Homework 1

Pattern Mining and Social Network Analysis

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Classification

Overall

Classification algorithms have categorical responses. In classification we build a function f(X) that takes a vector of input variables X and predicts its class membership, such that Y in C.

Possibilities of models

There are classifiers as logistic regression, Decision trees, Perceptron / Neural networks, K-nearest-neighbors, linear and quadratic logistic regression, Bayes classifiers ...

Some indicators to analyze the results

Sensitivity and recall

The sensitivity (also named recall) is the percentage of true defaulters that are identified (True positive tests). For example, probability of predicting disease given true state is disease.

$$sensitivity = recall = \frac{TruePositiveTests}{PositivePopulation}$$

Specificity

The specificity is the percentage of non-defaulters that are correctly identified (True negative tests). 1 - specificity is the Type 1 error, it is the false positive rate. For example, probability of predicting non-disease given true state is non- disease.

$$specificity = \frac{TrueNegativeTests}{NegativePopulation}$$

Precision

The precision is the proportion of true positive tests among the positive tests.

$$precision = \frac{TruePositiveTests}{PositiveTests}$$

F-Mesure

The traditional F measure is calculated as follows:

$$F_{M}easure = \frac{(2*Precision*Recall)}{(Precision+Recall)}$$

Rand index

The rand index is a mesure of similarity between two partitions from a single set.

Given two partitions π_1 and π_2 in E :

- a, the number of elements in π_1 and π_2
- b, the number of elements in π_1 and not in π_2
- c, the number of elements in π_2 and not in π_1
- d, the number of elements not in both π_1 and π_2

	in π_2	not in π_2
in π_1	a	b
not in π_1	c	d

$$RI(\pi_1,\pi_2) = \frac{a+d}{a+b+c+d}$$

Mutual information

Mutual information is calculated between two variables and measures the reduction in uncertainty for one variable given a known value of the other variable. The mutual information between two random variables X and Y can be stated formally as follows:

$$MI = I(X;Y) = H(X) - H(X|Y)$$

Cross Entropy(log loss)

Cross-entropy loss, or log loss, measures the performance of a classification model whose output is a probability value between 0 and 1. Cross-entropy loss increases as the predicted probability diverges from the actual label. So predicting a probability of .012 when the actual observation label is 1 would be bad and result in a high loss value. A perfect model would have a log loss of 0.

In binary classification, where the number of classes M equals 2, cross-entropy can be calculated as:

$$CE = -(y \log(p) + (1 - y) \log(1 - p))$$

If M>2 (i.e. multiclass classification), we calculate a separate loss for each class label per observation and sum the result.

$$CE = -\sum_{c=1}^b y_{o,c} \log(p_{o,c})$$

Accuracy of a model

How do we determine which model is best? Various statistics can be used to judge the quality of a model. These include Mallow's C_p , Akaike information criterion (AIC), Bayesian information criterion (BIC), and adjusted \mathbb{R}^2 .

MSE: Mean Squarred Error

Let's define the mean squared error or MSE.

$$MSE = \frac{1}{n} \sum_{i} 1_{y_i - \hat{f}(x_i)}$$

where:

$$1_{y_i - \hat{f}(x_i)} = \left\{ \begin{array}{ll} 1 & \text{if } y_i! = \hat{f}(x_i) \\ 0 & \text{otherwise} \end{array} \right.$$

Recall:

$$RSS = MSE * n$$

RSS and \mathbb{R}^2 are not suitable for selecting the best model among a collection of models with different numbers of predictors.

Mallow's Cp

Mallows's Cp addresses the issue of overfitting, in which model selection statistics such as the residual sum of squares always get smaller as more variables are added to a model

If there are d predictors:

$$C_p = \frac{RSS + 2d\hat{\sigma}^2}{n}$$

AIC: Akaike information criterion

The Akaike information criterion (AIC) is an estimator of in-sample prediction error and thereby relative quality of statistical models for a given set of data.[1] In-sample prediction error is the expected error in predicting the resampled response to a training sample

The AIC criterion is defined for a large class of models fit by maximum likelihood.

$$AIC = \frac{RSS + 2d\hat{\sigma}^2}{n\hat{\sigma}^2}$$

To use AIC for model selection, we simply choose the model giving small- est AIC over the set of models considered.

BIC: Bayesian information criterion

In statistics, the Bayesian information criterion or Schwarz information criterion is a criterion for model selection among a finite set of models; the model with the lowest BIC is preferred. It is based, in part, on the likelihood function and it is closely related to the Akaike information criterion

BIC is derived from a Bayesian point of view, but ends up looking similar to Cp (and AIC) as well. For the least squares model with d predictors, the BIC is, up to irrelevant constants, given by

$$BIC = \frac{RSS + log(n)d\hat{\sigma}^2}{n}$$

Adjusted R statistic

The adjusted R-squared is a modified version of R-squared that has been adjusted for the number of predictors in the model. The adjusted R-squared increases only if the new term improves the model more than would be expected by chance. It decreases when a predictor improves the model by less than expected by chance

$$AdustedR^{2} = 1 - \frac{\frac{RSS}{n-d-1}}{\frac{TSS}{n-1}}$$

Logistic Regression

How it works

In logistic regression, for covariates (X_1 , . . . , X_p), we want to estimate $p_i = P_r(Y_i = 1 | X_1, ..., X_p)$

$$p_i = \frac{e^{\beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \beta_4 x_{i4} + \dots}}{1 + e^{\beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \beta_4 x_{i4} + \dots}}$$

To come back to linear regression we define the logistic function as follow.

$$logit(p_i) = log(\frac{p_i}{1 - p_i}) = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \beta_4 x_{i4} + \dots$$

We can define the odds :

$$\frac{odds(Y_i=1|X1=x_{i1}+1)}{odds(Y_i=1|X1=x_{i1})}=e^{\beta_1}$$

Which indicator to construct the model?

We use Maximum Likehood:

$$L(\beta) = \prod_{i=1}^{n} p_i^{y_i} * (1 - p_i)^{y_i}$$

The goal is to maximise it by adjusting β vector.

Example on the Wimbledon tennis tournament

We use a dataset from the Wimbledon tennis tournament for Women in 2013. We will predict the result for player 1 (win=1 or loose=0) based on the number of aces won by each player and the number of unforced errors committed by both players. The data set is a subset of a data set from https://archive.ics.uci.edu/ml/datasets/Tennis+Major+Tournament+Match+Statistics.

##		Player1	Player2	${\tt Result}$	ACE.1	UFE.1	ACE.2	UFE.2	${\tt ACEdiff}$	${\tt UFEdiff}$
##	1	M.Koehler	V.Azarenka	0	2	18	3	14	-1	4
##	2	E.Baltacha	F.Pennetta	0	0	10	4	14	-4	-4
##	3	S-W.Hsieh	T.Maria	1	1	13	2	29	-1	-16
##	4	A.Cornet	V.King	1	4	30	0	45	4	-15
##	5	Y.Putintseva	K.Flipkens	0	2	28	6	19	-4	9
##	6	A.Tomljanovic	B.Jovanovski	0	6	42	11	40	-5	2

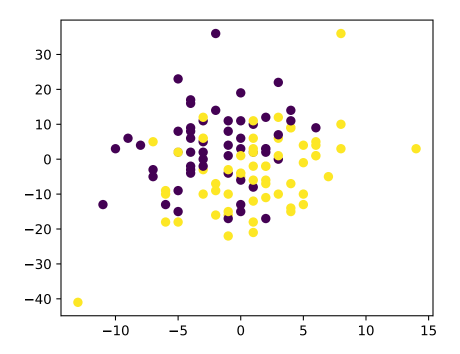
Looking if the dataset is balanced :

1 63 ## 0 55

Name: Result, dtype: int64

##	Result	ACE.1	UFE.1	 UFE.2	ACEdiff	UFEdiff
## count	118.000000	118.000000	118.000000	 118.000000	118.000000	118.000000
## mean	0.533898	2.974576	20.177966	 20.466102	-0.296610	-0.288136
## std	0.500977	2.835857	10.248728	 11.444912	4.356564	11.410822
## min	0.000000	0.000000	4.000000	 2.000000	-13.000000	-41.000000
## 25%	0.000000	1.000000	13.000000	 12.000000	-3.000000	-7.750000
## 50%	1.000000	2.000000	18.000000	 18.000000	0.000000	1.000000
## 75%	1.000000	4.000000	25.750000	 27.000000	2.000000	6.000000
## max	1.000000	14.000000	54.000000	 55.000000	14.000000	36.000000
##						

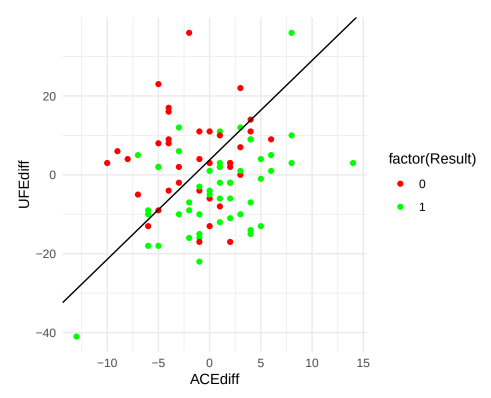
[8 rows x 7 columns]



On R

```
##
## Call:
## glm(formula = Result ~ ACEdiff + UFEdiff, family = "binomial",
       data = tennisTrain)
##
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                           Max
   -2.1204 -0.9994
                      0.5662
                               0.8918
                                        1.8714
##
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) 0.31318
                           0.24439
                                     1.281 0.20004
## ACEdiff
                0.20856
                           0.06575
                                     3.172 0.00151 **
## UFEdiff
               -0.08272
                                   -3.371 0.00075 ***
                           0.02454
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
   (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 120.352
                              on 87 degrees of freedom
## Residual deviance: 99.102 on 85 degrees of freedom
## AIC: 105.1
##
## Number of Fisher Scoring iterations: 4
```

With the model, we can draw the slope which indicates the category of a point.



We can write:

$$logit(p_i) = log(\frac{p_i}{1 - p_i}) = 0,31318 + 0,20856 * ACEDiff - 0,08272 * UFEDiff$$

We can observe AIC = 105.1

The confusion matrix is :

glm.Result_pred 0 1 ## 0 15 5 ## 1 2 8

The accuracy rate is $\frac{15+8}{30} = 0.7667$.

The sensitivity is the percentage of true output giving Player1-winner among the population of true Player1-winner:

[1] 0.6153846

The specificity is the percentage of true output giving Player2-winner (= Player1-looser) among the population of true Player2-winner:

[1] 0.8823529

The precision is the percentage of true output giving Player1-winner among all the outputs giving Player1-winner (even if not winner):

[1] 0.8

So the F_Mesure is :

[1] 0.6956522

On Python with Scikit-learn

```
## Mean accuracy is :
## 0.7499502537460511
## Mean performance is :
## [9.53, 3.38, 4.07, 12.73]
## The Sensitivity is : 0.700735294117647
```

The specificity is : 0.7901924270639354 ## The precision is : 0.7381874515879163

So, we can deduce that the F-mesure is: 0.7189739720860052

Using the Logistic Regression model in scikit we obtain an accuracy of 0.74. In average, this model has 9.48 True Positive, 3.23 False Positive, 4.4 False Negative, 12.63 True Positive.

Decision trees and Random Forest

Decision Tree

Decision Tree algorithm belongs to the family of supervised learning algorithms. Unlike other supervised learning algorithms, the decision tree algorithm can be used for solving regression and classification problems too.

The goal of using a Decision Tree is to create a training model that can use to predict the class or value of the target variable by learning simple decision rules inferred from prior data(training data).

Decision trees use multiple algorithms to decide to split a node into two or more sub-nodes. The creation of sub-nodes increases the homogeneity of resultant sub-nodes. In other words, we can say that the purity of the node increases with respect to the target variable. The decision tree splits the nodes on all available variables and then selects the split which results in most homogeneous sub-nodes.

Random Forest

Random forests or random decision forests are an ensemble learning method for classification, regression and other tasks that operate by constructing a multitude of decision trees at training time and outputting the class that is the mode of the classes or mean/average prediction of the individual trees.

Which indicator to construct the model?

Entropy Entropy is a measure of the randomness in the information being processed. The higher the entropy, the harder it is to draw any conclusions from that information. Flipping a coin is an example of an action that provides information that is random.

Mathematically Entropy for 1 attribute is represented as:

$$E(s) = \sum_{i=1}^c -p_i \log_2(p_i)$$

Mathematically Entropy for multiple attributes is represented as:

$$E(T,X) = \sum_{c \in X} P(c)E(c)$$

Gini You can understand the Gini index as a cost function used to evaluate splits in the dataset. It is calculated by subtracting the sum of the squared probabilities of each class from one. It favors larger partitions and easy to implement whereas information gain favors smaller partitions with distinct values.

$$Gini = 1 - \sum_{i=1}^{c} (p_i)^2$$

Example on the Wimbledon tennis tournament

On R

[1] 2

MSE R2 RMSE MAE ## 1 0.2074194 0.1835982 0.4554331 0.4063749

The confusion matrix is:

##

model.Result_pred 0 1 ## 0 10 4 ## 1 7 9

The accuracy rate is:

[1] 0.6333333

The sensitivity is the percentage of true output giving Player1-winner among the population of true Player1-winner:

[1] 0.6923077

The specificity is the percentage of true output giving Player2-winner (= Player1-looser) among the population of true Player2-winner:

[1] 0.5882353

The precision is the percentage of true output giving Player1-winner among all the outputs giving Player1-winner (even if not winner):

[1] 0.5625

So the F_Mesure is :

[1] 0.6206897

On Python with Scikit-learn

0.6875003547642073

[8.69, 4.14, 5.19, 11.82]

The Sensitivity is : 0.6260806916426513 ## The specificity is : 0.7406015037593985

The precision is : 0.6773187840997662

So, we can deduce that the F-mesure is : 0.6506926244852115

KNN: K-nearest neighbors

How does it work?

To make predictions for an observation x, the KNN use the training observations to find the k closest training observations to x. Then x is assigned to the class to which the most of these observations belong. We just need to definine the notion of distance which can be euclidean.

Comments

KNN is very good for non-linear classification.

However it doesn't give the coefficient for the predictors so we can't see their impacts.

It also needs a lot od training observations well-balanced. Otherwise if a class is over-represented, it would assign too much this label.

Choosing k

To choose k, it can be useful to use cross-validation. We test multiple times the model with different k and we choose the one with the smallest error.

Discriminant Analysis

Discriminant analysis is a technique that helps classifying 2 or more groups into clusters. This technique uses - the Bayes' theorem - the covariance (the shape of one group of point) - and the center of this group.

The aim is to use those 3 components to create boundaries between the classes. The boundaries are composed by a set of points that have the same chance to belong to either classes.

There is two types of discriminant Analysis. The first one is the Linear Discriminant Analysis (LDA). To use that model, each class need to be drawn from a multivariate Gaussian distribution and the covariance for each class needs to be equals (the groups of points need to have the same shape).

Quadratic Discriminant Analysis (QDA) is a bit similar to (LDA) because it assumes that each class are drawn from a Gaussian distribution. However, the covariance doesn't need to be the same for each class.

The advantages of such models is that they have a good accuracy and they are robust to outliers (because it is based on statistical distribution of all the observations).

Support Vector machines

Maximal margin classifier

A hyperplane in p dimensions can be written as follow:

$$\beta_0+\beta_1*x_1+\beta_2*x_2+\ldots+\beta_p*x_p=\beta_0+X^T\beta=0$$

It is a a p-1 dimensional subspace of R^p

The hyperplane leads to a natural classifier, depending on the side of the hyperplane where the new observation lies.

If $-0 + X^T > 0$: it lies on one side of the hyperplane meaning it belongs to class labelled 1. If $-0 + X^T < 0$: it lies on the opposite side of the hyperplane meaning it belongs to class labelled -1.

Thus $y_i * (\beta_0 + X^T \beta) > 0$

The distance of any point x to the hyperplane is given by:

$$d = \frac{1}{||\beta||}(\beta_0 + X^T\beta)$$

The best hyperplane is the maximal margin hyperplane which maximises the distance d from the training observations.

It is an optimization problem: we want to classify well all observations and maximize d.

Let's call M, the minimal distance from the observations to the hyperplan. We write the problem:

$$\max_{\beta_0,\beta} M$$
 (1) subject to $\Sigma_j \beta_j^2 = 1$

for all training observations $y_i * (\beta_0 + X^T \beta) > M$

(2)

(7)

Support vector classifier

For some data sets a separating hyperplane does not exist, the data set is non-separable.

To obtain a support vector classifier we relax the conditions that we had for the maximal margin hyperplane by allowing for a "budget" C of misclassifications.

We write the new problem:

$$\max_{\beta_0,\beta} M \tag{3}$$

subject to
$$\Sigma_j \beta_j^2 = 1$$
 (4)

for all training observations
$$y_i*(\beta_0+X^T\beta)>M*(1-\epsilon_i) \eqno(5)$$

$$0 \le \epsilon_i \text{and} \Sigma_i \epsilon_i \le C \tag{6}$$

Once we have the hyperplane, we can deduce to which class the observation belong by projection.

Support Vector Machines

For some datasets a non-linear decicion boundary between the classes is more suitable than a linear decision boundary.

To compute the boundary, we use interactions terms between predictors or functions on predictors.

An example can be:

$$\beta_0 + \beta_1 * x_1^2 + \beta_2 * x_2 * x_1 + \dots + \beta_p * x_p^3 = 0$$

$$f(X_i) = \beta_0 + \Sigma_i \beta_i * K(x_i, x_i)$$

where K is a function between x_i and x_j

We write the new problem:

$$\max_{\beta_0,\beta} M \tag{8}$$

for all training observations
$$y_i * f(X_i) > M * (1 - \epsilon_i)$$
 (9)

$$0 \le \epsilon_i \text{and} \Sigma_i \epsilon_i \le C \tag{10}$$

(11)

Once we have the hyperplane, we can deduce to which class the observation belong by projection.

Neural networks

Neural network structure

Neural networks can be decomposed as three parts the input the hidden layers and the output. Those parts are in fact nodes link together with edges that are given weights describing the strength of the connection. With a given input the value of the first node of the hidden layer will correspond to the inputs time the weights on the edges going to the node. This product will give a result that needs to go through an activation function before giving the value of the current node. This process of computing the values of the nodes from the inputs to the output is called feedforward.

Once, we have computed the output, we compare our prediction to the answer. Then, we backpropagate the error. It means that depending on the error, the value of the nodes and the learning rate, we are going to updates the weights.

Once the weights are updated we iterate until the amount of errors become acceptable.

Regression

Overall

Regression is a statistical method used in finance, investing, and other disciplines that attempts to determine the strength and character of the relationship between one dependent variable (usually denoted by Y) and a series of other variables (known as independent variables).

Accuracy of a model

MSE: Mean Squarred Error

The MSE mesures the mean accuracy of the predicted responses values for given observations. There are two MSE: the train MSE and the test MSE.

The train MSE is use to fit a model while training.

The test MSE is use to choose between models already trained.

Let's define the mean squared error or MSE.

$$MSE = \frac{1}{n} \sum_{i} (y_i - \hat{f}(x_i))^2$$

where $\hat{f}(x_i)$ is the prediction of y_i obtained with the model.

Then the expected test MSE refers to the average test MSE that we would obtain if we repeatedly estimated f using a large number of training sets, and tested each at x_0 . So that the expected test MSE is:

$$E(y_0 - \hat{f}(x_0))^2$$

$$E(y_0 - \hat{f}(x_0))^2 = Var(\hat{f}(x_0)) + (f(x_0) - E(\hat{f}(x_0)))^2 + Var(\varepsilon)$$

 $Var(\varepsilon)$ represents the irreductible error. This term can not be reduced regardless how well our statistical model fits the data.

 $(f(x_0) - E(\hat{f}(x_0))^2 = [Bias(\hat{f}(x_0))]^2$ is the squared Bias and refers to the error that is introduced by approximating a real-life problem, which may be extremely complicated, by a much simpler model. If the bias is low the model gives a prediction which is close to the true value.

 $Var(\hat{f}(x_0))$ is the Variance of the prediction at $\hat{f}(x_0)$ and refers to the amount by which \hat{f} would change if we estimated it using a different training data set. If the variance is high, there is a large uncertainty associated with the prediction.

RMSE: Root Mean Squared Error

Root Mean Squared Error (RMSE), which measures the average prediction error made by the model in predicting the outcome for an observation. That is, the average difference between the observed known outcome values and the values predicted by the model. The lower the RMSE, the better the model.

$$RMSE = \sqrt{MSE} = \sqrt{\frac{1}{n}\sum_i (y_i - \hat{f}(x_i))^2}$$

RSS: Residual Sum of Squares

We define the residual sum of squares (RSS) as the sum of the squares of residuals (deviations predicted from actual empirical values of data):

Since $\hat{f}(x_i) = a + b * x_i$ and $y_i = a + b * x_i + \epsilon_i$

$$RSS = \Sigma \epsilon_i^2 = \Sigma (y_i - \hat{f}(x_i))^2 = n * MSE$$

We want to minimize the RSS.

RSE: Residual Standard Error

The residual standard error is the square root of the residual sum of squares divided by the residual degrees of freedom. Mean Square Error. The mean square error is the mean of the sum of squared residuals, i.e. it measures the average of the squares of the errors. Lower values (closer to zero) indicate better fit.

$$RSE = \sqrt{\frac{1}{n-2}RSS}$$

R squared statistic

In statistics, the coefficient of determination, denoted R² or r² and pronounced "R squared", is the proportion of the variance in the dependent variable that is predictable from the independent variable

$$R^2 = 1 - \frac{RSS}{TSS}$$

$$TSS = \Sigma (y_i - \bar{y}_i)^2$$

is the total sum of squares. TSS measures the total variance in the response Y.

TSS – RSS measures the amount of variability in the response that is explained.

 R^2 measures the proportion of variability in Y that can be explained using X.

MAE : Mean Absolute Error

Mean Absolute Error (MAE), an alternative to the RMSE that is less sensitive to outliers. It corresponds to the average absolute difference between observed and predicted outcomes. The lower the MAE, the better the model is.

$$MAE = \frac{1}{n} \sum_i |y_i - \hat{f}(x_i)|$$

Simple Linear Regression

Definition

In statistics, linear regression is a linear approach to modeling the relationship between a scalar response (or dependent variable) and one or more explanatory variables (or independent variables). The case of one explanatory variable is called simple linear regression.

Simple linear regression lives up to its name: it is a very straightforward approach for predicting a quantitative response Y on the basis of a single predictor variable X. It assumes that there is approximately a linear relationship between X and Y. Mathematically, we can write this linear relationship as

$$Y \approx \beta_0 + \beta_1 * X$$

Multiple Linear Regression

Definition

Multiple linear regression (MLR), also known simply as multiple regression, is a statistical technique that uses several explanatory variables to predict the outcome of a response variable. Multiple regression is an extension of linear (OLS) regression that uses just one explanatory variable

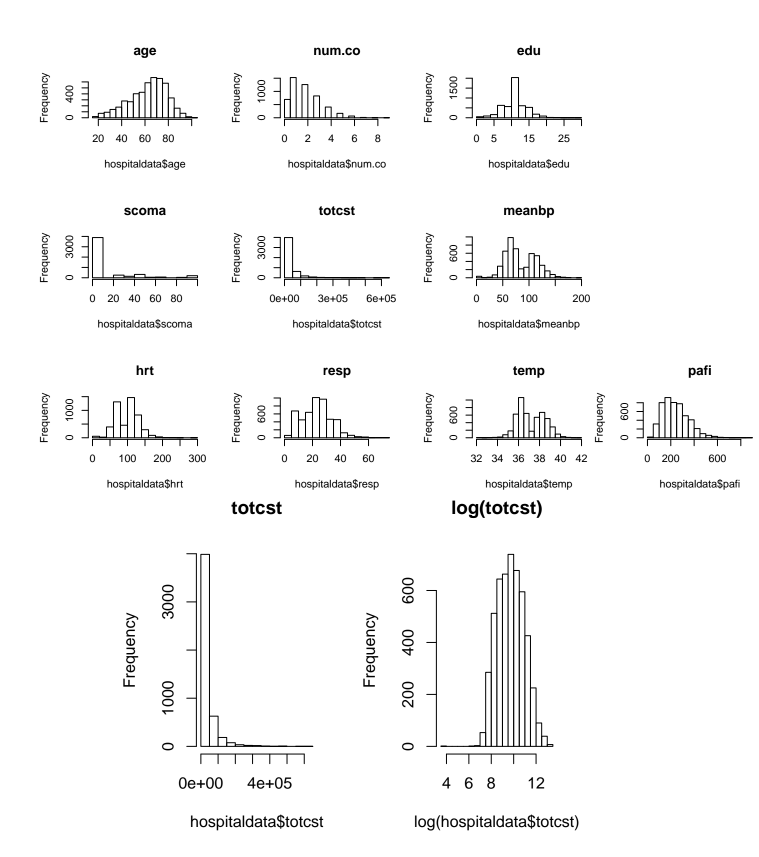
Formula and Calcualtion of Multiple Linear Regression

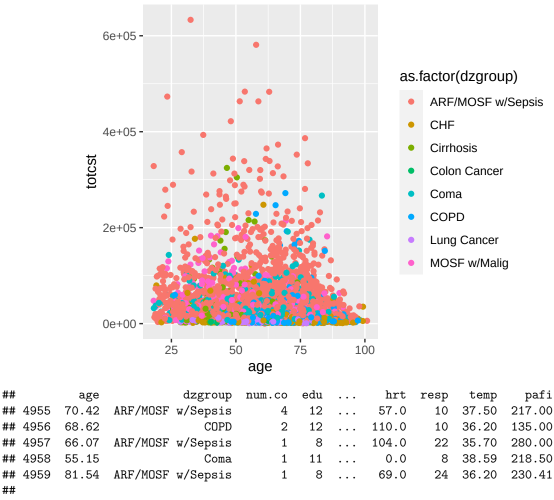
$$y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_p x_{ip} + \epsilon$$

Hospital Costs dataset

The next dataset (source F. E. Harrell, Regression Modeling Strategies) contains the total hospital costs of 9105 patients with certain diseases in American hospitals between 1989 and 1991. The different variables are:

##		age)	C	lzgroup	num.co	edu	iı	ncome	scoma	totcst	race	meanbp	hrt
##	1	62.85)	Lung	Cancer	0	11	\$11-	-\$25k	0	NA	other	97	69
##	2	60.34	ŀ	Cii	rhosis	2	12	\$11-	-\$25k	44	NA	white	43	112
##	3	52.75	j	Cii	rhosis	2	12	under	\$11k	0	NA	white	70	88
##	4	42.38	3	Lung	Cancer	2	11	under	\$11k	0	NA	white	75	88
##	5	79.88	ARF/N	OSF w	'Sepsis	1	NA			26	NA	white	59	112
##	6	93.02	2		Coma	1	14			55	NA	white	110	101
##		resp	temp	pafi	Ĺ									
##	1	22	36.00	388.00)									
##	2	34	34.59	98.00)									
##	3	28	37.40	231.66	3									
##	4	32	35.00	NA	Α									
##	5	20	37.90	173.31	L									
##	6	44	38.40	266.63	3									





[5 rows x 13 columns]

We can see that there is a lot of NaN values.

Remove NaN and null data:

##		age	num.co	 temp	pafi
##	count	4960.000000	4960.000000	 4960.000000	4960.000000
##	mean	62.668768	1.912298	 37.218907	241.739718
##	std	15.892083	1.397348	 1.301758	110.397054
##	min	18.120000	0.000000	 32.000000	12.000000
##	25%	52.597500	1.000000	 36.200000	156.977500
##	50%	65.090000	2.000000	 36.900000	226.660000
##	75%	74.472500	3.000000	 38.300000	308.000000
##	max	100.850000	9.000000	 41.700000	890.380000
##					
##	[8 row	s x 10 column	s]		

On R We would like to build models that help us to understand which predictors are mostly driving the total cost.

Looking at the distribution of the cost we see we should apply a log transformation for a better distribution.

We can calculate the MSE on the test set to evaluate the simple linear regression model.

```
##
                       (Intercept)
                                                                 age
                       7.848450651
                                                       -0.006610500
##
##
                              temp
                                                        0.026642892
                       0.075106886
##
##
                              resp
                                                             num.co
                      -0.004025088
                                                       -0.043124831
##
##
            as.factor(dzgroup)CHF
                                       as.factor(dzgroup)Cirrhosis
##
                      -1.405057960
                                                       -0.923605089
   as.factor(dzgroup)Colon Cancer
                                            as.factor(dzgroup)Coma
##
##
                      -1.458633036
                                                       -0.452564283
##
           as.factor(dzgroup)COPD
                                     as.factor(dzgroup)Lung Cancer
                      -1.242045040
                                                       -1.688831933
##
   as.factor(dzgroup)MOSF w/Malig
##
##
                      -0.256302673
           MSE
                       R2
                               RMSE
                                           MAE
## 1 0.8903145 0.3742044 0.9435648 0.7440975
```

On Python with Scikit-learn

```
## Ridge(alpha=0.5)
## [-0.00775316 -0.16374392 -0.11901479 0.02506879 -0.00197573 0.1099544 ]
## 6.48346400118024
```

Linear regression with interaction terms

Definition

It is a regression which introduces an operation between two predictors like multiplication, division ...

Hospital Costs dataset

On ${\bf R}$ We use the same example than for simple linear regression.

```
##
                           (Intercept)
                                                                        age
##
                          7.6621089858
                                                              -0.0042188616
##
                as.factor(dzgroup)CHF
                                               as.factor(dzgroup)Cirrhosis
##
                         -1.1077199267
                                                              -0.5713961342
##
       as.factor(dzgroup)Colon Cancer
                                                    as.factor(dzgroup)Coma
                         -1.3449083302
                                                               0.3636117761
##
##
               as.factor(dzgroup)COPD
                                             as.factor(dzgroup)Lung Cancer
##
                         -1.2782357031
                                                              -2.1009737479
       as.factor(dzgroup)MOSF w/Malig
##
                                                                        temp
##
                          0.3648729712
                                                               0.0762561613
##
                                   edu
                                                                       resp
                          0.0263008321
##
                                                              -0.0039559477
##
                                num.co
                                                 age:as.factor(dzgroup)CHF
##
                         -0.0431664138
                                                              -0.0046449011
##
      age:as.factor(dzgroup)Cirrhosis age:as.factor(dzgroup)Colon Cancer
##
                         -0.0062251497
                                                              -0.0018857174
##
           age:as.factor(dzgroup)Coma
                                                age:as.factor(dzgroup)COPD
##
                         -0.0129436877
                                                               0.0002001618
    age:as.factor(dzgroup)Lung Cancer age:as.factor(dzgroup)MOSF w/Malig
##
##
                          0.0066143639
                                                              -0.0103087258
```

We can calculate the MSE on the test set to evaluate the multiple linear regression model.

MSE R2 RMSE MAE ## 1 0.8872275 0.3764346 0.9419275 0.7408729

The MSE-test for multiple linear regression is worst than for simple linear regression.

Simple linear regression is the best model so far for this problem.

K-nearest neighbor regression

It works the same way as the KNN for classification.

Given a value for k and a prediction point x_i , KNN regression identifies the k closest training observations to x_i represented by N_i .

Then it estimates $f(x_i)$ using the average of all the training responses in N_i .

$$\hat{y_i} = f(x_i) = \frac{1}{k} \Sigma_{x_j \in N_i} y_j$$

Validation techniques

Sampling

This consists in dividing the dataset into a training set and a test set.

Cross validation

R2, RMSE and MAE are used to measure the regression model performance during cross-validation.

Validation set approach

The Validation Set Approach is a type of method that estimates a model error rate by holding out a subset of the data from the fitting process (creating a testing dataset). The model is then built using the other set of observations (the training dataset)

Example on R

```
## MSE R2 RMSE MAE
## 1 0.8842554 0.3711666 0.9403485 0.7451318
```

Example on Python

```
## Ridge(alpha=0.5)
## {'fit_time': array([0.01695275, 0.01170397, 0.01566696, 0.01770115, 0.01134706]), 'score_time': array
```

Leave One out cross-validation

Leave-one-out cross-validation is a special case of cross-validation where the number of folds equals the number of instances in the data set.

This method works as follow:

Leave out one data point and build the model on the rest of the data set Test the model against the data point that is left out at step 1 and record the test error associated with the prediction Repeat the process for all data points Compute the overall prediction error by taking the average of all these test error estimates recorded at step 2.

Example on R

```
## Linear Regression
##
## 4960 samples
##
      6 predictor
##
## No pre-processing
## Resampling: Leave-One-Out Cross-Validation
## Summary of sample sizes: 4959, 4959, 4959, 4959, 4959, 4959, ...
## Resampling results:
##
##
     RMSE
                Rsquared
                           MAE
               0.3785367
##
     0.9348588
                           0.7467919
## Tuning parameter 'intercept' was held constant at a value of TRUE
```

k-Fold Cross-Validation

K-Fold Cross-Validation is where a given data set is split into a K number of sections/folds where each fold is used as a testing set at some point.

We divide the set of data in k equals part and we use k-1 parts to train the model and 1 to test. We do do that k times in order to use each part as a test part.

Here are the steps:

1. Split the dataset into k equal partitions (or "folds")

2. For each fold

One fold is used as the testing set and the union of the other folds as the training set

Calculate testing accuracy for this fold:

$$\begin{split} \hat{f}_i &= \frac{1}{K} \sum_{j \in N_0} (y_j) \\ MSE &= \frac{k}{n} \sum_i I(y_i \neq \hat{y_i}) \end{split}$$

3. Use the average testing accuracy as the estimate of out-of-sample accuracy :

We would use the cross-validation error:

$$CV_k = \frac{1}{k} \sum_i MSE_i$$

with $I(y_i \neq \hat{y_i}) = 1$ if $y_i \neq \hat{y_i}$, 0 else. So that we calculate the average of wrong predicted values.

Example on R

```
## Linear Regression
##
## 4960 samples
      6 predictor
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 4464, 4464, 4464, 4464, 4464, ...
## Resampling results:
##
##
     RMSE
                Rsquared
                           MAE
##
     0.9347368 0.3788688 0.7470814
## Tuning parameter 'intercept' was held constant at a value of TRUE
```