EasyqpcR: low-throughput real-time quantitative PCR data analysis

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1 Introduction

The package "EasyqpcR" has been created to facilitate the analysis of real-time quantitative RT-PCR data. This package contains five functions (badCt, nrmData, calData, totData, slope). In this manuscript, we describe and demonstrate how we can use this package. The last section presents how we can use the free R GUI RStudio and the gWidgets package created by John Verzani in order to facilitate the qPCR data analysis by a graphical user interface.

2 Amplification efficiency calculation

In this section, we describe how we can use the **slope** function of this package. As an example, we have 2 genes (gene 1 and gene 2) and 5 samples in triplicates (control group 1 and control group 2, treatment group 1 and treatment group 2, calibrator). We want to calculate the amplification efficiency of these two genes:

You can put the returned values into a vector to use it (without the need to type every amplification efficiency) in the next functions.

```
efficiency <- slope(data = Efficiency_calculation, q = c(1000, 100, 10, 1, 0.1), r = 3, r = TRUE
```

3 Calculation of the expression values from one or multiple qPCR run(s)

We describe the calculation of the normalization factors, the relative quantities, the normalized relative quantities and the normalized relative quantities scaled to the control group of your choice using the method of Hellemans et al (2007) [2]. We have a set of three qPCR runs, each run representing an independent biological replicate. The raw data of these runs can be found in the data folder of this package. The no template control and no amplification control have been discarded in order to facilitate the understanding of the workflow. The limiting step is that you need to put the control samples on the top of the data frame otherwise, the algorithm will not work correctly. Firstly, we load the datasets:

```
data(qPCR_run1, qPCR_run2, qPCR_run3)
str(c(qPCR_run1, qPCR_run2, qPCR_run3))
List of 15
$ Samples: Factor w/ 5 levels "Calibrator", "Control 1",..: 2 2 2 3 3 3 4 4 4 5 ...
          : num [1:15] 19.6 19.3 19.5 19.6 19.4 ...
 $ RG2
          : num [1:15] 18.5 18.4 18.5 18.6 18.7 ...
 $ TG
          : num [1:15] 26.3 26.1 26.2 26.1 26.1 ...
          : num [1:15] 16.3 16.5 16.7 16.8 17 ...
 $ TGb
$ Samples: Factor w/ 5 levels "Calibrator", "Control 1",..: 2 2 2 3 3 3 4 4 4 5 ...
          : num [1:15] 20.7 20.7 20.5 20.8 21 ...
 $ RG1
 $ RG2
          : num [1:15] 19.7 19.6 19.5 20.2 20.2 ...
 $ TG
          : num [1:15] 27.5 27.3 27.3 27.5 27.7 ...
          : num [1:15] 17.5 17.7 17.7 18.2 18.6 ...
 $ TGb
$ Samples: Factor w/ 5 levels "Calibrator", "Control 1",...: 2 2 2 3 3 3 4 4 4 5 ...
 $ RG1
          : num [1:15] 20.7 20.7 20.5 20.5 20.6 ...
 $ RG2
          : num [1:15] 19.7 19.6 19.5 19.8 19.8 ...
 $ TG
          : num [1:15] 27.5 27.3 27.3 27.2 27.3 ...
          : num [1:15] 17.5 17.7 17.7 17.8 18.1 ...
 $ TGb
```

Each dataset contains 15 observations: 5 samples (2 control groups, 2 treatment groups, 1 calibrator) in triplicates. There are 4 genes: 2 reference genes (RG1 and RG2) and 2 target genes (TG and Tgb). In order to facilitate the understanding of the nrmData function, I suggest you to read its man page by typing ?nrmData in your R session.

Concerning the reference genes, I suggest you to use the **selectHKgenes** function of the SLqPCR package from Matthias Kohl [1].

In order to avoid the inter-run variations, we have used a calibrator (one is the minimum recommended, more is better). Thus, we have to calculate the calibration factor for each

gene. We have to include the normalized relative quantities of our calibrator in an object:

```
## Isolate the calibrator NRQ values of the first biological replicate

aa <- nrmData(data = qPCR_run1, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02, 0.02, 0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = c(1, 1, 1, 1), CalPos = 5, trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]]

## Isolate the calibrator NRQ values of the first biological replicate

bb <- nrmData(data = qPCR_run2, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02, 0.02, 0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = c(1, 1, 1, 1), CalPos = 5, trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]]

## Isolate the calibrator NRQ values of the first biological replicate

cc <- nrmData(data = qPCR_run3, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02, 0.02, 0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = c(1, 1, 1, 1), CalPos = 5, trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]]</pre>
```

Now, we have to run the calData function.

4 Calculation of the calibration factors

Here, we describe how to use the calData function. In the continuation of what has been done before, we have three objects containing the NRQ values of the calibrator(s) and we now have to calculate the calibration factors for each gene:

```
## Calibration factor calculation
e <- calData(aa)
f <- calData(bb)
g <- calData(cc)</pre>
```

5 Attenuation of inter-run variations

Now, we have the calibration factors, we can calculate the expression value without the obsession of the inter-run variability:

```
nrmData(data = qPCR_run1, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02, 0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = e, CalPos = 5, trace = FALSE, geo = TRUE, na.rm = TRUE)
nrmData(data = qPCR_run2, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02, 0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = f, CalPos = 5, trace = FALSE, geo = TRUE, na.rm = TRUE)

nrmData(data = qPCR_run3, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02, 0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = g, CalPos = 5, trace = FALSE, geo = TRUE, na.rm = TRUE)
```

Remark: The validity of IRCs must be interpreted with care: two or more IRCs must be used to control if the IRCs measure the technical variation between the runs with the same extent the calData value divided by each calibrator NRQ value must be sensitively equal). If this ratio is really different, you must exclude the highly variable IRC in all the qPCR runs.

6 Aggregation of multiple independent biological replicates from the same experiment

In this section, we will discuss about the final function of this package totData. In some research fields, the reproducibility of an observation can be tough (notably in the stem cells field). An algorithm published by Willems et al. (2008) [3] attenuates the high variations between independent biological replicates which have the same tendency in order to draw relevant statistical conclusions. This algorithm has been inputed in this function for the scientists experiencing this kind of issue.

```
## Isolate the NROs scaled to control of the first biological replicate
a1 <- nrmData(data = qPCR_run1, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02,
   0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = e, CalPos = 5,
    trace = FALSE, geo = TRUE, na.rm = TRUE)[1]
## Isolate the NRQs scaled to control of the second biological replicate
b1 \leftarrow nrmData(data = qPCR_run2, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02, 1.00)
   0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = f, CalPos = 5,
    trace = FALSE, geo = TRUE, na.rm = TRUE)[1]
## Isolate the NRQs scaled to control of the third biological replicate
c1 < -nrmData(data = qPCR_run3, r = 3, E = c(2, 2, 2, 2), Eerror = c(0.02, 0.02, 0.02, 0.02)
   0.02, 0.02), nSpl = 5, nbRef = 2, Refposcol = 1:2, nCTL = 2, CF = g, CalPos = 5,
    trace = FALSE, geo = TRUE, na.rm = TRUE)[1]
## Data frame transformation
a2 <- as.data.frame(a1)</pre>
b2 <- as.data.frame(b1)
c2 <- as.data.frame(c1)</pre>
## Aggregation of the three biological replicates
d2 <- rbind(a2, b2, c2)
```

Finally, we use the final function **totData** and indicate that we want to use the transformation algorithm published by Willems et al. (2008) [3] followed by the linearization process:

```
totData(data = d2, r = 3, geo = TRUE, logarithm = TRUE, base = 2, transformation = TRUE,
    nSpl = 5, linear = TRUE, na.rm = TRUE)
$Mean of your gPCR runs
                RG1
                         RG2
                                   TG
                                          TGb
Calibrator -0.08222 0.08222 -0.06722 0.1878
          -0.06889 0.06889 -0.06611 0.1744
Control 1
Control 2
          0.06889 -0.06889 0.06611 -0.1744
Treatment 1 0.08222 -0.08222 5.80500 -0.6711
Treatment 2 0.02611 -0.02611 8.67444 0.2761
$Standard deviations of
       your qPCR runs
               RG1
                       RG2
                                TG
Calibrator 0.03221 0.03653 0.01568 0.03713
Control 1 0.01654 0.01818 0.01046 0.01698
Control 2 0.01818 0.01654 0.01140 0.01331
Treatment 1 0.03187 0.02837 2.77617 0.03137
Treatment 2 0.04769 0.04634 9.20333 0.03650
$Standard errors of your qPCR runs
                RG1
                         RG2
                                   TG
Calibrator 0.018597 0.021092 0.009054 0.021438
Control 1 0.009551 0.010496 0.006037 0.009801
Control 2 0.010496 0.009551 0.006583 0.007687
Treatment 1 0.018401 0.016381 1.602820 0.018113
Treatment 2 0.027534 0.026753 5.313544 0.021074
$Transformed data
               RG1
                      RG2
                               TG
                                     TGb
Control 1
            0.9669 1.0342 0.1284 1.1445
Control 2
            1.0268 0.9739 0.1432 0.9254
Treatment 1 1.0878 0.9193 7.5994 0.6438
Treatment 2 1.0605 0.9429 55.4285 1.2758
            0.9691 1.0318 0.1276 1.1419
Calibrator
Control 11
            0.9340 1.0707 0.1311 1.1794
Control 21 1.0630 0.9408 0.1402 0.8980
Treatment 11 1.0244 0.9762 7.2061 0.6162
Treatment 21 0.9658 1.0354 56.6586 1.2597
Calibrator1 0.9063 1.1033 0.1318 1.2154
```

```
Control 12 0.9492 1.0535 0.1290 1.1605
Control 22 1.0459 0.9561 0.1425 0.9126
Treatment 12 1.0532 0.9495 7.9588 0.6806
Treatment 22 1.0197 0.9807 54.1625 1.2042
Calibrator2 0.9492 1.0535 0.1290 1.1605
$Reordered transformed data
               RG1
                      RG2
                              TG
                                    TGb
            0.9691 1.0318 0.1276 1.1419
Calibrator
Calibrator1 0.9063 1.1033 0.1318 1.2154
Calibrator2 0.9492 1.0535 0.1290 1.1605
Control 1
            0.9669 1.0342 0.1284 1.1445
Control 11
           0.9340 1.0707 0.1311 1.1794
Control 12 0.9492 1.0535 0.1290 1.1605
Control 2
            1.0268 0.9739 0.1432 0.9254
Control 21 1.0630 0.9408 0.1402 0.8980
Control 22 1.0459 0.9561 0.1425 0.9126
Treatment 1 1.0878 0.9193 7.5994 0.6438
Treatment 11 1.0244 0.9762 7.2061 0.6162
Treatment 12 1.0532 0.9495 7.9588 0.6806
Treatment 2 1.0605 0.9429 55.4285 1.2758
Treatment 21 0.9658 1.0354 56.6586 1.2597
Treatment 22 1.0197 0.9807 54.1625 1.2042
```

7 RStudio and gWidgets: tools to facilitate your qPCR data analysis

To facilitate the use of R, a free software named RStudio has been created [4]. This interface allows (among other things) easy data importation/exportation. In the same spirit of having an interface for using R, John Verzani has published a package gWidgets which has the great advantage to easily create a graphical user interface [5] for the function you want. In this last section, we will present how we can use these tools to facilitate the qPCR data analysis.

To begin, we must choose our workspace directory by typing this in your R session: setwd(gfile(type='selectdir')). You will see the opening of a window and you will just need to define your workspace directory. Then, we have to import some datasets in our R session. RStudio allows an easy data importation (see 1).

This can be done by following these steps:

- 1. uncompress the csv file (qPCR run1.csv) in the data folder
- 2. move it to inst/extdata

Then, you just need to type this in your R session:

```
file <- system.file("extdata", "qPCR_run1.csv", package = "EasyqpcR")</pre>
qPCR_run1 <- read.table(file, header = TRUE, sep = "", dec = ".")</pre>
qPCR_run1
       Samples
                 RG1
                       RG2
                               TG
                                    TGb
1
     Control 1 19.62 18.54 26.32 16.32
2
     Control 1 19.32 18.35 26.12 16.48
3
     Control 1 19.48 18.49 26.24 16.68
4
     Control 2 19.63 18.63 26.08 16.75
     Control 2 19.40 18.72 26.12 16.98
5
6
     Control 2 19.35 18.51 26.23 16.89
  Treatment 1 19.21 18.45 20.32 17.12
  Treatment 1 19.32 18.49 20.15 17.23
 Treatment 1 19.09 18.66 20.26 17.33
10 Treatment 2 19.63 18.75 17.65 16.65
11 Treatment 2 19.46 18.69 17.75 16.54
12 Treatment 2 19.51 18.92 17.60 16.40
13 Calibrator 19.88 18.91 26.69 16.69
14 Calibrator 19.78 18.72 26.49 16.85
15 Calibrator 19.85 18.86 26.63 17.05
```

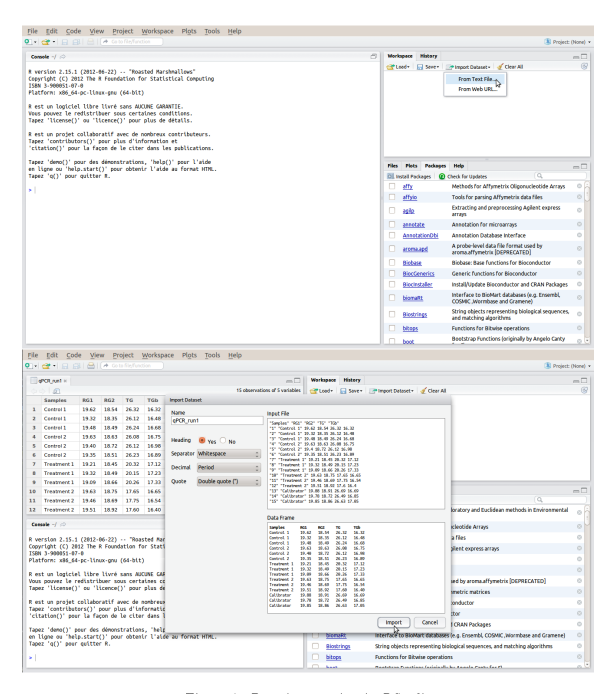


Figure 1: Data importation in RStudio

After data importation, you must control if your qPCR technical replicates satisfy your

threshold variation value (0.5 classically):

```
badCt(data = qPCR_run1, r = 3, threshold = 0.5, na.rm = TRUE)
$Bad replicates localization
     row col
$Mean of the Cq
              RG1
                    RG2
                           TG
                                TGb
Control 1
            19.47 18.46 26.23 16.49
          19.46 18.62 26.14 16.87
Control 2
Treatment 1 19.21 18.53 20.24 17.23
Treatment 2 19.53 18.79 17.67 16.53
Calibrator 19.84 18.83 26.60 16.86
$Standard error of the Cq
                RG1
                        RG2
                                 TG
                                        TGb
Control 1
            0.08667 0.05686 0.05812 0.10414
Control 2 0.08622 0.06083 0.04485 0.06692
Treatment 1 0.06642 0.06438 0.04978 0.06064
Treatment 2 0.05044 0.06888 0.04410 0.07234
Calibrator 0.02963 0.05686 0.05925 0.10414
```

Here, there is no bad replicates, but as an example, we will set the threshold value to 0.2 and see what it returns:

```
badCt(data = qPCR_run1, r = 3, threshold = 0.2, na.rm = TRUE)
$Bad replicates localization
            row col
Control 1
             1
                 2
             2
                 2
Control 2
Treatment 1
                 2
Control 2
             2
                 3
             3 3
Treatment 1
Treatment 2
             4 3
Calibrator
             5
                 4
Control 1
             1
                 5
             2
                 5
Control 2
Treatment 1 3
                 5
```

```
Treatment 2
                  5
Calibrator
                  5
$Mean of the Cq
                    RG2
                            TG
                                 TGb
Control 1
            19.47 18.46 26.23 16.49
Control 2
            19.46 18.62 26.14 16.87
Treatment 1 19.21 18.53 20.24 17.23
Treatment 2 19.53 18.79 17.67 16.53
Calibrator 19.84 18.83 26.60 16.86
$Standard error of the Cq
                RG1
                        RG2
                                  TG
                                         TGb
Control 1
            0.08667 0.05686 0.05812 0.10414
Control 2
            0.08622 0.06083 0.04485 0.06692
Treatment 1 0.06642 0.06438 0.04978 0.06064
Treatment 2 0.05044 0.06888 0.04410 0.07234
Calibrator 0.02963 0.05686 0.05925 0.10414
```

There are some bad replicates (according to the example threshold value). Now, we want to easily remove technical error (no more than one in qPCR technical triplicates), we just need to use the gdfnotebook function of the gWidgets package by typing this in the R session: gdfnotebook(cont=TRUE), and then choose which dataset you want to edit (qPCR_run1, here). After saving it (on an other name in order to easily reproduce your analysis with the same raw data, by example: qPCR_run1 is the raw dataset, and after removing technical replicates you can save it under the name qPCR_run1_cor). Or, you can edit your dataset directly in your spreadsheet and save it under an other name.

Finally, to easily analyze your qPCR data, you just will need to type this in your R session for each function of the package: **ggenericwidget(function,cont=TRUE)**, where function has to be replaced by nrmData, calData, totData, or badCt. This can also be done with the command lines described above.

8 How to analyse qPCR data with EasyqpcR when samples and genes are spread across runs?

All the previous examples showed how to perform qPCR data analysis when all the samples were present for each gene in each run. Here we present the procedure to follow when we have too much samples to be contained in each run, thus when samples and genes are spread across different runs.

Here are some examples of plate designs:

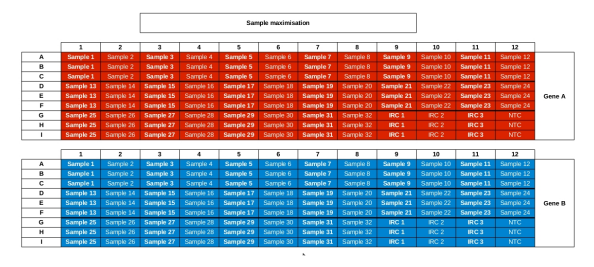


Figure 2: Sample maximisation strategy

										1			
			Sample maximisation										
										J			
	1	2	3	4	5	6	7	8	9	10	11	12	
Α	Sample 1	Sample 1	Sample 1	Sample 2	Sample 2	Sample 2	Sample 3	Sample 3	Sample 3	Sample 4	Sample 4	Sample 4	
В	IRC 1.1	IRC 1.1	IRC 1.1	IRC 2.1	IRC 2.1	IRC 2.1	IRC 3.1	IRC 3.1	IRC 3.1				
С	Sample 1	Sample 1	Sample 1		Sample 2	Sample 2	Sample 3	Sample 3	Sample 3	Sample 4	Sample 4	Sample 4	
D	IRC 1.1	IRC 1.1	IRC 1.1	IRC 2.1	IRC 2.1	IRC 2.1	IRC 3.1	IRC 3.1	IRC 3.1				Run 1
E	Sample 1	Sample 1	Sample 1	Sample 2	Sample 2	Sample 2	Sample 3	Sample 3	Sample 3	Sample 4	Sample 4	Sample 4	Run 1
F	IRC 1.1	IRC 1.1	IRC 1.1	IRC 2.1	IRC 2.1	IRC 2.1	IRC 3.1	IRC 3.1	IRC 3.1				
G	NTC 1.1	NTC 1.1	NTC 1.1	NTC 2.1	NTC 2.1	NTC 2.1	NTC 3.1	NTC 3.1	NTC 3.1				
н													
	1	2	3	4	5	6	7	8	9	10	11	12	
Α	Sample 5	Sample 5	Sample 5	Sample 6	Sample 6	Sample 6	Sample 7	Sample 7	Sample 7	Sample 8	Sample 8	Sample 8	
В	IRC 1.2	IRC 1.2	IRC 1.2	IRC 2.2	IRC 2.2	IRC 2.2	IRC 3.2	IRC 3.2	IRC 3.2				
С	Sample 5	Sample 5	Sample 5	Sample 6	Sample 6	Sample 6	Sample 7	Sample 7	Sample 7	Sample 8	Sample 8	Sample 8	
D	IRC 1.2	IRC 1.2	IRC 1.2	IRC 2.2	IRC 2.2	IRC 2.2	IRC 3.2	IRC 3.2	IRC 3.2				Run 2
E	Sample 5	Sample 5	Sample 5	Sample 6	Sample 6	Sample 6	Sample 7	Sample 7	Sample 7	Sample 8	Sample 8	Sample 8	Ruii 2
F	IRC 1.2	IRC 1.2	IRC 1.2	IRC 2.2	IRC 2.2	IRC 2.2	IRC 3.2	IRC 3.2	IRC 3.2				
G	NTC 1.2	NTC 1.2	NTC 1.2	NTC 2.2	NTC 2.2	NTC 2.2	NTC 3.2	NTC 3.2	NTC 3.2				
Н													
	1	2	3	4	5	6	7	8	9	10	11	12	
A	Sample 9	Sample 9	Sample 9	Sample 10	Sample 10	Sample 10	Sample 11	Sample 11	Sample 11	Sample 12	Sample 12	Sample 12	
В	IRC 1.3	IRC 1.3	IRC 1.3	IRC 2.3	IRC 2.3	IRC 2.3	IRC 3.3	IRC 3.3	IRC 3.3				
C	Sample 9	Sample 9	Sample 9	Sample 10	Sample 10	Sample 10	Sample 11	Sample 11	Sample 11	Sample 12	Sample 12	Sample 12	
D	IRC 1.3	IRC 1.3	IRC 1.3	IRC 2.3	IRC 2.3	IRC 2.3	IRC 3.3	IRC 3.3	IRC 3.3				
E	Sample 9	Sample 9	Sample 9	Sample 10	Sample 10	Sample 10	Sample 11	Sample 11	Sample 11	Sample 12	Sample 12	Sample 12	Run 3
F	IRC 1.3	IRC 1.3	IRC 1.3	IRC 2.3	IRC 2.3	IRC 2.3	IRC 3.3	IRC 3.3	IRC 3.3				
G	NTC 1.3	NTC 1.3	NTC 1.3	NTC 2.3	NTC 2.3	NTC 2.3	NTC 3.3	NTC 3.3	NTC 3.3				
н													
		L							L				

Figure 3: Gene maximisation strategy 1/3

	1	2	3	4	5	6	7	8	9	10	11	12	
Α	Sample 13	Sample 13	Sample 13	Sample 14	Sample 14	Sample 14	Sample 15	Sample 15	Sample 15	Sample 16	Sample 16	Sample 16	
В	IRC 1.4	IRC 1.4	IRC 1.4	IRC 2.4	IRC 2.4	IRC 2.4	IRC 3.4	IRC 3.4	IRC 3.4				
С	Sample 13	Sample 13	Sample 13	Sample 14	Sample 14	Sample 14	Sample 15	Sample 15	Sample 15	Sample 16	Sample 16	Sample 16	
D	IRC 1.4	IRC 1.4	IRC 1.4	IRC 2.4	IRC 2.4	IRC 2.4	IRC 3.4	IRC 3.4	IRC 3.4				
E	Sample 13	Sample 13	Sample 13	Sample 14	Sample 14	Sample 14	Sample 15	Sample 15	Sample 15	Sample 16	Sample 16	Sample 16	Run 4
F	IRC 1.4	IRC 1.4	IRC 1.4	IRC 2.4	IRC 2.4	IRC 2.4	IRC 3.4	IRC 3.4	IRC 3.4				
G	NTC 1.4	NTC 1.4	NTC 1.4	NTC 2.4	NTC 2.4	NTC 2.4	NTC 3.4	NTC 3.4	NTC 3.4				
н													
	1	2	3	4	5	6	7	8	9	10	11	12	
Α	Sample 17	Sample 17	Sample 17	Sample 18	Sample 18	Samplle 18	Sample 19	Sample 19	Sample 19	Sample 20	Sample 20	Sample 20	
В	IRC 1.5	IRC 1.5	IRC 1.5	IRC 2.5	IRC 25	IRC 2.5	IRC 3.5	IRC 3.5	IRC 3.5	Sample 20	Sample 20	Sample 20	
c	Sample 17	Sample 17	Sample 17	Sample 18	Sample 18	Sample 18	Sample 19	Sample 19	Sample 19	Sample 20	Sample 20	Sample 20	
D	IRC 1.5	IRC 1.5	IRC 1.5	IRC 2.5	IRC 2.5	IRC 2.5	IRC 3.5	IRC 3.5	IRC 3.5	Sample 20	Sample 20	Sample 20	
E	Sample 17	Sample 17	Sample 17	Sample 18	Sample 18	Sample 18	Sample 19	Sample 19	Sample 19	Sample 20	Sample 20	Sample 20	Run 5
	IRC 1.5	IRC 1.5	IRC 1.5	IRC 2.5	IRC 2.5	IRC 2.5	IRC 3.5	IRC 3.5	IRC 3.5	oumpie 20	Ottimpio 20	ouripie 20	
	NTC 1.5	NTC 1.5	NTC 1.5	NTC 2.5	NTC 2.5	NTC 2.5	NTC 3.5	NTC 3.5	NTC 3.5				
Н .	141 0 2.0	1110 2.0	1410 2.0	1110 2.0	111 0 2.0	1110 2.0	1110 0.0	11100.0	1110 0.0				
	1	2	3	4	5	6	7	8	9	10	11	12	
Α	Sample 21	Sample 21	Sample 21	Sample 22	Sample 22	Sample 22	Sample 23	Sample 23	Sample 23	Sample 24	Sample 24	Sample 24	
В В	IRC 1.6	IRC 1.6	IRC 1.6	IRC 2.6	IRC 2.6	IRC 2.6	IRC 3.6	IRC 3.6	IRC 3.6	Sample 24	Sample 24	Sample 24	
C B	Sample 21	Sample 21	Sample 21	Sample 22	Sample 22	Sample 22	Sample 23	Sample 23	Sample 23	Sample 24	Sample 24	Sample 24	
D	IRC 1.6	IRC 1.6	IRC 1.6	IRC 2.6	IRC 26	IRC 2.6	IRC 3.6	IRC 3.6	IRC 3.6	Sample 24	Sample 24	Sample 24	
E		Sample 21		Sample 22	Sample 22		Sample 23			Sample 24	Cample 24	Sample 24	Run 6
F	Sample 21 IRC 1.6	IRC 1.6	Sample 21 IRC 1.6	IRC 2.6	IRC 2.6	Sample 22 IRC 2.6	IRC 3.6	Sample 23 IRC 3.6	Sample 23 IRC 3.6	затріе 24	Sample 24	Sample 24	
G	NTC 1.6	NTC 1.6	NTC 1.6	NTC 2.6	NTC 2.6	NTC 2.6	NTC 3.6	NTC 3.6	NTC 3.6				
Н	NIC1.6	NIC 1.6	NIC 1.6	NIC 26	NI C 2.6	NTC 2.6	N1C 3.6	N1 C 3.6	N1C 3.6				
н	1	I	1	I	1	I	I	I	1	I	I	1	

Figure 4: Gene maximisation strategy 2/3

	1	2	3	4	5	6	7	8	9	10	11	12	
Α	Sample 25	Sample 25	Sample 25	Sample 26	Sample 26	Sample 26	Sample 27	Sample 27	Sample 27	Sample 28	Sample 28	Sample 28	
В	IRC 1.7	IRC 1.7	IRC 1.7	IRC 2.7	IRC 2.7	IRC 2.7	IRC 3.7	IRC 3.7	IRC 3.7				
С	Sample 25	Sample 25	Sample 25	Sample 26	Sample 26	Sample 26	Sample 27	Sample 27	Sample 27	Sample 28	Sample 28	Sample 28	
D	IRC 1.7	IRC 1.7	IRC 1.7	IRC 2.7	IRC 2.7	IRC 2.7	IRC 3.7	IRC 3.7	IRC 3.7				Run 7
E	Sample 25	Sample 25	Sample 25	Sample 26	Sample 26	Sample 26	Sample 27	Sample 27	Sample 27	Sample 28	Sample 28	Sample 28	Ruii /
F	IRC 1.7	IRC 1.7	IRC 1.7	IRC 2.7	IRC 2.7	IRC 2.7	IRC 3.7	IRC 3.7	IRC 3.7				
G	NTC 1.7	NTC 1.7	NTC 1.7	NTC 2.7	NTC 2.7	NTC 2.7	NTC 3.7	NTC 3.7	NTC 3.7				
Н													

	1	2	3	4	5	6	7	8	9	10	11	12	
Α	Sample 29	Sample 29	Sample 29	Sample 30	Sample 30	Sample 30	Sample 31	Sample 31	Sample 31	Sample 32	Sample 32	Sample 32	
В	IRC 1.8	IRC 1.8	IRC 1.8	IRC 2.8	IRC 2.8	IRC 2.8	IRC 3.8	IRC 3.8	IRC 3.8				
С	Sample 29	Sample 29	Sample 29	Sample 30	Sample 30	Sample 30	Sample 31	Sample 31	Sample 31	Sample 32	Sample 32	Sample 32	
D	IRC 1.8	IRC 1.8	IRC 1.8	IRC 2.8	IRC 2.8	IRC 2.8	IRC 3.8	IRC 3.8	IRC 3.8				Run 8
E	Sample 29	Sample 29	Sample 29	Sample 30	Sample 30	Sample 30	Sample 31	Sample 31	Sample 31	Sample 32	Sample 32	Sample 32	Ruito
F	IRC 1.8	IRC 1.8	IRC 1.8	IRC 2.8	IRC 2.8	IRC 2.8	IRC 3.8	IRC 3.8	IRC 3.8				
G	NTC 1.8	NTC 1.8	NTC 1.8	NTC 28	NTC 2.8	NTC 2.8	NTC 3.8	NTC 3.8	NTC 3.8				
Н													

Figure 5: Gene maximisation strategy 3/3

As described in the work of Hellemans et al (2007) [2], the sample maximisation strategy does not need inter-run calibration factor because there is no samples spread over runs (all the samples for each gene are contained in one plate). Unless you have to many samples to be contained in one run even with sample maximisation strategy, you will have to perform the inter-run variation correction presented below.

In gene maximisation strategy the samples and genes are spread across runs. Thus, it is necessary to calibrate the NRQs with a run-specific and gene-specific calibration factor.

We will now describe how to handle this issue. Follow these steps:

- 1. uncompress the csv file (Gene_maximisation.csv) in the data folder
- 2. move it to inst/extdata

Then, you just have to do this:

```
filebis <- system.file("extdata", "Gene_maximisation.csv", package = "EasyqpcR")
Gene_maximisation <- read.table(filebis, header = TRUE, sep = ";", dec = ",")</pre>
```

First, you have to analyze if there are some bad replicates:

```
badCt(data = Gene_maximisation, r = 3, threshold = 0.5, na.rm = FALSE)[1]
$Bad replicates localization
    row col
```

```
Sample 15
           15
                  2
NTC 1
                  2
            36
NTC 2
                  2
            40
NTC 3
                  2
            44
NTC 5
            52
                  2
NTC 7
            60
                  2
NTC 8
            64
                  2
Sample 18
            18
                  3
NTC 1
                  3
            36
NTC 2
            40
                  3
NTC 3
                  3
            44
NTC 4
            48
                  3
NTC 7
            60
                  3
NTC 8
            64
                  3
NTC 1
            36
                  4
NTC 2
            40
                  4
NTC 3
            44
                  4
NTC 4
            48
                  4
NTC 7
            60
                  4
NTC 8
            64
```

Do not worry about the NTC (no template control), they are not needed in qPCR data analysis (but they have to present at least a Ct difference > 5 Ct compared to the samples). Here, you can use the gdfnotebook function of the gWidgets package.

After having removed the two aberrant values (RG2, Sample 15, Ct=22.0691567571; and RG3, Sample 18, Ct=19.0232804823), rename the data frame as Gene_maximisation_cor. Now, we remove the NTC values:

```
fileter <- system.file("extdata", "Gene_maximisation_cor.csv", package = "EasyqpcR")

Gene_maximisation_cor <- read.table(fileter, header = TRUE, sep = ";", dec = ",")

Gene_maximisation_cor1 <- Gene_maximisation_cor[-c(106:108, 118:120, 130:132, 142:144, 154:156, 166:168, 178:180, 190:192), ]

rownames(Gene_maximisation_cor1) <- c(1:168)</pre>
```

Now, we will calculate run-specific calibration factors:

```
calr1 <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02,
    0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = c(1, 1)
    1, 1), CalPos = c(33:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]][1:3,
calr2 <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02,
    0.02, 0.02), nSp1 = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = c(1, 1)
    1, 1), CalPos = c(33:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]][4:6,
calr3 <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02,
    0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = c(1, 1)
    1, 1), CalPos = c(33:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]][7:9,
calr4 <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02,
    0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = c(1, 1)
   1, 1), CalPos = c(33:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]][10:12,
calr5 <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02,
    0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = c(1, 1)
   1, 1), CalPos = c(33:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]][13:15,
    calr6 <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02,
    0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = c(1, 1)
   1, 1), CalPos = c(33:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]][16:18,
    calr7 <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02,
    0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = c(1, 1)
   1, 1), CalPos = c(33:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]][19:21,
calr8 <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02,
    0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = c(1, 1)
   1, 1), CalPos = c(33:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[3]][22:24,
```

```
e <- calData(calr1)

f <- calData(calr2)

g <- calData(calr3)

h <- calData(calr4)

i <- calData(calr5)

j <- calData(calr6)

k <- calData(calr7)

1 <- calData(calr8)</pre>
```

To respect the calculation of the NRQs which need the whole samples to take into account the whole variability, we have to apply the nrmData function on the whole samples. But we will do it for each run with the correction by the run-specific calibration factor and after each inter-run variation correction we isolate the corresponding CNRQs (i.e. the NRQs corrected by the specific CF), for example:

We perform inter-run variation correction on the whole samples by the CF of the first run, which corresponds to the samples 1 to 4 and IRC 1.1, 2.1, 3.1. But, the CF of the first is not the correct CF for the second run and for any other run. Thus, we isolate (after inter-run variation correction on the whole samples by the CF of the first run) the samples concerned by this specific CF which are the samples 1 to 4 and IRC 1.1, 2.1, 3.1. And we do it for each run-specific CF. Then we isolate the NRQs of the samples 5 to 8 and IRC 1.2, 2.2, 3.2 corrected by the CF of the second run, etc...

```
m <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02, 0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = e, CalPos = c(33:35), trace = FALSE, geo = TRUE, na.rm = TRUE)[[2]][c(1:4, 33:35), ]</pre>
n <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02, 0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = f, CalPos = c(36:56), trace = FALSE, geo = TRUE, na.rm = TRUE)[[2]][c(5:8, 36:38), ]
```

```
o <- nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02, 1)
  0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = g, CalPos = c(36:56),
    trace = FALSE, geo = TRUE, na.rm = TRUE)[[2]][c(9:12, 39:41), ]
p < -nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02, 1)
  0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = h, CalPos = c(33:35),
    trace = FALSE, geo = TRUE, na.rm = TRUE)[[2]][c(13:16, 42:44), ]
q < -nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02, 1)
  0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = i, CalPos = c(36:56),
    trace = FALSE, geo = TRUE, na.rm = TRUE)[[2]][c(17:20, 45:47), ]
r < -nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02, 1)
  0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = j, CalPos = c(36:56),
    trace = FALSE, geo = TRUE, na.rm = TRUE)[[2]][c(21:24, 48:50), ]
s < -nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02, 1)
  0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = k, CalPos = c(33:35),
    trace = FALSE, geo = TRUE, na.rm = TRUE)[[2]][c(25:28, 51:53), ]
t < -nrmData(data = Gene_maximisation_cor1, r = 3, E = c(2, 2, 2), Eerror = c(0.02, 1)
  0.02, 0.02), nSpl = 56, nbRef = 2, Refposcol = 1:2, nCTL = 16, CF = 1, CalPos = c(36:56),
    trace = FALSE, geo = TRUE, na.rm = TRUE)[[2]][c(29:32, 54:56), ]
## Aggregation of all the CNRQs
u <- rbind(m, n, o, p, q, r, s, t)
```

Explanation of what we have done before: we have isolated the corresponding NRQs corrected by the specific CF (remember that after correction by the specific CF , we talk in terms of CNRQs and not anymore of NRQs):

- Samples 1 to 4 and IRC 1.1, 2.1, 3.1 are corrected by e
- Samples 5 to 8 and IRC 1.2, 2.2, 3.2 are corrected by f
- Samples 9 to 12 and IRC 1.3, 2.3, 3.3 are corrected by g
- Samples 13 to 16 and IRC 1.4, 2.4, 3.4 are corrected by h

- Samples 17 to 20 and IRC 1.5, 2.5, 3.5 are corrected by i
- Samples 21 to 24 and IRC 1.6, 2.6, 3.6 are corrected by j
- Samples 25 to 28 and IRC 1.7, 2.7, 3.7 are corrected by k
- Samples 29 to 32 and IRC 1.8, 2.8, 3.8 are corrected by l

Do not worry about the nCTL parameter, because in gene maximisation (or sample maximisation if there are too many samples), the CTL samples are not present in all the runs, so you will have to perform the scaling to control after.

Note that in this case where the control samples are not present in all the runs, the nCTL parameter is not relevant and we only take into account the NRQs corrected by the calibration factor (CNRQs). Thus, to have nicer graphs, you will need to perform a scaling to your control group by doing a geometric mean of the CNRQs of your control samples and divide all the CNRQs by this geometric mean. Here the control group is composed by the samples 1 to 16, thus:

```
ctlgroup <- u[c(1:4, 8:11, 15:18, 22:25), ]

ctlgeom <- colProds(ctlgroup)^(1/dim(ctlgroup)[1])
ctlgeom1 <- (as.data.frame(ctlgeom)[rep(1:(ncol(u)), each = nrow(u)), ])
ctlgeom2 <- as.data.frame(matrix(ctlgeom1, ncol = ncol(u), byrow = FALSE))

CNRQs_scaled_to_group <- u/ctlgeom2</pre>
```

References

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