Visualization and evaluation of RNA-seq data normalization – the Normalization Visualization Tool

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Abstract

Differential expression analysis between two samples, is a common task in the analysis of RNA-Seq data. In order to identify differential expressed genes, it is crucial that the two compared data sets are normalized. For the task of normalizing gene expression data there are multiple methods available. However all of these methods are based on certain assumptions, these can or can not be suitable for the data which should be normalized. For planning an adequate normalization strategy one need to answer, how the normalization affects the data sets and which normalization method and its underlying assumptions apply to the analyzed expression data. The *NVT* package provides a fast and simple way to analyze multiple normalization methods via visualization and correlation values. This vignette explains the use of the package and demonstrates a typical work flow.

NVT version: 0.9

If you use NVT in published research, please cite:

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Contents

1	Inst	allation
2	Star	ndard work flow
	2.1	Load data
		2.1.1 Load expression data
		2.1.2 Load gene length data
	2.2	Generate NVTdata objects
		2.2.1 Normalize the NVTdata
	2.3	Visualize expression data
		2.3.1 Simple plot
		2.3.2 Advanced plot
	2.4	Correlation values
		2.4.1 Pearson correlation
		2.4.2 RMSD and MEA correlation
	2.5	Test all methods
3	Sess	sion Info

1 Installation

The latest NVT package can be downloaded from: https://github.com/Edert/NVT/releases. The installation of the package can be done directly in R.

```
install.packages(file.path("/home/user/Downloads/","NVT_1.0.tar.gz"),
repos=NULL, type="source")
```

2 Standard work flow

Initially the library has to be loaded, which provides access to the data examples 2.1.1 and the NVT functions.

```
library("NVT")
```

2.1 Load data

For the demonstration of the NVT functions we need gene expression data, so we load the example data provided with the package. We use the two human expression data sets GSM1275862 and GSM1275863 from the airway[1] package.

2.1.1 Load expression data

We simply load the provided data sets *myexp1* from *GSM1275862*, *myexp2* from *GSM1275863* and the length data per gene as *mylen*.

```
data(mylen)
data(myexp1)
data(myexp2)
#show just the first four elements of the loaded data
head(mylen,4)
##
                   Length
## ENSG00000000003
                     3688
## ENSG00000000005
                     1881
## ENSG00000000419
                     4032
## ENSG0000000457 10784
head(myexp1,4)
##
                   GSM1275862
## ENSG0000000003
                           679
## ENSG00000000005
                             0
## ENSG0000000419
                           467
## ENSG0000000457
                           260
```

In order to evaluate the expression between the two RNA-seq expression samples, we have to define a list of human housekeeping genes *GAPDH*, *ALB*, *ACTB*, *QARS*, *PGK2*, *HPRT1*, *ADA* and *POLR2L*.

2.1.2 Load gene length data

Instead of using the length data generated directly in R or using a simple flat file, it is also possible to load the gene or exon length data directly from an annotation file in gtf or gff format. This function utilizes the GenomicRanges[2] and rtracklayer[3] packages.

- gff-version: version of the provided gff file [gff1, gff2, gff3, gtf]
- gff-feature: feature to use [default: exon]
- gff-name: name to use $[default:gene_id]$

```
#this line gets the path to the gff file provided in the NVT package
#(annotation from Chlamydia trachomatis, ACCESSION: NC_000117)
mygffpath <- system.file("extdata", "Ctr-D-UW3CX.gff", package = "NVT")</pre>
#this function loads the gff file from the gffpath
mylen1 <- NVTloadgff(mygffpath, "gff3", "gene", "locus_tag")</pre>
head(mylen1)
##
         Length
## CT001
            273
## CT002
            303
## CT003
           1476
## CT004
           1467
## CT005
           1092
## CT006
          570
```

2.2 Generate NVTdata objects

In the first step you need to generate an *NVTdata* object with the *NVTinit* function. Here you have to provide the list of housekeeping genes, the two gene expression samples and the normalization method. Optionally you can also add the gene or exon length data.

The normalization methods are:

- N = No normalization
- TC = Total count normalization
- Med = Median normalization
- TMM = Trimmed Mean of M-values normalization,
- UQ = Upper Quartile normalization
- UQ2 = Upper Quartile normalization (from NOISeq[4])
- Q = Quantile normalization implemented in limma[5]

- RPKM = Reads Per Kilobase per Million mapped reads normalization
- RPM = Reads Per Million mapped reads normalization
- TPM = Transcripts Per Million normalization
- DEQ = Relative log expression method included in the DESeq[6] package
- G = Use the provided genes to normalize

For the most methods no length information is required.

```
mynvt <- NVTinit(mylist1,myexp1,myexp2,"TMM")</pre>
```

But for RPKM, RPM and TPM normalization the length data has to be provided.

```
mynvt <- NVTinit(mylist1,myexp1,myexp2,"RPKM",mylen)</pre>
```

2.2.1 Normalize the NVTdata

The now initialized NVTdata object can be normalized in the next step.

```
mynorm <- NVTnormalize(mynvt)
## [1] "RPKM normalization!"</pre>
```

If required the now normalized data can be retrieved easily.

2.3 Visualize expression data

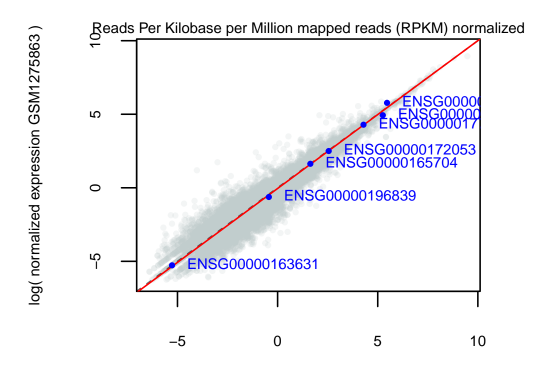
One of the key features of NVT is the plotting of the XY-Scatter-Plots. This can be done with two functions: NVT plot and NVT advanced plot.

2.3.1 Simple plot

The normalized *NVTdata* object can be visualized with the plot function. With the second parameter you can modify the size-ratio of the text and the data points. All expression values are shown as grey dots and the linear model as red line, it is calculated with the data from the housekeeping genes only. The dashed grey line highlights the diagonal, so the perfect correlation between the two data sets.

NVTplot(mynorm,0.6)

Scatter plot of: GSM1275862 vs. GSM1275863



log(normalized expression GSM1275862)

Figure 1: **XY-Scatter-Plot.** The resulting plot from the simple NVTplot function

2.3.2 Advanced plot

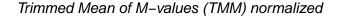
The *NVTadvancedplot* plots via ggplot2[7] and the size parameters control data points, text and labels separately. Here we use the default values of 1 for the data points and the text and increase the labels of the housekeeping genes to 2. Again the grey dots represent the expression data and their density is visualized by the rug in dark red for x- and y-axis. The linear model of the housekeeping gene data is shown in red and the diagonal is highlighted as grey dashed line. As we want to compare different methods we now normalize with the TMM method.

```
mynvt <- NVTinit(mylist1,myexp1,myexp2,"TMM",mylen)
mynorm <- NVTnormalize(mynvt)

## [1] "Trimmed Mean of M-values normalization!"

NVTadvancedplot(mynorm,1,1,2)</pre>
```

Scatter plot of: GSM1275862 vs GSM1275863



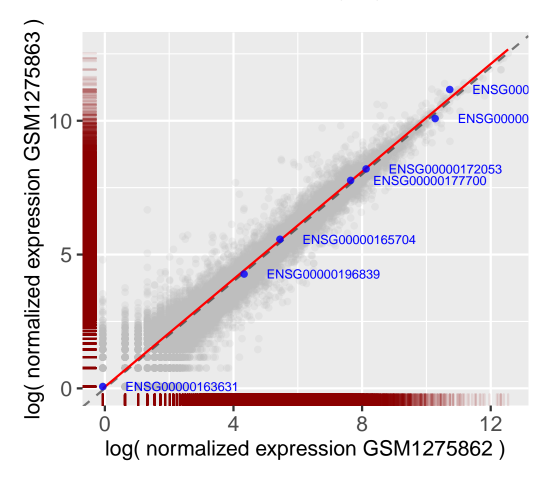


Figure 2: **Advanced XY-Scatter-Plot.** The NVTadvancedplot function results in a plot produced by ggplot2 with a rug on both axis, indicating the density of the data-points

2.4 Correlation values

In addition to the graphical representation of the gene expression data, the correlation coefficients of the housekeeping genes of the two samples can be calculated with the functions *NVTpearson*, *NVTrmsd* and *NVTmae*.

2.4.1 Pearson correlation

The Pearson correlation coefficient of the normalized expression of the housekeeping genes is calculated with the following command, using an already normalized *NVTdata* object.

```
NVTpearson(mynorm)
## pearson p-value
## 0.9636246988 0.0001170673
```

2.4.2 RMSD and MEA correlation

The root mean square error (RMSE) also called the root mean square deviation (RMSD) is calculated with the *NVTrmsd* function.

```
NVTrmsd(mynorm)

## RMSD

## 9097.041
```

And the mean absolute error (MAE) can be calculated and retrieved with the NVTmae function.

```
NVTmae(mynorm)
## MAE
## 3855.185
```

2.5 Test all methods

To test more normalization methods on the provided data sets in one single step, the correlation coefficients of all implemented normalization methods can be calculated with the *NVTtestall* function. It requires a normalized *NVTdata* object and the correlation coefficient you are interested in. The method can be defined with:

- p = Pearson correlation
- rmsd = root mean square deviation
- mae = mean absolute error

```
NVTtestall(mynorm, "p")
## [1] "No normalization!"
## [1] "Total count normalization!"
## [1] "Median normalization!"
## [1] "Trimmed Mean of M-values normalization!"
```

```
## [1] "Upper Quartile normalization!"
## [1] "Upper Quartile normalization (from NOISeq)!"
## [1] "Quantile normalization!"
## [1] "RPKM normalization!"
## [1] "RPM normalization!"
## [1] "DESeq normalization!"
## [1] "Using DESeq"
## [1] "Input counts are not integer, converting them!"
## [1] "TPM normalization!"
## [1] "Normalization by given gene-set!"
## [1] "ENSG00000111640" "ENSG00000163631" "ENSG00000075624" "ENSG00000172053"
## [5] "ENSG00000170950" "ENSG00000165704" "ENSG000000196839" "ENSG00000177700"
##
         pearson
                     p-value
       0.9716940 5.550234e-05
## Q
## DEQ 0.9636247 1.170673e-04
## TMM 0.9636247 1.170673e-04
       0.9636247 1.170673e-04
## G
       0.9636247 1.170673e-04
## TC
       0.9636247 1.170673e-04
## UQ
## N
       0.9636247 1.170673e-04
## Med 0.9636247 1.170673e-04
## UQ2 0.9636247 1.170673e-04
## RPM 0.9636247 1.170673e-04
## RPKM 0.9489857 3.193364e-04
## TPM 0.9489857 3.193364e-04
```

3 Session Info

- R version 3.2.2 (2015-08-14), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=de_AT.UTF-8, LC_COLLATE=en_US.UTF-8, LC_MONETARY=de_AT.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=de_AT.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=de_AT.UTF-8, LC_IDENTIFICATION=C
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- Other packages: knitr 1.12, NVT 0.9
- Loaded via a namespace (and not attached): annotate 1.48.0, AnnotationDbi 1.32.3, Biobase 2.30.0, BiocGenerics 0.16.1, BiocParallel 1.4.3, BiocStyle 1.8.0, Biostrings 2.38.3, bitops 1.0-6, colorspace 1.2-6, DBI 0.3.1, DESeq 1.22.0, evaluate 0.8, formatR 1.2.1, futile.logger 1.4.1, futile.options 1.0.0, genefilter 1.52.0, geneplotter 1.48.0, GenomeInfoDb 1.6.1, GenomicAlignments 1.6.3, GenomicRanges 1.22.3, ggplot2 2.0.0, grid 3.2.2, gtable 0.1.2, highr 0.5.1, IRanges 2.4.6, labeling 0.3, lambda.r 1.1.7, lattice 0.20-33, limma 3.26.5, magrittr 1.5, Matrix 1.2-3, munsell 0.4.2, NOISeq 2.14.0, parallel 3.2.2, plyr 1.8.3, RColorBrewer 1.1-2, Rcpp 0.12.3, RCurl 1.95-4.7, Rsamtools 1.22.0, RSQLite 1.0.0, rtracklayer 1.30.1, S4Vectors 0.8.7, scales 0.3.0, splines 3.2.2, stats4 3.2.2, stringi 1.0-1, stringr 1.0.0, SummarizedExperiment 1.0.2, survival 2.38-3, tools 3.2.2, XML 3.98-1.3, xtable 1.8-0, XVector 0.10.0, zlibbioc 1.16.0

References

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