

# Using MBatch Assessments: Boxplot\_\_AllSamplesData\_\_Structures

*Tod Casasent*

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## 1 Introduction

These instructions are aimed at people familiar with R and familiar with TCGA/GDC platforms and data types. They are intended to introduce the reader to producing the given assessment. These instructions will only rarely, if ever, touch on the appropriateness of the assessment algorithm or interpretation of output. See MBatch\_01\_InstallLinux for instructions on downloading test data.

## 2 Algorithm

Boxplot\_\_AllSamplesData\_\_Structures is a function used to perform batch effects assessments using the boxplots on all samples without modification.

## 3 Output

The primary output method for MBatch is to view results in the Batch Effects Website, described elsewhere. The PNG files are rough versions of the website output.

Graphical output is a set of boxplots where each boxplot (also called a box and whisper plot) represent a single sample. For datasets with many samples, the static PNG may be so dense as to be unusable.

Here is an example of a smallish dynamic boxplot. (See Batch Effects Viewer documentation for more details.)

Here is an example of the static plot for a medium-sized dataset.

## 4 Usage

Boxplot\_\_AllSamplesData\_\_Structures(theData, theTitle, theOutputPath, theBatchTypeAndValuePairsToRemove, theBatchTypeAndValuePairsToKeep, theMaxGeneCount=20000, theJavaParameters = “-Xms8000m”)

## 5 Arguments

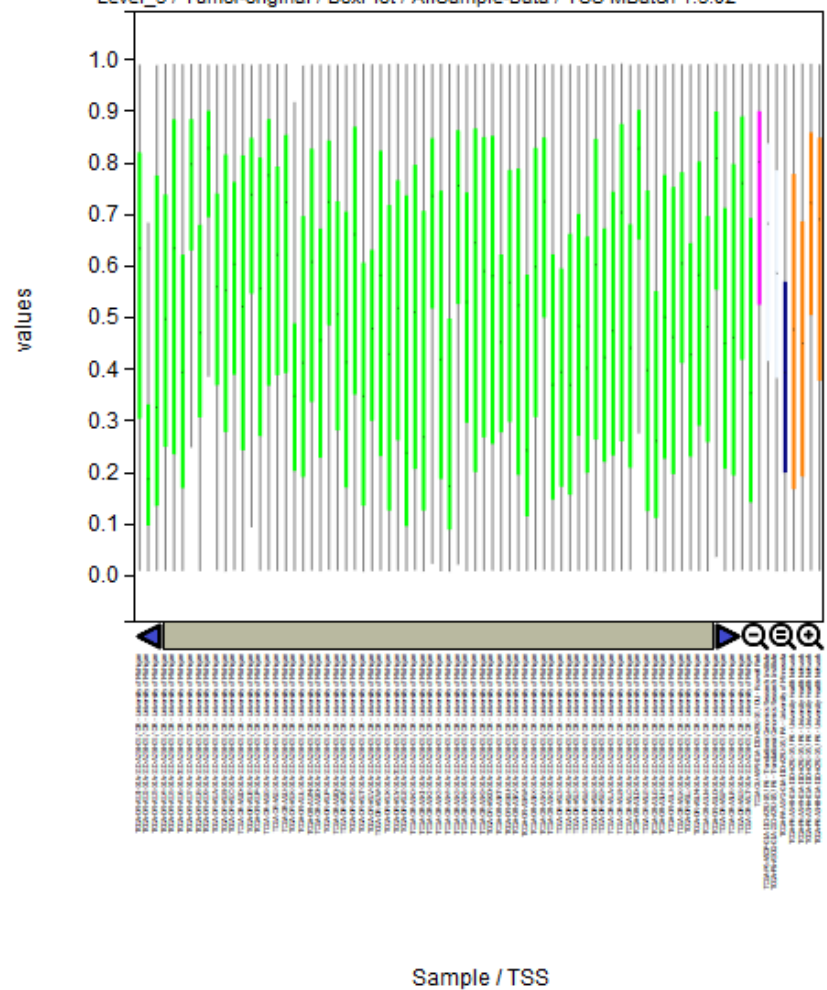
### 5.1 theData

An instance of BEA\_\_DATA.

BEA\_\_DATA objects can be created by calls of the form new(“BEA\_\_DATA”, theData, theBatches, theCovariates). If you have no covariate data, use an empty data.frame created with data.frame()

mData: Object of class “matrix” A matrix where the colnames are sample ids and the rownames are gene equivalents. All names should be strings, not factors.

2016\_06\_13\_0834-2016\_08\_16\_1052 / acc / methylation / humanmethylation450\_methWxy /  
Level\_3 / Tumor-original / BoxPlot / AllSample-Data / TSS MBatch 1.3.02



TCGA-OR-A5J1-01A-11D-A29J-05  
n=99988

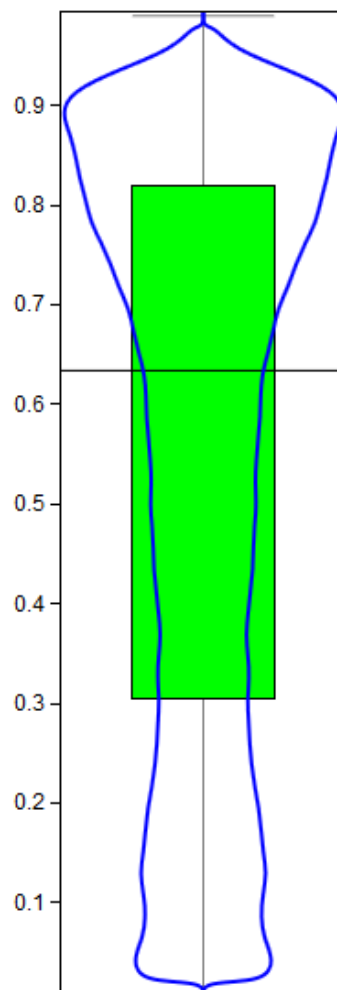


Figure 1: Dynamic Boxplot Example

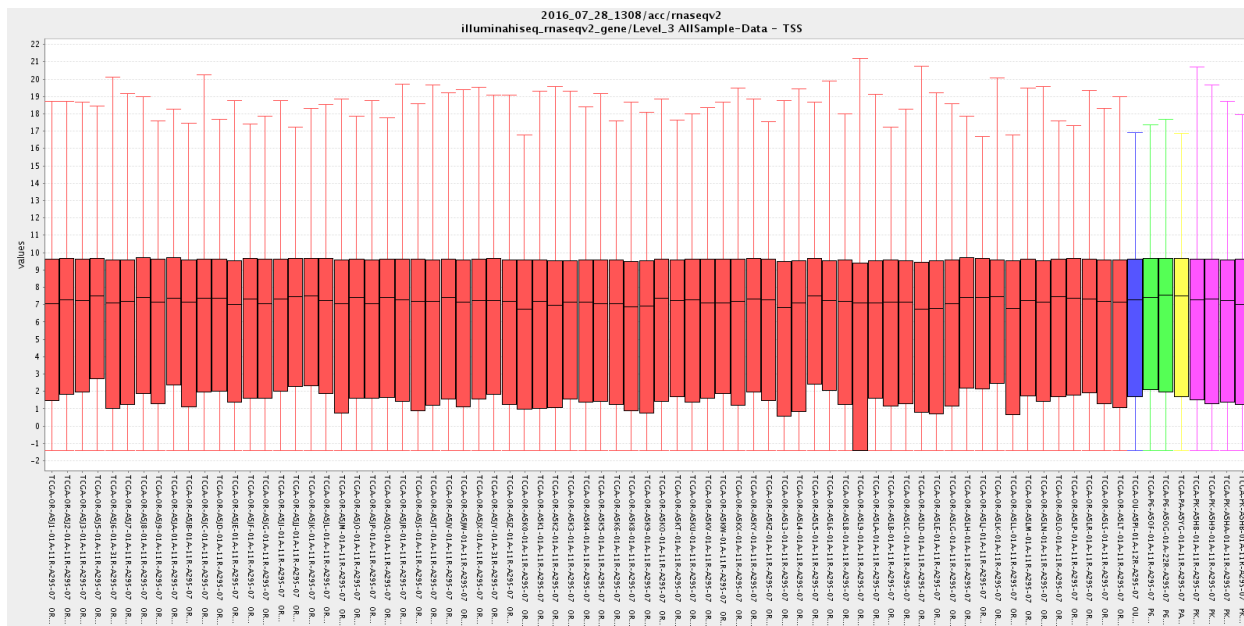


Figure 2: Static Boxplot Example

**mBatches:** Object of class “data.frame” A data.frame where the column “names” are batch types. The first batch “type” is “Sample”. All names and values should be strings, not factors or numeric.

**mCovariates:** Object of class “data.frame” A data.frame where the column “names” are covariate types. The first covariate “type” is “Sample”. All names and values should be strings, not factors or numeric.

## 5.2 theTitle

A string title to use in PNG files.

## 5.3 theOutputPath

String giving directory in which to place output PNG files.

## 5.4 theBatchTypeAndValuePairsToRemove

A list of vectors containing the batch type (or \* for all types) and the value to remove. list() indicates none while NULL will cause an error.

## 5.5 theBatchTypeAndValuePairsToKeep

A list of vectors containing the batch type (or \* for all types) and a vector of the the value(s) to keep. list() indicates none while NULL will cause an error.

## 5.6 theMaxGeneCount

Integer giving maximum number of features (genes) to keep. Default is 20000. 0 means keep all.

## 5.7 theJavaParameters

Object of class “character” String for initializing JVM. Defaults to -Xms8000m.

## 6 Example Call

The following code is adapted from the tests/Boxplot\_AllSamplesData\_Structures file. Data used is from the testing data as per the MBatch\_01\_InstallLinux document. In the future, we plan to make the output from MBatch more user friendly, but currently, this produces the following output at the command line.

This output can generally be skipped as very long and generally obscure. After the output is an explanation of files and directories created.

```
{
  library(MBatch)

  # set the paths
  theGeneFile="/bea_testing/MATRIX_DATA/matrix_data-Tumor.tsv"
  theBatchFile="/bea_testing/MATRIX_DATA/batches-Tumor.tsv"
  theOutputDir="/bea_testing/output/Boxplot_AllSamplesData_Structures"
  theRandomSeed=314

  # make sure the output dir exists and is empty
  unlink(theOutputDir, recursive=TRUE)
  dir.create(theOutputDir, showWarnings=FALSE, recursive=TRUE)

  # load the data and reduce the amount of data to reduce run time
  myData <- mbatchLoadFiles(theGeneFile, theBatchFile)
  myData@mData <- mbatchTrimData(myData@mData, 100000)

  # here, we take most defaults
  Boxplot_AllSamplesData_Structures(myData, "Disease/Data Type/Platform/Data Level", theOutputDir, list
}

## 2018 06 21 10:32:35.842 DEBUG megazone23 Changing LC_COLLATE to C for duration of run
## 2018 06 21 10:32:35.843 INFO megazone23 \ / \ / \ / \ / \ / \ / \ / \ / \ /
## 2018 06 21 10:32:35.844 INFO megazone23 Starting mbatchLoadFiles
## 2018 06 21 10:32:35.844 INFO megazone23 MBatch Version: 2017-09-19-1530
## 2018 06 21 10:32:35.844 INFO megazone23 read batch file= /bea_testing/MATRIX_DATA/batches-Tumor.tsv
## 2018 06 21 10:32:35.846 INFO megazone23 read gene file= /bea_testing/MATRIX_DATA/matrix_data-Tumor.t
## 2018 06 21 10:32:44.676 INFO megazone23 filter samples in batches using gene samples
## 2018 06 21 10:32:44.677 INFO megazone23 sort batches by gene file samples
## 2018 06 21 10:32:44.766 INFO megazone23 Finishing mbatchLoadFiles
## 2018 06 21 10:32:44.766 INFO megazone23 ~~~~~
## 2018 06 21 10:32:44.766 DEBUG megazone23 Changing LC_COLLATE to C for duration of run
## 2018 06 21 10:32:44.767 INFO megazone23 \ / \ / \ / \ / \ / \ / \ / \ / \ /
## 2018 06 21 10:32:44.767 INFO megazone23 mbatchTrimData Starting
## 2018 06 21 10:32:44.767 INFO megazone23 MBatch Version: 2017-09-19-1530
## 2018 06 21 10:32:52.584 INFO megazone23 mbatchTrimData Finishing
## 2018 06 21 10:32:52.584 INFO megazone23 ~~~~~
## 2018 06 21 10:32:52.586 DEBUG megazone23 Changing LC_COLLATE to C for duration of run
## 2018 06 21 10:32:52.587 INFO megazone23 \ / \ / \ / \ / \ / \ / \ / \ / \ /
## 2018 06 21 10:32:52.587 INFO megazone23 mbatchFilterData Starting
## 2018 06 21 10:32:52.587 INFO megazone23 MBatch Version: 2017-09-19-1530
```

```
## 2018 06 21 10:32:52.588 DEBUG megazone23 rows pre filter 1250
## 2018 06 21 10:32:52.844 DEBUG megazone23 rows post filter 1250
## 2018 06 21 10:32:52.844 DEBUG megazone23 mbatchFilterData Prefilter, gene data had 1250 while post
## 2018 06 21 10:32:52.845 DEBUG megazone23 mbatchFilterData Prefilter, batch data had 80 while post
## 2018 06 21 10:32:52.845 INFO megazone23 mbatchFilterData Finishing
## 2018 06 21 10:32:52.846 INFO megazone23 ~~~~~
## 2018 06 21 10:32:52.846 DEBUG megazone23 Changing LC_COLLATE to C for duration of run
## 2018 06 21 10:32:52.846 INFO megazone23 \ / \ / \ / \ / \ / \ / \ / \ / \ /
## 2018 06 21 10:32:52.846 INFO megazone23 mbatchTrimData Starting
## 2018 06 21 10:32:52.847 INFO megazone23 MBatch Version: 2017-09-19-1530
## 2018 06 21 10:32:52.847 INFO megazone23 mbatchTrimData Finishing
## 2018 06 21 10:32:52.847 INFO megazone23 ~~~~~
## 2018 06 21 10:32:52.847 DEBUG megazone23 checkCreateDir: /bea_testing/output/Boxplot_AllSamplesData
## 2018 06 21 10:32:52.849 DEBUG megazone23 boxplotJinit - Calling .jinit /home/linux/R/x86_64-pc-linux
## 2018 06 21 10:33:04.520 DEBUG megazone23 after allSampleData call

## [1] TRUE
```

## 7 Example File Output

The above code creates the following subdirectories and files. The subdirectories correspond to the run type were requested.

```
/bea_testing/output/Boxplot_AllSamplesData_Structures$ ls -l
total 44
drwxr-xr-x 2 linux linux 40960 Jun 19 11:41 AllSample-RLE
```

Looking at the “AllSample-RLE” subdirectory, it contains the diagram and legend files, and data usable with dynamic displays.

```
/bea_testing/output/Boxplot_AllSamplesData_Structures/AllSample-Data$ ls -l
total 6228
-rw-r--r-- 1 linux linux 3873 Jun 19 15:12 BoxPlot_AllSample-Data_Annotations-BatchId.tsv
-rw-r--r-- 1 linux linux 3873 Jun 19 15:13 BoxPlot_AllSample-Data_Annotations-PlateId.tsv
-rw-r--r-- 1 linux linux 3873 Jun 19 15:13 BoxPlot_AllSample-Data_Annotations-ShipDate.tsv
-rw-r--r-- 1 linux linux 3873 Jun 19 15:13 BoxPlot_AllSample-Data_Annotations-TSS.tsv
-rw-r--r-- 1 linux linux 15072 Jun 19 15:12 BoxPlot_AllSample-Data_BoxData-BatchId.tsv
-rw-r--r-- 1 linux linux 15072 Jun 19 15:13 BoxPlot_AllSample-Data_BoxData-PlateId.tsv
-rw-r--r-- 1 linux linux 15072 Jun 19 15:13 BoxPlot_AllSample-Data_BoxData-ShipDate.tsv
-rw-r--r-- 1 linux linux 15072 Jun 19 15:13 BoxPlot_AllSample-Data_BoxData-TSS.tsv
-rw-r--r-- 1 linux linux 9 Jun 19 15:12 BoxPlot_AllSample-Data_CatData-BatchId-TCGA-OR-A5J1-01A-11D-A2
-rw-r--r-- 1 linux linux 7552 Jun 19 15:12 BoxPlot_AllSample-Data_CatData-BatchId-TCGA-OR-A5J2-01A-11D-A2
#snipped out "CatData" files for each sample for each batch type
-rw-r--r-- 1 linux linux 6469 Jun 19 15:13 BoxPlot_AllSample-Data_CatData-TSS-TCGA-PK-A5HA-01A-11D-A2
-rw-r--r-- 1 linux linux 5461 Jun 19 15:13 BoxPlot_AllSample-Data_CatData-TSS-TCGA-PK-A5HB-01A-11D-A2
-rw-r--r-- 1 linux linux 70954 Jun 19 15:12 BoxPlot_AllSample-Data_Diagram-BatchId.png
-rw-r--r-- 1 linux linux 70493 Jun 19 15:13 BoxPlot_AllSample-Data_Diagram-PlateId.png
-rw-r--r-- 1 linux linux 70713 Jun 19 15:13 BoxPlot_AllSample-Data_Diagram-ShipDate.png
-rw-r--r-- 1 linux linux 66492 Jun 19 15:13 BoxPlot_AllSample-Data_Diagram-TSS.png
-rw-r--r-- 1 linux linux 916490 Jun 19 15:13 BoxPlot_AllSample-Data_Histogram-BatchId.png
-rw-r--r-- 1 linux linux 44432 Jun 19 15:12 BoxPlot_AllSample-Data_Histogram-BatchId.tsv
-rw-r--r-- 1 linux linux 916490 Jun 19 15:13 BoxPlot_AllSample-Data_Histogram-PlateId.png
-rw-r--r-- 1 linux linux 44432 Jun 19 15:13 BoxPlot_AllSample-Data_Histogram-PlateId.tsv
-rw-r--r-- 1 linux linux 916490 Jun 19 15:13 BoxPlot_AllSample-Data_Histogram-ShipDate.png
```

```
-rw-r--r-- 1 linux linux 44432 Jun 19 15:13 BoxPlot_AllSample-Data_Histogram-ShipDate.tsv
-rw-r--r-- 1 linux linux 916490 Jun 19 15:13 BoxPlot_AllSample-Data_Histogram-TSS.png
-rw-r--r-- 1 linux linux 44432 Jun 19 15:13 BoxPlot_AllSample-Data_Histogram-TSS.tsv
-rw-r--r-- 1 linux linux 4431 Jun 19 15:12 BoxPlot_AllSample-Data_Legend-BatchId.png
-rw-r--r-- 1 linux linux 4450 Jun 19 15:13 BoxPlot_AllSample-Data_Legend-PlateId.png
-rw-r--r-- 1 linux linux 4521 Jun 19 15:13 BoxPlot_AllSample-Data_Legend-ShipDate.png
-rw-r--r-- 1 linux linux 13135 Jun 19 15:13 BoxPlot_AllSample-Data_Legend-TSS.png
```

## 7.1 Files

Example data may not match output from above.

### 7.1.1 Annotations Files

Looking at BoxPlot\_AllSample-RLE\_Annotations-TSS.tsv, we see it is a tab-delimited file, with two columns with the headers “key” nad “value”. The first entry after that is the “Total-Data-Points”, and then for each sample, we have the number of points available for that sample that are not NA. These two numbers will not always be equal, since some samples may have NAs for genes or probes where the other samples have values.

```
key value
Total-Data-Points 1250
Non-NA-Points-TCGA-OR-A5J1-01A-11D-A29J-05 1250
Non-NA-Points-TCGA-OR-A5J2-01A-11D-A29J-05 1250
Non-NA-Points-TCGA-OR-A5J3-01A-11D-A29J-05 1250
Non-NA-Points-TCGA-OR-A5J4-01A-11D-A29J-05 1250
Non-NA-Points-TCGA-OR-A5J5-01A-11D-A29J-05 1250
```

### 7.1.2 BoxData Files

Looking at BoxPlot\_AllSample-RLE\_BoxData-TSS.tsv, we see it is a tab delimited file with headers indicating the Id (sample) and the different parts of the boxplot. Subsequent rows give the box settings for each sample. NAs are possible in this data.

Id	LowerOutMax	LowerOutMin	LowerNotch	LowerWhisker	LowerHinge	Median	UpperHinge	UpperWhisker
TCGA-OR-A5J1-01A-11D-A29J-05			NA	NA	-0.020527642802858643	-0.8467955227772493	-0.4056428985980960	
TCGA-OR-A5J2-01A-11D-A29J-05			NA	NA	-0.002911079872554134	-0.039930119705853896	-0.021222413369	
TCGA-OR-A5J3-01A-11D-A29J-05			NA	NA	-0.035001758602725926	-0.3988124487830225	-0.3498757664560811	
TCGA-OR-A5J4-01A-11D-A29J-05			NA	NA	-0.017185120892053492	-0.8247218460963763	-0.3183107584949181	
TCGA-OR-A5J5-01A-11D-A29J-05			NA	NA	-0.03364073791133153	-0.8079754846584357	-0.644252206301912	
TCGA-OR-A5J6-01A-31D-A29J-05			NA	NA	-0.0034328681890936023	-0.04187597080189986	-0.022337461512	
TCGA-OR-A5J7-01A-11D-A29J-05					-0.865878813052012	-0.18821669173503153	-0.003889412141825995	-0.

### 7.1.3 CatData Files

If we look at BoxPlot\_AllSample-RLE\_CatData-TSS-TCGA-PK-A5HB-01A-11D-A29J-05.tsv, we see it is a tab-delimited file with “id” and “value” as headers. The id is a feature (in this case a gene, probe, location) combination and then the value from the data for that id. This is used to populate the violin plot with a subset of outliers, if any.

```
id value
ADCY4-cg14287235-14-24804339 -0.7667974166463363
ASCL2-cg12499235-11-2293173 -0.7077020078715286
```

```
BAI1-cg09968723-8-143545789 -0.8074333452970504
BNC1-cg06523224-15-83953883 -0.7850694441252194
```

## 7.1.4 Histogram Data Files

Looking at BoxPlot\_AllSample-RLE\_Histogram-TSS.tsv, we see it is a tab-delimited file. The first row is headers, with “entry” and “size” being the first two, followed by pairs of headers of the form “xN” and “yN”, where they are pairs of X,Y coordinates for plotting the histogram. The entry column is the sample id and the size entry is the number of X,Y pairs.

entry	size	x0	y0	x1	y1	x2	y2	x3	y3	x4	y4	x5	y5	x6	y6	x7	y7	x8	y8	x9	y9	x10	y10
TCGA-OR-A5J1-01A-11D-A29J-05	12						-0.8064387185053226	193.0				-0.7257251099614688	44.0								-0.64501150		
TCGA-OR-A5J2-01A-11D-A29J-05	79						-0.033911616995144944					168.0			-0.02187461157372705						253.0		-0.
TCGA-OR-A5J3-01A-11D-A29J-05	7						-0.32982819164709853					520.0			-0.19185967737525045						68.0		-0.

## 7.2 Diagram

Here is a diagram generated from this code.

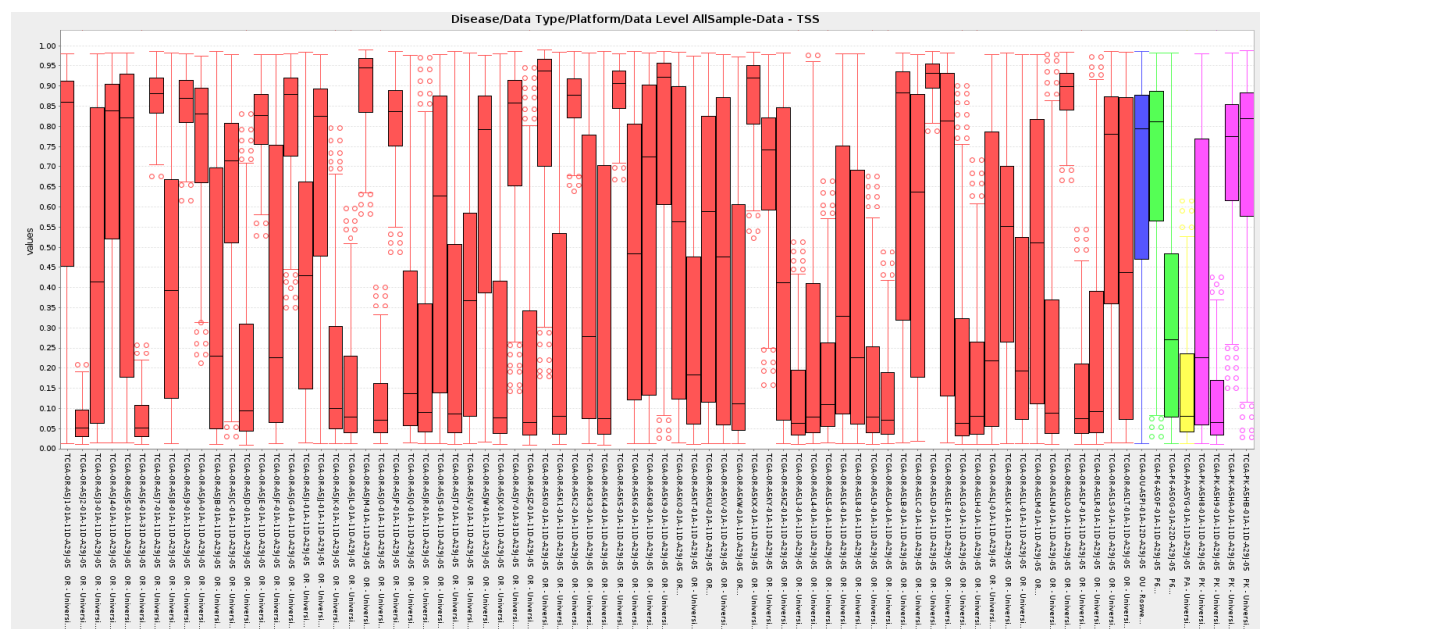


Figure 3: Boxplot All Samples Data Output