# Re-analysis of JQ1 lysate dataset by Savitski et al, 2018

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#### **Package**

TPP2D 1.3.10

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# 1 Step-by-step walk through the anlysis

```
# This script uses the development version of TPP2D
if(require("BiocManager"))
   install.packages("BiocManager")
BiocManager::install("nkurzaw/TPP2D")
```

#### Load required libraries

```
library(TPP2D)
## Loading required package: dplyr
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
## filter, lag
## The following objects are masked from 'package:base':
##
## intersect, setdiff, setequal, union
library(dplyr)
library(tidyr)
library(ggplot2)
library(readxl)
```

#### Define plot style

```
theme_paper <- theme_bw(base_size = 6) +
    theme(legend.background = element_blank(),
        legend.key = element_blank(),
        panel.background = element_blank(),
        panel.grid.major = element_line(colour = "grey92", size = 0.25),
        panel.grid.minor = element_line(colour = "grey92", size = 0.15),
        panel.border = element_blank(),
        strip.background = element_blank(),
        plot.background = element_blank(),
        complete = TRUE,
        axis.line = element_line(color = "black", size = 0.25),
        text = element_text(size = 7),
        axis.ticks = element_line(color = "black", size = 0.25),
        axis.title = element_text(size = 8),
        axis.text = element_text(size = 6))</pre>
```

Download the supplementary excel table (Supplementary Dataset S1) by Savitski et al. (2018)

```
if(!file.exists("Savitski_et_al_Figure_3/Supplementary Dataset 2_2D-TPP.xlsx")){
    download.file(
        url = "https://data.mendeley.com/datasets/8pzhg2tdyb/1/files/115f60c9-01d1-4213-9abb-aa095d70a626/Savidestfile = "Savitski_et_al_Figure_3.zip")
    unzip(zipfile = "Savitski_et_al_Figure_3.zip",
        exdir = "Savitski_et_al_Figure_3")
    system("rm Savitski_et_al_Figure_3.zip")
```

```
}
```

Read in the data and reformat to a data frame as would be obtained after import of the raw data:

```
jq1_lys_raw <- read_xlsx("Savitski_et_al_Figure_3/Supplementary Dataset 2_2D-TPP.xlsx", sheet = 3, skip = 1)</pre>
 dplyr::select(representative = `Accession No.`,
                clustername = `protein name`,
                qupm = QUPM,
                qusm = QUSM,
                temperature,
                matches("sumionarea"),
                -matches("total"),
                matches("rel_fc_protein"),
                -matches("transformed"),
                -matches("orig")) %>%
 gather(key, value, matches("sumionarea"), matches("rel_fc_protein")) %>%
 mutate(conc = as.numeric(gsub("uM", "", gsub(".+_protein_[0-9,H,L]+_[0-9,H,L]+_", "", key))),
         temperature = as.numeric(gsub("C", "", temperature)),
         key = case_when(grepl("sumionarea", key) ~ "raw_value",
                         grepl("rel_fc", key) ~ "rel_value")) %>%
 spread(key, value) %>%
 arrange(representative, temperature, conc) %>%
 group_by(clustername, temperature, conc) %>%
  filter(qupm == max(qupm),
         qusm == max(qusm),
         raw_value == max(raw_value)) %>%
  filter(!duplicated(clustername)) %>%
 ungroup %>%
 mutate(log2_value = log2(raw_value),
         log_conc = log10(conc/le6)) %>%
  filter(qupm > 1)
# resolve ambiguous protein names
jq1_lys_fil <- resolveAmbiguousProteinNames(jq1_lys_raw)</pre>
# recompute reporter ion signal from robust Isobarquant fold changes
jq1_lys_df <- recomputeSignalFromRatios(jq1_lys_fil)</pre>
```

Compute null and alternative model fits and extract parameters

```
jq1_params_df <- getModelParamsDf(jq1_lys_df, maxit = 500)
saveRDS(jq1_params_df, file = "../pre_run_data/jq1_params_df.rds")</pre>
```

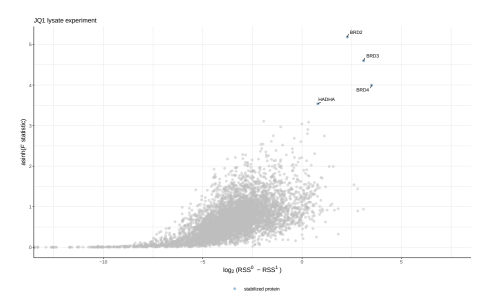
Compute F statistics

```
jq1_fstat_df <- computeFStatFromParams(jq1_params_df)</pre>
```

Get B datasets expected under the null model and perform model fitting and compute F statistics to obtain a null distribution for FDR calibration:

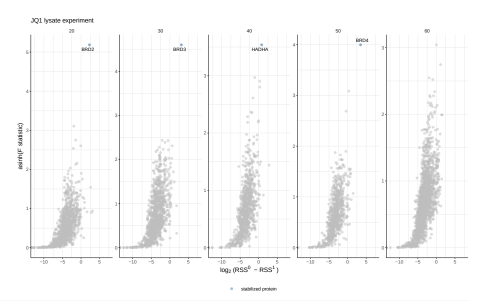
```
set.seed(12, kind = "L'Ecuyer-CMRG")
jq1_null_df <- bootstrapNullAlternativeModel(</pre>
  df = jq1_lys_df, params_df = jq1_params_df,
 maxit = 500, B = 100,
  BPPARAM = BiocParallel::MulticoreParam(workers = 20, progressbar = TRUE),
  verbose = FALSE)
saveRDS(jq1_null_df, file = "../pre_run_data/jq1_null_df.rds")
Compute FDR and find hits:
jq1_fdr_df <- getFDR(df_out = jq1_fstat_df,</pre>
                     df_null = jq1_null_df
jq1_hits_df <- findHits(jq1_fdr_df, alpha = 0.1)</pre>
ggplot(jq1_fdr_df %>%
           filter(dataset == "true") %>%
           mutate(group = case_when(slopeH1 > 0 ~ "stabilized protein",
                                     slopeH1 < 0 ~ "destabilized protein")),</pre>
       aes(log2(rssH0 - rssH1), asinh(F_statistic))) +
  geom_point(color = "gray", alpha = 0.5, size = 1) +
  geom_point(aes(color = group), alpha = 0.5,
             size = 1,
             data = jql_hits_df %>%
                 mutate(group = case_when(
                     slopeH1 > 0 ~ "stabilized protein",
                     slopeH1 < 0 ~ "destabilized protein"))) +</pre>
  ggrepel::geom_text_repel(
    aes(label = clustername),
    data = jq1_hits_df,
    size = 2, segment.size = 0.2, min.segment.length = unit(2, "pt")) +
  scale_color_manual("", values = c("steelblue", "orange")) +
  labs(x = expression('log'[2]~'(RSS'^0~' - RSS'^1~')'),
       y = expression('asinh('*italic(F)*' statistic)')) +
  ggtitle("JQ1 lysate experiment") +
  coord_cartesian(xlim = c(-12.5, 7.5)) +
  theme_paper +
  theme(legend.position = "bottom")
```

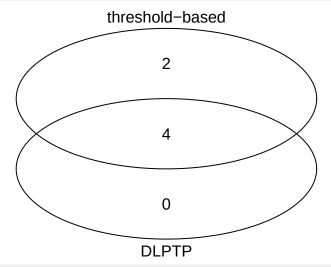
#### Re-analysis of JQ1 lysate dataset by Savitski et al, 2018



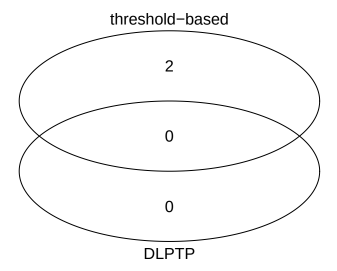
```
ggplot(jq1_fdr_df %>%
           filter(dataset == "true") %>%
           mutate(group = case_when(slopeH1 > 0 ~ "stabilized protein",
                                    slopeH1 < 0 ~ "destabilized protein")),</pre>
       aes(log2(rssH0 - rssH1), asinh(F_statistic))) +
 geom_point(color = "gray", alpha = 0.5, size = 1) +
 geom_point(aes(color = group), alpha = 0.5,
             size = 1,
             data = jq1_hits_df %>%
                 mutate(group = case_when(
                     slopeH1 > 0 \sim "stabilized protein",
                     slopeH1 < 0 ~ "destabilized protein"))) +</pre>
 ggrepel::geom_text_repel(
    aes(label = clustername),
    data = jq1_hits_df,
    size = 2, segment.size = 0.2, min.segment.length = unit(2, "pt")) +
 scale_color_manual("", values = c("steelblue", "orange")) +
  facet_wrap(~n0bsRound, scales = "free", ncol = 5) +
 labs(x = expression('log'[2]~'(RSS'^0~' - RSS'^1~')'),
       y = expression('asinh('*italic(F)*' statistic)')) +
 ggtitle("JQ1 lysate experiment") +
 coord\_cartesian(xlim = c(-12.5, 7.5)) +
  theme_paper +
  theme(legend.position = "bottom")
```

#### Re-analysis of JQ1 lysate dataset by Savitski et al, 2018

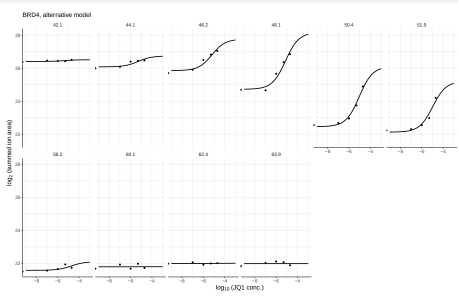




### Re-analysis of JQ1 lysate dataset by Savitski et al, 2018



#### BRD4 profile



```
sessionInfo()
## R version 3.6.1 (2019-07-05)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: macOS Mojave 10.14.6
## Matrix products: default
          /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRlapack.dylib
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## attached base packages:
                 graphics grDevices utils
## [1] stats
                                               datasets methods
                                                                   base
## other attached packages:
## [1] gplots_3.0.1.2
                        readxl_1.3.1
                                         ggplot2_3.2.1
                                                          tidyr_1.0.0
## [5] TPP2D_1.3.10
                        dplyr_0.8.3
                                         BiocStyle_2.12.0
## loaded via a namespace (and not attached):
## [1] gtools_3.8.1
                            tidyselect_0.2.5
                                                xfun_0.10
## [4] purrr_0.3.3
                            colorspace_1.4-1
                                                vctrs_0.2.0
## [7] htmltools_0.4.0
                            yaml_2.2.0
                                                rlang_0.4.1
## [10] pillar_1.4.2
                            glue_1.3.1
                                                withr_2.1.2
## [13] BiocParallel_1.18.1 foreach_1.4.7
                                                lifecycle_0.1.0
## [16] stringr_1.4.0
                            munsell_0.5.0
                                                gtable_0.3.0
## [19] cellranger_1.1.0
                            zip_2.0.4
                                                caTools_1.17.1.2
## [22] codetools_0.2-16
                            evaluate_0.14
                                                labeling_0.3
## [25] knitr_1.25
                            doParallel_1.0.15
                                                parallel_3.6.1
## [28] Rcpp_1.0.2
                            KernSmooth_2.23-16 scales_1.0.0
## [31] backports_1.1.5
                            BiocManager_1.30.9 gdata_2.18.0
## [34] digest_0.6.22
                            stringi_1.4.3
                                                openxlsx_4.1.0.1
## [37] bookdown_0.14
                            ggrepel_0.8.1
                                                qrid_3.6.1
## [40] tools_3.6.1
                            bitops_1.0-6
                                                magrittr_1.5
## [43] lazyeval_0.2.2
                            RCurl_1.95-4.12
                                                tibble_2.1.3
## [46] crayon_1.3.4
                            pkgconfig_2.0.3
                                                zeallot_0.1.0
## [49] MASS_7.3-51.4
                            ellipsis_0.3.0
                                                assertthat_0.2.1
## [52] rmarkdown_1.16
                                                R6_2.4.0
                            iterators_1.0.12
## [55] compiler_3.6.1
```

### References

Savitski, M.M., Zinn, N., Faelth-Savitski, M., Poeckel, D., Gade, S., Becher, I., Muelbaier, M., Wagner, A.J., Strohmer, K., Werner, T., et al. (2018). Multiplexed Proteome Dynamics Profiling Reveals Mechanisms Controlling Protein Homeostasis. Cell 1–15.