# Anaquin - Vignette 

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November 01, 2022

## Citation

- [1] Representing genetic variation with synthetic DNA standards. Nature Methods, 2017
- [2] Spliced synthetic genes as internal controls in RNA sequencing experiments. Nature Methods, 2016.
- [3] Reference standards for next-generation sequencing. Nature Reviews, 2017.
- [4] Anaquin: a software toolkit for the analysis of spike-in controls for next generation sequencing. Bioinformatics, 2017.


## Website

Visit our website to learn more about sequins: www.sequin.xyz.

## Overview

In this document, we show how to conduct statistical analysis that models the performance of sequin controls in next-generation-sequencing (NGS) experiment. We call the sequins RnaQuin for "RNA-Seq sequins", MetaQuin for "metagenomic sequins", VarQuin for "genomics variant sequins", and the statistical framework Anaquin.

This vignette is written for R-usage. However, Anaquin is a framework covering the entire NGS workflow. Consequently, the R-package (and it's documentation) is a subset of the overall Anaquin framework. We also distribute a detailed workflow guide on our website.

It is important to note Anaquin is both command-line tool and R-package. Our workflow guide has the details on how the command-line tool can be used with the R-package.

## Sequins

Next-generation sequencing (NGS) enables rapid, cheap and high-throughput determination of sequences within a user's sample. NGS methods have been applied widely, and have fuelled major advances in the life sciences and clinical health care over the past decade. However, NGS typically generates a large amount of sequencing data that must be first analyzed and interpreted with bioinformatics tools. There is no standard way to perform an analysis of NGS data; different tools provide different advantages in different situations. The complexity and variation of sequences further compound this problem, and there is little reference by which compare next-generation sequencing and analysis.

To address this problem, we have developed a suite of synthetic nucleic-acid sequins (sequencing spike-ins). Sequins are fractionally added to the extracted nucleic-acid sample prior to library preparation, so they are sequenced along with your sample of interest. We can use the sequins as an internal quantitative and qualitative control to assess any stage of the next-generation sequencing workflow.


Figure 1: NGS Workflow for sequins

## Mixture

Sequins are combined together across a range of concentrations to formulate a mixture. Mixture file (CSV) is a text file that specifies the concentration of each sequin within a mixture. Mixture files are often required as input to enable Anaquin to perform quantitative analysis. Mixture file can be downloaded from our website.

Let's demonstrate RnaQuin mixture A with a simple example. Load the mixture file (you can also download the file directly from our website):

```
library('Anaquin')
## Loading required package: ggplot2
data("RnaQuinIsoformMixture")
head(RnaQuinIsoformMixture)
```

| \#\# | Name | Length | MixA | MixB |
| :--- | ---: | ---: | ---: | ---: |
| \#\# 1 | R1_101_1 | 719 | 11.329650 | 0.472075 |
| \#\# 2 | R1_101_2 | 430 | 3.776550 | 1.416225 |
| \#\# 3 | R1_102_1 | 1490 | 13.217925 | 7.553100 |
| \#\# 4 | R1_102_2 | 1362 | 1.888275 | 52.871700 |
| \#\# 5 R1_103_1 | 1754 | 60.424806 | 453.186000 |  |
| \#\# 6 | R1_103_2 | 1856 | 906.372094 | 30.212400 |

Each row represents a sequin. Name gives the sequin names, Length is the length of the sequins in nucleotide bases, MixA gives the concentration level in attoml/ul for Mixture A.

Imagine we have two RNA-Seq experiments; a well-designed experiment and a poorly-designed experiment. We would like to quantify their isoform expression.
Let's simulate the experiments:

```
set.seed(1234)
sim1 <- 1.0 + 1.2*log2(RnaQuinIsoformMixture$MixA) + rnorm(nrow(RnaQuinIsoformMixture),0,1)
sim2 <- c(1.0 + rnorm(100,1,3), 1.0 +
    1.2*log2(tail(RnaQuinIsoformMixture,64)$MixA) +
    rnorm(64,0,1))
```

In the first experiment, sequins are expected to correlate linearly with the measured FPKM. Indeed, the variables are strongly correlated:

```
names <- row.names(RnaQuinIsoformMixture)
input <- log2(RnaQuinIsoformMixture$MixA)
title <- 'Isoform expression (Good)'
xlab <- 'Input concentration (log2)'
```

```
ylab <- 'Measured FPKM (log2)'
plotLinear(names, input, sim1, title=title, xlab=xlab, ylab=ylab)
## Warning: `guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> =
## "none")` instead.
## Warning: Use of `data$x` is discouraged. Use ` x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
## Warning: Use of `data$x` is discouraged. Use ` x` instead.
## Warning: Use of `data$y` is discouraged. Use ` y` instead.
```

Isoform expression (Good)


In our second experiment, the weakly expressed isoforms exhibit stochastic behavior and are clearly not linear with the input concentration. Furthermore, there is a limit of quantification (LOQ); below which accuracy of the experiment becomes questionable.

```
names <- row.names(RnaQuinIsoformMixture)
input <- log2(RnaQuinIsoformMixture$MixA)
title <- 'Isoform expression (Bad)'
xlab <- 'Input concentration (log2)'
ylab <- 'Measured FPKM (log2)'
plotLinear(names, input, sim2, title=title, xlab=xlab, ylab=ylab)
## Warning: `guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> =
## "none")` instead.
```

```
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
```

Isoform expression (Bad)


The primary observation is that the artificial scale imposed by sequins allow us to quantify our experiments.

## Quantifying transcriptome assembly

To quantify RNA-Seq transcriptome assembly, we need to run a transcriptome assember; a software that can assemble transcripts and estimates their abundances. Our workflow guide has the details.

Here, we use a data set generated by Cufflinks, described in Section 5.4.5.1 in the user guide:

```
data(UserGuideData_5.4.5.1)
head(UserGuideData_5.4.5.1)
## Input Sn
## R1_101_1 10.0708 0.990264
## R1_101_2 5.0354 0.393023
## R1_102_1 0.8886 0.519463
## R1_102_2 14.2176 0.902349
## R1_103_1 107.4220 0.995439
## R1_103_2 859.3750 0.904095
```

The first column gives the input concentration for each sequin in attomol/ul. The second column is the measured sensitivity. Run the following R-code to generate a sensitivity plot.

```
title <- 'Assembly Plot'
xlab <- 'Input Concentration (log2)'
ylab <- 'Sensitivity'
# Sequin names
names <- row.names(UserGuideData_5.4.5.1)
# Input concentration
x <- log2(UserGuideData_5.4.5.1$Input)
# Measured sensitivity
y <- UserGuideData_5.4.5.1$Sn
plotLogistic(names, x, y, title=title, xlab=xlab, ylab=ylab, showLOA=TRUE)
## Warning: `guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> =
## "none")` instead.
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
```



The fitted logistic curve reveals clear relationship between input concentration and sensitivity. Unsurprisingly, the assembler has higher sensitivity with highly expressed isoforms. The limit-of-assembly (LOA) is defined as the intersection of the curve to sensitivity of 0.70 .

## Quantifying gene expression

Quantifying gene/isoform expression involves building a linear model between input concentration and measured FPKM. In this section, we consider a dataset generated by Cufflinks, described in Section 5.4.5.1 of the user guide.

Load the data set:

```
data(UserGuideData_5.4.6.3)
head(UserGuideData_5.4.6.3)
## Input Observed1 Observed2 Observed3
## R1_101 15.1062 0.958838 1.456650 0.960190
## R1_102 15.1062 0.806596 0.604539 0.652783
## R1_103 966.7970 2.650470 2.890570 3.211090
## R1_11 241.6990 3.876010 
```



The first column gives input concentration for each sequin in attomol/ul. The other columns are the FPKM values for each replicate (three replicates in total). The following code will quantify the first replicate:

```
title <- 'Gene Expression'
xlab <- 'Input Concentration (log2)'
ylab <- 'FPKM (log2)'
# Sequin names
names <- row.names(UserGuideData_5.4.6.3)
# Input concentration
x <- log2(UserGuideData_5.4.6.3$Input)
# Measured FPKM
y <- log2(UserGuideData_5.4.6.3$Observed1)
plotLinear(names, x, y, title=title, xlab=xlab, ylab=ylab, showLOQ=TRUE)
## Warning: `guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> =
## "none")` instead.
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
```


## Gene Expression



Coefficient of determination is over 0.90 ; over $90 \%$ of the variation (e.g. technical bias) can be explained by the model. LOQ is 3.78 attomol/ul, this is the estimated emphirical detection limit.

We can also quantify multiple replicates:

```
title <- 'Gene Expression'
xlab <- 'Input Concentration (log2)'
ylab <- 'FPKM (log2)'
# Sequin names
names <- row.names(UserGuideData_5.4.6.3)
# Input concentration
x <- log2(UserGuideData_5.4.6.3$Input)
# Measured FPKM
y <- log2(UserGuideData_5.4.6.3[,2:4])
plotLinear(names, x, y, title=title, xlab=xlab, ylab=ylab, showLOQ=TRUE)
## Warning: `guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> =
## "none")` instead.
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
```

\#\# Warning: Use of ‘data\$x` is discouraged. Use ‘x` instead.
\#\# Warning: Use of `data\$ymin` is discouraged. Use `ymin` instead.
\#\# Warning: Use of ‘data\$x` is discouraged. Use ‘x` instead.
\#\# Warning: Use of `data\$ymax` is discouraged. Use `ymax` instead.
Gene Expression


## Differential analysis

In this section, we show how to quantify differential expression analysis between expected fold-change and measured fold-change. We apply our method to a data set described in Section 5.6.3 of the user guide.

```
data(UserGuideData_5.6.3)
head(UserGuideData_5.6.3)
```

| \#\# | ExpLFC | ObsLFC | SD | Pval | Qval | Mean | Label |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: | ---: |
| \#\# R1_101 | -3 | -1.890122 | 0.701723 | $7.069675 \mathrm{e}-03$ | $2.056337 \mathrm{e}-02$ | 9.953556 | TP |
| \#\# R1_102 | -4 | -2.051777 | 0.546374 | $1.731616 \mathrm{e}-04$ | $7.646243 \mathrm{e}-04$ | 17.285262 | TP |
| \#\# R1_103 | -1 | 3.837784 | 0.377602 | $2.883289 \mathrm{e}-24$ | $6.534028 \mathrm{e}-23$ | 1221.301532 | TP |
| \#\# R1_11 | -4 | -2.431582 | 0.591352 | $3.924117 \mathrm{e}-05$ | $1.974336 \mathrm{e}-04$ | 47.174250 | TP |
| \#\# R1_12 | 1 | 1.542757 | 0.425562 | $2.887104 \mathrm{e}-04$ | $1.214989 \mathrm{e}-03$ | 73.008720 | TP |
| \#\# R1_13 | 0 | 0.717701 | 0.242493 | $3.079564 \mathrm{e}-03$ | $1.000416 \mathrm{e}-02$ | 44053.259914 | FP |

For each of the sequin gene, we have expected log-fold change, measured log-fold change, standard deviation, $p$-value, $q$-value and mean. The estimation was done by DESeq2.

Run the following code to construct a folding plot:

```
title <- 'Gene Fold Change'
xlab <- 'Expected fold change (log2)'
ylab <- 'Measured fold change (log2)'
# Sequin names
names <- row.names(UserGuideData_5.6.3)
# Expected log-fold
x <- UserGuideData_5.6.3$ExpLFC
# Measured log-fold
y <- UserGuideData_5.6.3$ObsLFC
plotLinear(names, x, y, title=title, xlab=xlab, ylab=ylab, showAxis=TRUE,
    showLOQ=FALSE)
## Warning: `guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> =
## "none")` instead.
## Warning: Use of `data$x` is discouraged. Use `x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
## Warning: Use of `data$x` is discouraged. Use ` x` instead.
## Warning: Use of `data$y` is discouraged. Use `y` instead.
```

Gene Fold Change


Outliers are obvious throughout the reference scale. Overall, DESeq2 is able to account for $78 \%$ of the variation.

We can also construct a ROC plot. [1] has details on how the true-positives and false-positives are defined.

```
title <- 'ROC Plot'
# Sequin names
seqs <- row.names(UserGuideData_5.6.3)
# Expected ratio
ratio <- UserGuideData_5.6.3$ExpLFC
# How the ROC points are ranked (scoring function)
score <- 1-UserGuideData_5.6.3$Pval
# Classified labels (TP/FP)
label <- UserGuideData_5.6.3$Label
plotROC(seqs, score, ratio, label, title=title, refGroup=0)
```

ROC Plot


AUC statistics for LFC 3 and 4 are higher than LFC 1 and 2. Overall, all LFC ratios can be correctly classified relative to LFC 0.
Furthermore, we can construct limit of detection ratio (LOD) curves:

```
xlab <- 'Average Counts'
ylab <- 'P-value'
title <- 'LOD Curves'
# Measured mean
mean <- UserGuideData_5.6.3$Mean
```

```
# Expected log-fold
ratio <- UserGuideData_5.6.3$ExpLFC
# P-value
pval <- UserGuideData_5.6.3$Pval
qval <- UserGuideData_5.6.3$Qval
```

plotLOD(mean, pval, abs(ratio), qval=qval, xlab=xlab, ylab=ylab, title=title, FDR=0.05)

LOD Curves


Unsurprisingly, p-value is inverse quadratically related with average counts. All the LFC ratios systematically outperform LFC 0 . The function also estimates the empirical detection limits, [1] has the details.

