

Using MBatch Corrections: AN_Adjusted

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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

AN Adjusted performs an ANOVA Adjusted correction taking a BEA_DATA object (with data matrix and batch dataframe) and returning either a corrected matrix or a string containing the path to where the data file was written.

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

```
AN_Adjusted(theBeaData, theBatchType, thePath = NULL, theWriteToFile = FALSE)
```

5 Arguments

5.1 theBeaData

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.

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 theBatchType

A string identifying the batch type to correct.

5.3 thePath

Output path for any files.

5.4 theWriteToFile

TRUE to write the corrected data to file and return the file pathname instead of the corrected matrix.

6 Example Call

The following code is adapted from the tests/AN_Adjusted.R 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.

```
{
  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

  theGeneFile="/bea_testing/MATRIX_DATA/matrix_data-Tumor.tsv"
  theBatchFile="/bea_testing/MATRIX_DATA/batches-Tumor.tsv"
  theOutputDir="/bea_testing/output/AN_Adjusted"
  theRandomSeed=314
  theBatchType="TSS"

  # make sure the output dir exists and is empty
  unlink(theOutputDir, recursive=TRUE)
  dir.create(theOutputDir, showWarnings=FALSE, recursive=TRUE)
  # load data
  myData <- mbatchLoadFiles(theGeneFile, theBatchFile)
  myData@mData <- mbatchTrimData(myData@mData, 100000)
  # call
  outputFile <- AN_Adjusted(theBeaData=myData,
                           theBatchType=theBatchType,
                           thePath=theOutputDir,
                           theWriteToFile=TRUE)
  correctedMatrix <- readAsGenericMatrix(outputFile)
  print(correctedMatrix[1:4, 1:4])
}
```

```
## 2018 06 21 10:35:21.037 DEBUG megazone23 Changing LC_COLLATE to C for duration of run
## 2018 06 21 10:35:21.038 INFO megazone23 \ / \ / \ / \ / \ / \ / \ / \ / \ /
## 2018 06 21 10:35:21.038 INFO megazone23 Starting mbatchLoadFiles
## 2018 06 21 10:35:21.039 INFO megazone23 MBatch Version: 2017-09-19-1530
## 2018 06 21 10:35:21.040 INFO megazone23 read batch file= /bea_testing/MATRIX_DATA/batches-Tumor.tsv
## 2018 06 21 10:35:21.041 INFO megazone23 read gene file= /bea_testing/MATRIX_DATA/matrix_data-Tumor.t
## 2018 06 21 10:35:29.957 INFO megazone23 filter samples in batches using gene samples
## 2018 06 21 10:35:29.958 INFO megazone23 sort batches by gene file samples
```

```

## 2018 06 21 10:35:30.041 INFO megazone23 Finishing mbatchLoadFiles
## 2018 06 21 10:35:30.041 INFO megazone23 ~~~~~
## 2018 06 21 10:35:30.042 DEBUG megazone23 Changing LC_COLLATE to C for duration of run
## 2018 06 21 10:35:30.042 INFO megazone23 \ / \ / \ / \ / \ / \ / \ / \ / \ /
## 2018 06 21 10:35:30.042 INFO megazone23 mbatchTrimData Starting
## 2018 06 21 10:35:30.043 INFO megazone23 MBatch Version: 2017-09-19-1530
## 2018 06 21 10:35:38.397 INFO megazone23 mbatchTrimData Finishing
## 2018 06 21 10:35:38.398 INFO megazone23 ~~~~~
## 2018 06 21 10:35:38.398 INFO megazone23 AN_Internal - starting
## 2018 06 21 10:35:38.398 DEBUG megazone23 checkCreateDir: /bea_testing/output/AN_Adjusted
## 2018 06 21 10:35:38.689 DEBUG megazone23 starting BeaAN
## 2018 06 21 10:35:38.690 DEBUG megazone23 AN names
## 2018 06 21 10:35:38.690 DEBUG megazone23 convertDataFrameToSi start
## 2018 06 21 10:35:38.690 DEBUG megazone23 convertDataFrameToSi asmatrixWithIssues
## 2018 06 21 10:35:38.692 DEBUG megazone23 convertDataFrameToSi rownames
## 2018 06 21 10:35:38.693 DEBUG megazone23 convertDataFrameToSi colnames
## 2018 06 21 10:35:38.693 DEBUG megazone23 convertDataFrameToSi done
## 2018 06 21 10:35:38.694 DEBUG megazone23 AN all
## 2018 06 21 10:35:38.694 DEBUG megazone23 AN cbin
## 2018 06 21 10:35:38.694 DEBUG megazone23 AN function
## 2018 06 21 10:35:38.694 DEBUG megazone23 AN check number of batch
## 2018 06 21 10:35:38.695 DEBUG megazone23 AN Check for missing values
## 2018 06 21 10:35:38.695 DEBUG megazone23 AN Check for genes with whole batch missing or no variation
## 2018 06 21 10:35:38.853 DEBUG megazone23 AN design
## 2018 06 21 10:35:38.854 DEBUG megazone23 AN build.X
## 2018 06 21 10:35:38.855 DEBUG megazone23 AN NAs
## 2018 06 21 10:35:38.855 INFO megazone23 NAs & var.adj
## 2018 06 21 10:35:38.856 INFO megazone23 is.matrix dat TRUE
## 2018 06 21 10:35:40.281 INFO megazone23 transpose
## 2018 06 21 10:35:40.281 INFO megazone23 is.matrix ANdat TRUE
## 2018 06 21 10:35:40.282 INFO megazone23 check nulls
## 2018 06 21 10:35:40.498 INFO megazone23 sum nulls
## 2018 06 21 10:35:40.499 DEBUG megazone23 finishing BeaAN
## 2018 06 21 10:35:40.499 TIMING megazone23      1.77999999999975      1.809999999999767      ANAdjusted
## 2018 06 21 10:35:40.499 DEBUG megazone23 Write to file /bea_testing/output/AN_Adjusted/ANY_Correcti
## 2018 06 21 10:35:40.611 DEBUG megazone23 Finished write to file /bea_testing/output/AN_Adjusted/ANY
## 2018 06 21 10:35:40.611 INFO megazone23 AN_Internal - completed
##
##          TCGA-OR-A5J1-01A-11D-A29J-05
## ABR-cg06968724-17-1012579      0.03374963
## ABR-cg23568341-17-1011974      0.11227667
## ABR-cg24479027-17-1012576      0.03534335
## ACOT7-cg16034168-1-6336711     1.04323610
##
##          TCGA-OR-A5J2-01A-11D-A29J-05
## ABR-cg06968724-17-1012579      0.03561384
## ABR-cg23568341-17-1011974      0.11935979
## ABR-cg24479027-17-1012576      0.03927621
## ACOT7-cg16034168-1-6336711     0.19927875
##
##          TCGA-OR-A5J3-01A-11D-A29J-05
## ABR-cg06968724-17-1012579      0.8873027
## ABR-cg23568341-17-1011974      0.9059181
## ABR-cg24479027-17-1012576      0.9004336
## ACOT7-cg16034168-1-6336711     0.2014968
##
##          TCGA-OR-A5J4-01A-11D-A29J-05
## ABR-cg06968724-17-1012579      0.9119515

```

## ABR-cg23568341-17-1011974	0.9844174
## ABR-cg24479027-17-1012576	0.9078395
## ACOT7-cg16034168-1-6336711	1.0079096

7 Example File Output

The above code creates the following output file. File is named using the following naming convention: ANY_Corrections-ANAdjusted.tsv The TSV file with the corrected dataset is written by the MBatch package. The end of the output shows a snippet from the corrected matrix.