# Differential analyses for RNA-seq: transcript-level estimates improve gene-level inferences <br> Supplementary Material 

Charlotte Soneson, Michael I. Love, Mark D. Robinson

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## 1 Simulation details, sim2 data set

The $\operatorname{sim} 2$ data set consists of simulated sequencing reads from the human chromosome 1 . The sequencing parameters as well as underlying TPM values for the 15,677 transcripts in one of the two simulated conditions were estimated using RSEM v1.2.21 [6] from the ERS326990 sample from the ArrayExpress data set with accession number $E_{-} M T A B_{-} 1733$. We simulated three biological replicates from each of two conditions.

Three types of differential abundance were introduced between the two conditions, all assigned to genes with estimated TPMs between 0.1 and 1000 in the reference data set (the "eligible" genes):

- DGE (differential gene expression) was introduced for $420(11 \%)$ of the eligible genes. For these genes, the overall gene expression (TPM) was modified, but the relative isoform abundances within the genes were kept constant. The TPMs of the genes in condition 2 were obtained by shuffling the TPMs of the genes in condition 1, in order to create a range of fold changes and keep the total TPM for differential genes constant across samples. All genes with simulated DGE also show DTE (differential transcript expression), but no DTU (differential transcript usage).
- DTE (differential transcript expression) was introduced for $422(11 \%)$ of the eligible genes (selected among genes with at least two isoforms). For each of these genes, $10 \%$ of the isoforms were selected to be differential between the conditions, again by shuffling their TPMs from condition 1 to generate TPMs in condition 2. In total, 528 transcripts were differentially expressed in these genes. All genes with simulated DTE also show DGE (since the total TPM of the gene is modified) and DTU (since the relative usage of the isoforms is modified). However, the overall fold changes of the DTE genes is often lower than those of the DGE genes, since only a subset of the transcripts are affected.
- DTU (differential transcript usage) was introduced for $420(11 \%)$ of the eligible genes (selected among genes with at least two isoforms). For each of these genes, the total TPM was kept constant between the two conditions, but the relative isoform usage for a given gene with DTU in condition 2 was sampled randomly among the relative isoform usages for all genes in the data set with the same number of isoforms. All genes with simulated DTU also show DTE (since at least one isoform changes expression), but no DGE.

Based on the expected number of reads for each transcript in each condition, we simulated three biological replicates per condition from a negative binomial distribution, using a mean-dispersion relationship estimated from large experimental data sets (see [9] for more information on how these estimates were generated). The individual sample counts were then converted to TPMs and used as inputs to RSEM to simulate 10,000,000 paired-end reads of length 100 bp for each sample.

From the three types of differential genes in this data set, we can also define three sets of "truly differential" features:

- Differentially expressed transcripts
- Genes with between-condition differences in overall expression (the sum of the transcript TPMs)
- Genes with between-condition differences in at least one associated transcript

As we show in the manuscript, it is important to select the type of interest before choosing a quantification and representation method, since different methods have unequal power to detect different types of signals.

## 2 Experimental data sets

In addition to the two simulated data sets, the results presented in this manuscripts are based on the following four experimental data sets:

- Bottomly [1] This data set contains striatal tissue samples from 11 mice of the DBA/2J strain and 10 mice of the C57BL/6J strain. We downloaded the FASTQ files with sequencing reads from SRA (see Supplementary File 4 for exact file names and URLs) and used the GRCm38.82 Ensembl annotation for analysis.
- GSE64570 11 We use a subset of the data, consisting of 3 replicates of wildtype zebrafish injected with water, and 3 replicates of tlr5a morphants injected with Flagellin. We downloaded the FASTQ files from SRA (see Supplementary File 5 for exact file names and URLs) and used the GRz10 Ensembl annotation for analysis.
- GSE69244 4 Also here, we use a subset of the data, consisting of 3 replicates of 10-month old SAMP8 mice given vehicle diet for 7 months and 3 replicates of 10 -month old SAMP8 mice treated with the Alzheimer's disease drug candidate J147 for 7 months. We downloaded the FASTQ files from SRA (see Supplementary File 6 for exact file names and URLs) and used the GRCm38.82 Ensembl annotation for analysis.
- GSE72165 3 This data set contains 3 samples each from the carotid body and the adrenal medulla in mouse. We downloaded the FASTQ files from SRA (see Supplementary File 7 for exact file names and URLs) and used the GRCm38.82 Ensembl annotation for analysis.


## 3 Quantification

Abundances were quantified in two different ways:

- Transcript abundance quantification with Salmon [8]. We generated a Salmon index from the cDNA fasta file from Ensembl GRCh37.71 (for the human data sets), Ensembl GRCm38.82 (for the mouse data sets) or Ensembl GRz10 (for the zebrafish data set), and used the quasi-alignment approach to obtain TPM and count estimates for each transcript in each of the samples. In order to estimate the variability in the estimates, we generated 30 bootstrap replicates.
- Gene abundance quantification with featureCounts [7, after alignment of the reads to the genome using STAR v2.4.2a [5]. The gene models were defined by the Ensembl GTF files from the same release as the cDNA fasta files used for Salmon.

As an alternative to Salmon, we also applied kallisto [2] to obtain TPM and count estimates for transcripts. The results were very similar to those obtained with salmon and are not shown in the manuscript.

## 4 Deriving offsets from transcript lengths and abundances

Here we motivate using the average transcript length as an offset for performing differential expression on summarized fragment counts from all transcripts of a gene. We use the notation of the Methods Supplement of the Cuffdiff2 paper [10]. Suppose a single gene $G$ with transcripts indexed by $t$, two samples $a$ and $b$ with observed fragment counts for the gene $G$ given by $X_{a}$ and $X_{b}$, and underlying, unobserved fragment counts from each transcript $t$ given by $X_{t a}$ and $X_{t b}$. The effective length of transcript $t$, the number of positions along the transcript where a fragment can align, is given by $\bar{l}_{t}$. For notational simplicity, we suppose this is the same for samples, but the following holds for sample-specific effective lengths as well. The true expression of transcript $t$ is given by $\rho_{t a}$ and $\rho_{t b}$ for the two samples.

The fold change in expression for the gene is given by $\Delta_{\rho_{G}}$. If we knew the true fragment counts for each transcript, we could estimate this using the sum of length corrected counts for all transcripts for sample $a$ and dividing by the same quantity for sample $b$.

$$
\begin{equation*}
\hat{\Delta}_{\rho_{G}}=\frac{\sum_{t} X_{t a} / \bar{l}_{t}}{\sum_{t} X_{t b} / \bar{l}_{t}} \tag{1}
\end{equation*}
$$

We define $\theta_{t a}$ as the ratio of total expression for gene $G$ from transcript $t$, such that $\sum_{t} \theta_{t a}=1$

$$
\theta_{t a}=\frac{\rho_{t a}}{\sum_{t^{\prime}} \rho_{t^{\prime} a}}
$$

The expected number of fragments coming from transcript $t$ is then given by the following product:

$$
E\left(X_{t a}\right)=X_{a} \frac{\theta_{t a} \bar{l}_{t}}{\sum_{t^{\prime}} \theta_{t^{\prime} a} \bar{l}_{t^{\prime}}}
$$

We now return to the numerator of Eq. 1. The expected value of this quantity can be broken down as follows:

$$
\begin{aligned}
E\left(\sum_{t} X_{t a} / \bar{l}_{t}\right) & =\sum_{t} E\left(X_{t a}\right) / \bar{l}_{t} \\
& =\sum_{t} X_{a} \frac{\theta_{t a}}{\sum_{t^{\prime}} \theta_{t^{\prime} a} \bar{l}_{t^{\prime}}} \\
& =\frac{X_{a}}{\sum_{t^{\prime}} \theta_{t^{\prime} a} \bar{l}_{t^{\prime}}} \sum_{t} \theta_{t a} \\
& =\frac{X_{a}}{\sum_{t^{\prime}} \theta_{t^{\prime} a} \bar{l}_{t^{\prime}}}
\end{aligned}
$$

We can therefore estimate $\Delta_{\rho_{G}}$ using expected values for the numerator and denominator:

$$
\frac{E\left(\sum_{t} X_{t a} / \bar{l}_{t}\right)}{E\left(\sum_{t} X_{t b} / \bar{l}_{t}\right)}=\frac{X_{a} / \sum_{t^{\prime}} \theta_{t^{\prime} a} \bar{l}_{t^{\prime}}}{X_{b} / \sum_{t^{\prime}} \theta_{t^{\prime} b} \bar{l}_{t^{\prime}}}
$$

This final line shows that the ratio of summarized fragment counts for the gene can be corrected for bias due to changes in transcript usage by using the average transcript length for each sample, where the average is weighted by abundances $\theta_{t a}$. As we do not have access to the true abundances of each transcript, estimates of transcript abundance $\hat{\theta}_{t a}$ are used.

## 5 Supplementary Figures



Supplementary Figure 1: Accuracy of transcript and gene TPM estimates from Salmon and featureCounts, restricted to features with non-zero true and estimated TPMs (sim2). Spearman correlations are indicated in each panel.


Supplementary Figure 2: Accuracy of transcript and gene TPM estimates from Salmon and featureCounts ( $\operatorname{sim} 1$ ). Top: all features. Bottom: restricted to features with non-zero true and estimated TPMs. Spearman correlations are indicated in each panel.


Supplementary Figure 3: Distribution of coefficients of variation for gene and transcript TPM estimates from Salmon (sim1), estimated across 30 bootstrap replicates of one of the simulated samples (sample1).


Supplementary Figure 4: Distribution of coefficients of variation for gene and transcript TPM estimates from Salmon (Bottomly), estimated across 30 bootstrap replicates of the SRR099223 sample.


Supplementary Figure 5: Comparison of different approaches for finding DTE, using edgeR (sim2). Gene-level quantification followed by differential analysis (i.e., DGE) is less powerful than transcript-level quantification and statistical testing aggregated on gene-level (i.e., DTE). This highlights the importance of appropriate specification of the question of interest before starting an analysis.


Supplementary Figure 6: Correlation among counts obtained with different approaches in one of the simulated samples (sampleA1) from the $\operatorname{sim} 2$ data set. Top: all genes. Bottom: only genes with non-zero counts with all methods. For visualization and correlation calculations, the counts were log2-transformed after adding a pseudo-count of 1.


Supplementary Figure 7: Correlation among counts obtained with different approaches in one of the simulated samples (sample1) from the sim1 data set. Top: all genes. Bottom: only genes with non-zero counts with all methods. For visualization and correlation calculations, the counts were log2-transformed after adding a pseudo-count of 1 .


Supplementary Figure 8: Correlation among counts obtained with different approaches in the SRR099223 sample from the Bottomly data set. Top: all genes. Bottom: only genes with non-zero counts with all methods. For visualization and correlation calculations, the counts were log2-transformed after adding a pseudo-count of 1 . featureCountsMM is similar to featureCounts, but includes multi-mapping reads by distributing them evenly between the mapping locations.


Supplementary Figure 9: Distribution of nominal p-values obtained by performing DGE using edgeR (top) and DESeq2 (bottom) on gene-level count matrices obtained with different approaches (sim2).


Supplementary Figure 10: Distribution of nominal p-values obtained by performing DGE using edge $R$ (top) and DESeq2 (bottom) on gene-level count matrices obtained with different approaches (sim1).


Supplementary Figure 11: Distribution of nominal p-values obtained by performing DGE using edge $R$ (top) and DESeq2 (bottom) on gene-level count matrices obtained with different approaches (Bottomly).


Supplementary Figure 12: Mean-dispersion relationship obtained from edgeR (top) and DESeq2 (bottom) applied to gene-level count matrices obtained with different approaches (sim2).


Supplementary Figure 13: Mean-dispersion relationship obtained from edgeR (top) and DESeq2 (bottom) applied to gene-level count matrices obtained with different approaches (sim1).


Supplementary Figure 14: Mean-dispersion relationship obtained from edgeR (top) and DESeq2 (bottom) applied to gene-level count matrices obtained with different approaches (Bottomly).


Supplementary Figure 15: Stratification of differential gene expression detection performance by the presence of differential isoform usage, using DESeq2 (sim2).


Supplementary Figure 16: Overlaps among collections of significant genes found with edgeR (top) and DESeq2 (bottom) for three different experimental RNA-seq data sets, using three different counting pipelines. In these data sets, the majority of the significant genes are found with all counting methods, suggesting that for many real data sets, simple gene counting produces overall similar results as methods accounting for transcript-level expression.


Supplementary Figure 17: Accuracy of overall gene log-fold change estimates based on results from edge $R$, using different count matrices (sim2). Only genes with non-zero true log-fold changes are considered.

Supplementary Table 1: Spearman correlations quantifying the accuracy of the overall gene log-fold change estimates based on different count matrices (sim2), shown in Supplementary Figure 17

|  | featureCounts | featureCounts_avetxl | scaledTPM_salmon | simplesum_salmon | simplesum_salmon_avetxl |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |
| No DTU | 0.834 | 0.839 | 0.816 | 0.823 | 0.828 |
| DTU | 0.725 | 0.788 | 0.796 | 0.735 | 0.783 |



Supplementary Figure 18: Correlations between estimated log2-fold changes obtained with the scaledTPM and simplesum count matrices, using edgeR. The color indicates whether or not the gene is expressed in only one condition.


Supplementary Figure 19: Correlations between estimated log2-fold changes obtained with the scaledTPM and simplesum count matrices, using DESeq2. The color indicates whether or not the gene is expressed in only one condition.

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