## High-Throughput Sequencing Course Receiver Operator Characteristic

Biostatistics and Bioinformatics



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#### RECEIVER OPERATOR CHARACTERISTIC

► A graphical tool that can be used to evaluate the performance of a a binary classifier of the form

$$g(m) = \begin{cases} 1 & M \ge m \\ 0 & M < m, \end{cases}$$

where M is a quantitative marker

- ► Example: *M* could denote the predicted probability of a "positive" response from your genomic classifier
- ► The ROC curve is a representation of sensitivity as a function of the specificity of the classifier
- ► The ROC contrasts sensivity and specificity of the classifier as the treshold is varied

## CONFUSION MATRIX: TERMS

#### Among n samples

- $\triangleright$  P: The number of *positive* samples
- $\blacktriangleright$  N: The number of *negative* samples
- ightharpoonup TP: Number of true-positives
- ightharpoonup TN: Number of true-negatives

## Confusion Matrix: Relationship among terms

- ightharpoonup n = P + N
- ightharpoonup P = TP + FN
- ightharpoonup N = TN + FP
- $\qquad \qquad n = TP + FP + TN + FN$

## CONFUSION MATRIX: LAYOUT

		Prediction		
		Positive	Negative	
Truth	Positive	TP	FN	
	Negative	FN	TN	

## SENSITIVITY AND SPECIFICITY

► Sensitivity is defined as the proportion of "positive" samples correctly predicted to be "positive"

Sensitivity = 
$$\frac{TP}{P}$$

► Specificity is defined as the proportion of "negative" samples correctly predicted to be "negative"

Specificity = 
$$\frac{TN}{N}$$

## RECEIVER OPERATOR CHARACTERISTICS (ROC)

- $\blacktriangleright$  Let M be the value of the marker
- $\blacktriangleright$  In our case, M could be the predicted probability of a positive event from our classifier
- $\blacktriangleright$  For each of the *n* samples we have (Y, M)
- $\blacktriangleright$  For a given "cutoff" m let

$$\hat{Y} = g(m) = \begin{cases} 0 & M < m \\ 1 & M \ge m. \end{cases}$$

- ▶ In other words, "plug in" the observed value of the marker M into g(m) to get either a positive or negative prediction
- ► ROC : A plot of Sensitivity versus 1-Specificity at a given cutoff m

## EXAMPLE

Table: Toy Example

		m=1		m=2	
M	Y	Yhat	Res	Yhat	Res
1.1	0	1	FP	0	TN
2.3	1	1	TP	0	FN
0.9	0	0	TN	0	TN
3.1	0	1	FP	1	FP
2.1	1	1	TP	0	FN
2.5	1	1	TP	1	TP
0.1	1	0	FN	0	FN

Note: the prediction  $\hat{Y}$  does not require that the true state (Y) is known. The result (TP, TN, FP, FN) does.

#### SIMULATE DATA FROM LOGISTIC MODEL

This function will simulate n samples from the following logistic model

$$\log \frac{P(Y=1)}{1 - P(Y=1)} = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3$$

```
sim.binomial <- function(n, b0, b1, b2, b3) {
    x1 <- rnorm(n)
    x2 <- rnorm(n)
    x3 <- rnorm(n)
    1x <- b0 + b1 * x1 + b2 * x2 + b3 * x3
    px <- exp(1x)/(1 + exp(1x))
    y <- rbinom(n, 1, px)
    data.frame(y, px, x1, x2, x3)
}
# apply(replicate(1000,glm(y*x1+x2+x3,family='binomial',data=sim.binomial(100,0.1,0,0,0))£coef),1,mean)</pre>
```

## SIMULATE LOGISTIC MODEL

- ▶ The model has three features  $X_1, X_2$  and  $X_3$
- ▶ For each sample in the training set we have  $(Y, X_1, X_2, X_2)$  where Y = 1 (positive) or Y = 0 (negative)
- ightharpoonup The model is trained on the basis of data from n samples
- ► The "trained" model is of the form

$$\log \frac{\hat{P}(Y=1)}{1 - \hat{P}(Y=1)} = \hat{\beta}_0 + \hat{\beta}_1 x_1 + \hat{\beta}_2 x_2 + \hat{\beta}_3 x_3$$

ightharpoonup For a "new" sample, the marker M on the basis of this trained model is

$$M = \frac{\exp(\hat{\beta}_0 + \hat{\beta}_1 x_1 + \hat{\beta}_2 x_2 + \hat{\beta}_3 x_3)}{1 + \exp(\hat{\beta}_0 + \hat{\beta}_1 x_1 + \hat{\beta}_2 x_2 + \hat{\beta}_3 x_3)}$$

#### LOOCV FUNCTION FOR LOGISTIC REGRESSION

```
logistic.loocv <- function(simdat) {
    preds <- foreach(i = 1:nrow(simdat), .combine = rbind) %do% {
        ## remove the i-th sample from the training data set
        traindati <- simdat[-i, ]
        ## Get the data for the left out sample
        testi <- simdat[i, , drop = FALSE]
        ## Fit a logistic model based on this data set
        trainmodi <- glm(y ~ x1 + x2 + x3, family = binomial, data = traindati)
        ## Predicted probability the left out sample against the trained model
        probhat <- predict(trainmodi, newdata = testi, type = "response")</pre>
        ## Predicted outcome: If the predicted probability is < 0.5 predict as 0 or 1
        ## otherwise
        vhat <- ifelse(probhat < 0.5, 0, 1)</pre>
        ## Output data.frame
        data.frame(y = testi$y, yhat = yhat, probhat = probhat)
    return(preds)
```

# FUNCTIONS TO CALCULATE ROC AND AUC USING THE ROCR PACKAGE

```
### Helper function to produce ROC and AUC using the ROCR package The
### arguments are m: A positive marker y: The outcome (0 or 1)

ROC <- function(preddat, mlab = "probhat", ylab = "y") {
    m <- preddat[[mlab]]
    y <- preddat[[ylab]]
    performance(prediction(m, y), measure = "tpr", x.measure = "fpr")
}

AUC <- function(preddat, mlab = "probhat", ylab = "y") {
    m <- preddat[[mlab]]
    y <- preddat[[ylab]]
    performance(prediction(m, y), "auc") @y.values[[1]]
}</pre>
```

# FUNCTION TO CALCULATE CONFUSION MATRIX FOR A GIVEN CUTOFF

```
confusion <- function(cutoff, preddat, mlab = "probhat", ylab = "y") {</pre>
    m <- preddat[[mlab]]
    y <- preddat[[ylab]]
    vhat <- ifelse(m < cutoff, 0, 1)</pre>
    ### Calculate TN, FN, FP and TP
    TN <- sum(v == 0 & vhat == 0)
    FN \leftarrow sum(y == 1 & yhat == 0)
    FP \leftarrow sum(y == 0 \& yhat == 1)
    TP <- sum(y == 1 & yhat == 1)
    ### Get P and N
    P <- TP + FN
    N <- TN + FP
    ### get sensitivity, specificity, and 1-spec
    sens <- TP/P
    spec <- TN/N
    FNR <- 1 - spec
    data.frame(cutoff, n = P + N, P, N, sens, spec, FNR)
```

## ROC: Analysis

#### ► Simulate data

```
set.seed(31219)
mydat <- sim.binomial(100, 0.1, 0, 1, 0)</pre>
```

#### ► Perform LOOCV

```
mypreds <- logistic.loocv(mydat)</pre>
```

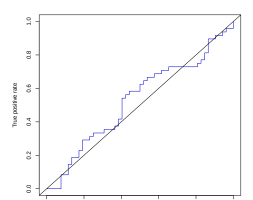
#### ► Calculate ROC

```
myroc <- ROC(mypreds)</pre>
```

## $\underset{\mathrm{Get}}{\mathrm{ROC}} : \underset{\mathrm{AUC}}{\mathrm{PLOT}}$

```
AUC(mypreds)
## [1] 0.5300481
```

```
plot(myroc, col = "blue3")
abline(0, 1)
```



## ADD RESUBSTITION ROC

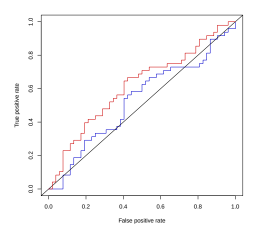
```
### Fit model using ALL n samples
trainmod <- glm(y ~ x1 + x2 + x3, family = binomial, data = mydat)
### Get the predicted probabilities
phat <- predict(trainmod, type = "response")
### Compare AUCs
AUC(mypreds)

## [1] 0.5300481

AUC(data.frame(probhat = phat, y = mydat$y))
## [1] 0.6169872</pre>
```

## SHOW RESUBSTITION ROC

```
### Plot LOCCV ROC
plot(myroc, col = "blue3")
abline(0, 1)
### Add resub ROC curve
plot(ROC(data.frame(probhat = phat, y = mydat$y)), col = "red3", add = TRUE)
```



#### LOOK AT THE PERFORMANCE OBJECT

## CALCULATE USING OUR OWN CONFUSION FUNCTION

```
perf3
       cutoff
##
                     tpr
                               fpr
## 1 0.6586603 0.06250000 0.07692308
## 2 0.6516634 0.08333333 0.07692308
## 3 0.6512297 0.08333333 0.09615385
confusion(perf3$cutoff[1], mypreds)
       cutoff n P N sens
                                              FNR
##
                                   spec
## 1 0.6586603 100 48 52 0.0625 0.9230769 0.07692308
confusion(perf3$cutoff[2], mypreds)
       cutoff n P N sens
                                                  FNR
                                       spec
## 1 0.6516634 100 48 52 0.08333333 0.9230769 0.07692308
confusion(perf3$cutoff[3], mypreds)
       cutoff n P N sens
                                       spec
## 1 0 6512297 100 48 52 0 08333333 0 9038462 0 09615385
```

## ROC: Analysis with larger effect size

#### ► Simulate data

```
set.seed(31219)
mydat <- sim.binomial(100, 0.1, 0, 2, 0)</pre>
```

#### ► Perform LOOCV

```
mypreds <- logistic.loocv(mydat)</pre>
```

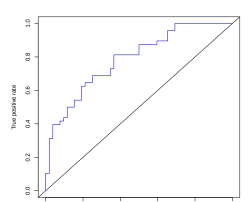
#### ► Calculate ROC

```
myroc <- ROC(mypreds)</pre>
```

## $\underset{\mathrm{Get}}{\mathrm{ROC}} : \underset{\mathrm{AUC}}{\mathrm{PLOT}}$

```
AUC(mypreds)
## [1] 0.7864583
```

```
plot(myroc, col = "blue3")
abline(0, 1)
```



## ADD RESUBSTITION ROC

```
### Fit model using ALL n samples
trainmod <- glm(y ~ x1 + x2 + x3, family = binomial, data = mydat)
### Get the predicted probabilities
phat <- predict(trainmod, type = "response")
### Compare AUCS
AUC(mypreds)

## [1] 0.7864583

AUC(data.frame(probhat = phat, y = mydat$y))
## [1] 0.8165064</pre>
```

## SHOW RESUBSTITION ROC

```
### Plot LOCCV ROC
plot(myroc, col = "blue3")
abline(0, 1)
### Add resub ROC curve
plot(ROC(data.frame(probhat = phat, y = mydat$y)), col = "red3", add = TRUE)
```

