# MBatch 05-04 Using MBatch Assessments: RBN\_Pseudoreplicates Tod Casasent 2017-11-10-1455

### 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.docx for instructions on downloading test data.

# 2 Algorithm

RBN\_Pseudoreplicatesis a function used to perform the RBN correction algorithm on two datasets, using pseudoreplicates. This function takes structures (matrices) and returns a corrected matrix of data.

# 3 Output

The primary output method for MBatch is to view results in the Batch Effects Website. Correction algorithms generally do not create graphical output and instead create TSV output files.

# 4 Usage

RBN\_Pseudoreplicates(theInvariantMatrix, theVariantMatrix, theInvariantReplicates, theVariantReplicates, theInvariantGroupId = "", theVariantGroupId = "", theMatchedReplicatesFlag = TRUE, theCombineOnlyFlag = FALSE, thePath = NULL, theWriteToFile = FALSE)

# 5 Arguments

**theInvariantMatrix** Matrix with sample names in colnames and features (like genes) in rownames. This matrix is invariant.

**theVariantMatrix** Matrix with sample names in colnames and features (like genes) in rownames. This matrix is variant.

**theInvariantReplicates** Vector of feature ids indicating replicates for variant data.

**theVariantReplicates** Vector of feature ids indicating replicates for invariant data.

**theInvariantGroupId** Group name used for labelling invariant features when combining matrixes. This defaults to "", but the user should generally provide a value.

**theVariantGroupId** Group name used for labelling variant features when combining matrixes. This defaults to "", but the user should generally provide a value.

**theMatchedReplicatesFlag** If TRUE, indicates that NAs should be added for missing replicates. Defaults to TRUE.

**theCombineOnlyFlag** If TRUE, only combined the matrixes, do not correct. Defaults to FALSE.

**thePath** Location for output. Defaults to NULL. If NULL, no output file is created.

**theWriteToFile** TRUE means write corrected data to thePath. Only works if thePath is given. Defaults to FALSE.

# 6 Example Call

The following code is taken from the tests/RBN\_Pseudoreplicates.R file. Data used is from the testing data as per the MBatch\_01\_InstallLinux.docx document.

```
library(MBatch)
# set the paths
invariantFile="/bea testing/MATRIX DATA/rbn-pseudo-iset.tsv"
variantFile="/bea testing/MATRIX DATA/rbn-pseudo-vset.tsv"
theOutputDir="/bea_testing/output/RBN_Pseudoreplicates"
theRandomSeed=314
resolveDuplicates <- function(theNames)
 # keep first instance of a name
 # number subsequent ones starting with .1
 make.unique(theNames)
readRPPAdataAsMatrix_WithTab <- function(theFile)</pre>
 # read RPPA data as a dataframe
 # column rppaDF[,1] contains row names that may contain duplicates
 rppaDF <- readAsGenericDataframe(theFile)</pre>
 # resolve duplicates in row names here
 myRownames <- rppaDF[,1]
 myRownames <- resolveDuplicates(myRownames)
 # convert to matrix
 myMatrix <- data.matrix(rppaDF[,-1])</pre>
 rownames(myMatrix) <- myRownames
 t(myMatrix)
# make sure the output dir exists and is empty
 unlink(theOutputDir, recursive=TRUE)
 dir.create(theOutputDir, showWarnings=FALSE, recursive=TRUE)
 message("Reading invariant file")
 invMatrix = readRPPAdataAsMatrix WithTab(invariantFile)
 message("Reading variant file")
 varMatrix = readRPPAdataAsMatrix WithTab(variantFile)
 invPseudo <- c("BN", "BO", "BP", "BQ", "BR", "BS") varPseudo <- c("AN", "AO", "AP", "AQ", "AR", "AS")
 filename <- RBN_Pseudoreplicates(theInvariantMatrix=invMatrix,
                     theVariantMatrix=varMatrix,
                     theInvariantReplicates = invPseudo,
                     theVariantReplicates = varPseudo,
                     theInvariantGroupId="Grp1",
                     theVariantGroupId="Grp2",
                     theMatchedReplicatesFlag=FALSE,
                     theCombineOnlyFlag=FALSE,
                     thePath=theOutputDir,
                     theWriteToFile=TRUE)
```

# 6.1 Command Line Output

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.

```
> library(MBatch)
> # set the paths
> invariantFile="/bea_testing/MATRIX_DATA/rbn-pseudo-iset.tsv"
> variantFile="/bea_testing/MATRIX_DATA/rbn-pseudo-vset.tsv"
> theOutputDir="/bea_testing/output/RBN_Pseudoreplicates"
> theRandomSeed=314
> resolveDuplicates <- function(theNames)
+ {
    # keep first instance of a name
    # number subsequent ones starting with .1
    make.unique(theNames)
+ }
>
> readRPPAdataAsMatrix WithTab <- function(theFile)
   # read RPPA data as a dataframe
   # column rppaDF[,1] contains row names that may contain duplicates
   rppaDF <- readAsGenericDataframe(theFile)</pre>
   # resolve duplicates in row names here
   myRownames <- rppaDF[,1]
   myRownames <- resolveDuplicates(myRownames)</pre>
   # convert to matrix
    myMatrix <- data.matrix(rppaDF[,-1])</pre>
    rownames(myMatrix) <- myRownames
    t(myMatrix)
+ }
> # make sure the output dir exists and is empty
> unlink(theOutputDir, recursive=TRUE)
> dir.create(theOutputDir, showWarnings=FALSE, recursive=TRUE)
> message("Reading invariant file")
Reading invariant file
> invMatrix = readRPPAdataAsMatrix WithTab(invariantFile)
> message("Reading variant file")
Reading variant file
> varMatrix = readRPPAdataAsMatrix_WithTab(variantFile)
> invPseudo <- c("BN", "BO", "BP", "BO", "BR", "BS")
> varPseudo <- c("AN", "AO", "AP", "AQ", "AR", "AS")
> filename <- RBN_Pseudoreplicates(theInvariantMatrix=invMatrix,
                     theVariantMatrix=varMatrix.
                     theInvariantReplicates = invPseudo,
```

```
theVariantReplicates = varPseudo,
theInvariantGroupId="Grp1",
theVariantGroupId="Grp2",
theMatchedReplicatesFlag=FALSE,
theCombineOnlyFlag=FALSE,
thePath=theOutputDir,
theWriteToFile=TRUE)
```

2017 10 18 12:51:16.693 DEBUG megazone23 please note: internally, RBN processes transposed the data, output (file and matrix) match the submitted data with samples across columns and features down the rows

2017 10 18 12:51:16.694 INFO megazone23 RBN\_internal - starting

2017 10 18 12:51:16.694 DEBUG megazone23 checkCreateDir:

/bea\_testing/output/RBN\_Pseudoreplicates

2017 10 18 12:51:16.742 WARN megazone23 Less than 30 replicates provided for Invariant Matrix.

RBN's performance may deteriorate.

2017 10 18 12:51:16.743 WARN megazone23 Less than 30 replicates provided for Variant Matrix.

RBN's performance may deteriorate.

2017 10 18 12:51:16.743 INFO megazone23 Found 213 common features in both matrices.

2017 10 18 12:51:16.763 DEBUG megazone23 Write to file

 $/bea\_testing/output/RBN\_Pseudoreplicates/ANY\_Corrections-RBN\_Pseudoreps.tsv$ 

2017 10 18 12:51:16.772 DEBUG megazone23 Finished write to file

/bea\_testing/output/RBN\_Pseudoreplicates/ANY\_Corrections-RBN\_Pseudoreps.tsv

2017 10 18 12:51:16.773 INFO megazone23 RBN\_internal - completed

# 6.2 Example File Output

The above code creates the following output files. Files are named using the following naming convention:

ANY Corrections-RBN Pseudoreps.tsv

The TSV file with the combined/corrected dataset is written by the MBatch package.