# High-Throughput Sequencing Course Supervised Learning

Biostatistics and Bioinformatics

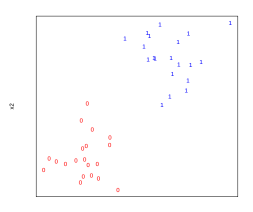


Summer 2018

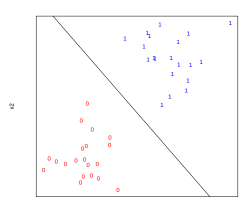




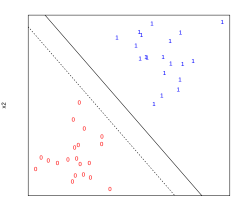
### CLASSIFICATION PROBLEM



## CLEAR-CUT CASE

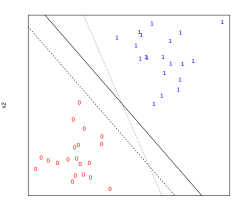


# CLEAR-CUT CASE?

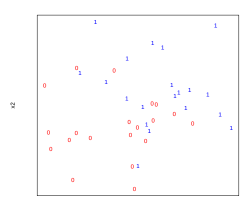


х1

# CLEAR-CUT CASE??

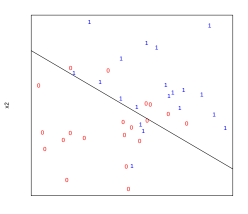


### LESS CLEAR-CUT CASE

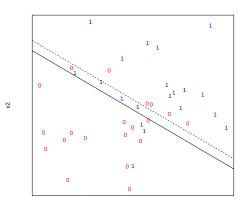


x1

### LESS CLEAR-CUT CASE

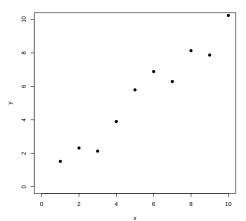


### LESS CLEAR-CUT CASE



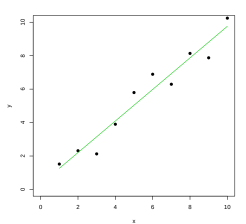
### REGRESSION PROBLEM

```
set.seed(10) x <- 1:10 y = x + rnorm(10, 0.5)
par(mfrow = c(1, 1), bg = "white") plot(x, y, xlim = c(0, 10), ylim = c(0, 10), pch = 19)</pre>
```

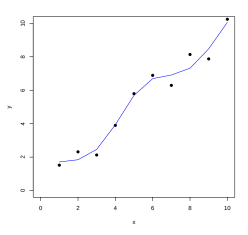


## LINEAR REGRESSION (LIN)

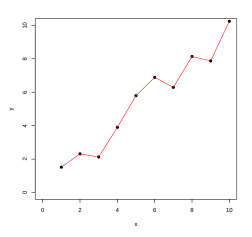
```
 par(mfrow = c(1, 1), bg = "white") \ plot(x, y, xlim = c(0, 10), ylim = c(0, 10), pch = 19) \ modlm = lm(y x) \ lines(x, predict(modlm), col = 3)
```



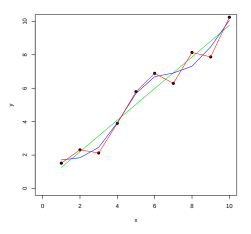
# Spline Regression (spl)



# CONNECT THE DOTS (CTD)



## WHICH APPROACH?



# SUPERVISED LEARNING (CLASSIFICATION)

- ▶ Goal: Predict a binary outcome (Y) on the basis of baseline information (X)
- ➤ Y assumes the value 0 or 1 (e.g., control vs case, or AML vs ALL)
- ► X could be single variable or be a vector of multiple variables
- ▶ Example: Can you predict Y on the basis of two genes say  $X_1$  and  $X_2$
- Note that a goal is to build a machine that will take on two values  $X_1$  and  $X_2$  and return a 0 or a 1
- ▶ You can denote this machine as a function  $g(x_1, x_2)$

#### Classifier

- We will denote the predictor or classifier by g(x)
- $x = (x_1, x_2)$  is the vector of gene expressions for genes 1 and 2
- $\blacktriangleright$  Based on x, the classifier g makes a prediction for the outcome
- ▶ Note that g(x) = 0 or g(x) = 1
- ► The prediction is *correct* if Y = 1 and  $g(x_1, x_2) = 1$ , or Y = 0 and  $g(x_1, x_2) = 0$
- ► The prediction is wrong if Y = 0 and  $g(x_1, x_2) = 1$ , or Y = 1 and  $g(x_1, x_2) = 0$

### PREDICTION ASSESSMENT

$$g(x_1, x_2) = 0$$
  $g(x_1, x_2) = 1$   
 $Y = 0$  True-Negative False-Positive  
 $Y = 1$  False-Negative True-Positive

### STEPS TO CONSTRUCT A CLASSIFIER

- ► Collect a random data set of size n to build (train) a classifer
- ► This is called the training data
- ▶ On the basis of these data, construct the classifier  $g_n$
- $\blacktriangleright$  It is subscripted by n to emphasize that it is trained on the basis of the training data
- ▶ Note that the final performance of  $g_n$  is not be judged on the basis of the training data
- ► It is to be judged on the basis of its performance on *future* data
- ► Called testing data

### STEPS IN NOTATION

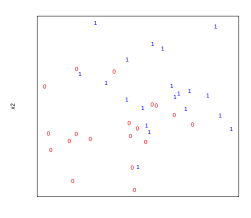
- ► Collect the training data  $(X_1, Y_1), \ldots, (X_n, Y_n)$
- ightharpoonup Construct a classifier  $g_n$  on the basis of the training data
- ▶ Apply  $g_n$  to a new data set  $X_1^*, \ldots, X_k^*$  to get
- ▶ k predictions:  $\hat{Y}_1^*, \dots, \hat{Y}_k^*$
- ► Compare the predictions to the observed outcomes  $Y_1^*, \ldots, Y_k^*$
- ▶ Note that at the testing stage, you are blinded to the  $Y_k^*$

## k-Nearest Neighborhood (Non-parametric)

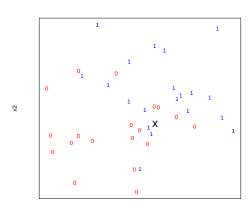
- $\blacktriangleright$  Generally, non-parametric methods (e.g., k-NN) are preferred
- ► These do not make strong assumptions on the specific shape of the underlying relationship, if there is one, between X and Y
- $\blacktriangleright$  For each x (point on the scatter plot), identify the k nearest neighbors
- ► Among the k neighbors, count the number of responders (say  $r_x$ )
- ► Set

$$\hat{\eta}(x) = \frac{r_x}{k}$$

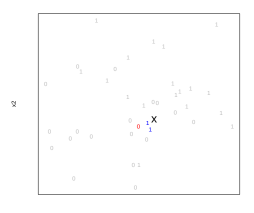
# K-Nearest Neighborhood (training data)



# K-Nearest Neighborhood (predict new sample)



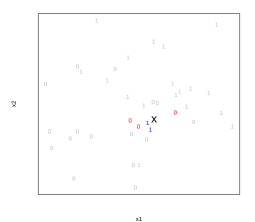
### 3-Nearest Neighborhood



x1

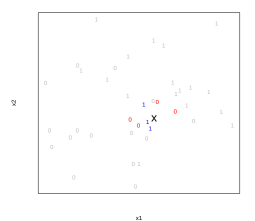
The prediction based on 3 nearest neighbors is  $\hat{Y}^* = 1$ 

### 5-Nearest Neighborhood



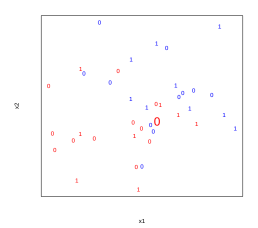
The prediction based on 5 nearest neighbors is  $\hat{Y}^* = 0$ 

#### 7-Nearest Neighborhood



The prediction based on 7 nearest neighbors is  $\hat{Y}^* = 0$ 

### PREDICTION VS TRUE STATE



Question: What should the parameter k be?

### MEAN REGRESSION MODEL

- ightharpoonup E(Y) is the unconditional (on X) mean of Y.
- $\blacktriangleright$  Model the mean relationship between Y and X

$$\eta(x) = E(Y|X=x)$$

 $\blacktriangleright \eta(x)$  is the *conditional* (on X) mean of Y given that X has realized the value x.

### Bayes Classifier

$$g(X) = \begin{cases} 1 & \text{if } \eta(X) \ge \frac{1}{2}, \\ 0 & \text{if } \eta(X) < \frac{1}{2} \end{cases}$$

- ▶ This classifier is "optimal" in the sense that there is no better classifier with respect to minimizing the error  $(P(g(X) \neq Y))$ .
- ▶ Suppose that  $g^*$  is another classifier. Then

$$P(g(X) \neq Y) \le P(g^*(X) \neq Y)$$

▶ Note that the optimality concerns  $\eta(x)$  and not  $\hat{\eta}(x)$ .

## LOGISTIC REGRESSION (PARAMETRIC)

► Most commonly used method for modeling the relationship between a binary response and a set of co-variables

$$logit(\eta(x)) = \beta_0 + \beta_1 x_1 + \beta_2 x_2,$$

where

$$logit(\eta(x)) = log\left(\frac{p}{1-p}\right),$$

for  $p \in (0,1)$  is called the "logit" function.

## Estimating $\eta(x)$

- ► Estimate the model parameters  $(\beta_0, \beta_1 \text{ and } \beta_2)$  using maximum-likelihood estimation to get  $\hat{\beta}_0, \hat{\beta}_1$  and  $\hat{\beta}_2$
- ► For the logistic model

$$\hat{\eta}(x) = \frac{\exp(\hat{\beta}_0 + \hat{\beta}_1 x_1 + \hat{\beta}_2 x_2)}{1 + \exp(\hat{\beta}_0 + \hat{\beta}_1 x_1 + \hat{\beta}_2 x_2)}$$

### OTHER CLASSIFICATION METHODS

- ► Fisher's Linear Discriminant
- ► Support Vector Machines (SVM)
- ► Classification and Regression Trees (CART)
- ► Random Forests (aggregated trees)
- ► Methods for "Deep" learning

### BIAS VERSUS VARIANCE

- ► A very important principle in statistical modeling is the so called bias-variance tradeoff
- ► The bias of  $\hat{\eta}(x)$  is

$$b(x) = \hat{\eta}(x) - \eta(x)$$

▶ The variance of  $\hat{\eta}(x)$  is

$$v(x) = E(\hat{\eta}(x) - \eta(x))^2)$$

- ► The bias-variance tradeoff implies that both cannot be minimized simultaneously
- $\blacktriangleright$  For example for the k-NN method increasing k increases bias while decreasing variance

### TRAINING AND TESTING

- ► In practice, the model is first estimated (trained) using an initial set of data
- ► This data set is usually called the "training" data
- ► Once the model is trained, then it is applied to an "independent" set of data
- ► This data set is usually called the "testing" (or validation) data set

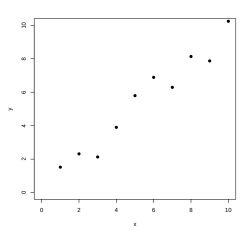
#### Parsimony

- ► The model should be parsimonious (less is more)
- ► Including too many noisy/unimportant features often degrades the performance of the classifier.
- ► Including highly dependent induces problems (e.g., multi-collinearity from simple linear regression).
- ► Additional complication: It is not practically/computationally feasible to include tens of thousands of features in the model.

#### **OVERFITTING**

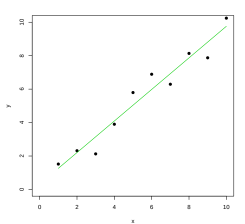
- ► Two many parameters compared to the number of data points in the training set
- ► A complicated model will fit the training set well
- $\blacktriangleright$  It will however perform poorly for an independent set.

### **OVERFITTING**

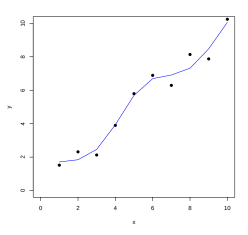


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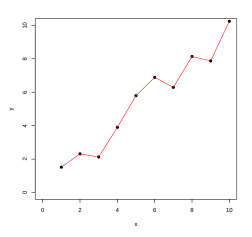
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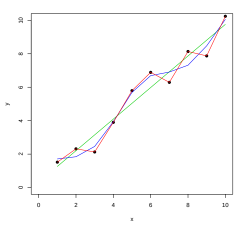
# Spline Regression (spl)



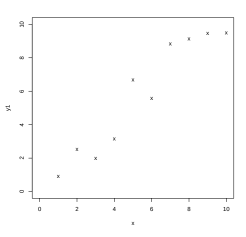
# CONNECT THE DOTS (CTD)



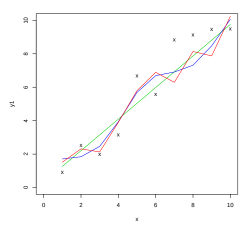
RSS: 4.1 (LIN) VS 1.9 (SPL) VS 0 (CTD)



### NEW DATA SET



RSS: 11 (LIN) VS 12.4 (SPL) VS 14 (CTD)



#### Two Challenges in Building a Classifier

#### 1. Feature Selection:

- It is neither feasible nor provident to build a classifier based on all available variables
- A subset of the variables has to be selected to build the model
- ▶ This is also called feature extraction

#### 2. Tuning Parameter Selection:

- Statistical methods may have one or more parameters that have to be set
- For example when using k-NN, one has to decide what k should be (e.g., 1, 3 or 5 or how about 8)?
- ► Choosing the defaults set by the software is inappropriate
- ► The feature selection method could also have tuning parameters that have to be set (e.g., the number of features to be selected)
- ► The performance of the method could be highly sensitive to the choice of these parameters

#### FEATURE SELECTION

- ► Reasonable Feature Selection is *critical* if not the most important component of model building.
- ➤ You cannot expect to build a good model if you select poor features.
- ► This is also called Feature Extraction
- ► We will talk about a few approaches that have been used in the literature.

# FEATURE SELECTION (RANKED BASED ON TEST-STATISTIC)

- ► Compute the two-sample t-test for all *m* features (based on the training set)
- ▶ Identify the top say 10 or 15 features (e.g, ranked based on the absolute value of the test statistic).
- ► Build a model on these "top" features (based on the training set)
- ► Alternatively, you could select all features for which the *P*-value is less than a certain threshold (say 0.001).
- ► You can also use the Wilcoxon rank sum statistic to protect against choosing features with outliers.

## FEATURE SELECTION (ORDINATION METHODS)

- ► A standard approach for reducing the dimension in the microarray setting is the method of Principal Components (PCs)
- ► The PCs are combinations of the original variables (gene expressions) that have maximum variability
- ► The are also constructed as to be uncorrelated with another
- ► This attempts to address the issue of high dimension and multi-collinearity simultaneously.
- ► One can use the principal components (say the first two or three) as the features
- ► Alternatively, one can first reduce the dimension by using the two-sample test-statistic approach and then get the PCs

#### TUNING

- ► You cannot expect to be able to build a model using default values provided by the software package.
- ► If you use k-NN you need to decide which k (e.g., 3 or 5 or 7) you want to use
- ► If you use the simple feature selection method you need to determine how many "top" features you want to use
- ► If you are doing PC dimension reduction, you need to determine how many PCs you want to use.
- ► In some books and articles, "tuning" only refers to the choice of the model parameter (e.g., k in k-NN)
- ► Must take a broader perspective as the choices in the FS part also affect the results.

#### VALIDATION

- ► Split the data into a training and a mutually exclusive testing set
- ► Build the model (including feature selection, tuning) on the *training* set
- ightharpoonup Evaluate the performance of the model on the  $testing\ set$
- ► IMPORTANT: The model is built based on the *training* set. The *testing* set should not contribute *any* information.
- ▶ Violating this principle will invariably result in bias

#### ERROR SUBSTITUTION VALIDATION

- ► Error Substitution Validation: The testing set is empty.
- ► Test the model you just built on the *training* set
- ► This approach cannot be recommended under any circumstance.
- ► Analogy: Assess the fit of the linear model by plotting the fitted (from the data) to the observed data.
- ► A bona-fide testing set is required.
- ▶ Will demonstrate how this can lead to noise discovery

#### HOLD-OUT METHOD

- ► Split the data into two parts
- $\blacktriangleright$  Keep the testing set locked up
- ▶ Better yet, ask an "honest" broker to keep it from you until you are ready to test the model
- ► This approach is reasonable if you have a large number of cases
- ▶ It may be problematic if the outcomes are sparse

#### k-fold Cross-Validation

- ► Many microarray experiments are from smaller (e.g., pilot) studies
- ► It is not impossible to get reasonably size training and testing sets this cases
- $\blacktriangleright$  A reasonable approach to get around this is k-fold cross-validation (CV)
- $\blacktriangleright$  Randomly split cases into k (nearly) equally sized subsets (folds).
- ▶ At each step take of these k portions as the *testing* set and construct the *training* set based on the other k-1 portions
- ▶ Special case is Leave-One-Out CV (LOOCV) where k = n
- ► For really small data sets, LOOCV is often the best (most practical) choice.

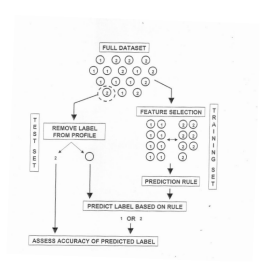
#### NAIVE CROSS-VALIDATION

- $\blacktriangleright$  Naive Validation: Do the feature selection once based on all n cases
- ► In each CV step use the same set of features.
- ► This will invariably make the results look better than they really are
- ► It should be avoided unless one feels *very* certain about the features (say biologically relevant gathered *a priori*

#### PROPER CROSS-VALIDATION

- ▶ Choose the first fold and set it aside the other k-1 folds
- ▶ Carry out Feature Selection on the other k-1 folds
- ▶ Train the model based the top features on the k-1 folds
- ► Test the model on the first fold left out
- ▶ Repeat the above for the second fold (set aside the second fold, leave in the first and the next k-2 folds).

# IMPORTANT ILLUSTRATION (FIG 8.5) FROM SIMON ET AL.



#### SIMULATE DATA FOR k-NN PREDICTION

► Simulate expression from 1000 genes for 40 patients. Let the first 20 be responders and the remaining 20 be non-responders

```
set.seed(123)
n = 20
m = 1000

EXPRS = matrix(rnorm(2 * n * m), 2 * n, m)
rownames(EXPRS) = paste("pt", 1:(2 * n), sep = "")
colnames(EXPRS) = paste("g", 1:m, sep = "")
grp = rep(0:1, c(n, n))
```

▶ Pick the top 10 features based on the two-sample t-test

```
stats = abs(rowttests(t(EXPRS), factor(grp))$statistic)
ii = order(-stats)
```

➤ Filter out all genes except the top 10

```
TOPEXPRS = EXPRS[, ii[1:10]]
```

#### ERROR RESUBSTITUTION AND NAIVE CV

► Error resubstitution (Training and Testing set are the same)

```
mod0 = knn(train = TOPEXPRS, test = TOPEXPRS, c1 = grp, k = 3)
table(mod0, grp)

## grp
## mod0 0 1
## 0 17 0
## 1 3 20
```

 Cross-validated predictions (the features selection is not part of the CV process)

```
mod1 = knn.cv(TOPEXPRS, grp, k = 3)
table(mod1, grp)

## grp
## mod1 0 1
## 0 16 0
## 1 4 20
```

▶ Note that in both examples, TOPEXPR not EXPR is used.

# R FUNCTION TO IMPLEMENT PROPER CV BASED ON k-NN

```
top.features <- function(EXP, resp, test, fsnum) {
    top.features.i <- function(i, EXP, resp, test, fsnum) {
        stats <- abs(mt.teststat(EXP[, -i], resp[-i], test = test))</pre>
        ii <- order(-stats)[1:fsnum]
       rownames(EXP)[ii]
    sapply(1:ncol(EXP), top.features.i, EXP = EXP, resp = resp, test = test,
        fsnim = fsnim)
# This function evaluates the knn
knn.loocv <- function(EXP, resp, test, k, fsnum, tabulate = FALSE, permute = FALSE) {
    if (permute)
        resp = sample(resp)
    topfeat = top.features(EXP, resp, test, fsnum)
    pids = rownames(EXP)
    EXP = t(EXP)
    colnames(EXP) = as.character(pids)
    knn.loocv.i = function(i, EXP, resp, k, topfeat) {
        ii = topfeat[, i]
        mod = knn(train = EXP[-i, ii], test = EXP[i, ii], cl = resp[-i], k = k)[1]
    out = sapply(1:nrow(EXP), knn.loocy.i, EXP = EXP, resp = resp, k = k, topfeat = topfeat)
    if (tabulate)
        out = ftable(pred = out, obs = resp)
    return(out)
```

#### PROPER CROSS-VALIDATION

- Finally, we conduct proper cross-validation using the previous R function
- ▶ At each iteration, the top 10 features are selected based on the data from the n-1 samples in the training set

- ► Note that EXPRS not TOPEXPR is used.
- ► The classification rate is 50% (as expected)

### NAIVE LOOCV: QUANTITATIVE TRAIT

- Repeat the last experiment with a noisy quantitative outcome
- First simulate a data matrix of dimension n = 50 (patients) and m (genes)
- Next draw the outcome for n = 50 patients from a standard normal distribution independent of the data matrix
- ► There is no relationship between the expressions and the outcome (by design)
- We consider m = 45 and m = 50000
- ► We conduct Naive LOOCV using the top 10 features

## NAIVE LOOCV: QUANTITATIVE TRAIT

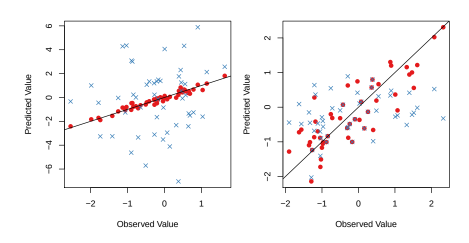


Figure taken from Owzar et al; Clin Transl Sci 2011.

## TRAINING, VALIDATION AND TESTING APPROACH

- ▶ Before you test the model, you must freeze it
- ► You may want to split the Training set further into a Training and Validation set
- ▶ Use the Validation set to "tune" the model.

#### FINAL REMARKS

- ► It is OK to try different methods (other classifiers, feature selection or tuning methods)
- ► Keep track of what you have done and report it (brief description in the paper and details in supplementary material)
- ▶ Be careful if you have too few responders
- ► You could have a model that will classify most patients as a non-responder.
- ▶ In this case a 00 (Y = 0 and g(X) = 0) may not be bona-fide true-negative
- ► The gold-standard for model validation, is to follow up the cross-validation by permutation resampling
- ► The R function provided can be used for this purpose

#### Pre-Processing Challenge

- ► The X profiles from the testing set need to be "compatible" to those used to train the model
- ► In classical experiments with a few biomarkers, the labs had internal controls to ensure that the measurements were properly normalized
- ► For RNA-Seq data, you observe counts (not expressions)
- ► Number of reads mapped to genes are *not* comparable
- ► Why?
- ► The current "state" of the art is to "normalize" the counts into expressions
- ► This is a practical but not rigorous solution
- ▶ One has to up the ante if the classifier is to be used for important decision (e.g., treating a patient with a toxic but potentially effective drug)

#### ON DATA AND ANSWERS

"The data may not contain the answer. The combination of some data and an aching desire for an answer does not ensure that a reasonable answer can be extracted from a given body of data."

John Wilder Tukey