- issues with selecting the proper amount of regularization;

# Meinshausen & Bühlmann (2010) suggested a procedure:

- based on subsampling (could work with bootstrapping as well);
- determines the amount of regularization to control the FWER;
- new structure estimation or variable selection scheme:
- here presented with  $L_1$ -penalty, works in general.

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#### Stability Selection: introduction

# Setting:

- β is a p-dimensional vector of coefficients;
- $S = \{j : \beta_j \neq 0\}, |S| < p;$
- $S^C = \{j : \beta_j = 0\};$
- $Z^{[i]} = (X^{[i]}, Y^{[i]}), i = 1, ..., N$ , are the i.i.d. data,
  - univariate response Y:
  - $N \times p$  covariate matrix X.
- consider a linear model

$$Y = X\beta + \epsilon$$

with  $\epsilon = (\epsilon_1, \dots, \epsilon_N)$  with i.i.d. components.

#### Stability Selection: introduction

The goal is to infer S from the data. We saw that lasso,

$$\hat{\beta}^{\lambda} = \operatorname{argmin}_{\beta \in \mathbb{R}^p} \left( ||Y - X\beta||_2^2 + \lambda \sum_{j=1}^p |\beta_j| \right)$$

provides an estimate of S,  $S^{\lambda} = \{j : \hat{\beta}_j \neq 0\} \subseteq \{1, \dots, p\}.$ 

#### Remember:

- λ ∈ ℝ<sup>+</sup> is the regularization factor;
- $||X_j||_2^2 = \sum_{i=1}^N (x_j^{[i]})^2 = 1;$

#### Stability Selection: selection probability

Stability selection is built on the concept of selection probability,

Definition 1: Let I be a random subsample of  $\{1,\ldots,N\}$  of size  $\lfloor N/2 \rfloor$  drawn without replacement. We define selection probability the probability for a variable  $X_j$  of being in  $S^{\lambda}(I)$ ,

$$\hat{\Pi}_j^{\lambda} = Pr^*[j \subseteq S^{\lambda}(I)]$$

#### Note:

- $Pr^*$  is with respect of both the random subsampling and other sources of randomness if  $S^{\lambda}$  is not deterministic:
- |N/2| is chosen for computational efficiency.

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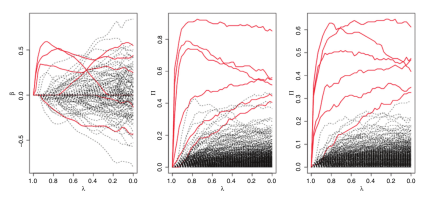
## Stability Selection: stability path

Once we have the selection probability, we can define the **stability** path, as the evolution of  $\hat{\Pi}_i^{\lambda}$  when  $\lambda \in \Lambda$  varies,

- similar to the learning path plot of lasso;
- it shows the selection probabilities for all variables;
- it is very useful for improved variable selection, especially in high-dimensional cases.

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## Stability Selection: stability path



- left: lasso learning path;
- center: stability path of the lasso;
- right: stability path of the randomized lasso.

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## Stability Selection: stability path

# Normally we would choose a specific $\lambda$ :

- it is a single element of the set  $\hat{S}^{\lambda}$ ,  $\lambda \in \Lambda$ ;
- S might not be a member of the set;
- even if it is, it is hard to find the right  $\lambda$  high-dimensions.

# With stability selection:

- we do not simply select one model in  $\hat{S}^{\lambda}$ ,  $\lambda \in \Lambda$ ;
- the data are perturbed (e.g. by subsampling) many times;
- we choose all variables that occur in a large fraction of the resulting selection sets.

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#### Stability Selection: stability selection

Definition 2: For a cut-off  $\pi_{thr}$ , with  $0 < \pi_{thr} < 1$ , and a set of regularization parameters  $\Lambda$ , the set of stable variables is defined as

$$\hat{S}^{\mathsf{stable}} = \left\{ j : \max_{\lambda \in \Lambda} (\Pi_j^{\lambda}) \geqslant \pi_{\mathsf{thr}} \right\}.$$

## In this way:

- we keep variables with a high selection probability;
- we disregard those with low selection probabilities;
- the exact cut-off  $\pi_{thr}$  is a tuning parameter;
- the results vary surprisingly little for sensible choices of  $\pi_{thr}$ ;
- results do not strongly depend on the choice of  $\lambda$  or  $\Lambda$ .

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#### Stability Selection: choice of regularization

#### Let:

- $S^{\Lambda} = \bigcup_{\lambda \in \Lambda} \hat{S}^{\lambda}$  be the set of selected variables  $\forall \lambda \in \Lambda$ ;
- $q_{\Lambda} = E[|\hat{S}^{\Lambda}(I)|]$  be the average number of selected variables;
- $V=|S^C \cap \hat{S}^{\rm stable}|$  the number of falsely selected variables with stability selection.

**Theorem (Meinshausen & Bühlmann, 2010)**: Assuming that the distribution of  $\{\mathbb{1}_{j\in \hat{S}^{\lambda}}\}$  is exchangeable  $\forall \lambda \in \Lambda$  and the procedure is not worse than a random guess, then

$$E[V] \leqslant \frac{1}{2\pi_{\rm thr} - 1} \frac{q_{\Lambda}^2}{p}$$

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## Stability Selection: choice of regularization

#### Therefore:

- $\pi_{thr}$  is a tuning parameter whose influence is very small;
  - sensible values are in (0.6, 0.9);
- once decided  $\pi_{thr}$ ,  $\Lambda$  is determined by the error control desired;
- specifically for  $\pi_{\text{thr}} = 0.9$ ,

  - $\Lambda: q_{\Lambda} = \sqrt{0.8\alpha p} \longrightarrow Pr[V > 0] \leqslant \alpha;$
- i.e., we need to find  $\Lambda$  that gives a specific  $q_{\Lambda}$ ,
  - q is given by the number of variables which enter in the model;
  - for lasso, find  $\lambda_{\min}: |\bigcup_{\lambda,\dots,\geq \lambda \geq \lambda-1} \hat{S}^{\lambda}| \leqslant q$

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#### Stability Selection: choice of regularization

#### Final remarks:

- without stability selection,  $\lambda$  depends on the unknown noise level of the observations:
- the advantages of stability selection are:
  - exact error control is possible;
  - the method works fine even though the noise level is unknown;
- real advantage when  $p \ge N$  (hard to estimate the noise level);
- consistency can be proved (see Meinshausen & Bühlmann, 2010, for the proof for randomized lasso);
- exchangeability in Theorem 1 is only need for the proof.

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#### References I

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- RIEGER, K. E., HONG, W.-J., TUSHER, V. G., TANG, J., TIBSHIRANI, R. & CHU, G. (2004). Toxicity from radiation therapy associated with abnormal transcriptional responses to dna damage. *Proceedings of the National Academy of Sciences* **101**, 6635–6640.

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