Variance All the Way Down: Quantifying the Uncertainty Introduced at each Stage of an End-to-End RNA-Seq Analysis

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Abstract

In the realm of RNA-Seq research, rigorous data preprocessing is a critical foundation for meaningful analysis. Despite its importance, this preprocessing involves numerous stages, each introducing potential sources of variance. While previous studies have examined the overall variance across entire RNA-Seq pipelines, (Arora et al. 2020) (Tong et al. 2020), (Vieth et al. 2019), the impact of individual stages remains less understood. We propose a comprehensive investigation into the variance introduced at each stage of RNA-Seq preprocessing. Our goal is to quantify these variances, study their distributions, and understand their statistical implications on downstream modeling. This will include exploring the multitude of decisions researchers face — from quality control to normalization and feature selection — and evaluating how these choices propagate uncertainty through the analysis. Of particular interest is whether variance amplifies due to interactions between decisions made at different stages. By modeling these interactions, we aim to identify cases where suboptimal combinations of preprocessing choices exacerbate variability, potentially distorting biological interpretations. Finally, we will assess various bias correction methods and uncertainty quantification strategies to incorporate into final models. This work aims to provide researchers with actionable insights and robust statistical tools to mitigate preprocessing-induced variance, ultimately enhancing the reliability and reproducibility of RNA-Seq studies.

Preliminary Results

Preliminary Methodology Section

Table 1: Basic RNA-Seq Differential Analysis End-to-End Pipeline

Pipeline Steps	Software	Options	Choices
1. Pull SRA data from the NIH.	prefetch	NA	NA
2. Compute quality scores.	fasterq-dump	$\begin{array}{l} -\mathrm{skip\text{-}technical} \\ -\mathrm{threads} \ X \end{array}$	Boolean Integer
3. Filter low quality reads.	fastp	$\begin{array}{l} -\text{qualified}_\text{quality}_\text{phred X} \\ -\text{length}_\text{required X} \end{array}$	Integer Integer
4. Trim excess bases.	fastp	-trim_poly_g -trim_ploy_x	Boolean Boolean
5. Align and count genes.	Various	Default	Salmon, Kallisto
6. Count normalization.	edgeR	calcNormFactors(method='X')	TMM, RLE, upperquartile

Statistical Model

Assume there are n samples of g gene counts. Let B_g denote the count for gene g report to the NIH database, and let C_{gX} denote the count obtained from pipeline with choices X. Similar let D_g and E_{gX} denote the p-values obtained from edgeR. Now,

$$Y_{1X} = \frac{1}{g} \sum_{i=1}^{g} (C_{gX} - B_g)^2 \tag{1}$$

and

$$Y_{2X} = \frac{1}{g} \sum_{i=1}^{g} (E_{gX} - D_g)^2 \tag{2}$$

Our primary analysis will focus on the two following regression models:

$$Y_{1X_{i}} = \beta_{0} + \sum_{i=1}^{p} \beta_{i} X_{i} + \sum_{1 \leq i < j \leq p} \beta_{ij} (X_{i} \times X_{j}) + \epsilon$$
(3)

and

$$Y_{2X_i} = \beta_0 + \sum_{i=1}^p \beta_i X_i + \sum_{1 \le i < j \le p} \beta_{ij} (X_i \times X_j) + \epsilon \tag{4}$$

where p is the number of pipeline choices from Table 1. The first model studies the effect of each pipeline choice, include all pairwise interactions, on the average square deviation from the official NIH count matrix. The second model does the same, but for the p-values from a differential expression analysis.

Simulated Regression Power Analysis

```
set.seed(33025)
num predictors <- 50
num_sig_predictors <- 5</pre>
alpha <- 0.05
effect_size_mean <- 0.1</pre>
effect_size_sd <- 0.05
iter <- 100
sample_sizes \leftarrow seq(from = 60, to = 200, by = 10)
simulate_reg <- function(sample_size, num_predictors, num_sig_predictors,</pre>
                             effect_size_mean, effect_size_sd){
    X <- matrix(rnorm(sample_size * num_predictors),</pre>
                        nrow = sample size, ncol = num predictors)
    colnames(X) <- paste0("X", 1:num_predictors)</pre>
    true_coef <- rnorm(num_sig_predictors, effect_size_mean, effect_size_sd)</pre>
    true_coef <- c(true_coef, rep(0, num_predictors - num_sig_predictors))</pre>
    Y <- X %*% true_coef + rnorm(sample_size)
    data <- data.frame(Y, X)</pre>
```

```
lm_fit \leftarrow lm(Y \sim X, data = data)
    p_values <- summary(lm_fit)$coefficient[, 4][-1]</pre>
    return(p_values)
}
power_reg <- function(sample_sizes, num_predictors, num_sig_predictors,</pre>
                        effect size mean, effect size sd, iter){
    df <- data.frame()</pre>
    for(sample_size in sample_sizes){
        type_2_errors <- 0</pre>
        type_1_errors <- 0</pre>
        for(i in 1:iter){
             p_values <- simulate_reg(</pre>
                  sample_size, num_predictors, num_sig_predictors,
                  effect_size_mean, effect_size_sd
             type_2_errors <- type_2_errors +</pre>
                  sum(p_values[num_sig_predictors] > alpha)
             type_1_errors <- type_1_errors +</pre>
                  sum(p_values[(num_sig_predictors + 1):num_predictors] < alpha)</pre>
        }
        power <- 1 - (type_2_errors / (iter * num_sig_predictors))</pre>
        type_1_error_rate <- (</pre>
             type_1_errors / (iter * (num_predictors - num_sig_predictors))
        df <- rbind(df, c(sample_size, power, type_1_error_rate))</pre>
    colnames(df) <- c("n", "Power", "Type I Error Rate")</pre>
    return(df)
power_df <- power_reg(sample_sizes, num_predictors, num_sig_predictors,</pre>
           effect_size_mean, effect_size_sd, iter)
```

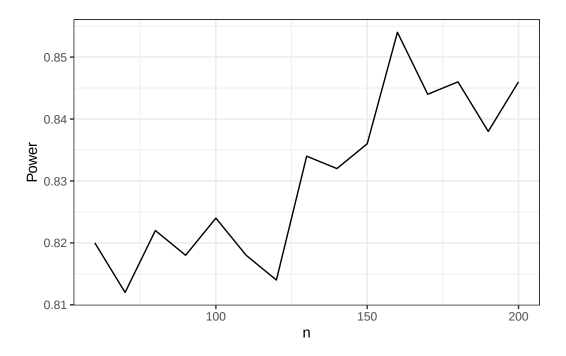


Figure 1: Multiple Linear Regression t-test Simulation Based Power Curve

Quality Score Variance Due to Fasterq-dump Options

```
if (!require("BiocManager", quietly = TRUE))
    install.packages("BiocManager")

BiocManager::install("ShortRead")

BiocManager::install("Rsubread")

library(ShortRead)

sample_1_fq_1 <- readFastq("./data/dump_1/SRR31476642.fastq")
    sample_1_fq_2 <- readFastq("./data/dump_2/SRR31476642.fastq")
    sample_1_fq_3 <- readFastq("./data/dump_3/SRR31476642.fastq")

sample_1_fq_2_qual <- as(quality(sample_1_fq_1), "matrix")
    sample_1_fq_2_qual <- as(quality(sample_1_fq_2), "matrix")
    sample_1_fq_3_qual <- as(quality(sample_1_fq_3), "matrix")

sample_1_fq_13_qual_diff <- sample_1_fq_1_qual - sample_1_fq_3_qual
    sample_1_fq_12_qual_diff <- sample_1_fq_1_qual - sample_1_fq_2_qual

mean(sample_1_fq_13_qual_diff)
mean(sample_1_fq_12_qual_diff)</pre>
```

Comparing Alignment Accuracy

```
fastq_files <- list.files(
    path = "./data/dump_1", pattern = "\\.fastq$", full.names = TRUE
)</pre>
```

```
library(Rsubread)
buildindex(basename="hg19_g1k",
           reference="./data/human_g1k_v37.fasta",
           memory=3600
)
align reads <- function(file, index base, output dir) {</pre>
  align(
    index = index_base,
    readfile1 = file,
    output_file = file.path(output_dir, paste0(basename(file), ".bam")),
    nthreads = 4
}
trim_reads <- function(file, quality_threshold = 20, min_length = 30) {</pre>
    fq <- readFastq(file)</pre>
    fq_filtered <- fq[
        alphabetScore(quality(fq)) >= quality_threshold & width(fq) >= min_length
    output_file <- sub(".fastq", "_trimmed.fastq", file)</pre>
    writeFastq(fq_filtered, output_file, compress = FALSE)
}
```

- Arora, S., Pattwell, S. S., Holland, E. C., and Bolouri, H. (2020), "Variability in estimated gene expression among commonly used RNA-seq pipelines," *Scientific reports*, Nature Publishing Group UK London, 10, 2734.
- Tong, L., Wu, P.-Y., Phan, J. H., Hassazadeh, H. R., Tong, W., and Wang, M. D. (2020), "Impact of RNA-seq data analysis algorithms on gene expression estimation and downstream prediction," *Scientific reports*, Nature Publishing Group UK London, 10, 17925.
- Vieth, B., Parekh, S., Ziegenhain, C., Enard, W., and Hellmann, I. (2019), "A systematic evaluation of single cell RNA-seq analysis pipelines," *Nature communications*, Nature Publishing Group UK London, 10, 4667.