

MVA_project.R

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```
#Importing the Hepatatis c Dataset
```

```
HCV= read.csv("HCV-Egy-Data.csv")
```

```
HCV
```

##	Age	Gender	BMI	Fever	Nausea.Vomting	Headache	Diarrhea
## 1	56	1	35	2	1	1	1
## 2	46	1	29	1	2	2	1
## 3	57	1	33	2	2	2	2
## 4	49	2	33	1	2	1	2
## 5	59	1	32	1	1	2	1
## 6	58	2	22	2	2	2	1
## 7	42	2	26	1	1	2	2
## 8	48	2	30	1	1	2	2
## 9	44	1	23	1	1	2	2
## 10	45	1	30	2	1	2	2
## 11	37	2	24	2	1	2	1
## 12	36	1	22	2	2	1	1
## 13	45	2	25	2	1	1	1
## 14	34	1	22	1	2	1	1
## 15	40	2	32	2	2	2	1
## 16	58	1	34	2	1	1	1
## 17	61	1	35	1	2	2	2
## 18	55	2	24	2	1	2	2
## 19	56	1	27	1	2	2	2
## 20	35	2	23	2	2	1	1
## 21	57	2	23	1	1	2	2
## 22	33	1	25	2	1	2	2
## 23	41	1	23	1	2	2	2
## 24	39	2	29	1	2	1	2
## 25	33	2	24	1	2	2	2
## 26	43	2	34	2	2	2	1
## 27	51	1	34	2	1	2	2
## 28	39	2	33	2	1	2	1
## 29	57	2	26	1	2	2	1
## 30	47	2	29	1	1	2	1
## 31	55	2	33	1	2	2	1
## 32	58	2	35	2	2	2	2
## 33	47	2	25	2	1	2	2
## 34	61	1	33	1	2	2	2
## 35	37	1	27	2	2	1	2
## 36	41	1	29	1	2	1	1

```
## 37      60      2  32      2      2      1      2
```

#Summary

attach(HCV)

summary(HCV)

```
##      Age      Gender      BMI      Fever
## Min.   :32.00  Min.    :1.00  Min.    :22.00  Min.     :1.000
## 1st Qu.:39.00  1st Qu.:1.00  1st Qu.:25.00  1st Qu.:1.000
## Median :46.00  Median :1.00  Median :29.00  Median :2.000
## Mean   :46.32  Mean    :1.49  Mean    :28.61  Mean     :1.516
## 3rd Qu.:54.00  3rd Qu.:2.00  3rd Qu.:32.00  3rd Qu.:2.000
## Max.   :61.00  Max.    :2.00  Max.    :35.00  Max.     :2.000
## Nausea.Vomting  Headache  Diarrhea
## Min.    :1.000  Min.    :1.000  Min.    :1.000
## 1st Qu.:1.000  1st Qu.:1.000  1st Qu.:1.000
## Median :2.000  Median :1.000  Median :2.000
## Mean    :1.503  Mean    :1.496  Mean    :1.503
## 3rd Qu.:2.000  3rd Qu.:2.000  3rd Qu.:2.000
## Max.    :2.000  Max.    :2.000  Max.    :2.000
## Fatigue...generalized.bone.ache  Jaundice  Epigastric.pain
## Min.    :1.000  Min.    :1.000  Min.    :1.000
## 1st Qu.:1.000  1st Qu.:1.000  1st Qu.:1.000
## Median :1.000  Median :2.000  Median :2.000
## Mean    :1.499  Mean    :1.501  Mean    :1.504
## 3rd Qu.:2.000  3rd Qu.:2.000  3rd Qu.:2.000
## Max.    :2.000  Max.    :2.000  Max.    :2.000
##      WBC      RBC      HGB      Plat
## Min.    : 2991  Min.    :3816422  Min.    :10.00  Min.    : 93013
## 1st Qu.: 5219  1st Qu.:4121374  1st Qu.:11.00  1st Qu.:124479
## Median : 7498  Median :4438465  Median :13.00  Median :157916
## Mean    : 7533  Mean    :4422130  Mean    :12.59  Mean    :158348
## 3rd Qu.: 9902  3rd Qu.:4721279  3rd Qu.:14.00  3rd Qu.:190314
## Max.    :12101  Max.    :5018451  Max.    :15.00  Max.    :226464
##      AST.1      ALT.1      ALT4      ALT.12
## Min.    : 39.00  Min.    : 39.00  Min.    : 39.00  Min.    : 39.00
## 1st Qu.: 60.00  1st Qu.: 62.00  1st Qu.: 61.00  1st Qu.: 60.00
## Median : 83.00  Median : 83.00  Median : 82.00  Median : 84.00
## Mean    : 82.77  Mean    : 83.92  Mean    : 83.41  Mean    : 83.51
## 3rd Qu.:105.00  3rd Qu.:106.00  3rd Qu.:107.00  3rd Qu.:106.00
## Max.    :128.00  Max.    :128.00  Max.    :128.00  Max.    :128.00
##      ALT.24      ALT.36      ALT.48      ALT.after.24.w
## Min.    : 39.00  Min.    :  5.00  Min.    :  5.00  Min.    :  5.00
## 1st Qu.: 61.00  1st Qu.: 61.00  1st Qu.: 61.00  1st Qu.:28.00
## Median : 83.00  Median : 84.00  Median : 83.00  Median :34.00
## Mean    : 83.71  Mean    : 83.12  Mean    : 83.63  Mean    :33.44
## 3rd Qu.:107.00  3rd Qu.:106.00  3rd Qu.:106.00  3rd Qu.:40.00
## Max.    :128.00  Max.    :128.00  Max.    :128.00  Max.    :45.00
##      RNA.Base      RNA.4      RNA.12      RNA.EOT
## Min.    :      11  Min.    :      5  Min.    :      5  Min.    :      5
## 1st Qu.: 269253  1st Qu.: 270893  1st Qu.:      5  1st Qu.:      5
```

```
## Median : 593103 Median : 597869 Median : 234359 Median :251376
## Mean : 590951 Mean : 600896 Mean : 288754 Mean :287660
## 3rd Qu.: 886791 3rd Qu.: 909093 3rd Qu.: 524819 3rd Qu.:517806
## Max. :1201086 Max. :1201715 Max. :3731527 Max. :808450
## RNA.EF Baseline.histological.Grading
## Min. : 5 Min. : 3.000
## 1st Qu.: 5 1st Qu.: 6.000
## Median :244049 Median :10.000
## Mean :291378 Mean : 9.762
## 3rd Qu.:527864 3rd Qu.:13.000
## Max. :810333 Max. :16.000
## Baselinehistological.staging
## Min. :1.000
## 1st Qu.:2.000
## Median :3.000
## Mean :2.536
## 3rd Qu.:4.000
## Max. :4.000
```

#Dimensions of the data set

NROW(HCV)

```
## [1] 1385
```

NCOL(HCV)

```
## [1] 29
```

#Displaying the column names of the dataset

colnames(HCV)

```
## [1] "Age" "Gender"
## [3] "BMI" "Fever"
## [5] "Nausea.Vomting" "Headache"
## [7] "Diarrhea"
"Fatigue...generalized.bone.ache"
## [9] "Jaundice" "Epigastric.pain"
## [11] "WBC" "RBC"
## [13] "HGB" "Plat"
## [15] "AST.1" "ALT.1"
## [17] "ALT4" "ALT.12"
## [19] "ALT.24" "ALT.36"
## [21] "ALT.48" "ALT.after.24.w"
## [23] "RNA.Base" "RNA.4"
## [25] "RNA.12" "RNA.EOT"
## [27] "RNA.EF" "Baseline.histological.Grading"
## [29] "Baselinehistological.staging"
```

#Another menthod for dimensions

dim(HCV)

```
## [1] 1385 29
```

#Preprocessing data was done but didn't find any discrepancies.

na= **is.na**(HCV)

na

[illegible]

```
## [49,] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
## [50,] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
## [51,] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
## [52,] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
## [53,] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
```

#Displaying the first six rows of the datasets
head(HCV)

```
## Age Gender BMI Fever Nausea.Vomting Headache Diarrhea
## 1 56 1 35 2 1 1 1
## 2 46 1 29 1 2 2 1
## 3 57 1 33 2 2 2 2
## 4 49 2 33 1 2 1 2
## 5 59 1 32 1 1 2 1
## 6 58 2 22 2 2 2 1
## Fatigue...generalized.bone.ache Jaundice Epigastric.pain WBC
RBC
## 1 2 2 2 7425
4248807
## 2 2 2 1 12101
4429425
## 3 1 1 1 4178
4621191
## 4 1 2 1 6490
4794631
## 5 2 2 2 3661
4606375
## 6 2 2 1 11785
3882456
## HGB Plat AST.1 ALT.1 ALT4 ALT.12 ALT.24 ALT.36 ALT.48
ALT.after.24.w
## 1 14 112132 99 84 52 109 81 5 5
5
## 2 10 129367 91 123 95 75 113 57 123
44
## 3 12 151522 113 49 95 107 116 5 5
5
## 4 10 146457 43 64 109 80 88 48 77
33
## 5 11 187684 99 104 67 48 120 94 90
30
## 6 15 131228 66 104 121 96 65 73 114
29
## RNA.Base RNA.4 RNA.12 RNA.EOT RNA.EF Baseline.histological.Grading
## 1 655330 634536 288194 5 5 13
## 2 40620 538635 637056 336804 31085 4
## 3 571148 661346 5 735945 558829 4
## 4 1041941 449939 585688 744463 582301 10
## 5 660410 738756 3731527 338946 242861 11
## 6 1157452 1086852 5 5 5 4
```

```
## Baselinehistological.staging
## 1 2
## 2 2
## 3 4
## 4 3
## 5 1
## 6 4
```

```
tail(HCV)
```

```
## Age Gender BMI Fever Nausea.Vomting Headache Diarrhea
## 1380 53 1 31 2 2 1 2
## 1381 44 1 29 1 2 2 2
## 1382 55 1 34 1 2 2 1
## 1383 42 1 26 2 2 1 1
## 1384 52 1 29 2 1 1 2
## 1385 55 2 26 1 2 2 2
## Fatigue...generalized.bone.ache Jaundice Epigastric.pain WBC
## 1380 2 2 1 4196
## 1381 1 1 1 7044
## 1382 1 1 1 6207
## 1383 1 2 1 4913
## 1384 2 2 1 7257
## 1385 1 2 1 11832
## RBC HGB Plat AST.1 ALT.1 ALT4 ALT.12 ALT.24 ALT.36 ALT.48
## 1380 4076324 12 150065 89 113 52 39 54 86 78
## 1381 4957174 15 202520 122 59 78 106 127 63 44
## 1382 4636951 10 115776 128 102 65 99 108 97 64
## 1383 4122038 14 128354 61 93 123 61 116 87 39
## 1384 4241990 10 205908 70 97 104 74 47 48 81
## 1385 4059176 14 136615 51 126 39 68 115 64 71
## ALT.after.24.w RNA.Base RNA.4 RNA.12 RNA.EOT RNA.EF
## 1380 36 886656 460080 591040 621014 20645
## 1381 45 387795 55938 5 5 5
## 1382 41 481378 152961 393339 73574 236273
## 1383 24 612664 572756 806109 343719 160457
## 1384 43 139872 76161 515730 2460 696074
## 1385 34 1190577 628730 5 5 5
## Baseline.histological.Grading Baselinehistological.staging
## 1380 13 4
## 1381 15 4
## 1382 10 2
## 1383 6 2
## 1384 15 3
## 1385 13 3
```

```
#CORRELATION, COVARIANCE AND DISTANCE
```

```
covariance<-cov(HCV[,c(11:16,23)]) #variamce-covariance matrix created
```

```
correlation<-cor(HCV[,c(11:16,23)]) #standardized
```

```
#colmeans
```

```
cm<-colMeans(HCV[,c(11:16,23)])
```

```
distance<-dist(scale(HCV[,c(11:16,23)],center=FALSE))
```

```
#Calculating di(generalized distance for all observations of our data)
```

```
#before that first extract all numeric variable in a dataframe
x<-HCV[,c(11:16,23)]
d <- apply(x, MARGIN = 1, function(x) + t(x - cm) %*% solve(covariance)
%*% (x - cm))
```

```
#Exploration of the data for high chances of HCV Infection
#Here RNA.base value if it is more than 700000 units then virus is
detected in high quantity.
#Here ALT.1 if value is greater than 57 then it is not normal.
#we sorted the data on these two components.
```

```
library(dplyr)
```

```
##
```

```
## Attaching package: 'dplyr'
```

```
## The following objects are masked from 'package:stats':
```

```
##
```

```
## filter, lag
```

```
## The following objects are masked from 'package:base':
```

```
##
```

```
## intersect, setdiff, setequal, union
```

```
HCV_male = HCV %>% filter(Gender == 1 & RNA.Base>= 700000 & ALT.1 >= 57)
HCV_male
```

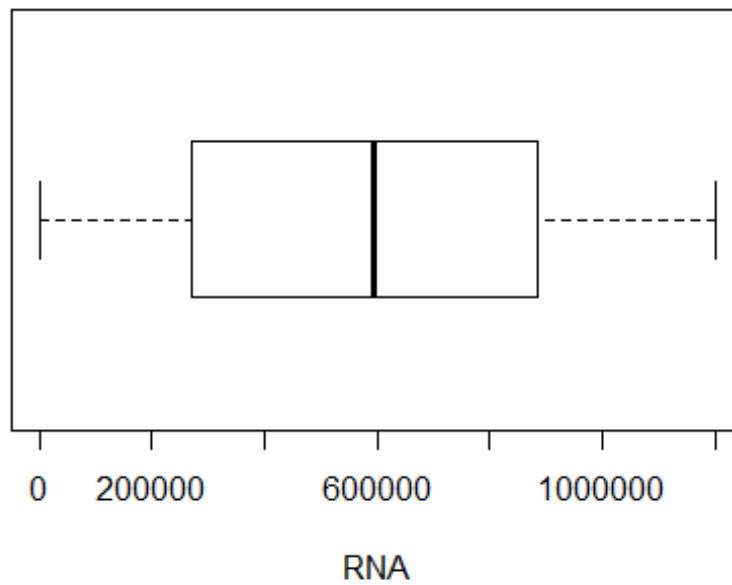
```
##      Age Gender BMI Fever Nausea.Vomting Headache Diarrhea
## 1     45      1  30     2          1          2          2
## 2     61      1  35     1          2          2          2
## 3     33      1  25     2          1          2          2
## 4     41      1  29     1          2          1          1
## 5     32      1  31     1          2          1          1
## 6     44      1  31     1          1          1          1
## 7     61      1  25     1          1          2          2
## 8     59      1  25     1          2          1          1
## 9     32      1  26     1          2          2          2
## 10    56      1  26     1          2          2          2
## 11    40      1  31     1          2          2          1
## 12    33      1  28     2          2          1          2
## 13    39      1  30     1          2          2          1
## 14    52      1  26     2          2          1          1
## 15    49      1  25     1          2          1          2
## 16    53      1  28     2          2          1          2
## 17    54      1  31     2          2          1          2
## 18    55      1  30     2          2          2          1
## 19    56      1  34     1          1          1          2
## 20    58      1  33     1          1          2          1
## 21    33      1  30     2          2          1          2
## 22    34      1  28     2          1          2          2
## 23    60      1  32     1          1          1          1
```

## 24	34	1	32	1	2	2	2
## 25	37	1	34	2	1	1	2
## 26	53	1	22	1	1	1	2
## 27	59	1	28	1	1	2	2
## 28	56	1	34	1	2	2	1
## 29	61	1	34	2	1	1	1
## 30	58	1	29	2	2	1	1
## 31	33	1	30	2	2	2	2
## 32	48	1	30	1	1	1	1
## 33	35	1	26	2	2	1	2
## 34	55	1	31	1	1	2	2
## 35	51	1	30	1	2	1	1
## 36	49	1	30	2	1	2	1
## 37	57	1	24	2	2	1	2
## 38	45	1	34	2	1	1	1
## 39	46	1	22	1	2	1	1
## 40	59	1	28	1	2	2	1
## 41	52	1	26	1	2	2	1
## 42	60	1	22	1	2	1	1
## 43	39	1	35	1	1	2	2
## 44	34	1	28	1	2	2	2
## 45	37	1	35	2	1	1	1
## 46	38	1	25	2	2	2	2
## 47	58	1	35	1	1	2	2
## 48	58	1	26	2	1	2	2
## 49	38	1	32	2	1	1	2
## 50	35	1	22	2	1	2	1
## 51	35	1	25	2	1	1	2
## 52	57	1	26	1	1	1	1
## 53	52	1	34	1	2	2	2
## 54	54	1	25	2	1	2	2

#Box Plot

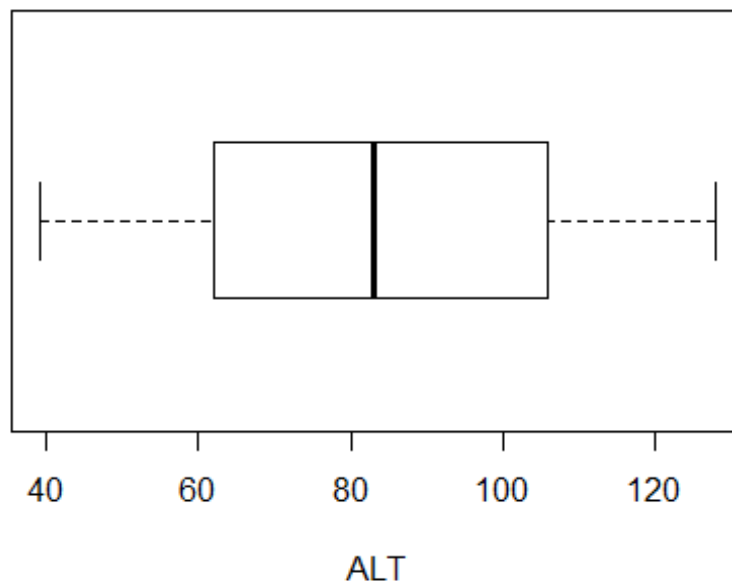
```
boxplot(RNA.Base, main="RNA.BASE Box plot",yaxt="n", xlab="RNA",
horizontal=TRUE)
```


RNA.BASE Box plot



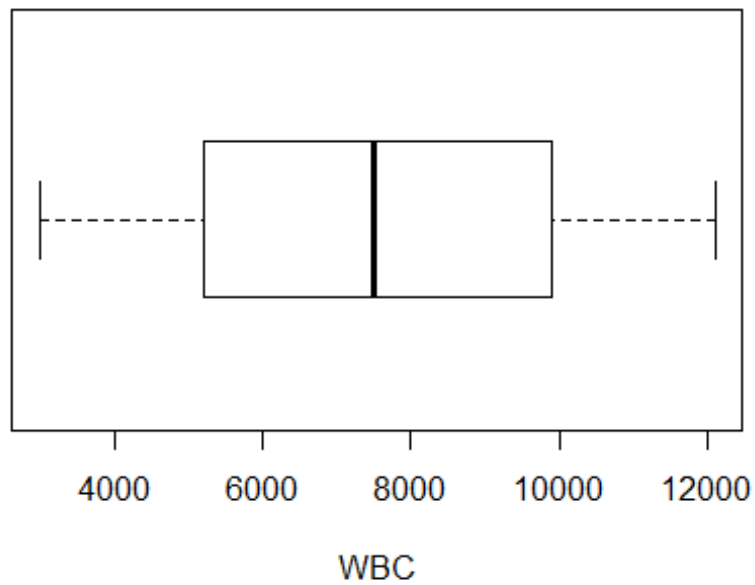
```
boxplot(ALT.1, main="ALT.1 Box plot", yaxt="n", xlab="ALT",  
horizontal=TRUE)
```

ALT.1 Box plot



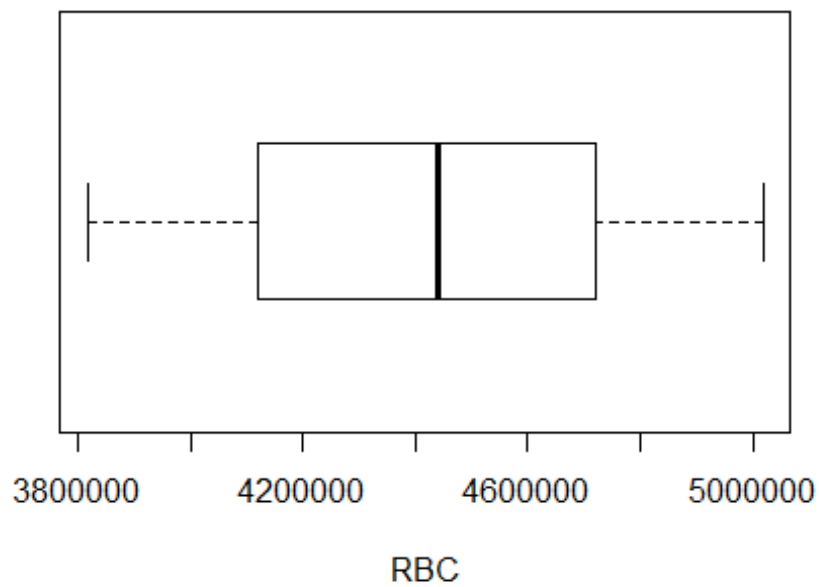
```
boxplot(WBC, main="WBC Box plot", yaxt="n", xlab="WBC", horizontal=TRUE)
```

WBC Box plot



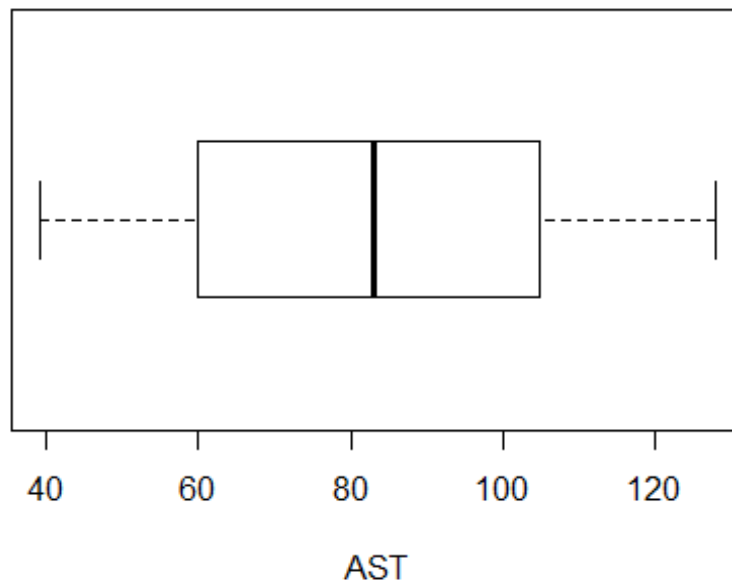
```
boxplot(WBC, main="WBC Box plot", yaxt="n", xlab="WBC", horizontal=TRUE)
```

WBC Box plot



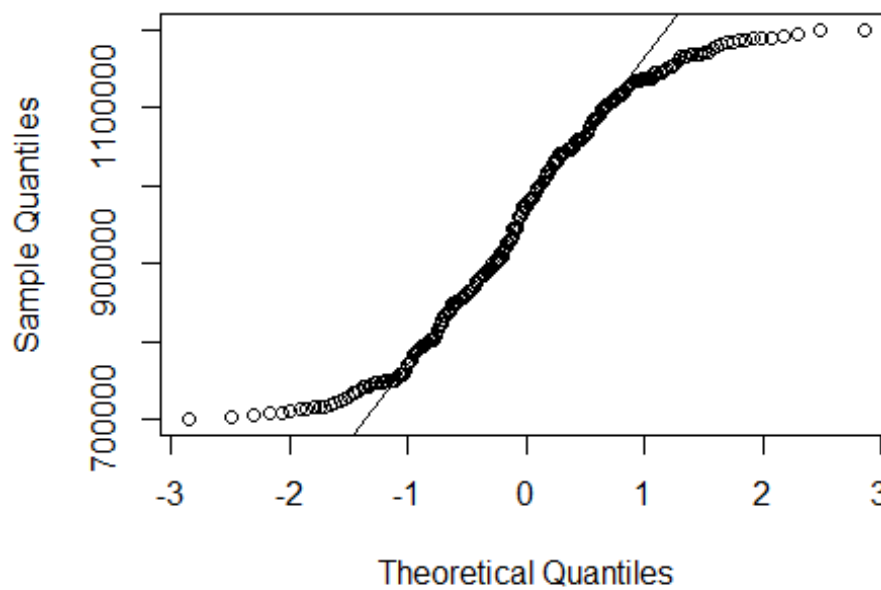
```
boxplot(RBC, main="RBC Box plot", yaxt="n", xlab="RBC", horizontal=TRUE)
```

AST.1 Box plot



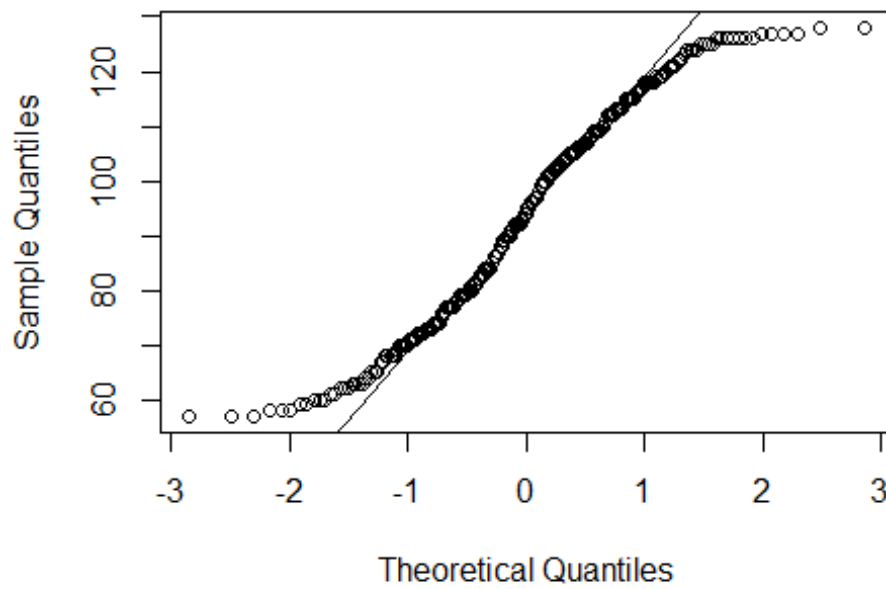
```
#plotting, Are they in a straight line.  
#Male Plotting of the dataset is done for five different attributes.  
qqnorm(HCV_male[, "RNA.Base"], main = "RNA.Base");  
qqline(HCV_male[, "RNA.Base"])
```

RNA.Base



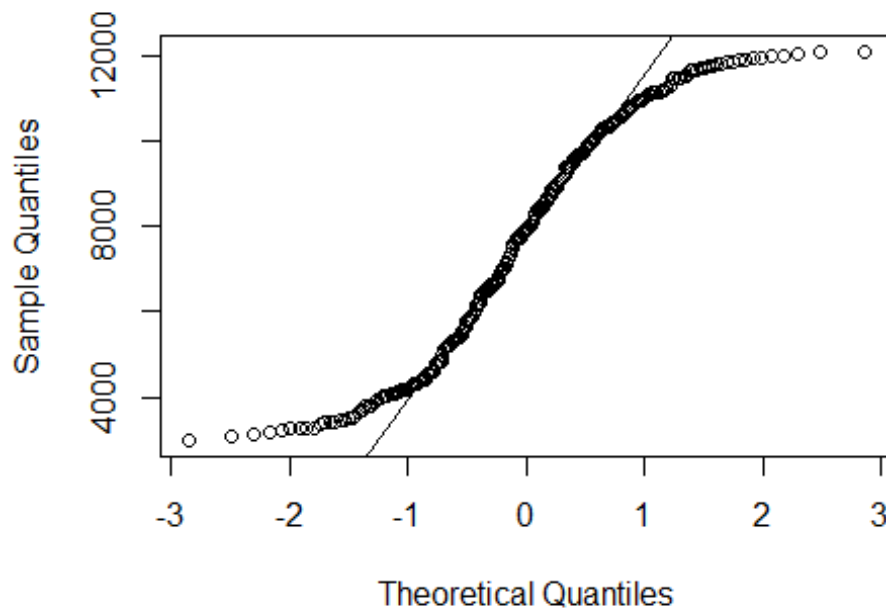
```
qqnorm(HCV_male[, "ALT.1"], main = "ALT.1"); qqline(HCV_male[, "ALT.1"])
```

ALT.1

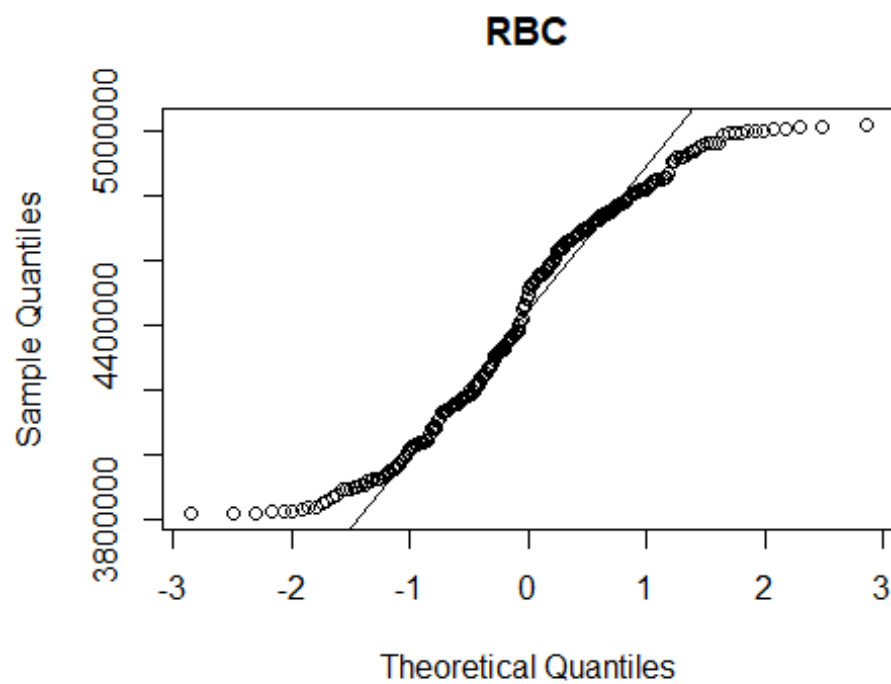


```
qqnorm(HCV_male[, "WBC"], main = "WBC"); qqline(HCV_male[, "WBC"])
```

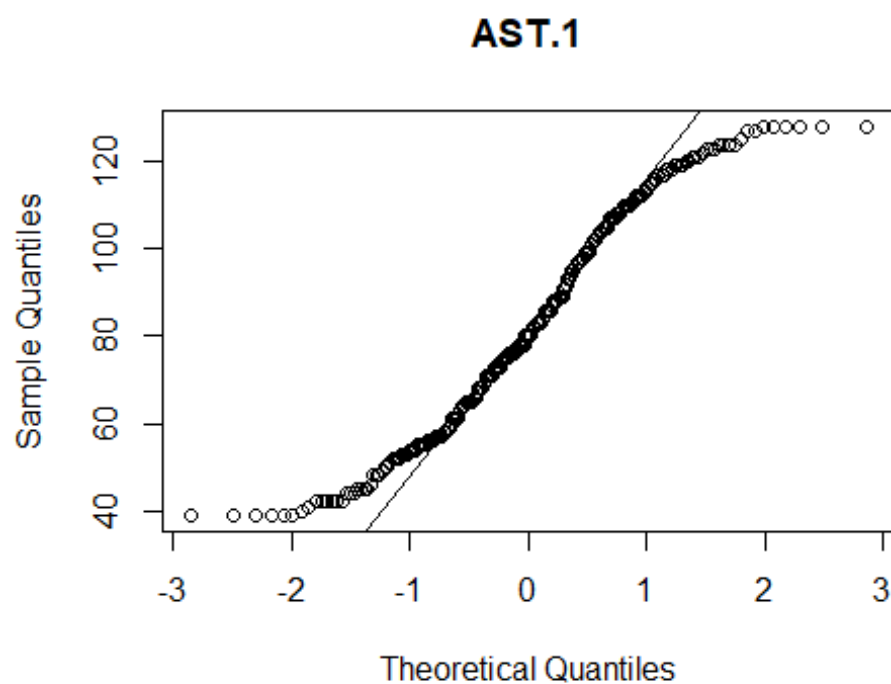
WBC



```
qqnorm(HCV_male[, "RBC"], main = "RBC"); qqline(HCV_male[, "RBC"])
```

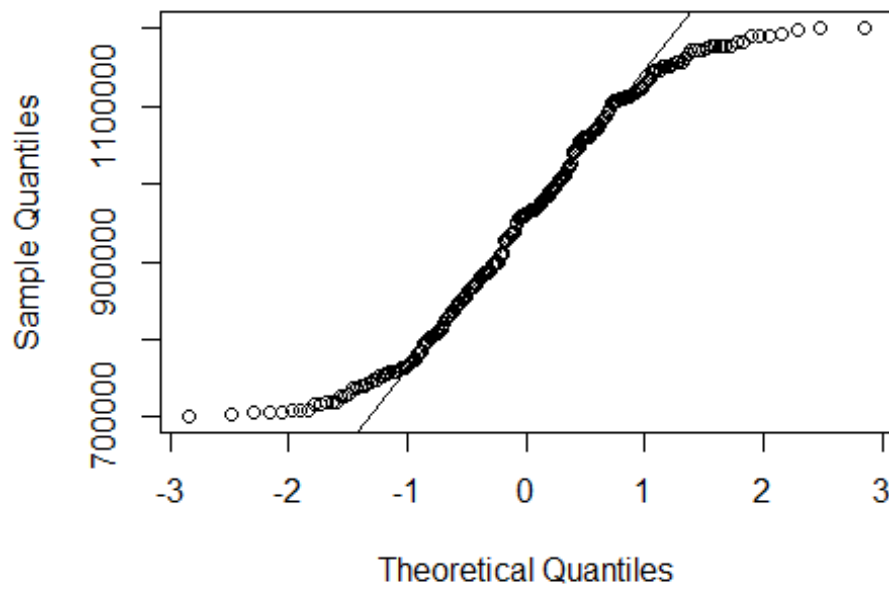


```
qqnorm(HCV_male[, "AST.1"], main = "AST.1"); qqline(HCV_male[, "AST.1"])
```



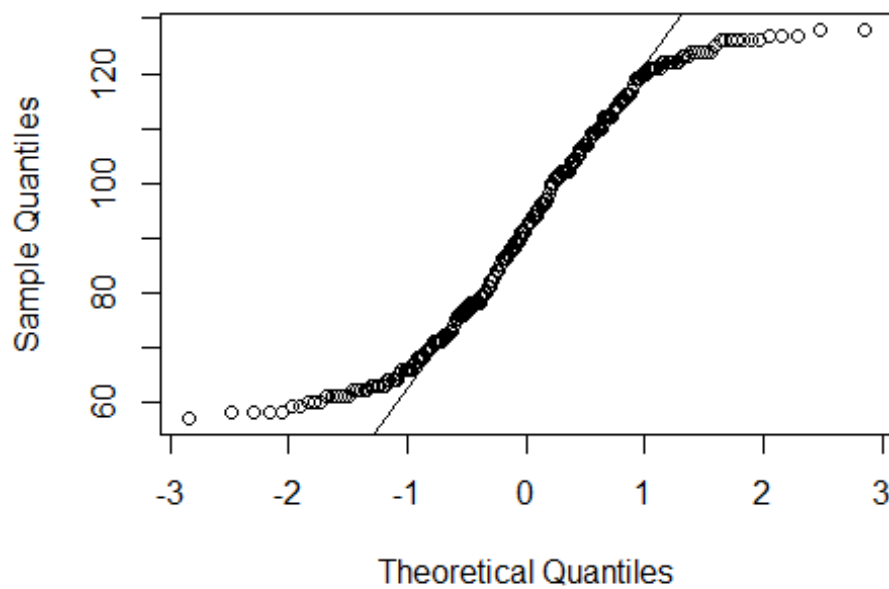
```
#Female, Are they in a straight line.  
#FeMale Plotting of the dataset is done for five different attributes.  
qqnorm(HCV_female[, "RNA.Base"], main = "RNA.Base");  
qqline(HCV_female[, "RNA.Base"])
```

RNA.Base

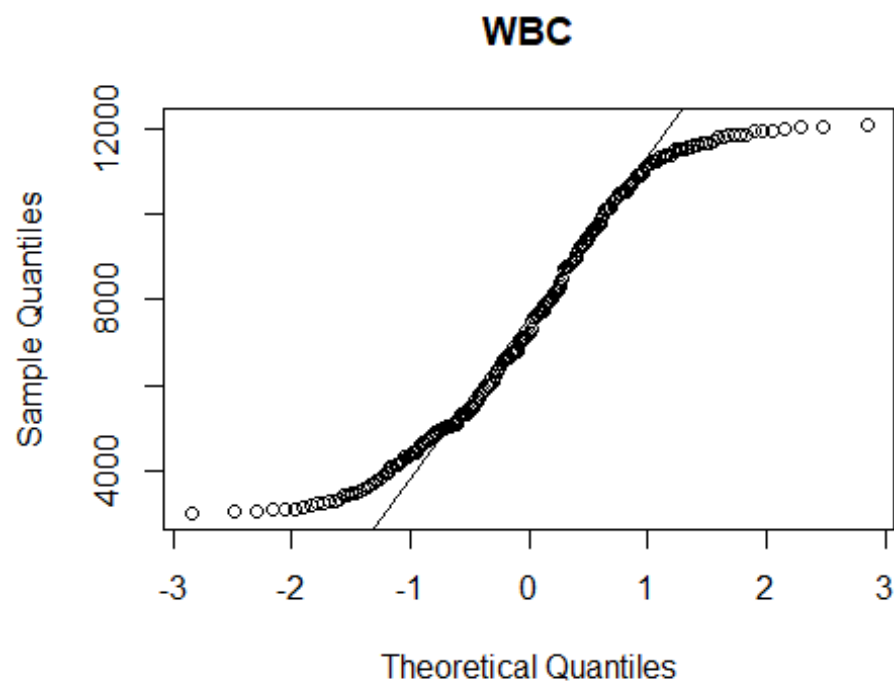


```
qqnorm(HCV_female[, "ALT.1"], main = "ALT.1"); qqline(HCV_female[, "ALT.1"])
```

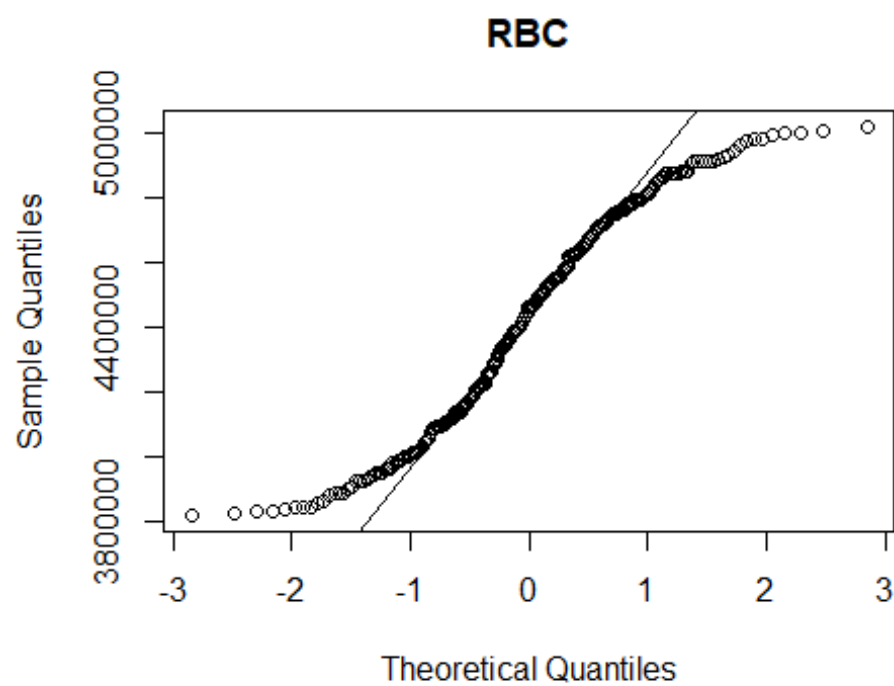
ALT.1



```
qqnorm(HCV_female[, "WBC"], main = "WBC"); qqline(HCV_female[, "WBC"])
```

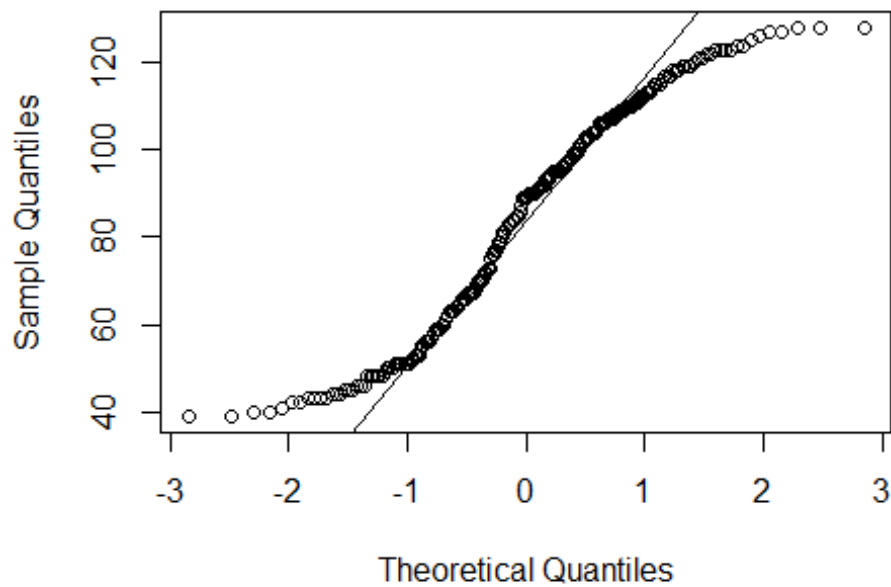


```
qqnorm(HCV_female[, "RBC"], main = "RBC"); qqline(HCV_female[, "RBC"])
```



```
qqnorm(HCV_female[, "AST.1"], main = "AST.1"); qqline(HCV_female[, "AST.1"])
```

AST.1

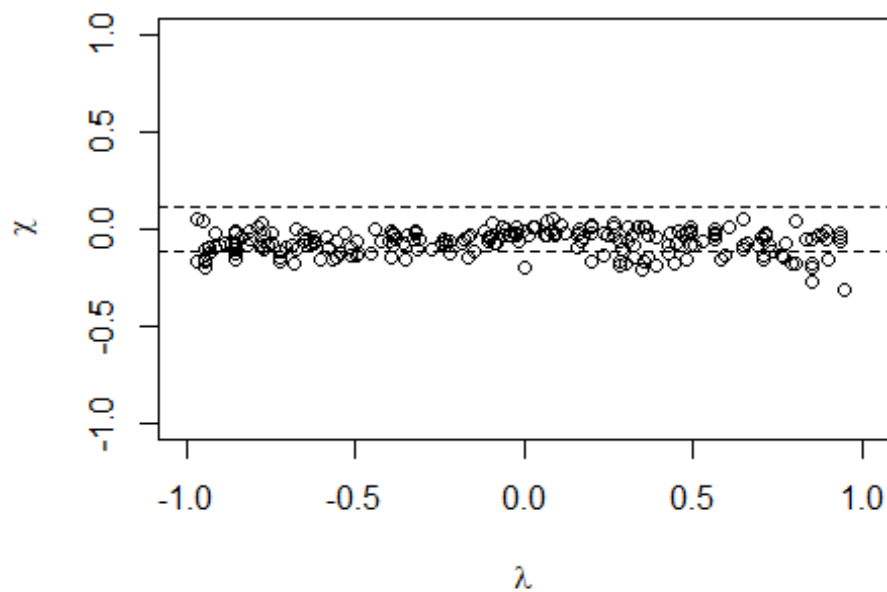


```
#Visualisation
#Chiplot
library(HSAUR2)

## Loading required package: tools

library(tools)
library(MVA)

#Chiplot
#For male data
with(HCV_male, chipplot(RNA.Base, ALT.1))
```

```
#For Female Data
with(HCV_female, chiplot(RNA.Base, ALT.1))

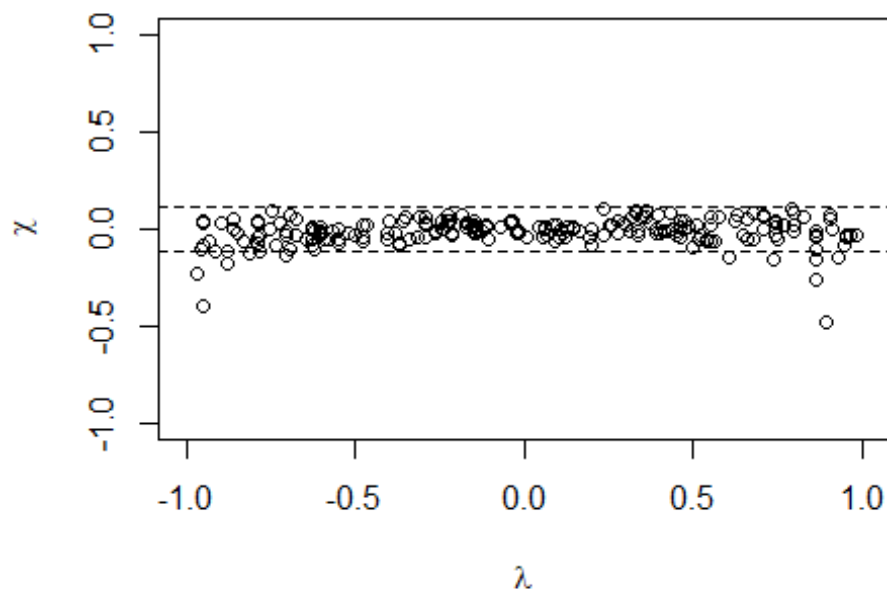
library(GGally)

## Loading required package: ggplot2

## Registered S3 method overwritten by 'GGally':
##   method from
##   +.gg      ggplot2

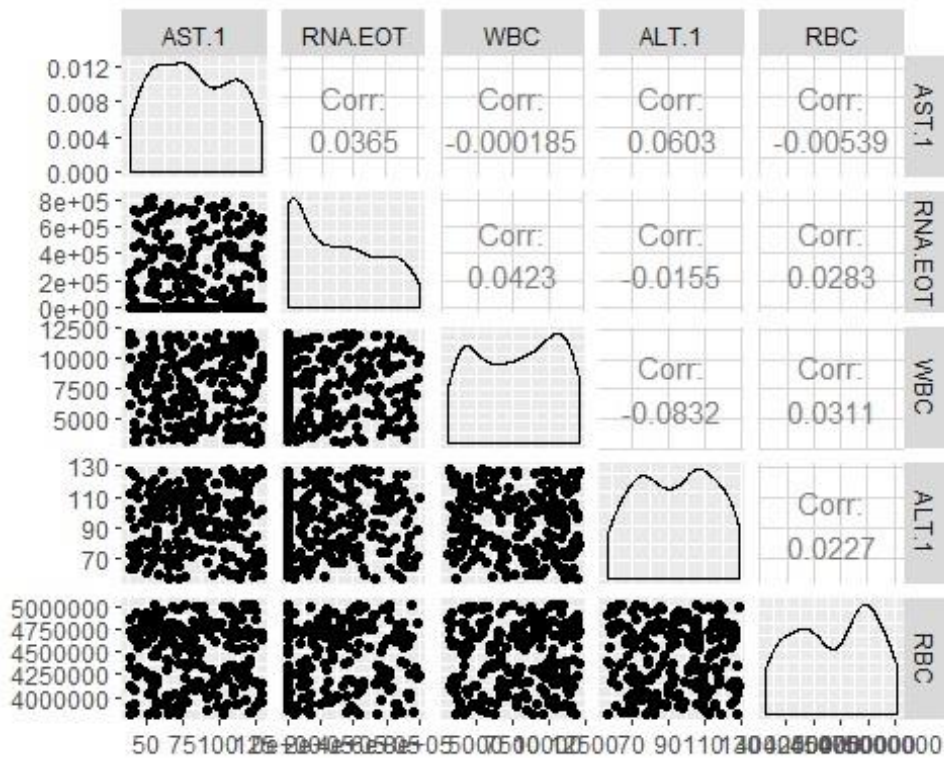
##
## Attaching package: 'GGally'

## The following object is masked from 'package:dplyr':
##
##   nasa
```



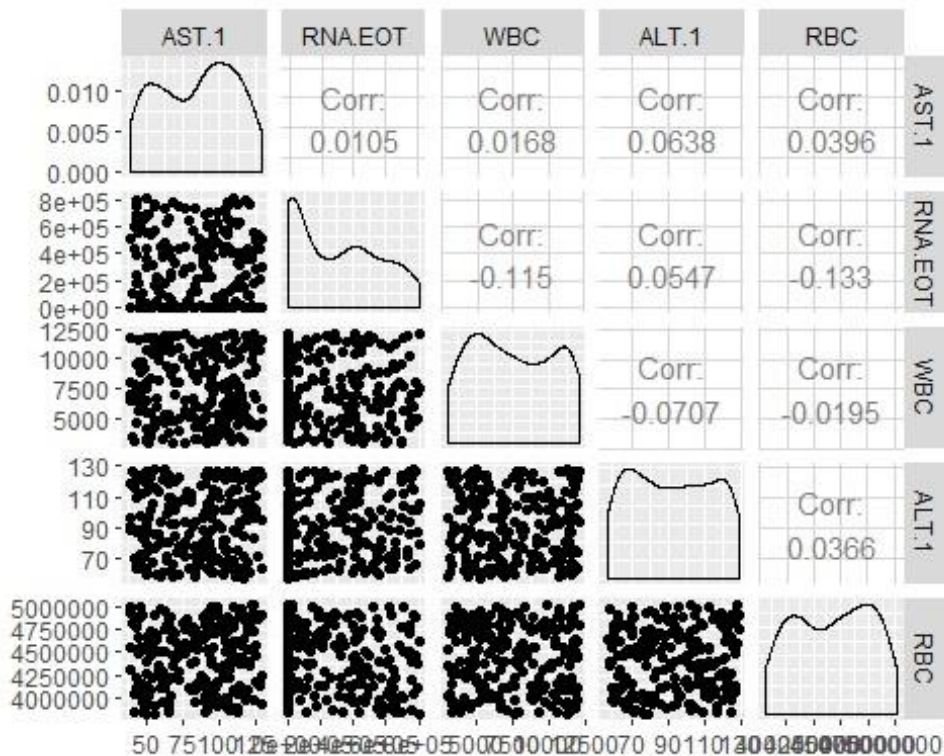
```
ggpairs(HCV_male, columns=c("AST.1", "RNA.EOT", "WBC", "ALT.1", "RBC"),  
color="Survivorship")
```

```
## Warning in warn_if_args_exist(list(...)): Extra arguments: "color" are  
## being ignored. If these are meant to be aesthetics, submit them using  
## the  
## 'mapping' variable within ggpairs with ggplot2::aes or  
## ggplot2::aes_string.
```



```
ggpairs(HCV_female, columns=c("AST.1", "RNA.EOT", "WBC", "ALT.1", "RBC"),
color="Survivorship")
```

```
## Warning in warn_if_args_exist(list(...)): Extra arguments: "color" are
## being ignored. If these are meant to be aesthetics, submit them using
the
## 'mapping' variable within ggpairs with ggplot2::aes or
ggplot2::aes_string.
```



```
summary(lm(data = HCV , RNA.EOT~Age))
```

```
##
## Call:
## lm(formula = RNA.EOT ~ Age, data = HCV)
##
## Residuals:
##      Min       1Q   Median       3Q      Max
## -308665 -271984 -35364  226006  533719
##
## Coefficients:
##              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 355622.3    38146.1   9.323  <2e-16 ***
## Age        -1467.3      809.1  -1.813    0.07 .
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 264300 on 1383 degrees of freedom
## Multiple R-squared:  0.002372,    Adjusted R-squared:  0.001651
## F-statistic: 3.288 on 1 and 1383 DF,  p-value: 0.07
```

```
summary(lm(data = HCV , RNA.EOT~Gender))
```

```
##
## Call:
## lm(formula = RNA.EOT ~ Gender, data = HCV)
##
## Residuals:
##      Min       1Q   Median       3Q      Max
## -297181 -278520 -34151  223816  529617
##
```

```
## Coefficients:
##           Estimate Std. Error t value Pr(>|t|)
## (Intercept)  259865     22338  11.634  <2e-16 ***
## Gender       18661     14217   1.313    0.19
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 264500 on 1383 degrees of freedom
## Multiple R-squared:  0.001244, Adjusted R-squared:  0.000522
## F-statistic: 1.723 on 1 and 1383 DF, p-value: 0.1896

summary(lm(data = HCV , RNA.EOT~WBC))

##
## Call:
## lm(formula = RNA.EOT ~ WBC, data = HCV)
##
## Residuals:
##      Min       1Q   Median       3Q      Max
## -296698 -280505 -38809  230123  527965
##
## Coefficients:
##           Estimate Std. Error t value Pr(>|t|)
## (Intercept) 302720.284  21302.853   14.21  <2e-16 ***
## WBC          -1.999     2.666   -0.75    0.453
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 264600 on 1383 degrees of freedom
## Multiple R-squared:  0.0004065, Adjusted R-squared: -0.0003163
## F-statistic: 0.5624 on 1 and 1383 DF, p-value: 0.4534

summary(lm(data = HCV , RNA.EOT~ALT.1))

##
## Call:
## lm(formula = RNA.EOT ~ ALT.1, data = HCV)
##
## Residuals:
##      Min       1Q   Median       3Q      Max
## -305707 -274761 -32847  226046  529732
##
## Coefficients:
##           Estimate Std. Error t value Pr(>|t|)
## (Intercept) 321386.0    24083.4  13.345  <2e-16 ***
## ALT.1        -401.9     274.2   -1.466    0.143
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 264400 on 1383 degrees of freedom
## Multiple R-squared:  0.001551, Adjusted R-squared:  0.0008288
## F-statistic: 2.148 on 1 and 1383 DF, p-value: 0.143
```

```
cor(HCV)
```

```
##                               Age          Gender          BMI
## Age          1.0000000000 -0.0099335141 -0.025353485
## Gender      -0.009933514  1.0000000000  0.006495822
## BMI         -0.025353485  0.0064958221  1.000000000
## Fever       -0.021208025  0.0216038570 -0.017581296
## Nausea.Vomting -0.025023686 -0.0367257983  0.005801303
## Headache     0.017561878 -0.0239990326 -0.007850790
## Diarrhea     0.043574033  0.0181605819 -0.026094093
## Fatigue...generalized.bone.ache -0.007817311  0.0454520816 -0.007295820
## Jaundice     0.010284829  0.0007675462 -0.071378670
## Epigastric.pain -0.010817131 -0.0251107556  0.008205190
## WBC          0.014310232  0.0272897099  0.036721434
## RBC         -0.002161729 -0.0009591287 -0.004922280
## HGB         -0.012810104 -0.0012462736  0.057987192
## Plat        -0.002225244  0.0192990586 -0.003093342
## AST.1       -0.016632599 -0.0130777126  0.001806482
## ALT.1        0.005722864  0.0224504101  0.034036533
## ALT4         0.030259565 -0.0087203690  0.001416015
## ALT.12       0.019046099  0.0099728383 -0.059658524
## ALT.24       0.002711517 -0.0159186857  0.007909135
## ALT.36      -0.008720825  0.0060865959 -0.029995451
## ALT.48       0.027556595 -0.0203239599 -0.007866295
## ALT.after.24.w 0.003865167  0.0101835990 -0.021412112
## RNA.Base     0.022775156 -0.0133707332 -0.016418030
## RNA.4        -0.012699117 -0.0233148008  0.036863599
## RNA.12       0.001578231 -0.0313789591 -0.009189913
## RNA.EOT      -0.048702492  0.0352722375 -0.021427851
## RNA.EF       -0.030296896 -0.0152924344 -0.044901060
## Baseline.histological.Grading -0.040705291  0.0145587520 -0.023045099
## Baselinehistological.staging -0.019599169  0.0119553382 -0.057258857
```

Principal Component Analysis

```
> hcv_pca <-prcomp(HCV[,1:29],scale=TRUE)
> hcv_pca
```

Standard deviations (1, .., p=29):

```
[1] 1.3769419 1.1174363 1.1113462 1.0916269 1.0782624 1.0707535 1.0626130
1.0521221 1.0306635 1.0292621
[11] 1.0172925 1.0141324 1.0121337 1.0049132 0.9926278 0.9870471 0.9805851
0.9792502 0.9631161 0.9530072
[21] 0.9405476 0.9304068 0.9283507 0.9201871 0.9127624 0.8919466 0.8754930
0.7514594 0.7276547
```


OUTPUT:

Rotation (n x k) = (29 x 29):

	PC1	PC2	PC3	PC4	PC5
Age	0.043992545	-0.035109552	0.15542945	-0.21681173	0.1846420065
Gender	0.008756739	-0.031538265	0.01225109	0.11432799	-0.2272447661
BMI	0.059438630	0.004938750	-0.21133076	0.43507918	0.2105290684
Fever	-0.018638143	-0.016959013	0.14647417	0.11272529	-0.0194418421
Nausea.Vomting	-0.018052038	0.523815972	0.02931082	0.03619724	0.0100755834
Headache	-0.007027392	0.124936603	0.20495296	-0.22055035	0.0328136750
Diarrhea	-0.037779605	0.112284699	0.31055207	0.09505129	0.1941946991
Fatigue...generalized.bone.ache	-0.039589819	-0.113283136	0.02431225	0.03182207	-0.5018034613
Jaundice	-0.060924334	-0.039975480	0.15722114	-0.28037289	0.0021353121
Epigastric.pain	-0.091593158	-0.314657318	0.08262704	0.13208431	-0.1562642350
WBC	0.079856632	-0.207631406	-0.09056514	0.08357598	0.0941100088
RBC	0.062330494	0.242506719	-0.14413534	-0.04346368	-0.0591444285
HGB	0.012151709	0.030902639	-0.39565562	0.16325012	0.0113733042
Plat	-0.051058807	0.378757510	0.01108177	0.28487255	-0.1382597553
AST.1	0.028793449	0.013713351	0.04919642	0.04294564	-0.1320669862
ALT.1	0.018974095	0.262467291	0.39485581	0.25382822	-0.0368534659
ALT4	-0.040780344	0.034435407	-0.08938033	-0.27015280	0.4057377396
ALT.12	0.047599395	-0.157661272	-0.16534036	-0.38085696	-0.2283790159
ALT.24	-0.018567737	0.212800327	-0.32270954	-0.24975390	0.0210555092
ALT.36	-0.054830139	0.088193685	0.08694682	-0.12844262	-0.2103541059
ALT.48	-0.026185451	-0.255155588	0.31754417	0.06703057	0.0024901089
ALT.after.24.w	-0.019391592	-0.145425335	0.17697872	0.01737588	0.0255244283
RNA.Base	-0.013683156	-0.002907261	0.30619803	-0.11191128	0.1894906165
RNA.4	0.056900757	-0.183428054	-0.04195740	0.16586510	0.3981169569
RNA.12	-0.552679994	0.001687528	-0.02336655	0.03313714	0.0093530466
RNA.EOT	-0.563868495	-0.039526114	-0.07119686	0.03930584	0.0771614060
RNA.EF	-0.570538298	-0.011466450	-0.04305053	-0.00047804	0.0008019717
Baseline.histological.Grading	0.051971451	-0.140448509	0.00159462	0.13066428	-0.1768391380
Baselinehistological.staging	-0.032348367	0.213485043	0.10947878	-0.19354490	-0.1009923507

```
> summary(hcv_pca)
```

Importance of components:

	PC1	PC2	PC3	PC4	PC5	PC6	PC7	PC8	PC9	PC10
Standard deviation	1.37694	1.11744	1.11135	1.09163	1.07826	1.07075	1.06261	1.05212	1.03066	1.02926
Proportion of Variance	0.06538	0.04306	0.04259	0.04109	0.04009	0.03953	0.03894	0.03817	0.03663	0.03653
Cumulative Proportion	0.06538	0.10844	0.15102	0.19212	0.23221	0.27174	0.31068	0.34885	0.38548	0.42201
	PC11	PC12	PC13	PC14	PC15	PC16	PC17	PC18	PC19	PC20
Standard deviation	1.01729	1.01413	1.01213	1.00491	0.99263	0.9870	0.98059	0.97925	0.96312	0.95301
Proportion of Variance	0.03569	0.03546	0.03532	0.03482	0.03398	0.0336	0.03316	0.03307	0.03199	0.03132
Cumulative Proportion	0.45770	0.49316	0.52848	0.56331	0.59728	0.6309	0.66404	0.69710	0.72909	0.76041
	PC21	PC22	PC23	PC24	PC25	PC26	PC27	PC28	PC29	
Standard deviation	0.9405	0.93041	0.92835	0.9202	0.91276	0.89195	0.87549	0.75146	0.72765	
Proportion of Variance	0.0305	0.02985	0.02972	0.0292	0.02873	0.02743	0.02643	0.01947	0.01826	
Cumulative Proportion	0.7909	0.82076	0.85048	0.8797	0.90841	0.93584	0.96227	0.98174	1.00000	

```
> |
```



```

> (eigen_hcv <- hcv_pca$sdev^2)
[1] 1.8959690 1.2486640 1.2350903 1.1916494 1.1626498 1.1465131 1.1291464 1.1069609 1.0622673 1.0593804
[11] 1.0348841 1.0284646 1.0244145 1.0098506 0.9853100 0.9742619 0.9615471 0.9589310 0.9275925 0.9082227
[21] 0.8846297 0.8656567 0.8618350 0.8467443 0.8331352 0.7955688 0.7664880 0.5646913 0.5294814
> names(eigen_hcv) <- paste("PC",1:29,sep="")
> eigen_hcv
      PC1      PC2      PC3      PC4      PC5      PC6      PC7      PC8      PC9      PC10
1.8959690 1.2486640 1.2350903 1.1916494 1.1626498 1.1465131 1.1291464 1.1069609 1.0622673 1.0593804
      PC11      PC12      PC13      PC14      PC15      PC16      PC17      PC18      PC19      PC20
1.0348841 1.0284646 1.0244145 1.0098506 0.9853100 0.9742619 0.9615471 0.9589310 0.9275925 0.9082227
      PC21      PC22      PC23      PC24      PC25      PC26      PC27      PC28      PC29
0.8846297 0.8656567 0.8618350 0.8467443 0.8331352 0.7955688 0.7664880 0.5646913 0.5294814
> sumlambdas <- sum(eigen_hcv)
> sumlambdas
[1] 29
> propvar <- eigen_hcv/sumlambdas
> propvar
      PC1      PC2      PC3      PC4      PC5      PC6      PC7      PC8      PC9
0.06537824 0.04305738 0.04258932 0.04109136 0.04009137 0.03953493 0.03893608 0.03817107 0.03662991
      PC10      PC11      PC12      PC13      PC14      PC15      PC16      PC17      PC18
0.03653036 0.03568566 0.03546430 0.03532464 0.03482243 0.03397621 0.03359524 0.03315680 0.03306659
      PC19      PC20      PC21      PC22      PC23      PC24      PC25      PC26      PC27
0.03198595 0.03131802 0.03050447 0.02985023 0.02971845 0.02919808 0.02872880 0.02743341 0.02643062
      PC28      PC29
0.01947211 0.01825798
> cumvar_hcv <- cumsum(propvar)
> cumvar_hcv
      PC1      PC2      PC3      PC4      PC5      PC6      PC7      PC8      PC9
0.06537824 0.10843562 0.15102494 0.19211630 0.23220767 0.27174261 0.31067869 0.34884975 0.38547966
      PC10      PC11      PC12      PC13      PC14      PC15      PC16      PC17      PC18
0.42201002 0.45769568 0.49315997 0.52848461 0.56330705 0.59728325 0.63087849 0.66403529 0.69710187
      PC19      PC20      PC21      PC22      PC23      PC24      PC25      PC26      PC27
0.72908782 0.76040585 0.79091032 0.82076055 0.85047900 0.87967708 0.90840588 0.93583929 0.96226991
      PC28      PC29
0.98174202 1.00000000
> |

```



```

> matlambdas <- rbind(eigen_hcv,propvar,cumvar_hcv)
> rownames(matlambdas) <- c("Eigenvalues","Prop. variance","Cum. prop. variance")
> round(matlambdas,29)

```

	PC1	PC2	PC3	PC4	PC5	PC6	PC7	PC8
Eigenvalues	1.89596904	1.24866396	1.23509034	1.19164939	1.16264975	1.14651308	1.12914639	1.10696093
Prop. variance	0.06537824	0.04305738	0.04258932	0.04109136	0.04009137	0.03953493	0.03893608	0.03817107
Cum. prop. variance	0.06537824	0.10843562	0.15102494	0.19211630	0.23220767	0.27174261	0.31067869	0.34884975

	PC9	PC10	PC11	PC12	PC13	PC14	PC15	PC16
Eigenvalues	1.06226726	1.05938037	1.03488407	1.0284646	1.02441453	1.00985059	0.98530999	0.97426192
Prop. variance	0.03662991	0.03653036	0.03568566	0.0354643	0.03532464	0.03482243	0.03397621	0.03359524
Cum. prop. variance	0.38547966	0.42201002	0.45769568	0.4931600	0.52848461	0.56330705	0.59728325	0.63087849

	PC17	PC18	PC19	PC20	PC21	PC22	PC23	PC24
Eigenvalues	0.9615471	0.95893098	0.92759253	0.90822269	0.88462975	0.86565673	0.86183502	0.84674429
Prop. variance	0.0331568	0.03306659	0.03198595	0.03131802	0.03050447	0.02985023	0.02971845	0.02919808
Cum. prop. variance	0.6640353	0.69710187	0.72908782	0.76040585	0.79091032	0.82076055	0.85047900	0.87967708

	PC25	PC26	PC27	PC28	PC29
Eigenvalues	0.8331352	0.79556880	0.76648805	0.56469127	0.52948135
Prop. variance	0.0287288	0.02743341	0.02643062	0.01947211	0.01825798
Cum. prop. variance	0.9084059	0.93583929	0.96226991	0.98174202	1.00000000

```

> summary(hcv_pca)
Importance of components:

```

	PC1	PC2	PC3	PC4	PC5	PC6	PC7	PC8	PC9	PC10
Standard deviation	1.37694	1.11744	1.11135	1.09163	1.07826	1.07075	1.06261	1.05212	1.03066	1.02926
Proportion of Variance	0.06538	0.04306	0.04259	0.04109	0.04009	0.03953	0.03894	0.03817	0.03663	0.03653
Cumulative Proportion	0.06538	0.10844	0.15102	0.19212	0.23221	0.27174	0.31068	0.34885	0.38548	0.42201

	PC11	PC12	PC13	PC14	PC15	PC16	PC17	PC18	PC19	PC20
Standard deviation	1.01729	1.01413	1.01213	1.00491	0.99263	0.9870	0.98059	0.97925	0.96312	0.95301
Proportion of Variance	0.03569	0.03546	0.03532	0.03482	0.03398	0.0336	0.03316	0.03307	0.03199	0.03132
Cumulative Proportion	0.45770	0.49316	0.52848	0.56331	0.59728	0.6309	0.66404	0.69710	0.72909	0.76041

	PC21	PC22	PC23	PC24	PC25	PC26	PC27	PC28	PC29
Standard deviation	0.9405	0.93041	0.92835	0.9202	0.91276	0.89195	0.87549	0.75146	0.72765
Proportion of Variance	0.0305	0.02985	0.02972	0.0292	0.02873	0.02743	0.02643	0.01947	0.01826
Cumulative Proportion	0.7909	0.82076	0.85048	0.8797	0.90841	0.93584	0.96227	0.98174	1.00000

```

> |

```



```
> hcv_pca$rotation
```

	PC1	PC2	PC3	PC4	PC5
Age	0.043992545	-0.035109552	0.15542945	-0.21681173	0.1846420065
Gender	0.008756739	-0.031538265	0.01225109	0.11432799	-0.2272447661
BMI	0.059438630	0.004938750	-0.21133076	0.43507918	0.2105290684
Fever	-0.018638143	-0.016959013	0.14647417	0.11272529	-0.0194418421
Nausea.Vomting	-0.018052038	0.523815972	0.02931082	0.03619724	0.0100755834
Headache	-0.007027392	0.124936603	0.20495296	-0.22055035	0.0328136750
Diarrhea	-0.037779605	0.112284699	0.31055207	0.09505129	0.1941946991
Fatigue...generalized.bone.ache	-0.039589819	-0.113283136	0.02431225	0.03182207	-0.5018034613
Jaundice	-0.060924334	-0.039975480	0.15722114	-0.28037289	0.0021353121
Epigastric.pain	-0.091593158	-0.314657318	0.08262704	0.13208431	-0.1562642350
WBC	0.079856632	-0.207631406	-0.09056514	0.08357598	0.0941100088
RBC	0.062330494	0.242506719	-0.14413534	-0.04346368	-0.0591444285
HGB	0.012151709	0.030902639	-0.39565562	0.16325012	0.0113733042
Plat	-0.051058807	0.378757510	0.01108177	0.28487255	-0.1382597553
AST.1	0.028793449	0.013713351	0.04919642	0.04294564	-0.1320669862
ALT.1	0.018974095	0.262467291	0.39485581	0.25382822	-0.0368534659
ALT4	-0.040780344	0.034435407	-0.08938033	-0.27015280	0.4057377396
ALT.12	0.047599395	-0.157661272	-0.16534036	-0.38085696	-0.2283790159
ALT.24	-0.018567737	0.212800327	-0.32270954	-0.24975390	0.0210555092
ALT.36	-0.054830139	0.088193685	0.08694682	-0.12844262	-0.2103541059
ALT.48	-0.026185451	-0.255155588	0.31754417	0.06703057	0.0024901089
ALT.after.24.w	-0.019391592	-0.145425335	0.17697872	0.01737588	0.0255244283
RNA.Base	-0.013683156	-0.002907261	0.30619803	-0.11191128	0.1894906165
RNA.4	0.056900757	-0.183428054	-0.04195740	0.16586510	0.3981169569
RNA.12	-0.552679994	0.001687528	-0.02336655	0.03313714	0.0093530466
RNA.EOT	-0.563868495	-0.039526114	-0.07119686	0.03930584	0.0771614060
RNA.EF	-0.570538298	-0.011466450	-0.04305053	-0.00047804	0.0008019717
Baseline.histological.Grading	0.051971451	-0.140448509	0.00159462	0.13066428	-0.1768391380
Baselinehistological.staging	-0.032348367	0.213485043	0.10947878	-0.19354490	-0.1009923507

```
> tabmeansPC <- aggregate(hcvtyp_pca[,2:30],by=list(Survivorship=HCV$Survivorship),mean)
```

```
> tabmeansPC
```

	Survivorship	PC1	PC2	PC3	PC4	PC5	PC6	PC7
1	C	0.6123744	0.02475479	0.06579593	-0.03430532	-0.04507313	0.01425565	-0.04281583
2	NC	-1.1472491	-0.04637671	-0.12326498	0.06426909	0.08444199	-0.02670717	0.08021306
	PC8	PC9	PC10	PC11	PC12	PC13	PC14	PC15
1	-0.004890105	0.06224251	-0.05949025	-0.004544576	-0.03920054	-0.002755935	-0.02361107	0.02266102
2	0.009161338	-0.11660786	0.11145164	0.008514008	0.07344002	0.005163090	0.04423401	-0.04245415
	PC16	PC17	PC18	PC19	PC20	PC21	PC22	PC23
1	0.02326687	0.05588213	0.002161106	0.02975834	-0.002715703	-0.04697310	-0.01407614	-0.001335249
2	-0.04358918	-0.10469204	-0.004048710	-0.05575059	0.005087717	0.08800146	0.02637085	0.002501515
	PC24	PC25	PC26	PC27	PC28	PC29		
1	-0.008750238	0.01021780	-0.02426390	-0.02104547	0.2054995	-0.2452732		
2	0.016393081	-0.01914247	0.04545705	0.03942750	-0.3849919	0.4595055		

```
> |
```



```
> tabmeansPC <- tabmeansPC[rev(order(tabmeansPC$Survivorship)),]
> tabmeansPC
Survivorship      PC1      PC2      PC3      PC4      PC5      PC6      PC7
2      NC -1.1472491 -0.04637671 -0.12326498 0.06426909 0.08444199 -0.02670717 0.08021306
1      C 0.6123744 0.02475479 0.06579593 -0.03430532 -0.04507313 0.01425565 -0.04281583
      PC8      PC9      PC10      PC11      PC12      PC13      PC14      PC15
2 0.009161338 -0.11660786 0.11145164 0.008514008 0.07344002 0.005163090 0.04423401 -0.04245415
1 -0.004890105 0.06224251 -0.05949025 -0.004544576 -0.03920054 -0.002755935 -0.02361107 0.02266102
      PC16      PC17      PC18      PC19      PC20      PC21      PC22      PC23
2 -0.04358918 -0.10469204 -0.004048710 -0.05575059 0.005087717 0.08800146 0.02637085 0.002501515
1 0.02326687 0.05588213 0.002161106 0.02975834 -0.002715703 -0.04697310 -0.01407614 -0.001335249
      PC24      PC25      PC26      PC27      PC28      PC29
2 0.016393081 -0.01914247 0.04545705 0.03942750 -0.3849919 0.4595055
1 -0.008750238 0.01021780 -0.02426390 -0.02104547 0.2054995 -0.2452732
> |
```

```
> tabfmeans <- t(tabmeansPC[, -1])
> tabfmeans
```

	2	1
PC1	-1.147249113	0.612374388
PC2	-0.046376709	0.024754788
PC3	-0.123264984	0.065795927
PC4	0.064269094	-0.034305319
PC5	0.084441992	-0.045073134
PC6	-0.026707165	0.014255652
PC7	0.080213059	-0.042815830
PC8	0.009161338	-0.004890105
PC9	-0.116607855	0.062242510
PC10	0.111451643	-0.059490246
PC11	0.008514008	-0.004544576
PC12	0.073440022	-0.039200543
PC13	0.005163090	-0.002755935
PC14	0.044234009	-0.023611066
PC15	-0.042454146	0.022661017
PC16	-0.043589184	0.023266873
PC17	-0.104692043	0.055882132
PC18	-0.004048710	0.002161106
PC19	-0.055750587	0.029758342
PC20	0.005087717	-0.002715703
PC21	0.088001462	-0.046973095
PC22	0.026370853	-0.014076137
PC23	0.002501515	-0.001335249
PC24	0.016393081	-0.008750238
PC25	-0.019142468	0.010217796
PC26	0.045457051	-0.024263896
PC27	0.039427504	-0.021045467
PC28	-0.384991894	0.205499549
PC29	0.459505511	-0.245273152

```
> |
```

```
> colnames(tabfmeans) <- t(as.vector(tabmeansPC[1]))
> tabfmeans
```

	NC	C
PC1	-1.147249113	0.612374388
PC2	-0.046376709	0.024754788
PC3	-0.123264984	0.065795927
PC4	0.064269094	-0.034305319
PC5	0.084441992	-0.045073134
PC6	-0.026707165	0.014255652
PC7	0.080213059	-0.042815830
PC8	0.009161338	-0.004890105
PC9	-0.116607855	0.062242510
PC10	0.111451643	-0.059490246
PC11	0.008514008	-0.004544576
PC12	0.073440022	-0.039200543
PC13	0.005163090	-0.002755935
PC14	0.044234009	-0.023611066
PC15	-0.042454146	0.022661017
PC16	-0.043589184	0.023266873
PC17	-0.104692043	0.055882132
PC18	-0.004048710	0.002161106
PC19	-0.055750587	0.029758342
PC20	0.005087717	-0.002715703
PC21	0.088001462	-0.046973095
PC22	0.026370853	-0.014076137
PC23	0.002501515	-0.001335249
PC24	0.016393081	-0.008750238
PC25	-0.019142468	0.010217796
PC26	0.045457051	-0.024263896
PC27	0.039427504	-0.021045467
PC28	-0.384991894	0.205499549
PC29	0.459505511	-0.245273152

```
> |
```

```

> tabsdsPC <- aggregate(hcvtyp_pca[,2:30],by=list(survivorship=HCV$survivorship),sd)
> tabfsds <- t(tabsdsPC[,-1])
> colnames(tabfsds) <- t(as.vector(tabsdsPC[1]))
> tabfsds

```

	C	NC
PC1	1.2298718	0.7719668
PC2	1.0967902	1.1548515
PC3	1.1099905	1.1045144
PC4	1.1015816	1.0709045
PC5	1.0739757	1.0823164
PC6	1.0599013	1.0914031
PC7	1.0790286	1.0274616
PC8	1.0597309	1.0387495
PC9	1.0180357	1.0450316
PC10	1.0153498	1.0468206
PC11	1.0196557	1.0138527
PC12	1.0407117	0.9591108
PC13	1.0061530	1.0242748
PC14	1.0201189	0.9752947
PC15	1.0217320	0.9352267
PC16	0.9916090	0.9779775
PC17	0.9823685	0.9696058
PC18	0.9770937	0.9842820
PC19	0.9740629	0.9407323
PC20	0.9417524	0.9747108
PC21	0.9223920	0.9685001
PC22	0.9380579	0.9162855
PC23	0.8997361	0.9806692
PC24	0.9241690	0.9134110
PC25	0.9266039	0.8868826
PC26	0.8808408	0.9115743
PC27	0.8760836	0.8739294
PC28	0.6719146	0.7418754
PC29	0.5891777	0.7402801

T-Tests

```
> t.test(PC1~HCV$Survivorship,data=hcvtyp_pca)
```

```
Welch Two Sample t-test
```

```
data: PC1 by HCV$Survivorship
t = 32.611, df = 1347.9, p-value < 2.2e-16
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
 1.653773 1.865474
sample estimates:
mean in group C mean in group NC
    0.6123744    -1.1472491
```

```
> t.test(PC2~HCV$Survivorship,data=hcvtyp_pca)
```

```
Welch Two Sample t-test
```

```
data: PC2 by HCV$Survivorship
t = 1.111, df = 939.52, p-value = 0.2669
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
-0.0545162 0.1967792
sample estimates:
mean in group C mean in group NC
    0.02475479    -0.04637671
```

```
> t.test(PC3~HCV$Survivorship,data=hcvtyp_pca)
```

```
Welch Two Sample t-test
```

```
data: PC3 by HCV$Survivorship
t = 3.0292, df = 986.5, p-value = 0.002516
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
 0.06658236 0.31153946
sample estimates:
mean in group C mean in group NC
    0.06570502    0.12226108
```

```
> t.test(PC4~HCV$Survivorship,data=hcvtyp_pca)
```

```
Welch Two Sample t-test
```

```
data: PC4 by HCV$Survivorship
t = -1.6155, df = 1006.5, p-value = 0.1065
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
-0.21831093 0.02116211
sample estimates:
mean in group C mean in group NC
   -0.03430532    0.06426909
```

```
> t.test(PC5~HCV$Survivorship,data=hcvtyp_pca)
```

```
Welch Two Sample t-test
```

```
data: PC5 by HCV$Survivorship
t = -2.127, df = 975.75, p-value = 0.03367
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
-0.24900621 -0.01002404
sample estimates:
mean in group C mean in group NC
   -0.04507313    0.08444199
```

F-Ratio Tests

```
> # F ratio tests
> var.test(PC1~HCV$Survivorship,data=hcvtyp_pca)

      F test to compare two variances

data:  PC1 by HCV$Survivorship
F = 2.5382, num df = 902, denom df = 481, p-value < 2.2e-16
alternative hypothesis: true ratio of variances is not equal to 1
95 percent confidence interval:
 2.166058 2.962996
sample estimates:
ratio of variances
    2.53818

> var.test(PC2~HCV$Survivorship,data=hcvtyp_pca)

      F test to compare two variances

data:  PC2 by HCV$Survivorship
F = 0.90198, num df = 902, denom df = 481, p-value = 0.1915
alternative hypothesis: true ratio of variances is not equal to 1
95 percent confidence interval:
 0.7697373 1.0529399
sample estimates:
ratio of variances
    0.9019758

> var.test(PC3~HCV$Survivorship,data=hcvtyp_pca)

      F test to compare two variances

data:  PC3 by HCV$Survivorship
F = 1.0099, num df = 902, denom df = 481, p-value = 0.9079
alternative hypothesis: true ratio of variances is not equal to 1
95 percent confidence interval:
 0.8618732 1.1789746
sample estimates:
ratio of variances
    1.00994



---


> var.test(PC4~HCV$Survivorship,data=hcvtyp_pca)

      F test to compare two variances

data:  PC4 by HCV$Survivorship
F = 1.0581, num df = 902, denom df = 481, p-value = 0.4858
alternative hypothesis: true ratio of variances is not equal to 1
95 percent confidence interval:
 0.9029828 1.2352093
sample estimates:
ratio of variances
    1.058113

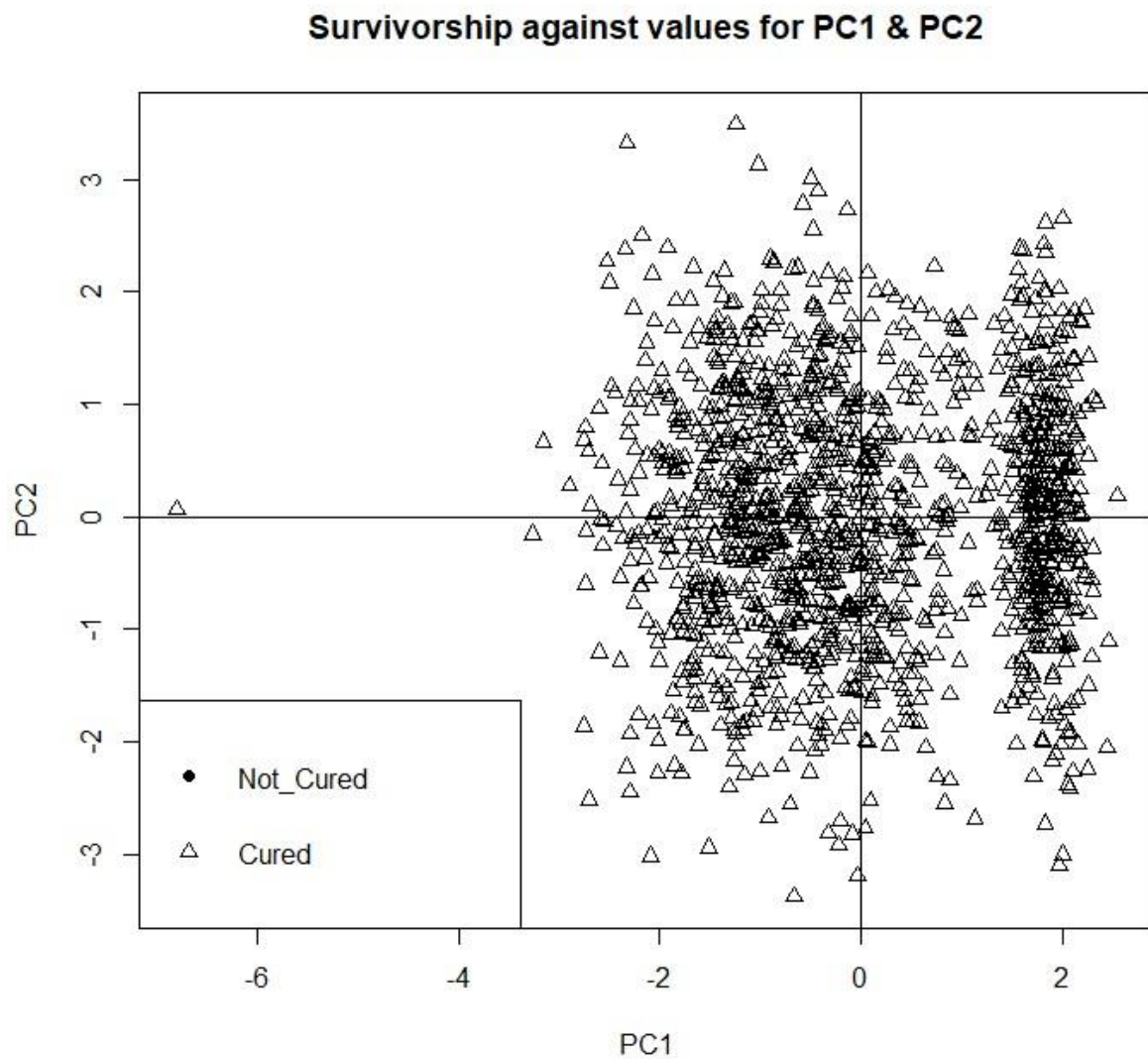
> var.test(PC5~HCV$Survivorship,data=hcvtyp_pca)

      F test to compare two variances

data:  PC5 by HCV$Survivorship
F = 0.98465, num df = 902, denom df = 481, p-value = 0.8399
alternative hypothesis: true ratio of variances is not equal to 1
95 percent confidence interval:
 0.8402878 1.1494474
sample estimates:
ratio of variances
    0.9846467
```

Plotting the scores for the first and second components

```
plot(hcvtyp_pca$PC1, hcvtyp_pca$PC2, pch=ifelse(hcvtyp_pca$Survivorship ==  
"S", 16, 2), xlab="PC1", ylab="PC2", main="49 HCV against values for PC1 & PC2")  
> abline(h=0)  
> abline(v=0)  
> legend("bottomleft", legend=c("Cured", "Not_Cured"), pch=c(16, 2),)
```




```

> plot(eigen_hcv, xlab = "Component number", ylab = "Component variance", type = "l", main = "Scree diagram")
> plot(log(eigen_hcv), xlab = "Component number", ylab = "log(Component variance)", type="l", main = "Log(eigenvalue) diagram")
> print(summary(hcv_pca))
Importance of components:

```

	PC1	PC2	PC3	PC4	PC5	PC6	PC7	PC8	PC9	PC10
Standard deviation	1.37694	1.11744	1.11135	1.09163	1.07826	1.07075	1.06261	1.05212	1.03066	1.02926
Proportion of Variance	0.06538	0.04306	0.04259	0.04109	0.04009	0.03953	0.03894	0.03817	0.03663	0.03653
Cumulative Proportion	0.06538	0.10844	0.15102	0.19212	0.23221	0.27174	0.31068	0.34885	0.38548	0.42201

	PC11	PC12	PC13	PC14	PC15	PC16	PC17	PC18	PC19	PC20
Standard deviation	1.01729	1.01413	1.01213	1.00491	0.99263	0.9870	0.98059	0.97925	0.96312	0.95301
Proportion of Variance	0.03569	0.03546	0.03532	0.03482	0.03398	0.0336	0.03316	0.03307	0.03199	0.03132
Cumulative Proportion	0.45770	0.49316	0.52848	0.56331	0.59728	0.6309	0.66404	0.69710	0.72909	0.76041

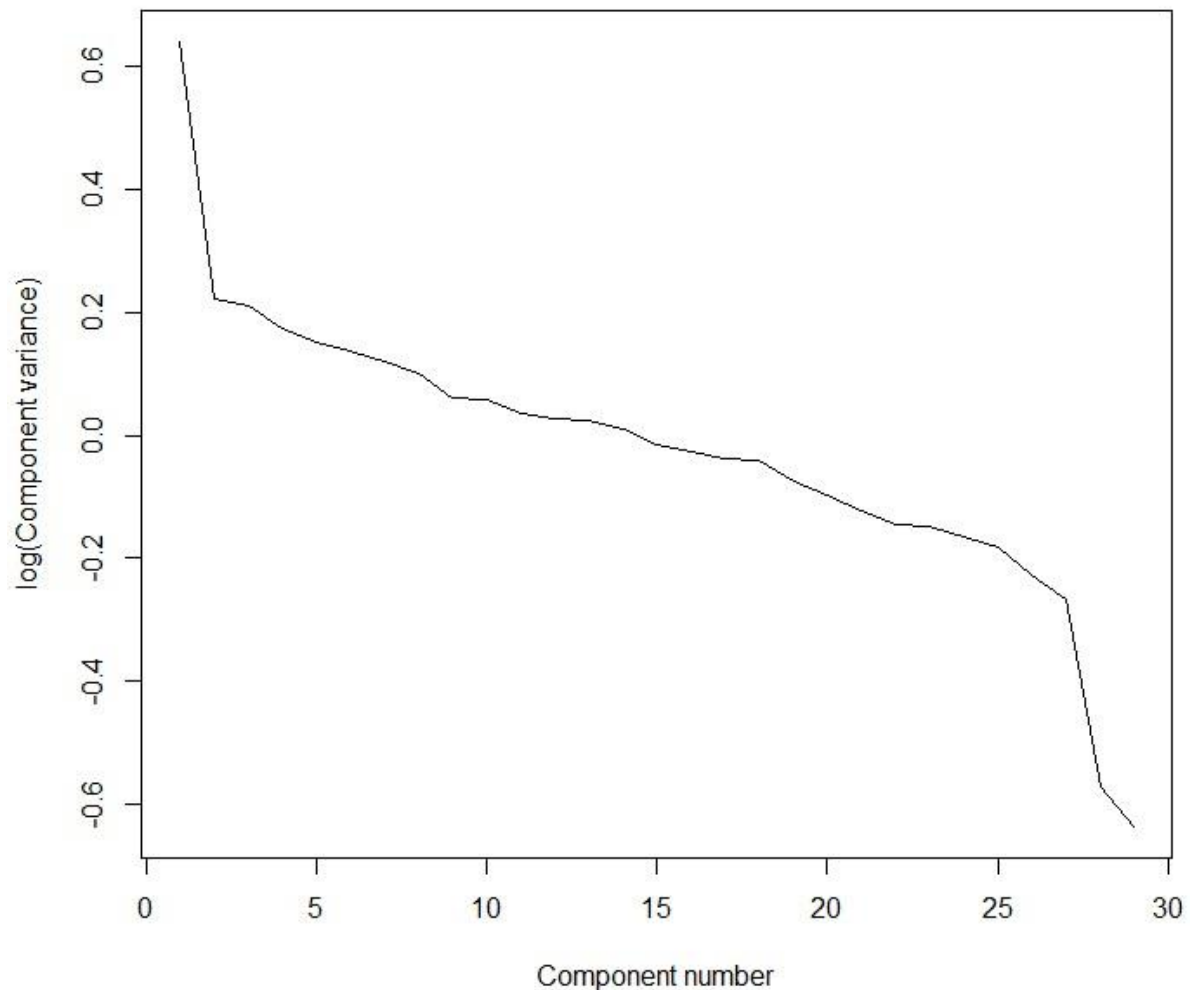
	PC21	PC22	PC23	PC24	PC25	PC26	PC27	PC28	PC29
Standard deviation	0.9405	0.93041	0.92835	0.9202	0.91276	0.89195	0.87549	0.75146	0.72765
Proportion of Variance	0.0305	0.02985	0.02972	0.0292	0.02873	0.02743	0.02643	0.01947	0.01826
Cumulative Proportion	0.7909	0.82076	0.85048	0.8797	0.90841	0.93584	0.96227	0.98174	1.00000

```

> |

```

Log(eigenvalue) diagram



```

> view(hcv_pca)
> diag(cov(hcv_pca$x))

```

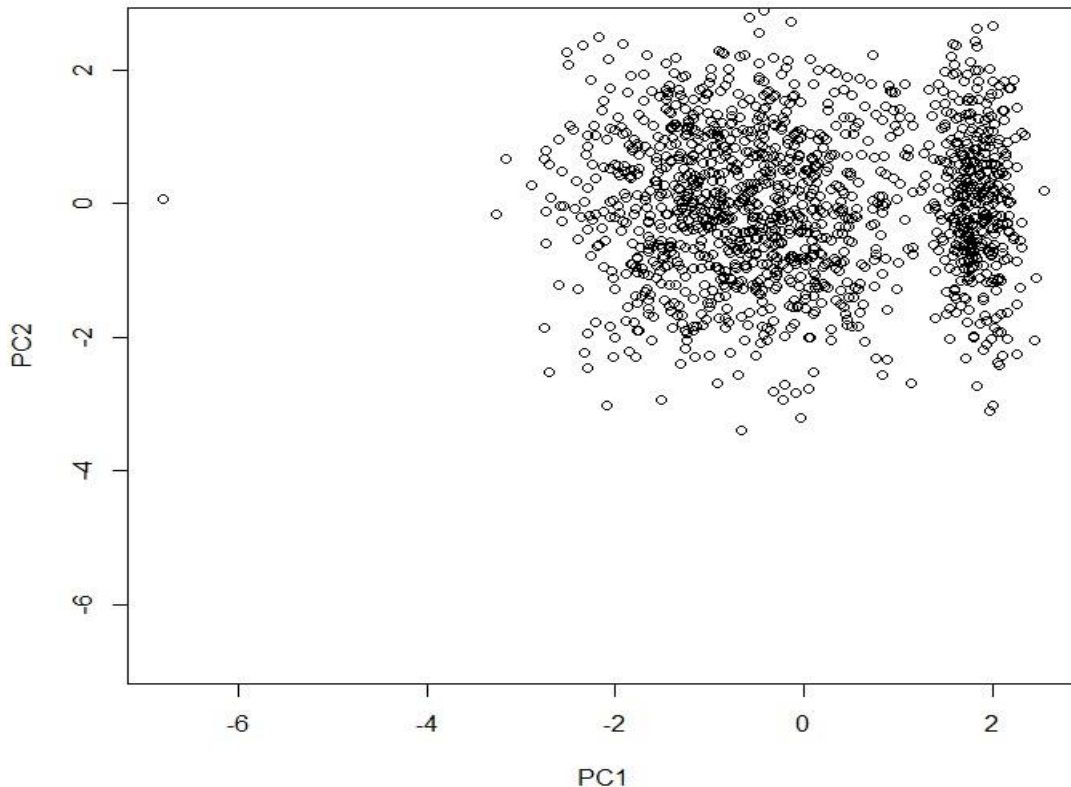
	PC1	PC2	PC3	PC4	PC5	PC6	PC7	PC8	PC9	PC10
1.8959690	1.2486640	1.2350903	1.1916494	1.1626498	1.1465131	1.1291464	1.1069609	1.0622673	1.0593804	
	PC11	PC12	PC13	PC14	PC15	PC16	PC17	PC18	PC19	PC20
1.0348841	1.0284646	1.0244145	1.0098506	0.9853100	0.9742619	0.9615471	0.9589310	0.9275925	0.9082227	
	PC21	PC22	PC23	PC24	PC25	PC26	PC27	PC28	PC29	
0.8846297	0.8656567	0.8618350	0.8467443	0.8331352	0.7955688	0.7664880	0.5646913	0.5294814		

```

> xlim <- range(hcv_pca$x[,1])
> hcv_pca$x[,1]

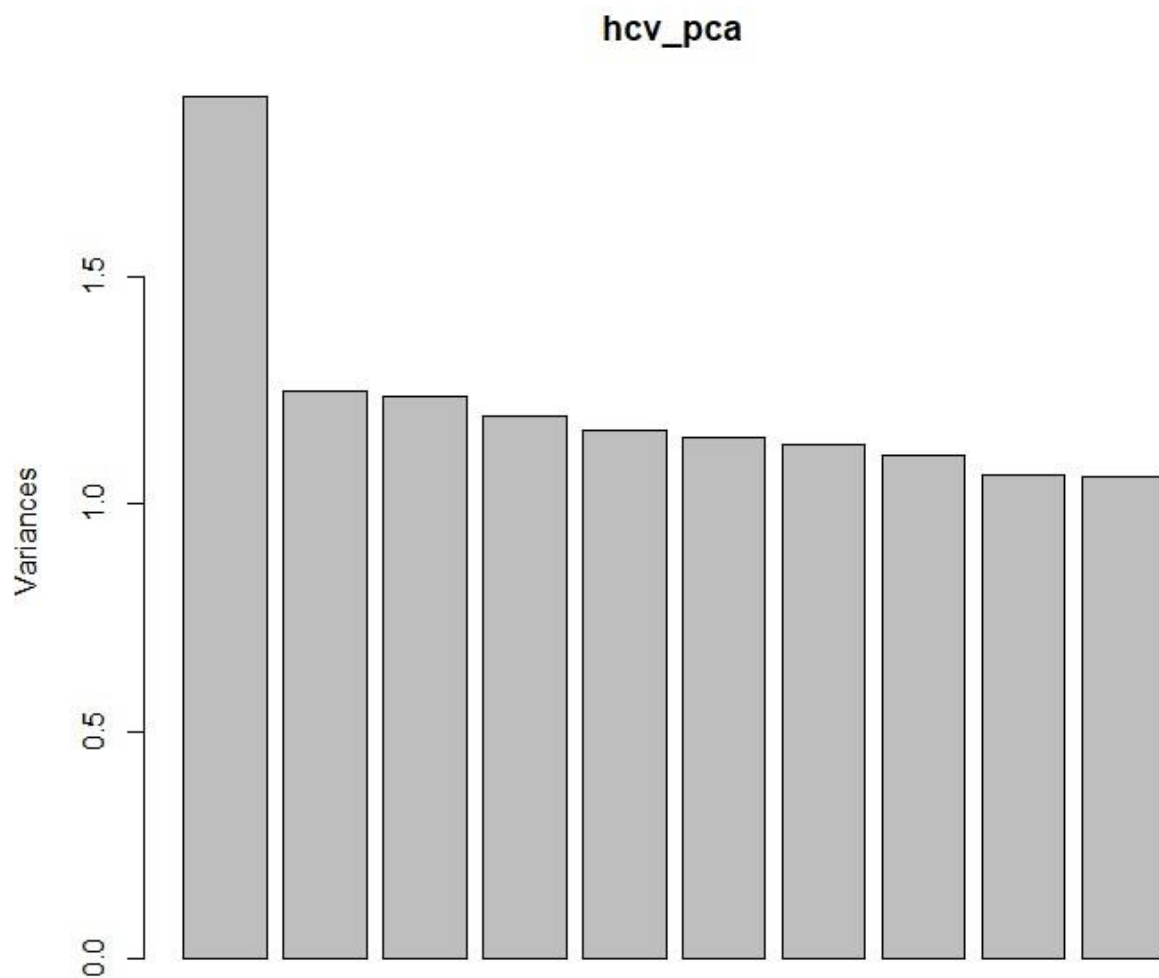
```

[1]	1.752028782	-0.125120259	-0.582399282	-2.085681979	-6.798188688	1.749113325	-0.590303483
[8]	-1.564717696	0.510283184	-0.588712397	-1.022320922	-0.962514349	1.707989062	-1.528713168
[15]	-0.049587298	0.432247099	1.852374878	-0.660406526	-1.127124155	-1.636602476	-1.088482877
[22]	1.666271696	1.064152892	-0.798078912	-0.969386323	2.021606955	-2.013528283	-0.017980495
[29]	1.760727933	-0.985329698	0.996146113	0.333530076	-1.780818574	-0.025899178	-1.417814487
[36]	-1.793894337	-0.390902443	-0.139473954	1.643574847	-0.669603195	-1.631512707	0.502948531
[43]	0.588390510	-2.586019373	0.116808274	-0.524253243	-0.778339505	-0.320301025	-2.294294180
[50]	-0.281952226	-0.510799747	0.456407698	-0.422122534	-1.374782140	-0.329563302	0.099821119
[57]	0.934665222	-0.438548488	-0.594170636	0.011562761	0.543354103	-1.773278006	-0.832846177
[64]	-2.127924184	-2.433998741	-0.427582983	0.734606924	-0.172389851	-0.842059433	-1.196245523
[71]	-1.179017676	-0.201355958	-0.579293636	-1.652600524	-1.469329779	1.320019225	-0.864879087
[78]	-1.170746846	-1.323100814	-0.025789209	-1.221369325	-0.488701234	-0.907145939	-0.766911350
[85]	-0.783371949	0.827179651	-0.606542502	-2.056249867	-1.799195517	-0.810106919	0.374473782
[92]	-1.337203727	1.750690841	-1.145751688	-1.534882120	2.075021466	-2.133537704	0.549400134
[99]	2.045395929	0.925674949	-2.049845810	-1.420305157	1.666908549	-1.107860127	2.079193403
[106]	-0.500317409	1.980382756	1.978614381	-0.970399214	1.939636252	1.489137414	-2.28822057
[113]	0.096747085	-2.310899960	-0.497826028	1.750726115	2.044411160	-2.295321121	0.823883639
[120]	-0.542571573	-0.033144043	1.657684088	-2.003172967	1.600796059	1.717756417	2.221271450
[127]	-2.604177500	1.817323300	-1.325819301	0.651781756	-1.287569137	-1.418202444	-1.180130962
[134]	-0.566850896	-0.100026377	-2.365308311	-2.072957921	-0.694508820	-1.222450327	0.185452086
[141]	1.634829190	-1.799663398	0.074848618	1.574529284	1.898653759	0.957344290	0.158573101
[148]	1.503510943	-1.873590833	-0.539976869	1.931465640	-1.573019968	-0.480779763	2.062468988
[155]	1.891818100	-1.850684355	-1.914856651	1.972169814	-1.826473077	1.839397182	1.915152600
[162]	0.135647552	0.112017231	0.521292370	1.681489545	0.110022689	-1.267206207	-0.031278772
[169]	-0.078879118	1.642842120	1.790709740	-2.551695792	-0.638398692	2.045832372	0.866146959
[176]	2.115080577	-0.848224403	-0.310674491	1.562725642	1.773527100	2.058737931	1.894057037
[183]	-2.243139605	-0.580830939	1.107961882	-0.328978421	-0.056139128	0.436533736	-1.279777144



```
> plot(hcv_pca$x,xlim=xlim,ylim=xlim)
> hcv_pca$rotation[,1]
```

Age	Gender	BMI
0.043992545	0.008756739	0.059438630
Fever	Nausea.Vomiting	Headache
-0.018638143	-0.018052038	-0.007027392
Diarrhea	Fatigue...generalized.bone.ache	Jaundice
-0.037779605	-0.039589819	-0.060924334
Epigastric.pain	WBC	RBC
-0.091593158	0.079856632	0.062330494
HGB	Plat	AST.1
0.012151709	-0.051058807	0.028793449
ALT.1	ALT4	ALT.12
0.018974095	-0.040780344	0.047599395
ALT.24	ALT.36	ALT.48
-0.018567737	-0.054830139	-0.026185451
ALT.after.24.w	RNA.Base	RNA.4
-0.019391592	-0.013683156	0.056900757
RNA.12	RNA.EOT	RNA.EF
-0.552679994	-0.563868495	-0.570538298
Baseline.histological.Grading	Baselinehistological.staging	
0.051971451	-0.032348367	




```
out <- sapply(1:5,
function(i){plot(HCV$Survivorship,hcv_pca$x[,i],xlab=paste("PC",i,sep=""),ylab="Survivorshi
p")})
```

```
pairs(hcv_pca$x[,1:5], ylim = c(-6,4),xlim = c(-
6,4),panel=function(x,y,...){text(x,y,HCV$Survivorship)})
```

