Supplementary material

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Introduction

Here we validate performance of the pulseR package on the simulated data. We will consider a pulse-experiment with no spike-ins added. The data include the total, the pull-out and the flow-through fractions for four time points in two replicates.

In the pulse-experiment, the unlabelled RNA is being degraded, starting from the steady-state amount at time t=0hr. The labelled fraction is being synthesised with the rate s, and it degrades with the same rate d as well. A system of ordinary differential equations, which describes this processes, is

$$\frac{d[\text{labelled RNA}]}{dt} = s - d[\text{labelled RNA}]$$

$$\frac{d[\text{unlabelled RNA}]}{dt} = -d[\text{unlabelled RNA}]$$

$$[\text{total}] = [\text{labelled}] + [\text{unlabelled}]$$

For constant rates of degradation d and synthesis s, the amounts of RNA at the time t are

$$[\text{total RNA}] = \frac{s}{d} \equiv \mu$$

$$[\text{labelled RNA}] = \mu(1 - e^{-dt})$$

$$[\text{unlabelled RNA}] = \mu e^{-dt},$$

where μ is the expression level of a gene in a steady state.

Definitions

Formulas

We can simulate count data using internal pulseR function pulseR:::generateTestDataFrom. For this, one needs to define the formulas for the mean read number of all three fractions:

Fractions

In reality, there is no pure labelled or unlabelled fractions in samples. That is why it may be important to model fraction cross-contamination. Here we assume that the labelled fraction consists of certain amount of the unlabelled one, and vice versa, the unlabelled one is contaminated with the labelled molecules. Hence, we distinguish the labelled RNA and the "labelled" pull-out fraction.

```
# important: the labels must be the same as in formulas,
# or they must be integer list indexes
formulaIndexes <- list(
  total_sample = 'total',
  label_sample = c('labelled', 'unlabelled'),</pre>
```

```
unlabel_sample = c('unlabelled', 'labelled')
)
```

This definition allows us to calculate mean read number in the fractions as a linear combination of the formulas for the labelled and unlabelled RNA amounts.

Conditions

Here we specify the condition data.frame, which includes information on the type of sample and the time points.

```
# create nReplicates * nTime * 3 (labelled, unlabelled, total) data points
conditions <- data.frame(
   condition = rep(names(formulaIndexes), each = nTime),
   time = rep(1:nTime, length(formulaIndexes) * nReplicates))
rownames(conditions) <- paste0("sample_", seq_along(conditions$condition))
knitr::kable(conditions)</pre>
```

	condition	time
sample_1	total_sample	1
$sample_2$	$total_sample$	2
$sample_3$	$total_sample$	3
$sample_4$	$total_sample$	4
$sample_5$	$label_sample$	1
$sample_6$	$label_sample$	2
$sample_7$	$label_sample$	3
$sample_8$	$label_sample$	4
$sample_9$	$unlabel_sample$	1
$sample_10$	$unlabel_sample$	2
$sample_11$	$unlabel_sample$	3
$sample_12$	$unlabel_sample$	4
$sample_13$	$total_sample$	1
$sample_14$	$total_sample$	2
$sample_15$	$total_sample$	3
$sample_16$	$total_sample$	4
$sample_17$	$label_sample$	1
$sample_18$	$label_sample$	2
$sample_19$	$label_sample$	3
$sample_20$	$label_sample$	4
$sample_21$	$unlabel_sample$	1
$sample_22$	$unlabel_sample$	2
$sample_23$	$unlabel_sample$	3
$sample_24$	unlabel_sample	4

Normalisation factors

Since we are going to simulate an experiments without spike-ins, we need to define the relations between different samples via *normalisation factors*. We divide samples into groups depending on their time point and fraction. Except the total fraction, other samples are considered to belong to different group if the time points are different:

```
fractions <- as.character(interaction(conditions))
# assume that the total RNA amount does not change with the time
# so all total fractions can be treated together
fractions[grep("total", fractions)] <- "total_sample"
knitr::kable(cbind(rownames(conditions), fractions))</pre>
```

	fractions
$sample_1$	$total_sample$
$sample_2$	$total_sample$
$sample_3$	$total_sample$
$sample_4$	$total_sample$
$sample_5$	$label_sample.1$
$sample_6$	$label_sample.2$
$sample_7$	$label_sample.3$
$sample_8$	$label_sample.4$
$sample_9$	$unlabel_sample.1$
$sample_10$	unlabel_sample.2
$sample_11$	$unlabel_sample.3$
$sample_12$	unlabel_sample.4
$sample_13$	$total_sample$
$sample_14$	$total_sample$
$sample_15$	$total_sample$
$sample_16$	$total_sample$
$sample_17$	$label_sample.1$
$sample_18$	$label_sample.2$
$sample_19$	$label_sample.3$
$sample_20$	$label_sample.4$
$sample_21$	$unlabel_sample.1$
$sample_22$	$unlabel_sample.2$
$sample_23$	$unlabel_sample.3$
$sample_24$	unlabel_sample.4

Here we use the inner function pulseR:::addKnownToFormulas in order to get the correct form of the list with the normalisation factors.

```
known <- pulseR:::addKnownToFormulas(formulas, formulaIndexes, conditions)
normFactors <- known$formulaIndexes[unique(names(known$formulaIndexes))]</pre>
```

```
# the total fractions have the reference factor = 1
normFactors <- normFactors[-grep("total", names(normFactors))]</pre>
normFactors <- c(list(total sample = 1), normFactors)</pre>
# linear combinations with the coefficient 0.1 for contaminating fraction
# and the coefficient 3 for the main fraction of the sample
normFactors <- relist(c(1, rep(3, length(unlist(normFactors)) - 1)),</pre>
                      normFactors)
normFactors[-1] <- lapply(normFactors[-1], '[[<-',2,.10)
str(normFactors)
## List of 9
## $ total sample : num 1
## $ label sample.1 : num [1:2] 3 0.1
## $ label_sample.2 : num [1:2] 3 0.1
## $ label sample.3 : num [1:2] 3 0.1
## $ label sample.4 : num [1:2] 3 0.1
## $ unlabel_sample.1: num [1:2] 3 0.1
## $ unlabel sample.2: num [1:2] 3 0.1
## $ unlabel sample.3: num [1:2] 3 0.1
## $ unlabel sample.4: num [1:2] 3 0.1
```

Simulation

We sample the mean expression level and the degradation rates from finite intervals:

```
# set size factor for the rnbinom function
par <- list(size = 1e2)
# set mean read number for every gene
# by sampling from the log-uniform distribution
par$mu <- exp(runif(nGenes, 0, log(1e5)))
# set the degradation rates from log-uniform distribution
par$d <- exp(runif(nGenes,log(0.01), log(2)))</pre>
```

And, finally, we simulate the data from the negative binomial distribution using the generateTestDataFrom function:

```
# the generateTestDataFrom needs for an input normFactors
# provided for every individual sample (i.e. we need to utilise
# the group information ourselves)
allNormFactors <- pulseR:::multiplyList(normFactors, fractions)

counts <- generateTestDataFrom(
   formulas, formulaIndexes, allNormFactors, par, conditions)</pre>
```

Fitting

This section describes the usual workflow for analysis of the count data coming from the experiment without spike-ins defined above, i.e. this is an entry point of the package usage on the real data.

Create the PulseData object

We define a PulseData object on the basis of the generated counts, the condition matrix, relations between formulas and samples. We also provide the information on how to split the samples for normalisation.

```
pd <- PulseData(
  counts = counts,
  conditions = conditions,
  formulas = formulas,
  formulaIndexes = formulaIndexes,
  groups = fractions
)</pre>
```

Set fitting options

In order to use the fitting function, one need to provide the boundaries of the parameters. In addition, we set here the relative tolerance thresholds.

```
# lower and upper boundaries for the normalisation factors.
# The total one will be ignored, because the first coefficient
# is always the reference one with the value 1.
# The reason is that we can identify the means of read numbers only
# up to an unknown multiplier.
lbNormFactors <- list(</pre>
 total_sample
 label sample = c(.1,.010),
 unlabel sample = c(.1,.010))
ubNormFactors <- list(</pre>
 total sample
 label sample
              = c(10, 2),
 unlabel sample = c(10, 2)
# set lower and upper boundaries for the rest of the parameters
opts <- setBoundaries(</pre>
 list(mu = c(1, 1e6), d = range(par$d) * c(1 / 5, 5)),
 normFactors = list(lbNormFactors, ubNormFactors))
# set the tolerance for the gene-specific parameters and
```

```
# the normalisation factors
opts <- setTolerance(params = .01, normFactors = .001, options = opts)</pre>
```

Fit

Before the fitting, one need to initialise the first guess for the parameters. The outcome of the fitting procedure may depend on this initial values a lot.

```
# initialise mu and d as gene-specific parameters, i.e. they must
# be sampled for every single gene individually
initPars <- initParameters(par = NULL, geneParams = c("mu", "d"), pd, opts)</pre>
str(initPars)
## List of 4
##
   $ mu
                 : num [1:530] 224008 871258 876756 302339 915588 ...
                 : num [1:530] 8.6 3.77 2.83 4.78 9.04 ...
##
    $ d
                 : num 7.31e+09
##
   $ size
   $ normFactors:List of 9
##
     ..$ : num 1
##
     ..$ : num [1:2] 1 0.01
##
##
     ..$ : num [1:2] 1 0.01
##
     ..$: num [1:2] 1 0.01
     ..$: num [1:2] 1 0.01
##
     ..$ : num [1:2] 1 0.01
##
     ..$ : num [1:2] 1 0.01
##
     ..$: num [1:2] 1 0.01
##
     ..$ : num [1:2] 1 0.01
Now we are ready to start the fitting:
```

```
res <- fitModel(pd, initPars, opts)</pre>
```

Results

Normalisation factors

Since the normalisation factors are fitted together with the rest of the parameters, they affect the identifiability of the last. However, the normalisation of the main fraction in the samples is recovered well (i.e. the coefficient 3).

```
str(res$normFactors)
## List of 9
## $ : num 1
## $ : num [1:2] 2.997 0.079
```

```
## $ : num [1:2] 3.0179 0.0581

## $ : num [1:2] 3.0185 0.0399

## $ : num [1:2] 3.05 0.01

## $ : num [1:2] 3.0621 0.0827

## $ : num [1:2] 3.0837 0.0885

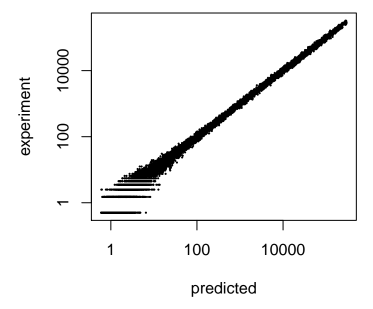
## $ : num [1:2] 3.0865 0.0972

## $ : num [1:2] 3.0995 0.0964
```

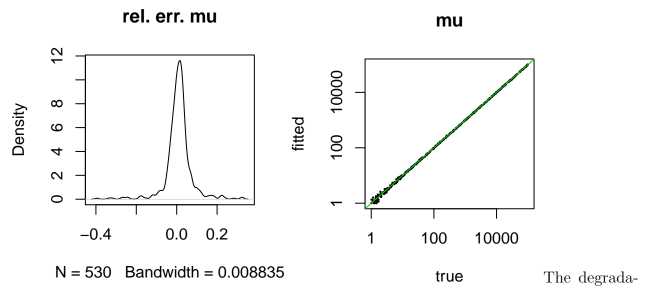
Gene-specific parameters

The estimated expected read mean numbers are well consistent with the raw simulated data:

Counts vs. means

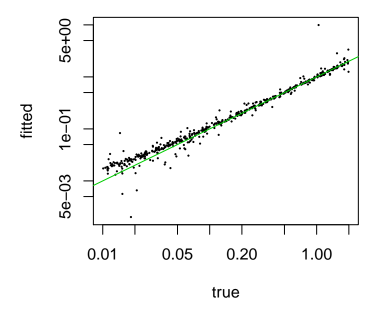


Due to high number of total fraction samples, the parameter of the mean μ is also well recovered from the data.



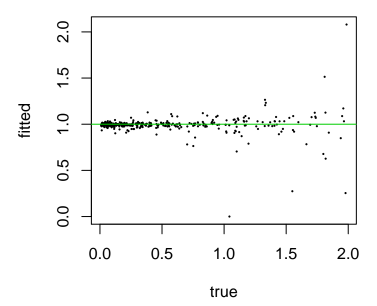
tion rate estimation is based on the less number of the samples (total provide the information only on the expression level). Hence, the quality of the fit is worse, than for the μ parameter. We see that the error is higher for the less degrading genes:

d



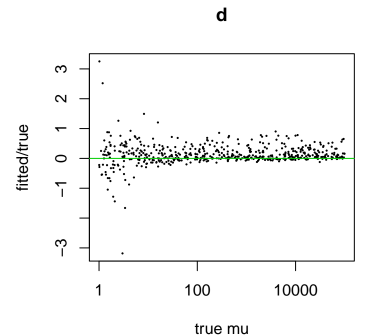
However, the misfit in d for lowly degrading genes has a minor effect on the error in predictions of the read number in comparison to the misfit in the higher ones, because it has an exponential impact:

$\exp(d_{true} - d_{fitted})$



Hence, the genes with the higher turn-over get a higher weight during optimisation of the degradation rate in the frame of the generated data.

As it was mentioned before, the normalisation of the samples has an impact on the genespecific parameters, and in our experiment the degradation rates are over-estimated for the majority of the genes. The estimation of d for the less expressed genes are less precise, because the have higher noise/signal ratio:



summary(res\$d/par\$d)

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.1100 0.9939 1.0620 1.1440 1.2080 9.5230
```

Conclusion

Here we presented a validation of the pulseR package performance on the simulated data. As a result, we show that the values of the gene-specific parameters can be recovered by the maximum likelihood estimation used by the package. The quality of the estimations directly depends on the data, i.e. on the amount of information and noise in the read numbers.

The pulseR package enables to perform estimations even if there are no spike-ins used in the experiment. However, this approach is limited to the systems, which has structurally and practically identifiable parameters. That depends on the number of replicates, the type of the fractions provided and the cross-contamination rate.