Relative Abundance Normalization Method Comparison

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Comparison of relative abundance error rate for different normalization methods

library(tidyverse)

```
## Loading tidyverse: ggplot2
## Loading tidyverse: tibble
## Loading tidyverse: tidyr
## Loading tidyverse: readr
## Loading tidyverse: purrr
## Loading tidyverse: dplyr
## Conflicts with tidy packages ------
## filter(): dplyr, stats
## lag():
            dplyr, stats
library(ggridges)
norm_count_df <- readRDS("~/Desktop/norm_count_df.RDS")</pre>
Mean variance relationship by normalization method - not sure if the difference is due to scaling or normal-
ization method.
filtered_norm <- norm_count_df %>%
    filter(mean_count != 0, var_count > 1e-10)
ggplot(filtered_norm) +
    geom_hex(aes(x = mean_count, y = var_count)) +
   geom_smooth(aes(x = mean_count, y = var_count)) +
   geom abline(aes(intercept = 0, slope = 1), color = "darkorange") +
   facet wrap(~norm method) +
   theme_bw() + scale_y_log10() + scale_x_log10() +
   labs(x = "Mean", y = "Variance") +
    # coord_equal() +
   theme(legend.position = "bottom", axis.text.x = element_text(angle = 315))
## `geom_smooth()` using method = 'gam'
Calculating Error Rate
pa summary anno df <- readRDS("~/Desktop/to file/mgtst RDS/pa summary anno df.RDS")
theta_est <- readRDS("~/Desktop/to_file/mgtst_RDS/bootstrap_theta_estimates.rds")</pre>
pre_post_prop <- norm_count_df %>%
      ungroup() %>%
     filter(t_fctr %in% c(0,20)) %>%
     mutate(end_point = if_else(t_fctr == 0 , "post", "pre")) %>%
      select(-t_fctr, -var_count) %>%
      ## setting values to 0 when one or more of the PCR replicates are 0 for titration end-points
      spread(end_point,mean_count, fill = 0)
```

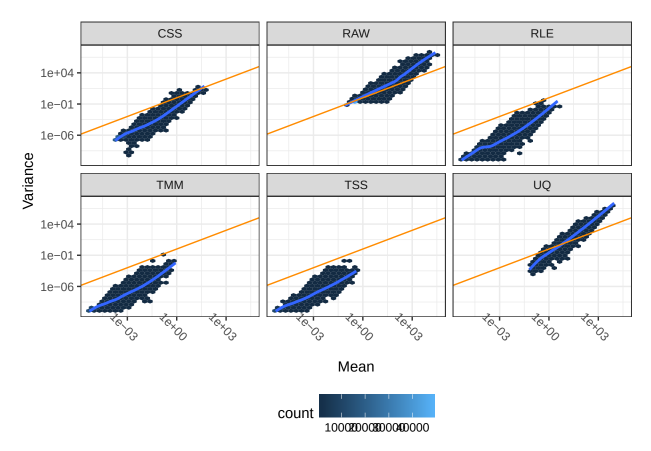
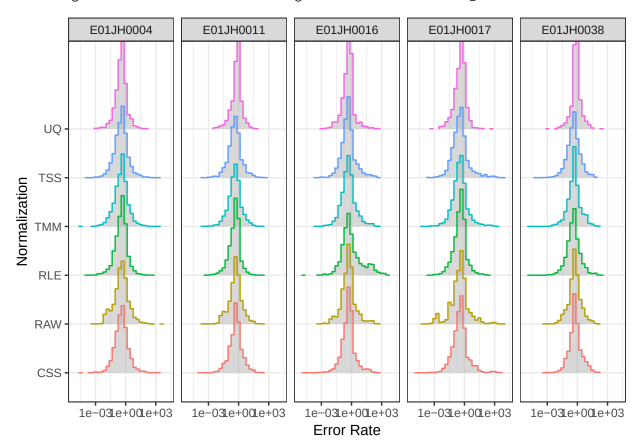


Figure 1: Comparison of relative abundance mean and variance relationship for PCR replicates across normalization methods. RLE - relative log expression, TMM - weighted trim mean of M-values, RAW - unnormalized, CSS - cumulative sum scaling, TSS - total sum scaling, UQ - upper quartile. Points with means of zero and variance < 1e-10 were excluded from the plot

```
prop_inferred <- theta_est %>%
      filter(pipe == "unclustered") %>%
      ungroup() %>%
      mutate(t fctr = factor(t fctr, levels = c(0:5, 10, 15, 20))) %>%
      select(biosample_id, theta_hat_mean, t_fctr) %>%
      right_join(norm_count_df) %>%
   right_join(pre_post_prop) %>%
      filter(t fctr %in% c(1:5,10,15)) %>%
      ## Using inferred theta estimates to calculate expected values
      mutate(inferred_prop = post * theta_hat_mean + pre * (1 - theta_hat_mean))
## Joining, by = c("biosample_id", "t_fctr")
## Warning: Column `t_fctr` joining factors with different levels, coercing to
## character vector
## Joining, by = c("biosample_id", "norm_method", "feature_id", "pipe")
## Excluding mix and unmix specific features
## Only including features observed in all or none of the four pre- post- PCR replicates
## Features with relative abundance estimates less than 1e-7, these are features that we would not expe
pa_filter <- pa_summary_anno_df %>%
      filter(pa_specific == "unspecific") %>%
     select(biosample_id, pipe, feature_id, full_pre, T00, T20, pa_mixed) %>%
      filter(T00 %in% c(0,4), T20 %in% c(04))
# prop_inferred <- prop_inferred %>%
       right_join(pa_filter) %>%
       filter(nb_prop > 1e-7)
#### Error Rate Calculations
rel_abu_error <- prop_inferred %>%
      mutate(t_fctr = factor(t_fctr, levels = c(1:5, 10, 15))) %>%
      mutate(inferred_error = abs(mean_count - inferred_prop),
             inferred_error_rate = inferred_error/inferred_prop)
rel_abu_ridge_df <- rel_abu_error %>%
    mutate(inferred_error_rate = if_else(inferred_error_rate < 1e-10, 0, inferred_error_rate)) %>%
    filter(inferred_error_rate != 0 & mean_count > 1e-10) %>%
    mutate(inferred_error_rate = if_else(inferred_prop == 0, NaN, inferred_error_rate))
rel_abu_med <- rel_abu_ridge_df %>%
      group_by(biosample_id, norm_method) %>%
      summarise(med_error = median( inferred_error_rate,na.rm = TRUE))
rel_abu_ridge_df %>%
      ggplot() +
      geom_density_ridges(aes(x = inferred_error_rate, y = norm_method, color = norm_method),
                          alpha = 0.5, stat = "binline", bins = 30, draw_baseline = FALSE)
      # geom_text(data = rel_abu_med,
                 aes(x = 100, y = norm\_method, label = round(med\_error, 2)), nudge_y = 0.1) +
      facet_wrap(~biosample_id, nrow = 1) + theme_bw() +
    scale_x_log10() +
      labs(x = "Error Rate", y = "Normalization", color = "Normalization") +
```

```
theme(legend.position = "none")
```

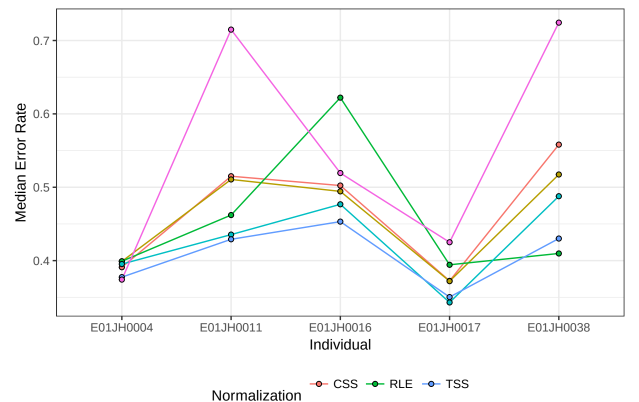
Warning: Removed 189663 rows containing non-finite values (stat_binline).



rel_abu_med %>% spread(norm_method, med_error) %>% knitr::kable(digits = 3)

| biosample_id | CSS | RAW | RLE | TMM | TSS | UQ |
|--------------|-------|-------|-------|-------|-------|-------|
| E01JH0004 | 0.391 | 0.399 | 0.399 | 0.395 | 0.378 | 0.374 |
| E01JH0011 | 0.515 | 0.511 | 0.462 | 0.435 | 0.429 | 0.715 |
| E01JH0016 | 0.502 | 0.494 | 0.622 | 0.477 | 0.453 | 0.519 |
| E01JH0017 | 0.372 | 0.372 | 0.394 | 0.343 | 0.350 | 0.425 |
| E01JH0038 | 0.558 | 0.517 | 0.410 | 0.488 | 0.430 | 0.724 |

```
rel_abu_med %>% ungroup() %>%
    mutate(biosample_id = factor(biosample_id)) %>%
    ggplot(aes(x = biosample_id, y = med_error)) +
    geom_blank() +
    geom_path(aes(x = as.numeric(biosample_id), y = med_error, color = norm_method)) +
    geom_point(aes(x = biosample_id, y = med_error, fill = norm_method), shape = 21) +
    theme_bw() +
    labs(x = "Individual", y = "Median Error Rate", fill = "Normalization", color = "Normalization") +
    theme(legend.position = "bottom")
```



```
fit <- aov(med_error ~ norm_method + biosample_id, data = rel_abu_med)
```

Significant difference between normalization methods when including biosample in the model. Need to used a mixed effects model to account for

summary(fit)

```
## Df Sum Sq Mean Sq F value Pr(>F)

## norm_method 5 0.06089 0.01218 2.956 0.037128 *

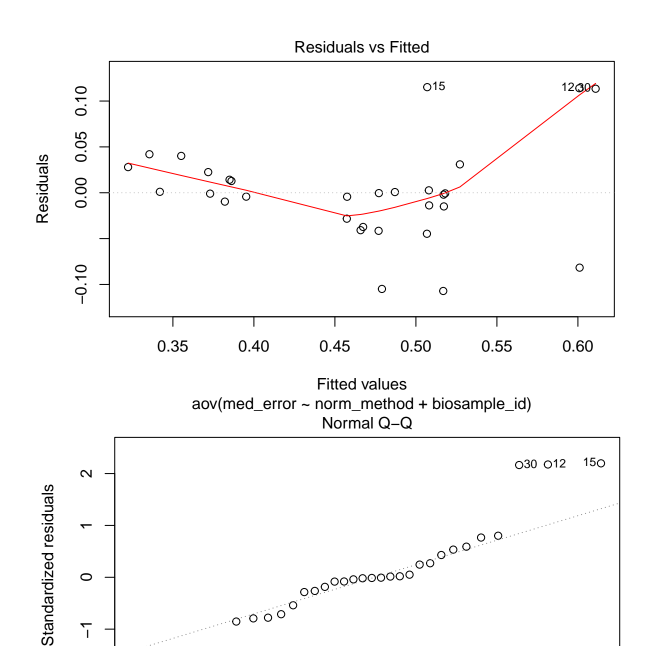
## biosample_id 4 0.12607 0.03152 7.650 0.000656 ***

## Residuals 20 0.08240 0.00412

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

plot(fit)
```

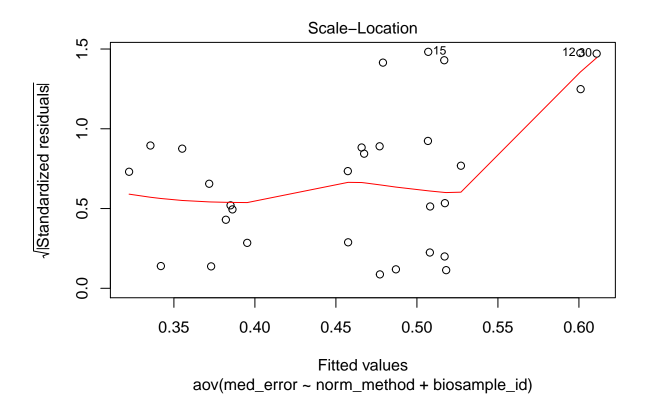


Theoretical Quantiles aov(med_error ~ norm_method + biosample_id)

hat values (leverages) are all = 0.3333333
and there are no factor predictors; no plot no. 5

-1

-2



Only two pairs normalization methods are significantly different from each other. Upper quartile is significantly different from TMM and TSS.

TukeyHSD(fit)

```
##
     Tukey multiple comparisons of means
##
       95% family-wise confidence level
##
## Fit: aov(formula = med_error ~ norm_method + biosample_id, data = rel_abu_med)
##
## $norm_method
##
                   diff
                                 lwr
                                            upr
                                                    p adj
## RAW-CSS -0.009103820 -0.136707088 0.11849945 0.9999085
## RLE-CSS -0.010287103 -0.137890371 0.11731616 0.9998331
## TMM-CSS -0.040116754 -0.167720023 0.08748651 0.9163956
## TSS-CSS -0.059738967 -0.187342236 0.06786430 0.6850444
## UQ-CSS
            0.083786623 -0.043816646 0.21138989 0.3439791
## RLE-RAW -0.001183283 -0.128786551 0.12641999 1.0000000
## TMM-RAW -0.031012934 -0.158616202 0.09659033 0.9705377
## TSS-RAW -0.050635147 -0.178238415 0.07696812 0.8087322
           0.092890443 -0.034712826 0.22049371 0.2442483
## UQ-RAW
## TMM-RLE -0.029829651 -0.157432919 0.09777362 0.9750577
## TSS-RLE -0.049451864 -0.177055132 0.07815140 0.8230580
            0.094073726 -0.033529542 0.22167699 0.2329747
## UQ-RLE
## TSS-TMM -0.019622213 -0.147225481 0.10798106 0.9962439
## UQ-TMM
            0.123903377 -0.003699891 0.25150665 0.0601562
## UQ-TSS
           0.143525590 \quad 0.015922322 \quad 0.27112886 \quad 0.0219373
##
## $biosample_id
##
                                diff
                                                         upr
                                                                 p adi
## E01JH0011-E01JH0004 0.1218466191 0.01095283 0.23274041 0.0269479
## E01JH0016-E01JH0004 0.1220207352 0.01112694
                                                  0.23291453 0.0266792
## E01JH0017-E01JH0004 -0.0131667180 -0.12406051 0.09772708 0.9962975
## E01JH0038-E01JH0004 0.1319069014 0.02101311 0.24280070 0.0149831
## E01JH0016-E01JH0011 0.0001741161 -0.11071968 0.11106791 1.0000000
## E01JH0017-E01JH0011 -0.1350133371 -0.24590713 -0.02411954 0.0124641
## E01JH0038-E01JH0011 0.0100602823 -0.10083351 0.12095408 0.9987010
## E01JH0017-E01JH0016 -0.1351874532 -0.24608125 -0.02429366 0.0123357
## E01JH0038-E01JH0016 0.0098861662 -0.10100763 0.12077996 0.9987869
## E01JH0038-E01JH0017 0.1450736194 0.03417983 0.25596741 0.0068211
```