

Reanalysis Johnson2007 - figures and code

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The `sessionInfo()` is provided at the end of the document.

```
rm(list = ls())

library(GGally)
## Loading required package: ggplot2
## Registered S3 method overwritten by 'GGally':
##   method from
##   +.gg   ggplot2
library(ggplot2)
library(limma)
library(sva)
## Loading required package: mgcv
## Loading required package: nlme
## This is mgcv 1.8-31. For overview type 'help("mgcv-package")'.
## Loading required package: genefilter
## Loading required package: BiocParallel
library(qvalue)
library(randRotation)
```

File from <https://github.com/ous-uiو-bioinfo-core/batch-adjust-warning-figures/tree/master/reanalysis/Johnson2007/data> See also Nygaard, V., Rodland, E. A. & Hovig, E. Methods that remove batch effects while retaining group differences may lead to exaggerated confidence in downstream analyses. Biostatistics kxv027 (2015). doi:10.1093/biostatistics/kxv027

```
edata <- read.table(file = "dataExample2.txt", header = TRUE,
                     sep = "\t", dec = ".", stringsAsFactors = FALSE)
```

File from <https://github.com/ous-uiو-bioinfo-core/batch-adjust-warning-figures/tree/master/reanalysis/Johnson2007/data> See also Nygaard, V., Rodland, E. A. & Hovig, E. Methods that remove batch effects while retaining group differences may lead to exaggerated confidence in downstream analyses. Biostatistics kxv027 (2015). doi:10.1093/biostatistics/kxv027

```
pdata <- read.table(file = "sampleInfoExample2.txt", header = TRUE,
                     sep = "\t", dec = ".", stringsAsFactors = TRUE,
                     row.names = 1)

samps <- which(pdata$Type != "WT")

pdata <- pdata[samps,]
edata <- edata[,samps]
pdata$Type <- droplevels(pdata$Type)
pdata$Batch <- as.factor(pdata$Batch)
```

Reanalysis Johnson2007 - figures and code

```
all.equal(rownames(pdata), colnames(edata))
## [1] TRUE

# flooring to 1
edata[edata<1] <- 1
# take out data with too many low/missing values.
negativeprobesfilter <- rowSums(edata>1) >= (0.9*ncol(edata))
edata <- edata[negativeprobesfilter,]
# quantilenormalize
edata.quan <- normalizeBetweenArrays(log2(edata), method="quantile")

# for debugging
debug = FALSE
if(debug) edata <- edata[1:1000,]

##### ComBat - "p ComBat" values

mod.com <- model.matrix(~Type, pdata)
edata.com <- ComBat(edata.quan,
                     batch = pdata$Batch,
                     mod = mod.com)
## Found3batches
## Adjusting for covariate(s) or covariate level(s)
## Standardizing Data across genes
## Fitting L/S model and finding priors
## Finding parametric adjustments
## Adjusting the Data

mod.fit <- model.matrix(~Type, pdata)
fit1 <- lmFit(edata.com, design = mod.fit)
fit1 <- eBayes(fit1)

ps.com <- topTable(fit1, number = Inf, sort.by = "none")$P.Value
## Removing intercept from test coefficients
fdr.com <- topTable(fit1, number = Inf, sort.by = "none")$adj.P.Val
## Removing intercept from test coefficients

sum(fdr.com<0.05)
## [1] 649
sum(qvalue(ps.com)$qvalues < 0.05)
## [1] 814

##### limma batch as covariate - "p Limma (+batch)" values

mod.fit <- model.matrix(~Type + Batch, pdata)
fit2 <- lmFit(edata.quan, design = mod.fit)
fit2 <- eBayes(fit2)
```

Reanalysis Johnson2007 - figures and code

```
ps.lim <- topTable(fit2, coef = 2, number = Inf, sort.by = "none")$P.Value
fdr.lim <- topTable(fit2, coef = 2, number = Inf, sort.by = "none")$adj.P.Val

sum(fdr.lim<0.05)
## [1] 271
sum(qvalue(ps.lim)$qvalues < 0.05)
## [1] 392

##### ComBat with random rotations - "p ComBat - 2000 rot." values

mod.fit <- model.matrix(~Type, pdata)

rr1 <- initBatchRandrot(edata.quan, mod.fit, 2, pdata$Batch)
## Initialising batch "1"
## Initialising batch "2"
## Initialising batch "3"

statistic <- function(Y, batch, mod){
  edata.com <- sva::ComBat(Y,
                            batch = batch,
                            mod = mod, mean.only = FALSE)

  fit1 <- limma::lmFit(edata.com, design = mod)
  fit1 <- limma::eBayes(fit1)

  abs(limma::topTable(fit1, number = Inf, sort.by = "none")$t)
}

rs1 <- rotateStat(rr1, R = 2000, statistic = statistic, pdata$Batch,
                  mod.fit, parallel = TRUE)

ps.rot <- pFdr(rs1)
fdr.rot <- p.adjust(ps.rot, "BH")

sum(fdr.rot<0.05)
## [1] 431
sum(qvalue(ps.rot)$qvalues < 0.05)
## [1] 539

ps <- cbind(ps.com, ps.lim, ps.rot = ps.rot[,1])
colnames(ps) <- c("p ComBat", "p Limma (+batch)", "p ComBat - 2000 rot.")

apply(ps, 2, function(i)sum(qvalue::qvalue(i)$qvalues < 0.05))
##          p ComBat      p Limma (+batch) p ComBat - 2000 rot.
##                      814                 392                 539
apply(ps, 2, function(i)sum(p.adjust(i, "BH") < 0.05))
##          p ComBat      p Limma (+batch) p ComBat - 2000 rot.
##                      649                 271                 431
```

Reanalysis Johnson2007 - figures and code

```
#### p-vals scatterplot

df1 <- data.frame(ps)
colnames(df1) <- colnames(ps)

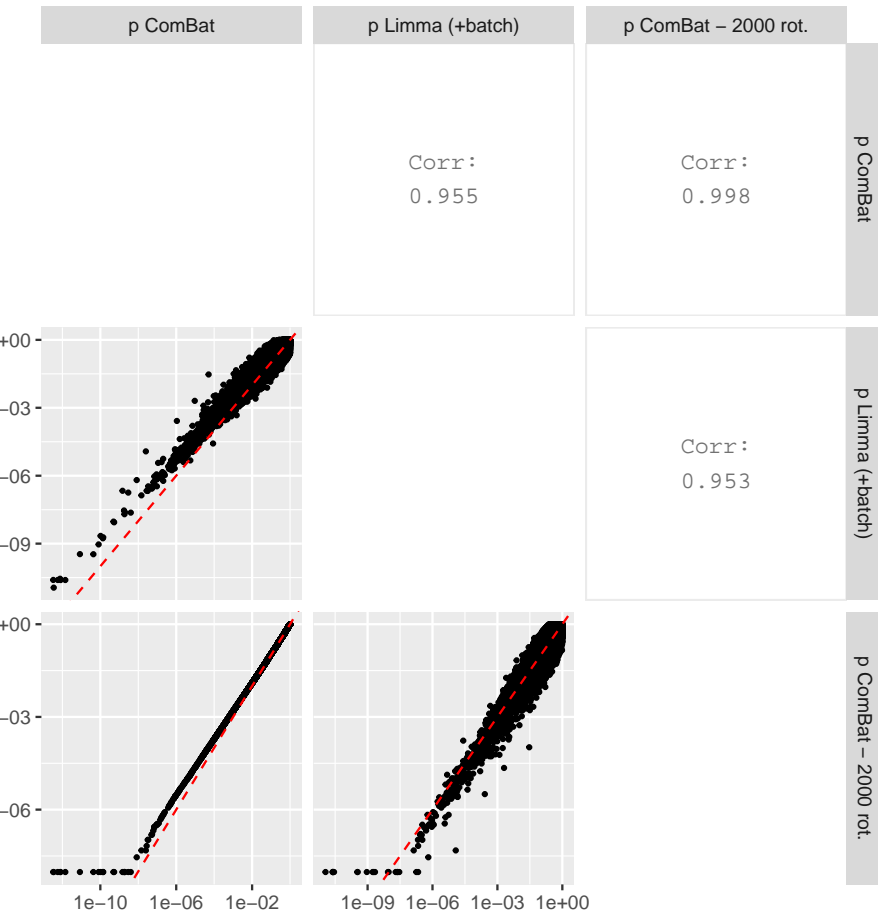
F1 <- function(...){
  ggally_points(..., size = 0.7) +
    scale_y_log10() +
    scale_x_log10() +
    geom_abline(slope = 1, intercept = 0, lty = 2, lwd = 0.5, col = "red")
}
lower.pan <- list(continuous = F1, combo = "facethist", discrete = "facetbar",
                   na = "na")

my.cor <- function(...)
  ggally_statistic(
    text_fn =
      function(x,y)formatC(cor(log(x),log(y)), digits = 3, format = "f"),
    title = "Corr",
    sep = ":\n",...)

upper.pan <- list(continuous = my.cor, combo = "box_no_facet",
                   discrete = "count", na = "na")

ggpairs(df1, lower = lower.pan, upper = upper.pan, diag = NULL)
```

Reanalysis Johnson2007 - figures and code

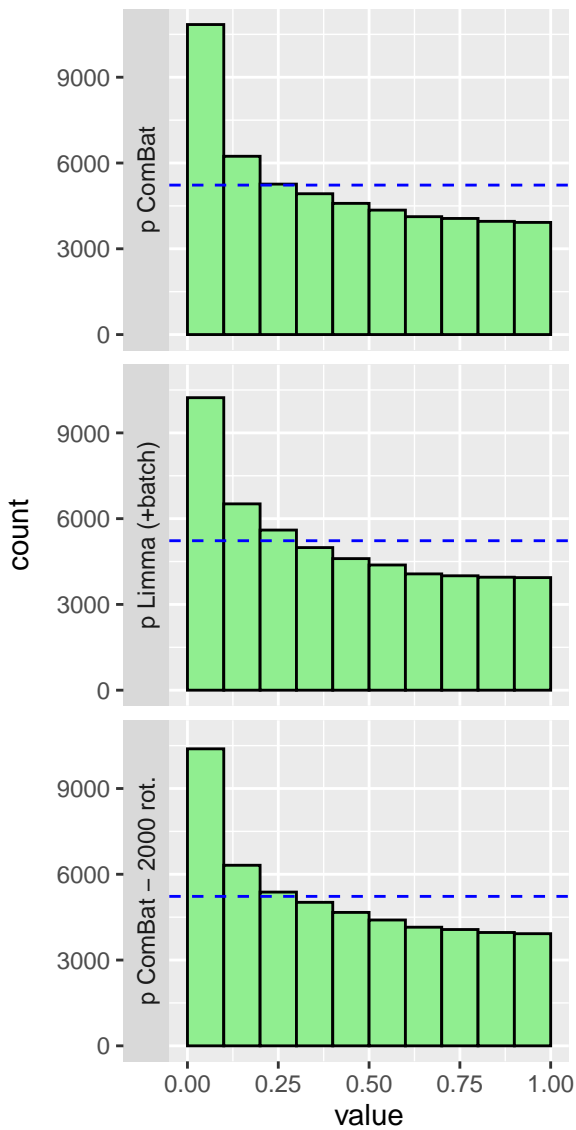


Histograms

```
df2 <- reshape2::melt(ps)

ggplot(df2, aes(x=value)) +
  geom_histogram(colour="black", fill="lightgreen", binwidth =0.1, boundary=0) +
  facet_grid(Var2 ~ ., switch = "y") +
  geom_abline(slope = 0, intercept = nrow(ps)/10, lty = 2, col = "blue") +
  theme(axis.title.y = element_text(vjust=+3.3))
```

Reanalysis Johnson2007 - figures and code



```
## Histograms 2

ind <- 1:15
h.com <- hist(ps.com, breaks = 100, plot = FALSE)
h.lim <- hist(ps.lim, breaks = 100, plot = FALSE)
h.rot <- hist(ps.rot, breaks = 100, plot = FALSE)

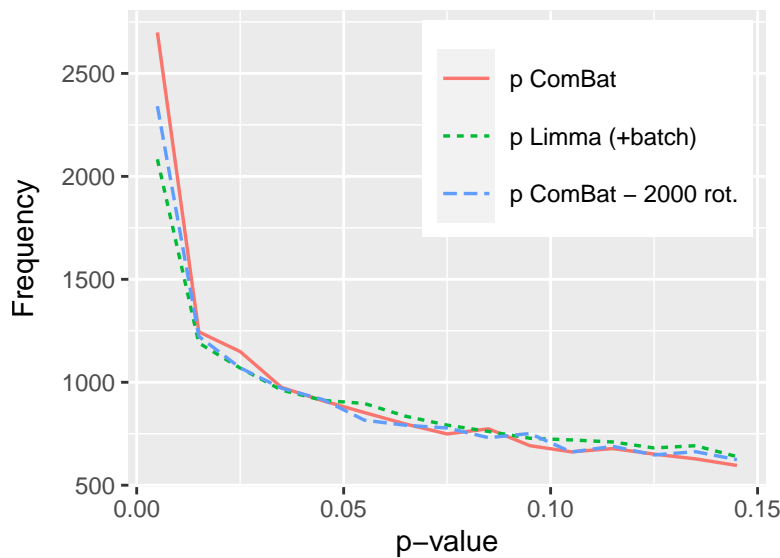
maxcount <- max(h.com$counts, h.lim$counts, h.rot$counts)

lab <- factor(rep(colnames(ps), rep(length(ind), 3)), levels = colnames(ps))

df1 <- data.frame(
  mids = c(h.com$mids[ind], h.lim$mids[ind], h.rot$mids[ind]),
  counts = c(h.com$counts[ind], h.lim$counts[ind], h.rot$counts[ind]),
  lab = lab)
```

Reanalysis Johnson2007 - figures and code

```
ggplot(df1, aes(x = mids, y = counts, colour = lab, lty = lab))+  
  geom_line(lwd = 0.6) +  
  xlab("p-value") + ylab("Frequency") +  
  theme(axis.title.y = element_text(vjust=+3.3)) +  
  theme(legend.justification=c(1,1), legend.position=c(0.98, 0.98),  
        legend.title = element_blank()) +  
  theme(legend.key.size = unit(1.5,"line"))
```



1 Session Info

```
sessionInfo()  
## R version 4.0.2 (2020-06-22)  
## Platform: x86_64-w64-mingw32/x64 (64-bit)  
## Running under: Windows 10 x64 (build 19041)  
##  
## Matrix products: default  
##  
## locale:  
## [1] LC_COLLATE=German_Austria.1252 LC_CTYPE=German_Austria.1252  
## [3] LC_MONETARY=German_Austria.1252 LC_NUMERIC=C  
## [5] LC_TIME=German_Austria.1252  
##  
## attached base packages:  
## [1] stats      graphics   grDevices    utils      datasets    methods     base  
##  
## other attached packages:  
## [1] randRotation_1.1.4    qvalue_2.21.0      sva_3.37.0  
## [4] BiocParallel_1.23.2   genefilter_1.71.0   mgcv_1.8-31  
## [7] nlme_3.1-148         limma_3.45.13    GGally_2.0.0  
## [10] ggplot2_3.3.2       BiocStyle_2.17.1
```

Reanalysis Johnson2007 - figures and code

```
##  
## loaded via a namespace (and not attached):  
## [1] Biobase_2.49.1      edgeR_3.31.4      bit64_4.0.5  
## [4] splines_4.0.2       Rdpack_1.0.0      BiocManager_1.30.10  
## [7] stats4_4.0.2        blob_1.2.1       yaml_2.2.1  
## [10] pillar_1.4.6       RSQLite_2.2.0     lattice_0.20-41  
## [13] glue_1.4.2         digest_0.6.25    RColorBrewer_1.1-2  
## [16] colorspace_1.4-1   htmltools_0.5.0   Matrix_1.2-18  
## [19] plyr_1.8.6         XML_3.99-0.5    pkgconfig_2.0.3  
## [22] bibtex_0.4.2.3    bookdown_0.20    purrr_0.3.4  
## [25] xtable_1.8-4      scales_1.1.1     snow_0.4-3  
## [28] tibble_3.0.3       annotate_1.67.1  generics_0.0.2  
## [31] farver_2.0.3       IRanges_2.23.10  ellipsis_0.3.1  
## [34] withr_2.3.0        BiocGenerics_0.35.4 survival_3.1-12  
## [37] magrittr_1.5        crayon_1.3.4     memoise_1.1.0  
## [40] evaluate_0.14      tools_4.0.2      gRd_0.4-11  
## [43] lifecycle_0.2.0    matrixStats_0.56.0 stringr_1.4.0  
## [46] S4Vectors_0.27.12  munsell_0.5.0    locfit_1.5-9.4  
## [49] AnnotationDbi_1.51.3 compiler_4.0.2   rlang_0.4.7  
## [52] grid_4.0.2         RCurl_1.98-1.2  bitops_1.0-6  
## [55] labeling_0.3       rmarkdown_2.3    gtable_0.3.0  
## [58] DBI_1.1.0          reshape_0.8.8   reshape2_1.4.4  
## [61] R6_2.4.1           knitr_1.30     dplyr_1.0.2  
## [64] bit_4.0.4          stringi_1.4.6  parallel_4.0.2  
## [67] Rcpp_1.0.5         vctrs_0.3.4    tidyselect_1.1.0  
## [70] xfun_0.16
```