

NormqPCR: Functions for normalisation of RT-qPCR data

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1 Introduction

The package "NormqPCR" provides methods for the normalization of real-time quantitative RT-PCR data. In this vignette we describe and demonstrate the available functions. Firstly we show how the user may combine technical replicates, deal with undetermined values and deal with values above a user-chosen threshold. The rest of the vignette is split into two distinct sections, the first giving details of different methods to select the best housekeeping gene/genes for normalisation, and the second showing how to use the selected housekeeping gene(s) to produce $2^{-\Delta Cq}$ normalised estimators and $2^{-\Delta\Delta Cq}$ estimators of differential expression.

2 Combining technical replicates

When a raw data file read in using `read.qPCR` contains technical replicates, they are dealt with by concatenating the suffix `_TechRep.n` to the detector name, where `n` in 1, 2...`N` is the number of the replication in the total number of replicates, `N`, based on order of appearance in the qPCR data file.

So if we read in a file with technical replicates, we can see that the detector/feature names are thus suffixed:

```
> library(ReadqPCR) # load the ReadqPCR library
> library(NormqPCR)
> path <- system.file("exData", package = "NormqPCR")
> qPCR.example.techReps <- file.path(path, "qPCR.techReps.txt")
> qPCRBatch.qPCR.techReps <- read.qPCR(qPCR.example.techReps)
> rownames(exprs(qPCRBatch.qPCR.techReps))[1:8]

[1] "gene_aj_TechReps.1" "gene_aj_TechReps.2" "gene_al_TechReps.1"
[4] "gene_al_TechReps.2" "gene_ax_TechReps.1" "gene_ax_TechReps.2"
[7] "gene_bo_TechReps.1" "gene_bo_TechReps.2"
```

It is likely that before continuing with the analysis, the user would wish to average the technical replicates by using the arithmetic mean of the raw Cq values. This can be achieved using the `combineTechReps` function, which will produce a new `qPCRBatch` object, with all tech reps reduced to one reading:

```
> combinedTechReps <- combineTechReps(qPCRBatch.qPCR.techReps)
> combinedTechReps
```

```
qPCRBatch (storageMode: lockedEnvironment)
assayData: 8 features, 3 samples
  element names: exprs
protocolData: none
phenoData
  sampleNames: one three two
  varLabels: sample
  varMetadata: labelDescription
featureData: none
experimentData: use 'experimentData(object)'
Annotation:
```

3 Dealing with undetermined values

When an RT-qPCR experiment does not produce a reading after a certain number of cycles (the cycle threshold), the reading is given as undetermined. These are represented in qPCRBatch objects as NA. Different users may have different ideas about how many cycles they wish to allow before declaring a detector as not present in the sample. There are two methods for the user to decide what to do with numbers above a given cycle threshold:

First the user might decide that anything above 38 cycles means there is nothing present in their sample, instead of the standard 40 used by the taqman software. They can replace the value of all readings above 38 as NA using the following:

Firstly read in the taqman example file which has 96 detectors, with 4 replicates for mia (case) and 4 non-mia (control):

```
> path <- system.file("exData", package = "NormqPCR")
> taqman.example <- file.path(path, "/example.txt")
> qPCRBatch.taqman <- read.taqman(taqman.example)
```

We can see that for the detector: Cc120.Rn00570287_m1 we have these readings for the different samples:

```
> exprs(qPCRBatch.taqman)["Cc120.Rn00570287_m1",]

      fp1.day3.v      fp2.day3.v      fp5.day3.mia      fp6.day3.mia      fp.3.day.3.v
              NA              NA             35.74190             34.05922             35.02052
fp.4.day.3.v fp.7.day.3.mia fp.8.day.3.mia
              NA             35.93689             36.57921
```

We can now use the `replaceAboveCutOff` method in order to replace anything above 35 with NA:

```
> qPCRBatch.taqman.replaced <- replaceAboveCutOff(qPCRBatch.taqman,
+                                                  newVal = NA, cutOff = 35)
> exprs(qPCRBatch.taqman.replaced)["Cc120.Rn00570287_m1",]

      fp1.day3.v      fp2.day3.v      fp5.day3.mia      fp6.day3.mia      fp.3.day.3.v
              NA              NA              NA             34.05922              NA
fp.4.day.3.v fp.7.day.3.mia fp.8.day.3.mia
              NA              NA              NA
```

It may also be the case that the user wants to get rid of all NA values, and replace them with an arbitrary number. This can be done using the `replaceNAs` method. So if the user wanted to replace all NAs with 40, it can be done as follows:

```
> qPCRBatch.taqman.replaced <- replaceNAs(qPCRBatch.taqman, newNA = 40)
> exprs(qPCRBatch.taqman.replaced)["Cc120.Rn00570287_m1",]

      fp1.day3.v    fp2.day3.v    fp5.day3.mia    fp6.day3.mia    fp.3.day.3.v
      40.00000      40.00000      35.74190      34.05922      35.02052
fp.4.day.3.v fp.7.day.3.mia fp.8.day.3.mia
      40.00000      35.93689      36.57921
```

In addition, the situation sometimes arises where some readings for a given detector are above a given cycle threshold, but some others are not. The user may decide for example that if a given number of readings are NAs, then all of the readings for this detector should be NAs. This is important because otherwise an unusual reading for one detector might lead to an inaccurate estimate for the expression of a given gene.

This process will necessarily be separate for the different sample types, since you might expect a given gene to show expression in one sample type compared to another. Therefore it is necessary to designate the replicates per sample type using a contrast matrix. It is also necessary to make a sampleMaxMatrix which gives a maximum number of NAs allowed for each sample type.

So in the example file above we two sample types, with 4 biological replicates for each, the contrastMatrix and sampleMaxMatrix might be constructed like this:

```
> sampleNames(qPCRBatch.taqman)

[1] "fp1.day3.v"      "fp2.day3.v"      "fp5.day3.mia"    "fp6.day3.mia"
[5] "fp.3.day.3.v"    "fp.4.day.3.v"    "fp.7.day.3.mia"  "fp.8.day.3.mia"

> a <- c(0,0,1,1,0,0,1,1) # one for each sample type, with 1 representing
> b <- c(1,1,0,0,1,1,0,0) # position of sample type in sampleNames vector
> contM <- cbind(a,b)
> colnames(contM) <- c("case","control") # set the names of each sample type
> rownames(contM) <- sampleNames(qPCRBatch.taqman) # set row names
> contM

      case control
fp1.day3.v      0      1
fp2.day3.v      0      1
fp5.day3.mia    1      0
fp6.day3.mia    1      0
fp.3.day.3.v    0      1
fp.4.day.3.v    0      1
fp.7.day.3.mia  1      0
fp.8.day.3.mia  1      0
```

```
> sMaxM <- t(as.matrix(c(3,3))) # now make the contrast matrix
> colnames(sMaxM) <- c("case","control") # make sure these line up with samples
> sMaxM
```

```
      case control
[1,]      3      3
```

More details on contrast matrices can be found in the limma manual, which requires a similar matrix when testing for differential expression between samples.

For example, if the user decides that if at least 3 out of 4 readings are NAs for a given detector, then all readings should be NA, they can do the following, using the `makeAllNewVal` method:

```
> qPCRBatch.taqman.replaced <- makeAllNewVal(qPCRBatch.taqman, contM,
+                                           sMaxM, newVal=NA)
```

Here you can see for the Ccl20.Rn00570287_m1 detector, the control values have been made all NA, whereas before 3 were NA and one was 35. However the case values have been kept, since they were all below the NA threshold. It is important to filter the data in this way to ensure the correct calculations are made downstream when calculating variation and other parameters.

```
> exprs(qPCRBatch.taqman.replaced)["Ccl20.Rn00570287_m1",]
```

fp1.day3.v	fp2.day3.v	fp5.day3.mia	fp6.day3.mia	fp.3.day.3.v
NA	NA	35.74190	34.05922	NA
fp.4.day.3.v	fp.7.day.3.mia	fp.8.day.3.mia		
NA	35.93689	36.57921		

4 Selection of most stable reference/housekeeping genes

This section contains two subsections containing different methods for the selection of appropriate housekeeping genes.

4.1 geNorm

We describe the selection of the best (most stable) reference/housekeeping genes using the method of Vandesompele et al (2002) [3] (in the sequel: Vand02) which is called *geNorm*. We first load the package and the data

```
> options(width = 68)
> data(geNorm)
> str(exprs(geNorm.qPCRBatch))
```

```

num [1:10, 1:85] 0.0425 0.0576 0.1547 0.1096 0.118 ...
- attr(*, "dimnames")=List of 2
..$ : chr [1:10] "ACTB" "B2M" "GAPD" "HMBS" ...
..$ : chr [1:85] "BM1" "BM2" "BM3" "BM4" ...

```

We start by ranking the selected reference/housekeeping genes. The geNorm algorithm implemented in function `selectHKs` proceeds stepwise; confer Section “Materials and methods” in Vand02. That is, the gene stability measure M of all candidate genes is computed and the gene with the highest M value is excluded. Then, the gene stability measure M for the remaining gene is calculated and so on. This procedure is repeated until two respectively, `minNrHK` genes remain.

```

> tissue <- as.factor(c(rep("BM", 9), rep("FIB", 20), rep("LEU", 13),
+                      rep("NB", 34), rep("POOL", 9)))
> res.BM <- selectHKs(geNorm.qPCRBatch[,tissue == "BM"], method = "geNorm",
+                     Symbols = featureNames(geNorm.qPCRBatch),
+                     minNrHK = 2, log = FALSE)

```

HPRT1	YWHAZ	RPL13A	UBC	GAPD	SDHA
0.5160313	0.5314564	0.5335963	0.5700961	0.6064919	0.6201470
TBP	HMBS	B2M	ACTB		
0.6397969	0.7206013	0.7747634	0.8498739		
HPRT1	RPL13A	YWHAZ	UBC	GAPD	SDHA
0.4705664	0.5141375	0.5271169	0.5554718	0.5575295	0.5738460
TBP	HMBS	B2M			
0.6042110	0.6759176	0.7671985			
HPRT1	RPL13A	SDHA	YWHAZ	UBC	GAPD
0.4391222	0.4733732	0.5243665	0.5253471	0.5403137	0.5560120
TBP	HMBS				
0.5622094	0.6210820				
HPRT1	RPL13A	YWHAZ	UBC	SDHA	GAPD
0.4389069	0.4696398	0.4879728	0.5043292	0.5178634	0.5245346
TBP					
0.5563591					
HPRT1	RPL13A	UBC	YWHAZ	GAPD	SDHA
0.4292808	0.4447874	0.4594181	0.4728920	0.5012107	0.5566762
UBC	RPL13A	HPRT1	YWHAZ	GAPD	
0.4195958	0.4204997	0.4219179	0.4424631	0.4841646	
RPL13A	UBC	YWHAZ	HPRT1		
0.3699163	0.3978736	0.4173706	0.4419220		
UBC	RPL13A	YWHAZ			
0.3559286	0.3761358	0.3827933			

```
RPL13A      UBC
0.3492712 0.3492712
```

```
> res.POOL <- selectHKs(geNorm.qPCRBatch[,tissue == "POOL"],
+                       method = "geNorm",
+                       Symbols = featureNames(geNorm.qPCRBatch),
+                       minNrHK = 2, trace = FALSE, log = FALSE)
> res.FIB <- selectHKs(geNorm.qPCRBatch[,tissue == "FIB"],
+                      method = "geNorm",
+                      Symbols = featureNames(geNorm.qPCRBatch),
+                      minNrHK = 2, trace = FALSE, log = FALSE)
> res.LEU <- selectHKs(geNorm.qPCRBatch[,tissue == "LEU"],
+                      method = "geNorm",
+                      Symbols = featureNames(geNorm.qPCRBatch),
+                      minNrHK = 2, trace = FALSE, log = FALSE)
> res.NB <- selectHKs(geNorm.qPCRBatch[,tissue == "NB"],
+                     method = "geNorm",
+                     Symbols = featureNames(geNorm.qPCRBatch),
+                     minNrHK = 2, trace = FALSE, log = FALSE)
```

We obtain the following ranking of genes (see Table 3 in Vand02)

```
> ranks <- data.frame(c(1, 1:9), res.BM$ranking, res.POOL$ranking,
+                      res.FIB$ranking, res.LEU$ranking,
+                      res.NB$ranking)
> names(ranks) <- c("rank", "BM", "POOL", "FIB", "LEU", "NB")
> ranks
```

	rank	BM	POOL	FIB	LEU	NB
1	1	RPL13A	GAPD	GAPD	UBC	GAPD
2	1	UBC	SDHA	HPRT1	YWHAZ	HPRT1
3	2	YWHAZ	HMBS	YWHAZ	B2M	SDHA
4	3	HPRT1	HPRT1	UBC	GAPD	UBC
5	4	GAPD	TBP	ACTB	RPL13A	HMBS
6	5	SDHA	UBC	TBP	TBP	YWHAZ
7	6	TBP	RPL13A	SDHA	SDHA	TBP
8	7	HMBS	YWHAZ	RPL13A	HPRT1	ACTB
9	8	B2M	ACTB	B2M	HMBS	RPL13A
10	9	ACTB	B2M	HMBS	ACTB	B2M

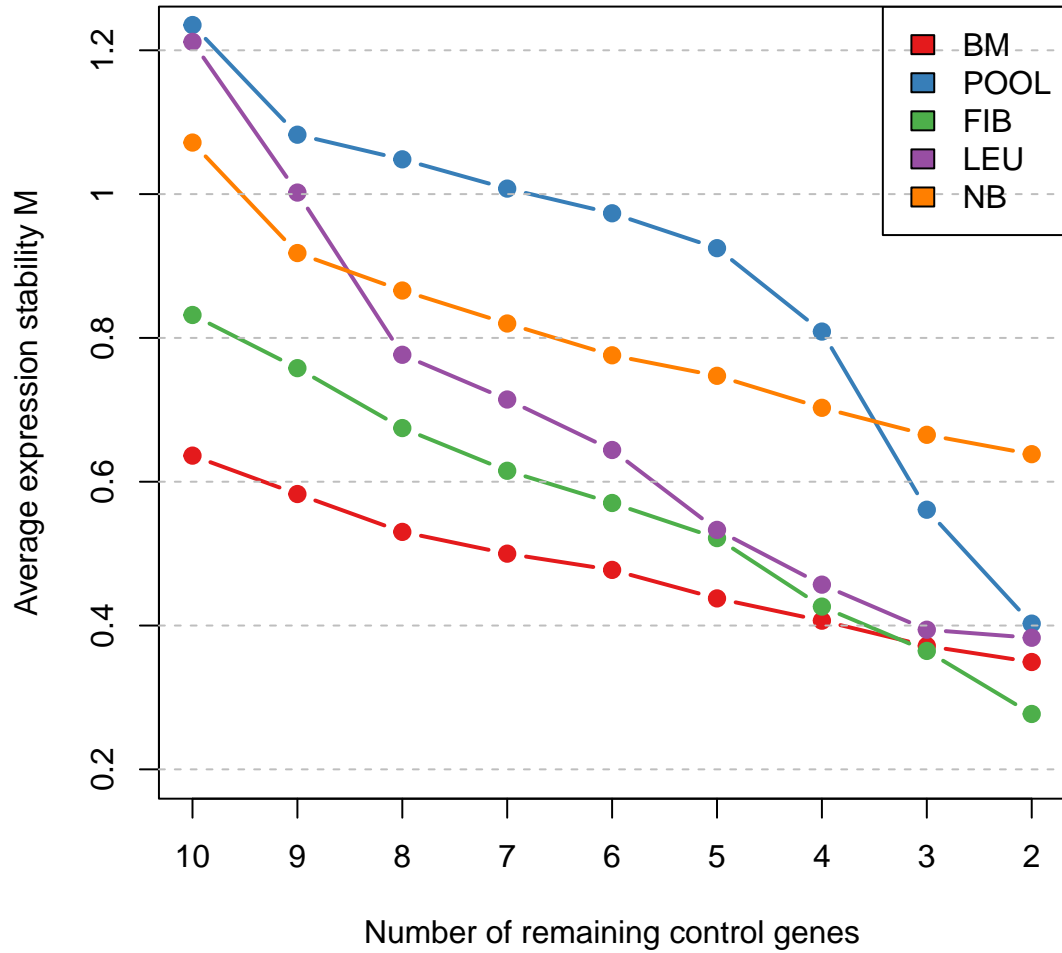
Remark 1:

Since the computation is based on gene ratios, the two most stable control genes in each cell type cannot be ranked.

We plot the average expression stability M for each cell type (see Figure 2 in Vand02).

```
> library(RColorBrewer)
> mypalette <- brewer.pal(5, "Set1")
> matplot(cbind(res.BM$meanM, res.POOL$meanM, res.FIB$meanM,
+               res.LEU$meanM, res.NB$meanM), type = "b",
+         ylab = "Average expression stability M",
+         xlab = "Number of remaining control genes",
+         axes = FALSE, pch = 19, col = mypalette,
+         ylim = c(0.2, 1.22), lty = 1, lwd = 2,
+         main = "Figure 2 in Vandesompele et al. (2002)")
> axis(1, at = 1:9, labels = as.character(10:2))
> axis(2, at = seq(0.2, 1.2, by = 0.2), labels = seq(0.2, 1.2, by = 0.2))
> box()
> abline(h = seq(0.2, 1.2, by = 0.2), lty = 2, lwd = 1, col = "grey")
> legend("topright", legend = c("BM", "POOL", "FIB", "LEU", "NB"),
+       fill = mypalette)
```


Figure 2 in Vandesompele et al. (2002)



Second, we plot the pairwise variation for each cell type (see Figure 3 (a) in Vand02)

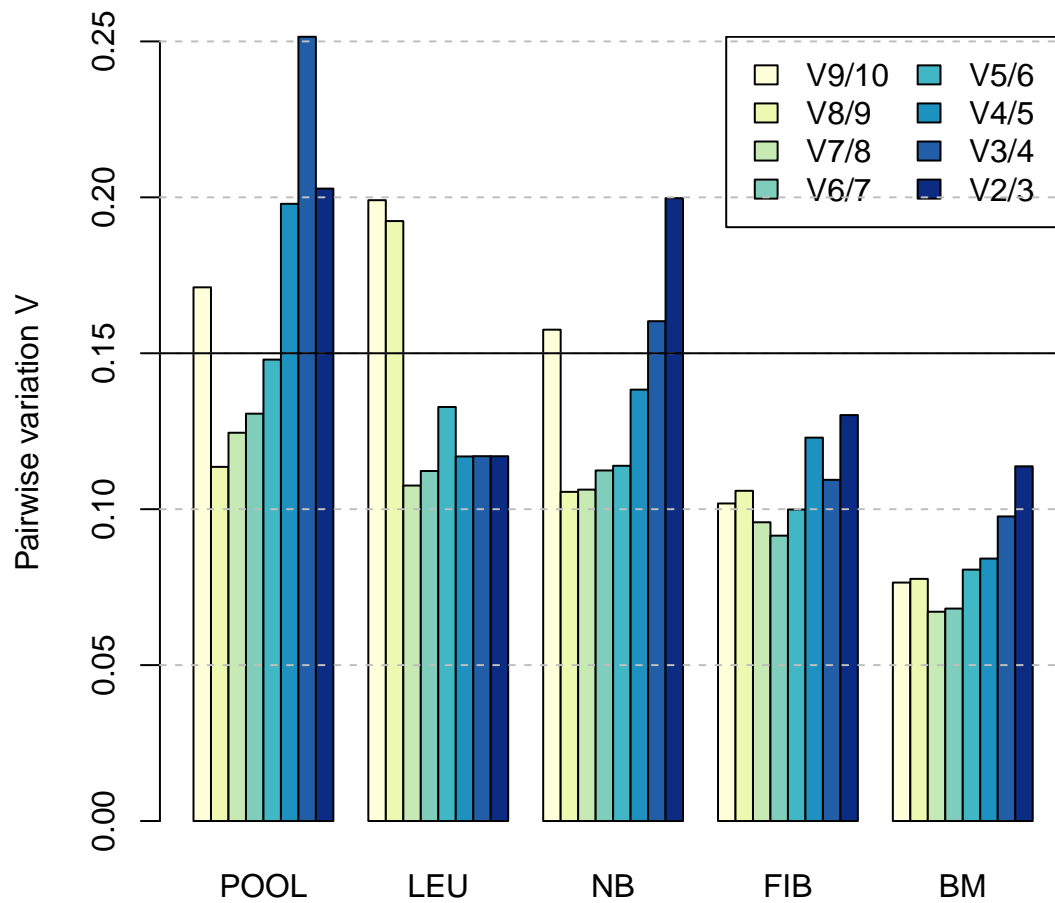
```
> mypalette <- brewer.pal(8, "YlGnBu")
> barplot(cbind(res.POOL$variation, res.LEU$variation, res.NB$variation,
+               res.FIB$variation, res.BM$variation), beside = TRUE,
+         col = mypalette, space = c(0, 2),
+         names.arg = c("POOL", "LEU", "NB", "FIB", "BM"),
+         ylab = "Pairwise variation V",
+         main = "Figure 3(a) in Vandesompele et al. (2002)")
```

```

> legend("topright", legend = c("V9/10", "V8/9", "V7/8", "V6/7",
+                               "V5/6", "V4/5", "V3/4", "V2/3"),
+       fill = mypalette, ncol = 2)
> abline(h = seq(0.05, 0.25, by = 0.05), lty = 2, col = "grey")
> abline(h = 0.15, lty = 1, col = "black")

```

Figure 3(a) in Vandesompele et al. (2002)



Remark 2:

Vand02 recommend a cut-off value of 0.15 for the pairwise variation. Below this bound the inclusion of an additional housekeeping gene is not required.

4.2 NormFinder

The second method for selection reference/housekeeping genes implemented in package is the method derived by [1] (in the sequel: And04) called *NormFinder*.

The ranking contained in Table 3 of And04 can be obtained via

```
> data(Colon)
> Colon

qPCRBatch (storageMode: lockedEnvironment)
assayData: 13 features, 40 samples
  element names: exprs
protocolData: none
phenoData
  sampleNames: I459N 90 ... I-C1056T (40 total)
  varLabels: Sample.no. Classification
  varMetadata: labelDescription
featureData
  featureNames: UBC UBB ... TUBA6 (13 total)
  fvarLabels: Symbol Gene.name
  fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
Annotation: Table 1 in Andersen et al. (2004)

> Class <- pData(Colon)[,"Classification"]
> res.Colon <- stabMeasureRho(Colon, group = Class, log = FALSE)
> sort(res.Colon) # see Table 3 in Andersen et al (2004)

      UBC      GAPD      TPT1      UBB      TUBA6      RPS13
0.1821707 0.2146061 0.2202956 0.2471573 0.2700641 0.2813039
      NACA      CFL1      SUI1      ACTB      CLTC      RPS23
0.2862397 0.2870467 0.3139404 0.3235918 0.3692880 0.3784909
      FLJ20030
0.3935173

> data(Bladder)
> Bladder

qPCRBatch (storageMode: lockedEnvironment)
assayData: 14 features, 28 samples
  element names: exprs
protocolData: none
phenoData
```

```

sampleNames: 335-6 1131-1 ... 1356-1 (28 total)
varLabels: Sample.no. Grade
varMetadata: labelDescription
featureData
  featureNames: ATP5B HSPCB ... FLJ20030 (14 total)
  fvarLabels: Symbol Gene.name
  fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
Annotation: Table 1 in Andersen et al. (2004)

> grade <- pData(Bladder)[,"Grade"]
> res.Bladder <- stabMeasureRho(Bladder, group = grade,
+                               log = FALSE)
> sort(res.Bladder)

```

HSPCB	TEGT	ATP5B	UBC	RPS23	RPS13
0.1539598	0.1966556	0.1987227	0.2033477	0.2139626	0.2147852
CFL1	FLJ20030	TPT1	UBB	FLOT2	GAPD
0.2666129	0.2672918	0.2691553	0.2826051	0.2960429	0.3408742
S100A6	ACTB				
0.3453435	0.3497295				

Of course, we can also reproduce the geNorm ranking also included in Table 3 of And04.

```

> selectHKs(Colon, log = FALSE, trace = FALSE,
+           Symbols = featureNames(Colon))$ranking

```

1	1	3	4	5	6
"RPS23"	"TPT1"	"RPS13"	"SUI1"	"UBC"	"GAPD"
7	8	9	10	11	12
"TUBA6"	"UBB"	"NACA"	"CFL1"	"CLTC"	"ACTB"
13					
"FLJ20030"					

```

> selectHKs(Bladder, log = FALSE, trace = FALSE,
+           Symbols = featureNames(Bladder))$ranking

```

1	1	3	4	5	6
"CFL1"	"UBC"	"ATP5B"	"HSPCB"	"GAPD"	"TEGT"
7	8	9	10	11	12
"RPS23"	"RPS13"	"TPT1"	"FLJ20030"	"FLOT2"	"UBB"
13	14				
"ACTB"	"S100A6"				

As we are often interested in more than one reference/housekeeping gene we also implemented a step-wise procedure of the NormFinder algorithm explained in Section “Average control gene” in the supplementary information of And04. This procedure is available via function `selectHKs`.

```
> Class <- pData(Colon)[,"Classification"]
> selectHKs(Colon, group = Class, log = FALSE, trace = TRUE,
+           Symbols = featureNames(Colon), minNrHKs = 12,
+           method = "NormFinder")$ranking
```

UBC	GAPD	TPT1	UBB	TUBA6	RPS13
0.1821707	0.2146061	0.2202956	0.2471573	0.2700641	0.2813039
NACA	CFL1	SUI1	ACTB	CLTC	RPS23
0.2862397	0.2870467	0.3139404	0.3235918	0.3692880	0.3784909
FLJ20030					
0.3935173					
GAPD	TPT1	UBB	NACA	CFL1	RPS13
0.1375298	0.1424519	0.1578360	0.1657364	0.1729069	0.1837057
TUBA6	SUI1	ACTB	RPS23	FLJ20030	CLTC
0.1849021	0.2065531	0.2131651	0.2188277	0.2359623	0.2447588
TPT1	NACA	UBB	RPS13	CFL1	TUBA6
0.1108474	0.1299802	0.1356690	0.1411173	0.1474242	0.1532953
FLJ20030	SUI1	ACTB	RPS23	CLTC	
0.1583031	0.1586250	0.1682972	0.1686139	0.1926907	
UBB	TUBA6	ACTB	CFL1	RPS13	SUI1
0.09656546	0.09674897	0.10753445	0.10830099	0.11801680	0.12612399
CLTC	NACA	FLJ20030	RPS23		
0.12773131	0.13422958	0.14609897	0.16530522		
RPS13	SUI1	TUBA6	NACA	FLJ20030	CFL1
0.09085973	0.09647829	0.09943424	0.10288912	0.11097074	0.11428399
ACTB	RPS23	CLTC			
0.11495336	0.12635109	0.13286210			
ACTB	TUBA6	CFL1	FLJ20030	NACA	CLTC
0.09215478	0.09499893	0.09674032	0.10528784	0.10718604	0.10879846
SUI1	RPS23				
0.11368091	0.13134766				
SUI1	NACA	FLJ20030	RPS23	TUBA6	CFL1
0.08281504	0.08444905	0.08922236	0.09072667	0.10559279	0.10993755
CLTC					
0.13142181					
NACA	CFL1	TUBA6	FLJ20030	CLTC	RPS23
0.08336046	0.08410148	0.09315528	0.09775742	0.10499056	0.10554332

CFL1	TUBA6	CLTC	FLJ20030	RPS23	
0.07222968	0.07722737	0.08440691	0.09831958	0.12735605	
FLJ20030	TUBA6	CLTC	RPS23		
0.08162006	0.08189011	0.10705192	0.11430674		
TUBA6	CLTC	RPS23			
0.06978897	0.08069582	0.13702726			
CLTC	RPS23				
0.1199009	0.1245241				
1	2	3	4	5	6
"UBC"	"GAPD"	"TPT1"	"UBB"	"RPS13"	"ACTB"
7	8	9	10	11	12
"SUI1"	"NACA"	"CFL1"	"FLJ20030"	"TUBA6"	"CLTC"

In case of the Bladder dataset the two top ranked genes are HSPCB and RPS13; see Figure 1 in And04.

```
> grade <- pData(Bladder)[,"Grade"]
> selectHKs(Bladder, group = grade, log = FALSE, trace = FALSE,
+           Symbols = featureNames(Bladder), minNrHKs = 13,
+           method = "NormFinder")$ranking
```

1	2	3	4	5	6
"HSPCB"	"RPS13"	"UBC"	"RPS23"	"ATP5B"	"TEGT"
7	8	9	10	11	12
"UBB"	"FLJ20030"	"CFL1"	"S100A6"	"FLOT2"	"ACTB"
13					
"TPT1"					

5 Normalization by means of reference/housekeeping genes

5.1 ΔCq method using a single housekeeper

The ΔCq method normalises detectors within a sample by subtracting the cycle time value of the housekeeper gene from the other genes. This can be done in NormqPCR as follows:

for the example dataset from "ReadqPCR" we must first read in the data:

```
> path <- system.file("exData", package = "NormqPCR")
> taqman.example <- file.path(path, "example.txt")
> qPCR.example <- file.path(path, "qPCR.example.txt")
> qPCRBatch.taqman <- read.taqman(taqman.example)
```

We then need to supply a housekeeper gene to be subtracted:

```
> hkg<-"Actb-Rn00667869_m1"
> qPCRBatch.norm <- deltaCq(qPCRBatch = qPCRBatch.taqman, hkg = hkg, calc="arith")
> head(exprs(qPCRBatch.norm))
```

	fp1.day3.v	fp2.day3.v	fp5.day3.mia
Actb.Rn00667869_m1	0.000000	0.000000	0.000000
Adipoq.Rn00595250_m1	0.016052	-0.116520	2.933523
Adrbk1.Rn00562822_m1	NA	NA	6.566628
Agtrl1.Rn00580252_s1	4.899380	5.035841	6.397364
Alpl.Rn00564931_m1	12.531942	11.808657	13.035166
B2m.Rn00560865_m1	0.741558	0.890717	2.040470

	fp6.day3.mia	fp.3.day.3.v	fp.4.day.3.v
Actb.Rn00667869_m1	0.000000	0.000000	0.000000
Adipoq.Rn00595250_m1	2.540987	-0.178971	-0.563263
Adrbk1.Rn00562822_m1	6.642561	NA	NA
Agtrl1.Rn00580252_s1	5.680837	5.220796	4.425364
Alpl.Rn00564931_m1	12.239549	12.394802	11.772896
B2m.Rn00560865_m1	2.234605	0.505516	0.877598

	fp.7.day.3.mia	fp.8.day.3.mia
Actb.Rn00667869_m1	0.000000	0.000000
Adipoq.Rn00595250_m1	2.458509	2.736475
Adrbk1.Rn00562822_m1	3.737100	6.873568
Agtrl1.Rn00580252_s1	4.794776	5.345202
Alpl.Rn00564931_m1	12.110000	12.255186
B2m.Rn00560865_m1	1.927563	1.903269

This returns a new `qPCRBatch`, with new values in the `exprs` slot. This will be compatible with many other bioconductor and R packages, such as `heatmap`.

Note these numbers might be negative. For further analysis requiring positive values only, $2^{\Delta C_T}$ can be used to transform the data into $2^{\Delta C_T}$ values.

5.2 ΔC_T method using a combination of housekeeping genes

If the user wishes to normalise by more than one housekeeping gene, for example if they have found a more than one housekeeping gene using the `NormFinder`/`geNorm` algorithms described above, they can. This is implemented by calculating the average of these values to form a "pseudo-housekeeper" which is subtracted from the other values. So using the same dataset as above, using housekeeping genes `GAPDH`, `Beta-2-microglobulin` and `Beta-actin`, the following steps would be taken:

```
> hkg<-c("Actb-Rn00667869_m1", "B2m-Rn00560865_m1", "Gapdh-Rn99999916_s1")
> qPCRBatch.norm <- deltaCq(qPCRBatch = qPCRBatch.taqman, hkg = hkg, calc="arith")
> head(exprs(qPCRBatch.norm))
```

	fp1.day3.v	fp2.day3.v	fp5.day3.mia
Actb.Rn00667869_m1	-1.2998917	-1.2816963	-1.380296
Adipoq.Rn00595250_m1	-1.2838397	-1.3982163	1.553227
Adrbk1.Rn00562822_m1	NA	NA	5.186332
Agtrl1.Rn00580252_s1	3.5994883	3.7541447	5.017068
Alpl.Rn00564931_m1	11.2320503	10.5269607	11.654870
B2m.Rn00560865_m1	-0.5583337	-0.3909793	0.660174

	fp6.day3.mia	fp.3.day.3.v	fp.4.day.3.v
Actb.Rn00667869_m1	-1.5106197	-1.1644617	-1.1714227
Adipoq.Rn00595250_m1	1.0303673	-1.3434327	-1.7346857
Adrbk1.Rn00562822_m1	5.1319413	NA	NA
Agtrl1.Rn00580252_s1	4.1702173	4.0563343	3.2539413
Alpl.Rn00564931_m1	10.7289293	11.2303403	10.6014733
B2m.Rn00560865_m1	0.7239853	-0.6589457	-0.2938247

	fp.7.day.3.mia	fp.8.day.3.mia
Actb.Rn00667869_m1	-1.323712	-1.286277
Adipoq.Rn00595250_m1	1.134797	1.450198
Adrbk1.Rn00562822_m1	2.413388	5.587291
Agtrl1.Rn00580252_s1	3.471064	4.058925
Alpl.Rn00564931_m1	10.786288	10.968909
B2m.Rn00560865_m1	0.603851	0.616992

5.3 $2^{-\Delta\Delta Cq}$ method using a single housekeeper

It is possible to use the $2^{-\Delta\Delta Cq}$ method for calculating relative gene expression between two sample types. Both the same well and the separate well methods as detailed in [2] can be used for this purpose, and will produce the same answers, but with different levels of variation. By default detectors in the same sample will be paired with the housekeeper, and the standard deviation used will be that of the differences between detectors and the housekeepers. However, if the argument `paired=FALSE` is added, standard deviation between case and control will be calculated as $s = \sqrt{s_1^2 + s_2^2}$, where s_1 is the standard deviation for the detector readings and s_2 is the standard deviation the housekeeper gene readings. The latter approach is not recommended when the housekeeper and genes to be compared are from the same sample, as is the case when using the taqman cards, but is included for completeness and for situations where readings for the housekeeper might be taken from a separate biological replicate (for example in a *post hoc* manner due to the originally designated housekeeping genes not performing well), or for when NormqPCR is used for more traditional qPCR where the products undergo amplifications from separate wells.

for the example dataset from "ReadqPCR" we must first read in the data:

```
> path <- system.file("exData", package = "NormqPCR")
```



```
> taqman.example <- file.path(path, "example.txt")
> qPCR.example <- file.path(path, "qPCR.example.txt")
> qPCRBatch.taqman <- read.taqman(taqman.example)
```

deltaDeltaCq also requires a contrast matrix. This is to contain columns which will be used to specify the samples representing `case` and `control` which are to be compared, in a similar way to the "limma" package. these columns should contain 1s or 0s which refer to the samples in either category:

```
> contM <- cbind(c(0,0,1,1,0,0,1,1),c(1,1,0,0,1,1,0,0))
> colnames(contM) <- c("interestingPhenotype","wildTypePhenotype")
> rownames(contM) <- sampleNames(qPCRBatch.taqman)
> contM
```

	interestingPhenotype	wildTypePhenotype
fp1.day3.v	0	1
fp2.day3.v	0	1
fp5.day3.mia	1	0
fp6.day3.mia	1	0
fp.3.day.3.v	0	1
fp.4.day.3.v	0	1
fp.7.day.3.mia	1	0
fp.8.day.3.mia	1	0

We can now normalise each sample by a given housekeeping gene and then look at the ratio of expression between the case and control samples. Results show (by column): 1) Name of gene represented by detector. 2) Case ΔCq for the detector: the average cycle time for this detector in the samples denoted as "case" - the housekeeper cycle time. 3) the standard deviation for the cycle times used to calculate the value in column 2). 4) Control ΔCq for the detector: the average cycle time for this detector in the samples denoted as "controller", or the "calibrator" samples - the housekeeper cycle time. 5) The standard deviation for the cycle times used to calculate the value in column 4). 6) $2^{-\Delta\Delta Cq}$ - The difference between the ΔCq values for case and control. We then find 2^{-} of this value. 7) and 8) correspond to 1 s.d. either side of the mean value, as detailed in [2].

```
> hkg <- "Actb-Rn00667869_m1"
> ddCq.taqman <- deltaDeltaCq(qPCRBatch = qPCRBatch.taqman, maxNACase=1, maxNAControl=1,
+                               hkg=hkg, contrastM=contM, case="interestingPhenotype",
+                               control="wildTypePhenotype", statCalc="geom", hkgCalc="arith")
> head(ddCq.taqman)
```

	ID	2 ⁻ -dCt.interestingPhenotype
1	Actb.Rn00667869_m1	1.000e+00

2	Adipoq.Rn00595250_m1	1.587e-01
3	Adrbk1.Rn00562822_m1	2.602e-02
4	Agtrl1.Rn00580252_s1	2.300e-02
5	Alpl.Rn00564931_m1	1.892e-04
6	B2m.Rn00560865_m1	2.464e-01
	interestingPhenotype.sd 2 ⁻ dCt.wildTypePhenotype	
1	0.000e+00	1.000e+00
2	2.280e-02	1.171e+00
3	3.266e-02	NA
4	1.014e-02	3.434e-02
5	4.770e-05	2.298e-04
6	2.498e-02	5.965e-01
	wildTypePhenotype.sd 2 ⁻ -ddCt 2 ⁻ -ddCt.min 2 ⁻ -ddCt.max	
1	0.000e+00 1	NA NA
2	2.131e-01 0.135541545192243	NA NA
3	NA +	NA NA
4	8.584e-03 0.669721905042939	NA NA
5	6.107e-05 0.823327272466571	NA NA
6	7.668e-02 0.413128242070071	NA NA

We can also average the taqman data using the separate samples/wells method . Here standard deviation is calculated separately and then combined, as described above. Therefore the pairing of housekeeper with the detector value within the same sample is lost. This can potentially increase variance.

```
> hkg <- "Actb-Rn00667869_m1"
> ddCqAvg.taqman <- deltaDeltaCq(qPCRBatch = qPCRBatch.taqman, maxNACase=1, maxNAControl=1,
+                               hkg=hkg, contrastM=contM, case="interestingPhenotype",
+                               control="wildTypePhenotype", paired=FALSE, statCalc="geom",
+                               hkgCalc="arith")
> head(ddCqAvg.taqman)
```

	ID	2 ⁻ -dCt.interestingPhenotype
1	Actb.Rn00667869_m1	1.000e+00
2	Adipoq.Rn00595250_m1	1.587e-01
3	Adrbk1.Rn00562822_m1	2.602e-02
4	Agtrl1.Rn00580252_s1	2.300e-02
5	Alpl.Rn00564931_m1	1.892e-04
6	B2m.Rn00560865_m1	2.464e-01
	interestingPhenotype.sd 2 ⁻ -dCt.wildTypePhenotype	
1	0.000e+00	1.000e+00
2	2.280e-02	1.171e+00

3	3.266e-02	NA
4	1.014e-02	3.434e-02
5	4.770e-05	2.298e-04
6	2.498e-02	5.965e-01
	wildTypePhenotype.sd	2 ^{-ddCt} 2 ^{-ddCt.min} 2 ^{-ddCt.max}
1	0.000e+00 1	NA NA
2	2.131e-01 0.135541545192243	NA NA
3	NA +	NA NA
4	8.584e-03 0.669721905042939	NA NA
5	6.107e-05 0.823327272466571	NA NA
6	7.668e-02 0.413128242070071	NA NA

5.4 $2^{\Delta\Delta Cq}$ method using a combination of housekeeping genes

If the user wishes to normalise by more than one housekeeping gene, for example if they have found a more than one housekeeping gene using the NormFinder/geNorm algorithms described above, they can. This is implemented by calculating the average of these values using the geometric mean to form a "pseudo-housekeeper" which is subtracted from the other values. For the dataset above, using housekeeping genes GAPDH, Beta-2-microglobulin and Beta-actin:

```
> qPCRBatch.taqman <- read.taqman(taqman.example)
> contM <- cbind(c(0,0,1,1,0,0,1,1),c(1,1,0,0,1,1,0,0))
> colnames(contM) <- c("interestingPhenotype","wildTypePhenotype")
> rownames(contM) <- sampleNames(qPCRBatch.taqman)
> hkg<-c("Actb-Rn00667869_m1", "B2m-Rn00560865_m1", "Gapdh-Rn99999916_s1")
> ddCq.gM.taqman <- deltaDeltaCq(qPCRBatch = qPCRBatch.taqman, maxNACase=1, maxNAControl=1,
+                               hkg=hkg, contrastM=contM, case="interestingPhenotype",
+                               control="wildTypePhenotype", statCalc="arith", hkgCalc="ari
> head(ddCq.gM.taqman)
```

	ID	2 ^{-dCt.interestingPhenotype}
1	Actb.Rn00667869_m1	2.594e+00
2	Adipoq.Rn00595250_m1	4.083e-01
3	Adrbk1.Rn00562822_m1	4.182e-02
4	Agtrl1.Rn00580252_s1	5.520e-02
5	Alpl.Rn00564931_m1	4.767e-04
6	B2m.Rn00560865_m1	6.367e-01
	interestingPhenotype.sd	2 ^{-dCt.wildTypePhenotype}
1	0.09819	2.345e+00
2	0.24929	2.713e+00
3	1.45844	NA

4	0.63719	7.878e-02
5	0.42589	5.242e-04
6	0.05413	1.390e+00
	wildTypePhenotype.sd	2 ⁻ -ddCt 2 ⁻ -ddCt.min 2 ⁻ -ddCt.max
1	0.071373 1.10638851325547	1.034e+00 1.184310
2	0.201905 0.150497255530234	1.266e-01 0.178884
3	NA +	NA NA
4	0.333840 0.700597907024805	4.505e-01 1.089636
5	0.386280 0.909381199520663	6.769e-01 1.221662
6	0.163975 0.457939394245865	4.411e-01 0.475448

There is also the option of using the mean housekeeper method using shared variance between the samples being compared, similar to the second `deltaDeltaCq` method shown above.

```
> qPCRBatch.taqman <- read.taqman(taqman.example)
> contM <- cbind(c(0,0,1,1,0,0,1,1),c(1,1,0,0,1,1,0,0))
> colnames(contM) <- c("interestingPhenotype","wildTypePhenotype")
> rownames(contM) <- sampleNames(qPCRBatch.taqman)
> hkg<-c("Actb-Rn00667869_m1", "B2m-Rn00560865_m1", "Gapdh-Rn99999916_s1")
> ddAvgCq.gM.taqman <-deltaDeltaCq(qPCRBatch = qPCRBatch.taqman, maxNACase=1, maxNAControl=1,
+                               hkg=hkg, contrastM=contM, case="interestingPhenotype",
+                               control="wildTypePhenotype", paired=FALSE, statCalc="arith",
+                               hkgCalc="arith")
> head(ddAvgCq.gM.taqman)
```

	ID	2 ⁻ -dCt.interestingPhenotype
1	Actb.Rn00667869_m1	2.594e+00
2	Adipoq.Rn00595250_m1	4.083e-01
3	Adrbk1.Rn00562822_m1	4.182e-02
4	Agtrl1.Rn00580252_s1	5.520e-02
5	Alpl.Rn00564931_m1	4.767e-04
6	B2m.Rn00560865_m1	6.367e-01
	interestingPhenotype.sd	2 ⁻ -dCt.wildTypePhenotype
1	0.3849	2.345e+00
2	0.4822	2.713e+00
3	1.4545	NA
4	0.6905	7.878e-02
5	0.5846	5.242e-04
6	0.2777	1.390e+00
	wildTypePhenotype.sd	2 ⁻ -ddCt 2 ⁻ -ddCt.min 2 ⁻ -ddCt.max
1	0.3574 1.10638851325547	8.473e-01 1.444684

2	0.2495	0.150497255530234	1.077e-01	0.210221
3	NA	+	NA	NA
4	0.2813	0.700597907024805	4.341e-01	1.130625
5	0.3689	0.909381199520663	6.064e-01	1.363762
6	0.4576	0.457939394245865	3.778e-01	0.555126

TO SHOW EXAMPLE USING GENORM/NORMFINDER DATA

5.5 Compute NRQs

THIS FUNCTION IS STILL EXPERIMENTAL!

We load a dataset including technical replicates.

```
> path <- system.file("exData", package = "ReadqPCR")
> qPCR.example <- file.path(path, "qPCR.example.txt")
> Cq.data <- read.qPCR(qPCR.example)
```

We combine the technical replicates and in addition compute standard deviations.

```
> Cq.data1 <- combineTechRepsWithSD(Cq.data)
```

We load efficiencies for the dataset and add them to the dataset.

```
> Effs <- file.path(path, "Efficiencies.txt")
> Cq.effs <- read.table(file = Effs, row.names = 1, header = TRUE)
> rownames(Cq.effs) <- featureNames(Cq.data1)
> effs(Cq.data1) <- as.matrix(Cq.effs[, "efficiency", drop = FALSE])
> se.effs(Cq.data1) <- as.matrix(Cq.effs[, "SD.efficiency", drop = FALSE])
```

Now we can compute normalized relative quantities for the dataset where we consider two of the included features as reference/housekeeping genes.

```
> res <- ComputeNRQs(Cq.data1, hkgs = c("gene_az", "gene_gx"))
> ## NRQs
> exprs(res)
```

	caseA	caseB	controlA	controlB
gene_ai	1.9253072	1.3586729	0.6479659	0.8749479
gene_az	1.0567118	1.1438982	1.0331980	0.9134997
gene_bc	1.1024935	0.7193500	0.7030487	1.2140836
gene_by	1.5102316	0.9573047	0.7527082	1.6008850
gene_dh	1.2982037	1.0722522	0.9623335	0.9392871
gene_dm	0.6590246	1.1690720	1.2475372	0.9366210
gene_dq	0.7541955	0.7036408	0.8327917	1.6165326

```

gene_dr 2.2192305 1.0581211 0.7026411 0.6900584
gene_eg 0.9366671 0.5800339 0.8313720 1.1848856
gene_er 0.5269062 0.9375427 0.6953326 2.3195978
gene_ev 1.4622280 2.3457021 0.9038912 1.1454535
gene_fr 1.4954763 1.6200792 0.9641192 0.7295680
gene_fw 0.6944248 0.8051075 1.5698382 0.7978611
gene_gx 0.9463318 0.8742037 0.9678687 1.0946911
gene_hl 1.0009372 1.4015267 0.7683665 0.7713712
gene_il 1.4632019 1.2595559 0.7216891 0.9318860
gene_iv 1.7263335 1.2275001 1.5464212 0.8881605
gene_jr 0.8984351 0.9834026 0.8754813 0.6637941
gene_jw 1.4655948 0.9340184 1.0505200 1.5504136
gene_qs 0.6730225 0.7610418 1.0665938 3.5329891
gene_qy 0.5287127 1.5722670 1.0615326 3.3252907
gene_rz 0.8690600 1.5588299 0.7287288 1.4812753
gene_sw 0.5975288 1.2406438 0.6982954 1.6007333
gene_vx 0.6942254 0.7168408 2.0253177 1.3190943
gene_xz 0.7668030 1.0218209 0.6136038 1.6729352

```

```

> ## SD of NRQs
> se.exprs(res)

```

```

           caseA      caseB  controlA  controlB
gene_ai 1.3996554 0.8787290 0.4855882 1.0034912
gene_az 0.6832730 0.7601966 0.8971054 0.6031927
gene_bc 0.7225348 0.4746146 0.9626570 1.0478385
gene_by 1.1522746 0.6116269 0.6088836 2.0409211
gene_dh 1.2483072 0.7889984 0.6165041 0.9767947
gene_dm 0.4711409 0.7780238 0.8476053 0.7294405
gene_dq 0.7023561 0.4849899 0.5813310 1.4067670
gene_dr 1.4407662 1.0804211 0.4543153 0.5149367
gene_eg 0.7355269 0.5497433 0.5588801 1.0938601
gene_er 0.4301195 0.6119514 0.4471454 1.5115897
gene_ev 1.0094209 2.4267114 0.6337126 0.7782519
gene_fr 1.6760391 1.1119157 0.6226081 0.5040967
gene_fw 0.5041070 0.9131565 1.1153268 0.7234551
gene_gx 0.6046042 0.9027816 0.6713914 1.7394961
gene_hl 0.7633174 0.9123997 1.0000329 0.5005813
gene_il 1.4621406 0.9540445 0.5678634 0.6067147
gene_iv 1.2668346 0.8039841 0.9995225 0.8171996
gene_jr 0.5749672 0.6786989 0.5595295 0.4405919

```

gene_jw 0.9626606 0.7890401 0.7194378 1.1512512
 gene_qs 0.5830335 0.5309990 0.6828952 2.8253799
 gene_qy 0.5947918 1.1294199 0.6794829 2.1713340
 gene_rz 0.5846751 1.7926435 0.4911506 2.1424580
 gene_sw 0.6284440 0.8062083 0.9638307 1.8398593
 gene_vx 0.5285361 1.0126959 1.3861226 0.8683886
 gene_xz 0.5231477 0.9270275 0.3972901 1.3643840

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